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# SLEPc for Python

Release 3.24.2

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### Abstract

This document describes [slepc4py](#), a [Python](#) wrapper to the [SLEPc](#) libraries.

[SLEPc](#) is a software package for the parallel solution of large-scale eigenvalue problems. It can be used for computing eigenvalues and eigenvectors of large, sparse matrices, or matrix pairs, and also for computing singular values and vectors of a rectangular matrix. It also provides more advanced functionality such as solvers for nonlinear eigenvalue problems (either polynomial or general) and matrix functions.

[SLEPc](#) relies on [PETSc](#) for basic functionality such as the representation of matrices and vectors, and the solution of linear systems of equations. Thus, [slepc4py](#) must be used together with its companion [petsc4py](#).

## 1 Discussion and Support

- You can write to the [PETSc Users Mailing List](#)
- See also the [contact information for SLEPc](#)

- Issue Tracker: <https://gitlab.com/slepc/slepc/-/issues>
- Git Repository: <https://gitlab.com/slepc/slepc.git> (the source code is in `src/binding/slepc4py`)
- Current release in PyPI: <https://pypi.org/project/slepc4py/>

## 2 Acknowledgments

L. Dalcin was partially supported by the Extreme Computing Research Center (ECRC), Division of Computer, Electrical, and Mathematical Sciences & Engineering (CEMSE), King Abdullah University of Science and Technology (KAUST).

See additional acknowledgements related to [the SLEPc project](#).

## 3 Contents

### 3.1 Overview

*SLEPc for Python* (slepc4py) is a Python package that provides convenient access to the functionality of SLEPc.

SLEPc<sup>1,2</sup> implements algorithms and tools for the numerical solution of large, sparse eigenvalue problems on parallel computers. It can be used for linear eigenvalue problems in either standard or generalized form, with real or complex arithmetic. It can also be used for computing a partial SVD of a large, sparse, rectangular matrix, and to solve nonlinear eigenvalue problems (polynomial or general). Additionally, SLEPc provides solvers for the computation of the action of a matrix function on a vector.

SLEPc is intended for computing a subset of the spectrum of a matrix (or matrix pair). One can for instance approximate the largest magnitude eigenvalues, or the smallest ones, or even those eigenvalues located near a given region of the complex plane. Interior eigenvalues are harder to compute, so SLEPc provides different methodologies. One such method is to use a spectral transformation. Cheaper alternatives are also available.

### Features

Currently, the following types of eigenproblems can be addressed:

- Standard eigenvalue problem,  $Ax = \lambda x$ , either for Hermitian or non-Hermitian matrices.
- Generalized eigenvalue problem,  $Ax = \lambda Bx$ , either Hermitian positive-definite or not.
- Partial singular value decomposition of a rectangular matrix,  $Au = \sigma v$ .
- Generalized singular values of a matrix pair,  $Ax = cu$ ,  $Bx = sv$ .
- Polynomial eigenvalue problem,  $P(\lambda) = 0$ .
- Nonlinear eigenvalue problem,  $T(\lambda) = 0$ .
- Computing the action of a matrix function on a vector,  $w = f(\alpha A)v$ .

For the linear eigenvalue problem, the following methods are available:

- Krylov eigensolvers, particularly Krylov-Schur, Arnoldi, and Lanczos.
- Davidson-type eigensolvers, including Generalized Davidson and Jacobi-Davidson.
- Subspace iteration and single vector iterations (inverse iteration, RQI).
- Conjugate gradient methods such as LOBPCG.

<sup>1</sup> J. E. Roman, C. Campos, L. Dalcin, E. Romero, A. Tomas. SLEPc Users Manual. DSIC-II/24/02 - Revision 3.24. D. Sistemas Informaticos y Computacion, Universitat Politecnica de Valencia. 2025.

<sup>2</sup> Vicente Hernandez, Jose E. Roman and Vicente Vidal. SLEPc: A Scalable and Flexible Toolkit for the Solution of Eigenvalue Problems, ACM Trans. Math. Softw. 31(3), pp. 351-362, 2005. <https://doi.org/10.1145/1089014.1089019>

- A contour integral solver using high-order moments.

For singular value computations, the following alternatives can be used:

- Use an eigensolver via the cross-product matrix  $A^*A$  or the cyclic matrix  $\begin{bmatrix} 0 & A \\ A^* & 0 \end{bmatrix}$ .
- Explicitly restarted Lanczos bidiagonalization.
- Implicitly restarted Lanczos bidiagonalization (thick-restart Lanczos).
- A basic randomized solver.

For polynomial eigenvalue problems, the following methods are available:

- Use an eigensolver to solve the generalized eigenvalue problem obtained after linearization.
- TOAR and Q-Arnoldi, memory efficient variants of Arnoldi for polynomial eigenproblems.
- Jacobi-Davidson for polynomial eigenproblems.
- A contour integral solver using high-order moments.

For general nonlinear eigenvalue problems, the following methods can be used:

- Solve a polynomial eigenproblem obtained via polynomial interpolation.
- Rational interpolation and linearization (NLEIGS).
- Newton-type methods such as SLP or RII.
- A subspace projection method (nonlinear Arnoldi).
- A contour integral solver using high-order moments.

Computation of interior eigenvalues is supported by means of the following methodologies:

- Spectral transformations, such as shift-and-invert. This technique implicitly uses the inverse of the shifted matrix  $A - \sigma I$  in order to compute eigenvalues closest to a given target value,  $\sigma$ .
- Harmonic extraction, a cheap alternative to shift-and-invert that also tries to approximate eigenvalues closest to a target,  $\sigma$ , but without requiring a matrix inversion.

Other remarkable features include:

- High computational efficiency, by using NumPy and SLEPc under the hood.
- Data-structure neutral implementation, by using efficient sparse matrix storage provided by PETSc. Implicit matrix representation is also available by providing basic operations such as matrix-vector products as user-defined Python functions.
- Run-time flexibility, by specifying numerous setting at the command line.
- Ability to do the computation in parallel and/or using GPUs.

## Components

SLEPc provides the following components, which are mirrored by slepc4py for its use from Python. The first six components are solvers for different classes of problems, while the rest can be considered auxiliary object.

### EPS

The Eigenvalue Problem Solver is the component that provides all the functionality necessary to define and solve an eigenproblem. It provides mechanisms for completely specifying the problem: the problem type (e.g. standard symmetric), number of eigenvalues to compute, part of the spectrum of interest. Once the problem has been defined, a collection of solvers can be used to compute the required solutions. The behavior of the solvers can be tuned by means of a few parameters, such as the maximum dimension of the subspace to be used during the computation.

#### **SVD**

This component is the analog of EPS for the case of Singular Value Decompositions. The user provides a rectangular matrix and specifies how many singular values and vectors are to be computed, whether the largest or smallest ones, as well as some other parameters for fine tuning the computation. Different solvers are available, as in the case of EPS.

#### **PEP**

This component is the analog of EPS for the case of Polynomial Eigenvalue Problems. The user provides the coefficient matrices of the polynomial. Several parameters can be specified, as in the case of EPS. It is also possible to indicate whether the problem belongs to a special type, e.g., symmetric or gyroscopic.

#### **NEP**

This component covers the case of general nonlinear eigenproblems,  $T(\lambda) = 0$ . The user provides the parameter-dependent matrix  $T$  via the split form or by means of callback functions.

#### **MFN**

This component provides the functionality for computing the action of a matrix function on a vector. Given a matrix  $A$  and a vector  $b$ , the call `MFNSolve(mfn,b,x)` computes  $x = f(A)b$ , where  $f$  is a function such as the exponential.

#### **LME**

This component provides the functionality for solving linear matrix equations such as Lyapunov or Sylvester where the solution has low rank.

#### **ST**

The Spectral Transformation is a component that provides convenient implementations of common spectral transformations. These are simple transformations that map eigenvalues to different positions, in such a way that convergence to wanted eigenvalues is enhanced. The most common spectral transformation is shift-and-invert, that allows for the computation of eigenvalues closest to a given target value.

#### **BV**

This component encapsulates the concept of a set of Basis Vectors spanning a vector space. This component provides convenient access to common operations such as orthogonalization of vectors. The BV component is usually not required by end-users.

#### **DS**

The Dense System (or Direct Solver) component, used internally to solve dense eigenproblems of small size that appear in the course of iterative eigensolvers.

#### **FN**

A component used to define mathematical functions. This is required by the end-user for instance to define function  $T(\cdot)$  when solving nonlinear eigenproblems with NEP in split form.

#### **RG**

A component used to define a region of the complex plane such as an ellipse or a rectangle. This is required by end-users in some cases such as contour-integral eigensolvers.

In addition to the above components, some extra functionality is provided in the `Sys` and `Util` sections.

## **3.2 Tutorial**

Below we provide links to tutorial examples intended for learning the basics of `slepc4py`. For additional information, the reader is referred to:

- `slepc4py` [Reference](#) (manual pages for all `slepc4py` classes and methods).
- [SLEPc documentation](#).
- [petsc4py documentation](#).

## Commented source of demo examples

- To get started, we recommend having a look at the source code of example `demo/ex1.py`, that has comments inserted inline. It is also [available online here](#). This example solves a standard symmetric eigenvalue problem.
- Demo examples for other problems, including linear and nonlinear eigenvalue problems and singular value problems, are available in the section *[slepc4py demos](#)*.

## 3.3 Installation

### Install from PyPI using pip

You can use **pip** to install `slepc4py` and its dependencies.

If you have a working MPI implementation and the `mpicc` compiler wrapper is on your search path, it is highly recommended to install `mpi4py` first:

```
$ python -m pip install mpi4py
```

Ensure you have `NumPy` and `petsc4py` installed:

```
$ python -m pip install numpy petsc4py
```

Finally, install `slepc4py`:

```
$ python -m pip install slepc slepc4py
```

If you already have working PETSc and SLEPc installs, set environment variables `SLEPC_DIR` and `PETSC_DIR` (and perhaps `PETSC_ARCH` for non-prefix installs) to appropriate values and next use **pip**:

```
$ export SLEPC_DIR=/path/to/slepc
$ export PETSC_DIR=/path/to/petsc
$ export PETSC_ARCH=arch-linux2-c-opt
$ python -m pip install petsc4py slepc4py
```

### Install from the SLEPc source tree

If you also want to install `petsc4py` from the PETSc source tree, follow the instructions in the [petsc4py installation](#) page.

Set the `PETSC_DIR` and `PETSC_ARCH` environment variables, as well as `SLEPC_DIR`. Follow the instructions to [build SLEPc](#). Then `cd` to the top of the SLEPc source tree and run:

```
$ python -m pip install src/binding/slepc4py
```

The installation of `slepc4py` supports multiple `PETSC_ARCH` in the form of a colon separated list:

```
$ PETSC_ARCH='arch-0:...:arch-N' python -m pip install src/binding/slepc4py
```

If you are cross-compiling, and the `numpy` module cannot be loaded on your build host, then before invoking **pip**, set the `NUMPY_INCLUDE` environment variable to the path that would be returned by `import numpy; numpy.get_include()`:

```
$ export NUMPY_INCLUDE=/usr/lib/pythonX/site-packages/numpy/core/include
```

## 3.4 Citations

If SLEPc for Python has been significant to a project that leads to an academic publication, please acknowledge that fact by citing the project.

- L. Dalcin, P. Kler, R. Paz, and A. Cosimo, *Parallel Distributed Computing using Python*, Advances in Water Resources, 34(9):1124-1139, 2011. <https://doi.org/10.1016/j.advwatres.2011.04.013>
- V. Hernandez, J.E. Roman, and V. Vidal, *SLEPc: A scalable and flexible toolkit for the solution of eigenvalue problems*, ACM Transactions on Mathematical Software, 31(3):351-362, 2005. <https://doi.org/10.1145/1089014.1089019>

There is also a list of [SLEPc-related references](#).

## 3.5 CHANGES

### Release 3.24

- Update to SLEPc 3.24.
- Support (opt-in via setting the environment variable `SLEPC4PY_BUILD_PYSABI=1`) for building with `Py_LIMITED_API` (Python Stable ABI) under Python 3.10+ (requires Cython 3.1+).
- Add support for standard Python operators for *BV* and *FN* classes.
- Add new *LME* class.

### Release 3.23

- Update to SLEPc 3.23.

### Release 3.22

- Update to SLEPc 3.22.
- In `slep4py` now *EPS.getEigenpair()* and *EPS.getEigenvalue()* will return a real value instead of a complex, if the problem is of Hermitian or generalized Hermitian type.

### Release 3.21

- Update to SLEPc 3.21.

### Release 3.20

- Update to SLEPc 3.20.

### Release 3.19

- Update to SLEPc 3.19.

### Release 3.18

- Update to SLEPc 3.18.

### Release 3.17

- Update to SLEPc 3.17.

### **Release 3.16**

- Update to SLEPc 3.16.

### **Release 3.15**

- Update to SLEPc 3.15.
- Updates in installation scripts.

### **Release 3.14**

- Update to SLEPc 3.14.

### **Release 3.13**

- Update to SLEPc 3.13.

### **Release 3.12**

- Update to SLEPc 3.12.

### **Release 3.11**

- Update to SLEPc 3.11.

### **Release 3.10**

- Update to SLEPc 3.10.

### **Release 3.9**

- Update to SLEPc 3.9.

### **Release 3.8**

- Update to SLEPc 3.8.

### **Release 3.7**

- Update to SLEPc 3.7.

### **Release 3.6**

- Update to SLEPc 3.6.

### **Release 3.5**

- Update to SLEPc 3.5.
- Add RG class introduced in SLEPc 3.5 release.
- Add PySlepcXXX\_New/Get C API functions.
- Fix compilation problem with complex scalars on OS X.
- Fix outdated SWIG interface file.

### Release 3.4

- Update to SLEPc 3.4.

### Release 3.3

- Update to SLEPc 3.3.
- Regenerate the wrappers using Cython 0.18 and fix binary compatibility issues with petsc4py 3.3.1.

### Release 1.2

- Update to SLEPc 3.2.

### Release 1.1

- Support for new QEP quadratic eigenproblem solver in SLEPc.
- Support for `pip install slepc4py` to download and install SLEPc.
- Support for PETSc/SLEPc static library builds (Linux-only).
- Preliminary support for Python 3.

### Release 1.0.0

- This is the first release of the all-new, Cython-based, implementation of *SLEPc for Python*.

## 3.6 Reference

<code>slepc4py</code>	SLEPc for Python
<code>slepc4py.typing</code>	Typing support.
<code>slepc4py.SLEPc</code>	Scalable Library for Eigenvalue Problem Computations

### slepc4py

#### SLEPc for Python

This package is an interface to [SLEPc](#) libraries.

[SLEPc](#) (the Scalable Library for Eigenvalue Problem Computations) is a software library for the solution of large scale sparse eigenvalue problems on parallel computers. It is an extension of [PETSc](#) and can be used for either standard or generalized eigenproblems, with real or complex arithmetic. It can also be used for computing a partial SVD of a large, sparse, rectangular matrix, and to solve nonlinear eigenvalue problems (polynomial or general). Additionally, SLEPc provides solvers for the computation of the action of a matrix function on a vector.

#### Functions

<code>get_config()</code>	Return a dictionary with information about SLEPc.
<code>get_include()</code>	Return the directory in the package that contains header files.
<code>init([args, arch, comm])</code>	Initialize SLEPc.



## slepc4py.get\_config

slepc4py.get\_config()

Return a dictionary with information about SLEPc.

### Return type

dict[str, str]

## slepc4py.get\_include

slepc4py.get\_include()

Return the directory in the package that contains header files.

Extension modules that need to compile against slepc4py should use this function to locate the appropriate include directory.

### Example

Using Python distutils or NumPy distutils:

```
import petsc4py, slepc4py
Extension('extension_name', ...
        include_dirs=[...,
                      petsc4py.get_include(),
                      slepc4py.get_include(),])
```

### Return type

str

## slepc4py.init

slepc4py.init(args=None, arch=None, comm=None)

Initialize SLEPc.

### Parameters

- **args** (str | list[str] | None) – Command-line arguments, usually the `sys.argv` list.
- **arch** (str | None) – Specific configuration to use.
- **comm** (Intracomm | None) – MPI communicator.

### Return type

None

### Notes

This function should be called only once, typically at the very beginning of the bootstrap script of an application.

## slepc4py.typing

Typing support.

## Attributes

<code>Scalar</code>	Scalar type.
<code>ArrayInt</code>	Array of <code>int</code> .
<code>ArrayReal</code>	Array of <code>float</code> .
<code>ArrayComplex</code>	Array of <code>complex</code> .
<code>ArrayScalar</code>	Array of <code>Scalar</code> numbers.
<code>LayoutSizeSpec</code>	<code>int</code> or 2-tuple of <code>int</code> describing the layout sizes.
<code>EPSStoppingFunction</code>	<code>EPS</code> stopping test callback.
<code>EPSArbitraryFunction</code>	<code>EPS</code> arbitrary selection callback.
<code>EPSEigenvalueComparison</code>	<code>EPS</code> eigenvalue comparison callback.
<code>EPSMonitorFunction</code>	<code>EPS</code> monitor callback.
<code>PEPStoppingFunction</code>	<code>PEP</code> stopping test callback.
<code>PEPEigenvalueComparison</code>	<code>PEP</code> eigenvalue comparison callback.
<code>PEPMonitorFunction</code>	<code>PEP</code> monitor callback.
<code>NEPStoppingFunction</code>	<code>NEP</code> stopping test callback.
<code>NEPEigenvalueComparison</code>	<code>NEP</code> eigenvalue comparison callback.
<code>NEPMonitorFunction</code>	<code>NEP</code> monitor callback.
<code>NEPFunction</code>	<code>NEP</code> Function callback.
<code>NEPJacobian</code>	<code>NEP</code> Jacobian callback.
<code>SVDStoppingFunction</code>	<code>SVD</code> stopping test callback.
<code>SVDMonitorFunction</code>	<code>SVD</code> monitor callback.
<code>MFNMonitorFunction</code>	<code>MFN</code> monitor callback.
<code>LMEMonitorFunction</code>	<code>LME</code> monitor callback.

### `slepc4py.typing.Scalar`

`slepc4py.typing.Scalar = float | complex`

Scalar type.

Scalars can be either `float` or `complex` (but not both) depending on how PETSc was configured (`./configure --with-scalar-type=real|complex`).

### `slepc4py.typing.ArrayInt`

`slepc4py.typing.ArrayInt`

Array of `int`.

alias of `ndarray[tuple[Any, ...], dtype[int]]`

### `slepc4py.typing.ArrayReal`

`slepc4py.typing.ArrayReal`

Array of `float`.

alias of `ndarray[tuple[Any, ...], dtype[float]]`

### `slepc4py.typing.ArrayComplex`

`slepc4py.typing.ArrayComplex`

Array of `complex`.

alias of `ndarray[tuple[Any, ...], dtype[complex]]`

### **slepc4py.typing.ArrayScalar**

`slepc4py.typing.ArrayScalar`

Array of *Scalar* numbers.

alias of `ndarray[tuple[Any, ...], dtype[float | complex]]`

### **slepc4py.typing.LayoutSizeSpec**

`slepc4py.typing.LayoutSizeSpec = int | tuple[int, int]`

`int` or 2-`tuple` of `int` describing the layout sizes.

A single `int` indicates global size. A `tuple` of `int` indicates (`local_size`, `global_size`).

### **slepc4py.typing.EPSStoppingFunction**

`slepc4py.typing.EPSStoppingFunction`

*EPS* stopping test callback.

alias of `Callable[[EPS, int, int, int, int], ConvergedReason]`

### **slepc4py.typing.EPSArbitraryFunction**

`slepc4py.typing.EPSArbitraryFunction`

*EPS* arbitrary selection callback.

alias of `Callable[[float | complex, float | complex, Vec, Vec, float | complex, float | complex], [float | complex, float | complex]]`

### **slepc4py.typing.EPSEigenvalueComparison**

`slepc4py.typing.EPSEigenvalueComparison`

*EPS* eigenvalue comparison callback.

alias of `Callable[[float | complex, float | complex, float | complex, float | complex], int]`

### **slepc4py.typing.EPSMonitorFunction**

`slepc4py.typing.EPSMonitorFunction`

*EPS* monitor callback.

alias of `Callable[[EPS, int, int, ndarray[tuple[Any, ...], dtype[float | complex]], ndarray[tuple[Any, ...], dtype[float | complex]], ndarray[tuple[Any, ...], dtype[float]], int, None]`

### **slepc4py.typing.PEPStoppingFunction**

`slepc4py.typing.PEPStoppingFunction`

*PEP* stopping test callback.

alias of `Callable[[PEP, int, int, int, int], ConvergedReason]`

### **slepc4py.typing.PEPEigenvalueComparison**

`slepc4py.typing.PEPEigenvalueComparison`

*PEP* eigenvalue comparison callback.

alias of `Callable[[float | complex, float | complex, float | complex, float | complex], int]`

### **slepc4py.typing.PEPMonitorFunction**

`slepc4py.typing.PEPMonitorFunction`

*PEP* monitor callback.

alias of `Callable[[PEP, int, int, ndarray[tuple[Any, ...], dtype[float | complex]], ndarray[tuple[Any, ...], dtype[float | complex]], ndarray[tuple[Any, ...], dtype[float]], int], None]`

### **slepc4py.typing.NEPStoppingFunction**

`slepc4py.typing.NEPStoppingFunction`

*NEP* stopping test callback.

alias of `Callable[[NEP, int, int, int, int], ConvergedReason]`

### **slepc4py.typing.NEPEigenvalueComparison**

`slepc4py.typing.NEPEigenvalueComparison`

*NEP* eigenvalue comparison callback.

alias of `Callable[[float | complex, float | complex, float | complex, float | complex], int]`

### **slepc4py.typing.NEPMonitorFunction**

`slepc4py.typing.NEPMonitorFunction`

*NEP* monitor callback.

alias of `Callable[[NEP, int, int, ndarray[tuple[Any, ...], dtype[float | complex]], ndarray[tuple[Any, ...], dtype[float | complex]], ndarray[tuple[Any, ...], dtype[float]], int], None]`

### **slepc4py.typing.NEPFunction**

`slepc4py.typing.NEPFunction`

*NEP* Function callback.

alias of `Callable[[NEP, float | complex, Mat, Mat], None]`

### **slepc4py.typing.NEPJacobian**

`slepc4py.typing.NEPJacobian`

*NEP* Jacobian callback.

alias of `Callable[[NEP, float | complex, Mat], None]`

## slepc4py.typing.SVDStoppingFunction

slepc4py.typing.SVDStoppingFunction

*SVD* stopping test callback.

alias of `Callable[[SVD, int, int, int, int], ConvergedReason]`

## slepc4py.typing.SVDMonitorFunction

slepc4py.typing.SVDMonitorFunction

*SVD* monitor callback.

alias of `Callable[[SVD, int, int, ndarray[tuple[Any, ...], dtype[float]], ndarray[tuple[Any, ...], dtype[float]], int], None]`

## slepc4py.typing.MFNMonitorFunction

slepc4py.typing.MFNMonitorFunction

*MFN* monitor callback.

alias of `Callable[[MFN, int, float], None]`

## slepc4py.typing.LMEMonitorFunction

slepc4py.typing.LMEMonitorFunction

*LME* monitor callback.

alias of `Callable[[LME, int, float], None]`

## slepc4py.SLEPc

Scalable Library for Eigenvalue Problem Computations

### Classes

<i>BV</i>	Basis Vectors.
<i>DS</i>	Direct Solver (or Dense System).
<i>EPS</i>	Eigenvalue Problem Solver.
<i>FN</i>	Mathematical Function.
<i>LME</i>	Linear Matrix Equation.
<i>MFN</i>	Matrix Function.
<i>NEP</i>	Nonlinear Eigenvalue Problem Solver.
<i>PEP</i>	Polynomial Eigenvalue Problem Solver.
<i>RG</i>	Region.
<i>ST</i>	Spectral Transformation.
<i>SVD</i>	Singular Value Decomposition Solver.
<i>Sys</i>	System utilities.
<i>Util</i>	Other utilities such as the creation of structured matrices.

## slepc4py.SLEPc.BV

**class** slepc4py.SLEPc.BV

Bases: `Object`

Basis Vectors.

The *BV* package provides the concept of a block of vectors that represent the basis of a subspace. It is a convenient way of handling a collection of vectors that often operate together, rather than working with an array of `petsc4py.PETSc.Vec`.

## Enumerations

<i>MatMultType</i>	BV mat-mult types.
<i>OrthogBlockType</i>	BV block-orthogonalization types.
<i>OrthogRefineType</i>	BV orthogonalization refinement types.
<i>OrthogType</i>	BV orthogonalization types.
<i>SVDMethod</i>	BV methods for computing the SVD.
<i>Type</i>	BV type.

## `slepc4py.SLEPc.BV.MatMultType`

**class** `slepc4py.SLEPc.BV.MatMultType`

Bases: `object`

BV mat-mult types.

- *VECS*: Perform a matrix-vector multiply per each column.
- *MAT*: Carry out a Mat-Mat product with a dense matrix.

### See also

`BVMatMultType`

## Attributes Summary

<i>MAT</i>	Constant MAT of type <code>int</code>
<i>VECS</i>	Constant VECS of type <code>int</code>

## Attributes Documentation

**MAT:** `int` = *MAT*

Constant MAT of type `int`

**VECS:** `int` = *VECS*

Constant VECS of type `int`

## `slepc4py.SLEPc.BV.OrthogBlockType`

**class** `slepc4py.SLEPc.BV.OrthogBlockType`

Bases: `object`

BV block-orthogonalization types.

- *GS*: Gram-Schmidt, column by column.
- *CHOL*: Cholesky QR method.
- *TSQR*: Tall-skinny QR method.

- *TSQRCHOL*: Tall-skinny QR, but computing the triangular factor only.
- *SVQB*: SVQB method.

#### See also

[BVOrthogBlockType](#)

### Attributes Summary

<i>CHOL</i>	Constant CHOL of type <code>int</code>
<i>GS</i>	Constant GS of type <code>int</code>
<i>SVQB</i>	Constant SVQB of type <code>int</code>
<i>TSQR</i>	Constant TSQR of type <code>int</code>
<i>TSQRCHOL</i>	Constant TSQRCHOL of type <code>int</code>

### Attributes Documentation

**CHOL:** `int` = CHOL

Constant CHOL of type `int`

**GS:** `int` = GS

Constant GS of type `int`

**SVQB:** `int` = SVQB

Constant SVQB of type `int`

**TSQR:** `int` = TSQR

Constant TSQR of type `int`

**TSQRCHOL:** `int` = TSQRCHOL

Constant TSQRCHOL of type `int`

### `slepc4py.SLEPc.BV.OrthogRefineType`

**class** `slepc4py.SLEPc.BV.OrthogRefineType`

Bases: `object`

BV orthogonalization refinement types.

- *IFNEEDED*: Reorthogonalize if a criterion is satisfied.
- *NEVER*: Never reorthogonalize.
- *ALWAYS*: Always reorthogonalize.

#### See also

[BVOrthogRefineType](#)

## Attributes Summary

<i>ALWAYS</i>	Constant ALWAYS of type <code>int</code>
<i>IFNEEDED</i>	Constant IFNEEDED of type <code>int</code>
<i>NEVER</i>	Constant NEVER of type <code>int</code>

## Attributes Documentation

**ALWAYS:** `int` = ALWAYS

Constant ALWAYS of type `int`

**IFNEEDED:** `int` = IFNEEDED

Constant IFNEEDED of type `int`

**NEVER:** `int` = NEVER

Constant NEVER of type `int`

## slepc4py.SLEPc.BV.OrthogType

**class** slepc4py.SLEPc.BV.OrthogType

Bases: `object`

BV orthogonalization types.

- *CGS*: Classical Gram-Schmidt.
- *MGS*: Modified Gram-Schmidt.

### See also

BVOrthogType

## Attributes Summary

<i>CGS</i>	Constant CGS of type <code>int</code>
<i>MGS</i>	Constant MGS of type <code>int</code>

## Attributes Documentation

**CGS:** `int` = CGS

Constant CGS of type `int`

**MGS:** `int` = MGS

Constant MGS of type `int`

## slepc4py.SLEPc.BV.SVDMethod

**class** slepc4py.SLEPc.BV.SVDMethod

Bases: `object`

BV methods for computing the SVD.

- *REFINE*: Based on the SVD of the cross product matrix  $S^*S$ , with refinement.



- *QR*: Based on the SVD of the triangular factor of  $qr(S)$ .
- *QR\_CAA*: Variant of QR intended for use in communication-avoiding. Arnoldi.

#### See also

*BVSVDMethod*

### Attributes Summary

<i>QR</i>	Constant QR of type <i>int</i>
<i>QR_CAA</i>	Constant QR_CAA of type <i>int</i>
<i>REFINE</i>	Constant REFINE of type <i>int</i>

### Attributes Documentation

**QR:** *int* = QR

Constant QR of type *int*

**QR\_CAA:** *int* = QR\_CAA

Constant QR\_CAA of type *int*

**REFINE:** *int* = REFINE

Constant REFINE of type *int*

### slepc4py.SLEPc.BV.Type

**class** slepc4py.SLEPc.BV.Type

Bases: *object*

BV type.

- *MAT*: A *BV* stored as a dense *Mat*.
- *SVEC*: A *BV* stored as a single *Vec*.
- *VECS*: A *BV* stored as an array of independent *Vec*.
- *CONTIGUOUS*: A *BV* stored as an array of *Vec* sharing a contiguous array of scalars.
- *TENSOR*: A special *BV* represented in compact form as  $V = (I \otimes U)S$ .

#### See also

*BVType*

### Attributes Summary

<i>CONTIGUOUS</i>	Object CONTIGUOUS of type <i>str</i>
<i>MAT</i>	Object MAT of type <i>str</i>
<i>SVEC</i>	Object SVEC of type <i>str</i>
<i>TENSOR</i>	Object TENSOR of type <i>str</i>
<i>VECS</i>	Object VECS of type <i>str</i>

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Table 11 – continued from previous page

**Attributes Documentation****CONTIGUOUS:** `str` = CONTIGUOUSObject CONTIGUOUS of type `str`**MAT:** `str` = MATObject MAT of type `str`**SVEC:** `str` = SVECObject SVEC of type `str`**TENSOR:** `str` = TENSORObject TENSOR of type `str`**VECS:** `str` = VECSObject VECS of type `str`**Methods Summary**

<code>appendOptionsPrefix([prefix])</code>	Append to the prefix used for searching for all BV options in the database.
<code>applyMatrix(x, y)</code>	Multiply a vector with the matrix associated to the bilinear form.
<code>copy([result])</code>	Copy a basis vector object into another one.
<code>copyColumn(j, i)</code>	Copy the values from one of the columns to another one.
<code>copyVec(j, v)</code>	Copy one of the columns of a basis vectors object into a vector.
<code>create([comm])</code>	Create the BV object.
<code>createFromMat(A)</code>	Create a basis vectors object from a dense matrix.
<code>createMat()</code>	Create a new dense matrix and copy the contents of the BV.
<code>createVec()</code>	Create a vector with the type and dimensions of the columns of the BV.
<code>destroy()</code>	Destroy the BV object.
<code>dot(Y)</code>	Compute the 'block-dot' product of two basis vectors objects.
<code>dotColumn(j)</code>	Dot products of a column against all the column vectors of a BV.
<code>dotVec(v)</code>	Dot products of a vector against all the column vectors of the BV.
<code>duplicate()</code>	Duplicate the BV object with the same type and dimensions.
<code>duplicateResize(m)</code>	Create a BV object of the same type and dimensions as an existing one.
<code>getActiveColumns()</code>	Get the current active dimensions.
<code>getArray([readonly])</code>	Return the array where the data is stored.
<code>getColumn(j)</code>	Get a vector with the entries of the column of the BV object.
<code>getDefiniteTolerance()</code>	Get the tolerance to be used when checking a definite inner product.

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<code>getLeadingDimension()</code>	Get the leading dimension.
<code>getMat()</code>	Get a matrix of dense type that shares the memory of the BV object.
<code>getMatMultMethod()</code>	Get the method used for the <code>matMult()</code> operation.
<code>getMatrix()</code>	Get the matrix representation of the inner product.
<code>getNumConstraints()</code>	Get the number of constraints.
<code>getOptionsPrefix()</code>	Get the prefix used for searching for all BV options in the database.
<code>getOrthogonalization()</code>	Get the orthogonalization settings from the BV object.
<code>getRandomContext()</code>	Get the <code>petsc4py.PETSc.Random</code> object associated with the BV.
<code>getSizes()</code>	Get the local and global sizes, and the number of columns.
<code>getType()</code>	Get the BV type of this object.
<code>getVecType()</code>	Get the vector type used when creating vectors via <code>createVec()</code> .
<code>insertConstraints(C)</code>	Insert a set of vectors as constraints.
<code>insertVec(j, w)</code>	Insert a vector into the specified column.
<code>insertVecs(s, W[, orth])</code>	Insert a set of vectors into the specified columns.
<code>matMult(A[, Y])</code>	Compute the matrix-vector product for each column, $Y = AV$ .
<code>matMultColumn(A, j)</code>	Mat-vec product for a column, storing the result in the next column.
<code>matMultHermitianTranspose(A[, Y])</code>	Pre-multiplication with the conjugate transpose of a matrix.
<code>matMultHermitianTransposeColumn(A, j)</code>	Conjugate-transpose matrix-vector product for a specified column.
<code>matMultTranspose(A[, Y])</code>	Pre-multiplication with the transpose of a matrix.
<code>matMultTransposeColumn(A, j)</code>	Transpose matrix-vector product for a specified column.
<code>matProject(A, Y)</code>	Compute the projection of a matrix onto a subspace.
<code>mult(delta, gamma, X, Q)</code>	Compute $Y = \gamma Y + \delta X Q$ .
<code>multColumn(delta, gamma, j, q)</code>	Compute $y = \gamma y + \delta X q$ .
<code>multInPlace(Q, s, e)</code>	Update a set of vectors as $V(:, s : e - 1) = VQ(:, s : e - 1)$ .
<code>multVec(delta, gamma, y, q)</code>	Compute $y = \gamma y + \delta X q$ .
<code>norm([norm_type])</code>	Compute the matrix norm of the BV.
<code>normColumn(j[, norm_type])</code>	Compute the vector norm of a selected column.
<code>orthogonalize([R])</code>	Orthogonalize all columns (except leading ones) (QR decomposition).
<code>orthogonalizeColumn(j)</code>	Orthogonalize a column vector with respect to the previous ones.
<code>orthogonalizeVec(v)</code>	Orthogonalize a vector with respect to all active columns.
<code>orthonormalizeColumn(j[, replace])</code>	Orthonormalize a column vector with respect to the previous ones.
<code>resize(m[, copy])</code>	Change the number of columns.
<code>restoreColumn(j, v)</code>	Restore a column obtained with <code>getColumn()</code> .
<code>restoreMat(A)</code>	Restore the matrix obtained with <code>getMat()</code> .
<code>scale(alpha)</code>	Multiply the entries by a scalar value.
<code>scaleColumn(j, alpha)</code>	Scale a column of a BV.

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Table 12 – continued from previous page

<code>setActiveColumns(l, k)</code>	Set the columns that will be involved in operations.
<code>setDefiniteTolerance(deftol)</code>	Set the tolerance to be used when checking a definite inner product.
<code>setFromOptions()</code>	Set BV options from the options database.
<code>setLeadingDimension(ld)</code>	Set the leading dimension.
<code>setMatMultMethod(method)</code>	Set the method used for the <code>matMult()</code> operation.
<code>setMatrix(B[, indef])</code>	Set the bilinear form to be used for inner products.
<code>setNumConstraints(nc)</code>	Set the number of constraints.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all BV options in the database.
<code>setOrthogonalization([otype, refine, eta, block])</code>	Set the method used for the (block-)orthogonalization of vectors.
<code>setRandom()</code>	Set the active columns of the BV to random numbers.
<code>setRandomColumn(j)</code>	Set one column of the BV to random numbers.
<code>setRandomCond(condn)</code>	Set the columns of a BV to random numbers.
<code>setRandomContext(rnd)</code>	Set the <code>petsc4py.PETSc.Random</code> object associated with the BV.
<code>setRandomNormal()</code>	Set the active columns of the BV to normal random numbers.
<code>setRandomSign()</code>	Set the entries of a BV to values 1 or -1 with equal probability.
<code>setSizes(sizes, m)</code>	Set the local and global sizes, and the number of columns.
<code>setSizesFromVec(w, m)</code>	Set the local and global sizes, and the number of columns.
<code>setType(bv_type)</code>	Set the type for the BV object.
<code>setVecType(vec_type)</code>	Set the vector type to be used when creating vectors via <code>createVec()</code> .
<code>view([viewer])</code>	Print the BV data structure.

## Attributes Summary

<code>column_size</code>	Basis vectors column size.
<code>local_size</code>	Basis vectors local size.
<code>size</code>	Basis vectors global size.
<code>sizes</code>	Basis vectors local and global sizes, and the number of columns.

## Methods Documentation

### `appendOptionsPrefix(prefix=None)`

Append to the prefix used for searching for all BV options in the database.

Logically collective.

#### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all BV option requests.

#### Return type

*None*

#### See also

[`setOptionsPrefix`](#), [`getOptionsPrefix`](#), [`BVAppendOptionsPrefix`](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:618](#) <[slepc4py/SLEPc/BV.pyx#L618](#)>

**applyMatrix**(*x*, *y*)

Multiply a vector with the matrix associated to the bilinear form.

Neighbor-wise collective.

#### Parameters

- **x** ([`Vec`](#)) – The input vector.
- **y** ([`Vec`](#)) – The result vector.

#### Return type

[`None`](#)

#### Notes

If the bilinear form has no associated matrix this function copies the vector.

#### See also

[`setMatrix`](#), [`BVApplyMatrix`](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:863](#) <[slepc4py/SLEPc/BV.pyx#L863](#)>

**copy**(*result=None*)

Copy a basis vector object into another one.

Logically collective.

#### Returns

The copy.

#### Return type

[`BV`](#)

#### Parameters

**result** ([`BV`](#) / [`None`](#)) – The copy.

#### Notes

Both objects must be distributed in the same manner; local copies are done. Only active columns (excluding the leading ones) are copied. In the destination BV, columns are overwritten starting from the leading ones. Constraints are not copied.

#### See also

[`BVCopy`](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:354](#) <[slepc4py/SLEPc/BV.pyx#L354](#)>

### **copyColumn**(*j*, *i*)

Copy the values from one of the columns to another one.

Logically collective.

#### **Parameters**

- **j** (*int*) – The index of the source column.
- **i** (*int*) – The index of the destination column.

#### **Return type**

*None*

#### **See also**

*copy*, *copyVec*, *BVCopyColumn*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1235 <slepc4py/SLEPc/BV.pyx#L1235>](#)

### **copyVec**(*j*, *v*)

Copy one of the columns of a basis vectors object into a vector.

Logically collective.

#### **Parameters**

- **j** (*int*) – The column index to be copied.
- **v** (*Vec*) – A vector.

#### **Return type**

*None*

#### **Notes**

The BV and v must be distributed in the same manner; local copies are done.

#### **See also**

*copy*, *copyColumn*, *BVCopyVec*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1210 <slepc4py/SLEPc/BV.pyx#L1210>](#)

### **create**(*comm=None*)

Create the BV object.

Collective.

#### **Parameters**

**comm** (*Comm* / *None*) – MPI communicator; if not provided, it defaults to all processes.

#### **Return type**

*Self*

#### **See also**

*createFromMat*, *BVCreate*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:225 <slepc4py/SLEPc/BV.pyx#L225>`

#### **createFromMat(A)**

Create a basis vectors object from a dense matrix.

Collective.

##### **Parameters**

**A** (*Mat*) – A dense tall-skinny matrix.

##### **Return type**

*Self*

#### **Notes**

The matrix values are copied to the *BV* data storage, memory is not shared.

The communicator of the *BV* object will be the same as *A*, and so will be the dimensions.

##### **See also**

*create*, *createMat*, *BVCreateFromMat*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:247 <slepc4py/SLEPc/BV.pyx#L247>`

#### **createMat()**

Create a new dense matrix and copy the contents of the *BV*.

Collective.

##### **Returns**

The new matrix.

##### **Return type**

`petsc4py.PETSc.Mat`

#### **Notes**

The matrix contains all columns of the *BV*, not just the active columns.

##### **See also**

*createFromMat*, *createVec*, *getMat*, *BVCreateMat*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:275 <slepc4py/SLEPc/BV.pyx#L275>`

#### **createVec()**

Create a vector with the type and dimensions of the columns of the *BV*.

Collective.

##### **Returns**

New vector.

##### **Return type**

`petsc4py.PETSc.Vec`

**See also**[createMat](#), [setVecType](#), [BVCreateVec](#)**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1147 <slepc4py/SLEPc/BV.pyx#L1147>](#)**destroy()**

Destroy the BV object.

Collective.

**See also**[BVDestroy](#)**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:211 <slepc4py/SLEPc/BV.pyx#L211>](#)**Return type***Self***dot(Y)**

Compute the ‘block-dot’ product of two basis vectors objects.

Collective.

 $M = Y^*X$  ( $m_{ij} = y_i^*x_j$ ) or  $M = Y^*BX$ **Parameters****Y (BV)** – Left basis vectors, can be the same as self, giving  $M = X^*X$ .**Returns**

The resulting matrix.

**Return type**[petsc4py.PETSc.Mat](#)**Notes**

This is the generalization of `Vec.dot()` for a collection of vectors,  $M = Y^*X$ . The result is a matrix  $M$  whose entry  $m_{ij}$  is equal to  $y_i^*x_j$  (where  $y_i^*$  denotes the conjugate transpose of  $y_i$ ).

$X$  and  $Y$  can be the same object.

If a non-standard inner product has been specified with [setMatrix\(\)](#), then the result is  $M = Y^*BX$ . In this case, both  $X$  and  $Y$  must have the same associated matrix.

Only rows (resp. columns) of  $M$  starting from  $l_y$  (resp.  $l_x$ ) are computed, where  $l_y$  (resp.  $l_x$ ) is the number of leading columns of  $Y$  (resp.  $X$ ).

**See also**[dotVec](#), [dotColumn](#), [setActiveColumns](#), [setMatrix](#), [BVDot](#)**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1500 <slepc4py/SLEPc/BV.pyx#L1500>](#)



### **dotColumn(*j*)**

Dot products of a column against all the column vectors of a BV.

Collective.

#### **Parameters**

**j** (*int*) – The index of the column.

#### **Returns**

The computed values.

#### **Return type**

*ArrayScalar*

### **Notes**

This operation is equivalent to [dotVec\(\)](#) but it uses column *j* of the BV rather than taking a vector as an argument. The number of active columns of the BV is set to *j* before the computation, and restored afterwards. If the BV has leading columns specified, then these columns do not participate in the computation. Therefore, the length of the returned array will be *j* minus the number of leading columns.

#### **See also**

[dot](#), [dotVec](#), [BVDotColumn](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1344 <slepc4py/SLEPc/BV.pyx#L1344>](#)

### **dotVec(*v*)**

Dot products of a vector against all the column vectors of the BV.

Collective.

#### **Parameters**

**v** (*Vec*) – A vector.

#### **Returns**

The computed values.

#### **Return type**

*ArrayScalar*

### **Notes**

This is analogue to [Vec.mDot\(\)](#), but using [BV](#) to represent a collection of vectors *X*. The result is  $m = X^*v$ , so  $m_i$  is equal to  $x_j^*v$ . Note that here *X* is transposed as opposed to [dot\(\)](#).

If a non-standard inner product has been specified with [setMatrix\(\)](#), then the result is  $m = X^*Bv$ .

#### **See also**

[dot](#), [dotColumn](#), [setMatrix](#), [BVDotVec](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1304 <slepc4py/SLEPc/BV.pyx#L1304>](#)

### **duplicate()**

Duplicate the BV object with the same type and dimensions.

Collective.

**Returns**

The new object.

**Return type**

*BV*

**Notes**

This function does not copy the entries, it just allocates the storage for the new *BV*. Use *copy()* to copy the content.

**See also**

*duplicateResize*, *BVDuplicate*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:299 <slepc4py/SLEPc/BV.pyx#L299>](#)

**duplicateResize(*m*)**

Create a *BV* object of the same type and dimensions as an existing one.

Collective.

**Parameters**

**m** (*int*) – The number of columns.

**Returns**

The new object.

**Return type**

*BV*

**Notes**

This is equivalent to a call to *duplicate()* followed by *resize()* with possibly different number of columns. The contents of this *BV* are not copied to the new one.

**See also**

*duplicate*, *resize*, *BVDuplicateResize*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:323 <slepc4py/SLEPc/BV.pyx#L323>](#)

**getActiveColumns()**

Get the current active dimensions.

Not collective.

**Returns**

- **l** (*int*) – The leading number of columns.
- **k** (*int*) – The active number of columns.

**Return type**

*tuple*[*int*, *int*]

#### See also

[setActiveColumns](#), [BVGetActiveColumns](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:924 <slepc4py/SLEPc/BV.pyx#L924>](#)

**getArray**(*readonly=False*)

Return the array where the data is stored.

Not collective.

#### Parameters

**readonly** (*bool*) – Enable to obtain a read only array.

#### Returns

The array.

#### Return type

[ArrayScalar](#)

#### See also

[BVGetArray](#), [BVGetArrayRead](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:546 <slepc4py/SLEPc/BV.pyx#L546>](#)

**getColumn**(*j*)

Get a vector with the entries of the column of the BV object.

Logically collective.

#### Parameters

**j** (*int*) – The index of the requested column.

#### Returns

The vector containing the *j*-th column.

#### Return type

[petsc4py.PETSc.Vec](#)

#### Notes

Modifying the returned vector will change the BV entries as well.

The returned vector must not be destroyed, [restoreColumn\(\)](#) must be called when it is no longer needed. At most, two columns can be fetched, that is, this function can only be called twice before the corresponding [restoreColumn\(\)](#) is invoked.

A negative index *j* selects the *i*-th constraint, where *i*=-*j*. Constraints should not be modified.

#### See also

[restoreColumn](#), [insertConstraints](#), [BVGetColumn](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1384 <slepc4py/SLEPc/BV.pyx#L1384>](#)

### **getDefiniteTolerance()**

Get the tolerance to be used when checking a definite inner product.

Not collective.

#### **Returns**

The tolerance.

#### **Return type**

`float`

#### **See also**

`setDefiniteTolerance`, `BVGetDefiniteTolerance`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1285 <slepc4py/SLEPc/BV.pyx#L1285>`

### **getLeadingDimension()**

Get the leading dimension.

Not collective.

#### **Returns**

The leading dimension.

#### **Return type**

`int`

### **Notes**

The returned value may be different in different processes.

The leading dimension must be used when accessing the internal array via `getArray()`.

#### **See also**

`setLeadingDimension`, `BVGetLeadingDimension`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:520 <slepc4py/SLEPc/BV.pyx#L520>`

### **getMat()**

Get a matrix of dense type that shares the memory of the BV object.

Collective.

#### **Returns**

The matrix.

#### **Return type**

`petsc4py.PETSc.Mat`

### **Notes**

The returned matrix contains only the active columns. If the content of the matrix is modified, these changes are also done in the BV object. The user must call `restoreMat()` when no longer needed.

This operation implies a call to `getArray()`, which may result in data copies.

**See also**

`restoreMat`, `createMat`, `getArray`, `BVGetMat`

**:sources:** ``Source code at slepc4py/SLEPc/BV.pyx:1447 <slepc4py/SLEPc/BV.pyx#L1447>``

**getMatMultMethod()**

Get the method used for the `matMult()` operation.

Not collective.

**Returns**

The method for the `matMult()` operation.

**Return type**

`MatMultType`

**See also**

`matMult`, `setMatMultMethod`, `BVGetMatMultMethod`

**:sources:** ``Source code at slepc4py/SLEPc/BV.pyx:760 <slepc4py/SLEPc/BV.pyx#L760>``

**getMatrix()**

Get the matrix representation of the inner product.

Not collective.

**Returns**

- `B (petsc4py.PETSc.Mat)` – The matrix of the inner product.
- `indef (bool)` – Whether the matrix is indefinite.

**Return type**

`tuple[Mat, bool] | tuple[None, bool]`

**See also**

`setMatrix`, `BVGetMatrix`

**:sources:** ``Source code at slepc4py/SLEPc/BV.pyx:799 <slepc4py/SLEPc/BV.pyx#L799>``

**getNumConstraints()**

Get the number of constraints.

Not collective.

**Returns**

The number of constraints.

**Return type**

`int`

**See also**

`insertConstraints`, `setNumConstraints`, `BVGetNumConstraints`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1128 <slepc4py/SLEPc/BV.pyx#L1128>`

#### **getOptionsPrefix()**

Get the prefix used for searching for all BV options in the database.

Not collective.

##### **Returns**

The prefix string set for this BV object.

##### **Return type**

`str`

##### **See also**

`setOptionsPrefix`, `appendOptionsPrefix`, `BVGetOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:637 <slepc4py/SLEPc/BV.pyx#L637>`

#### **getOrthogonalization()**

Get the orthogonalization settings from the BV object.

Not collective.

##### **Returns**

- **type** (*OrthogType*) – The type of orthogonalization technique.
- **refine** (*OrthogRefineType*) – The type of refinement.
- **eta** (`float`) – Parameter for selective refinement (used when the refinement type is *IFNEEDED*).
- **block** (*OrthogBlockType*) – The type of block orthogonalization.

##### **Return type**

`tuple[OrthogType, OrthogRefineType, float, OrthogBlockType]`

##### **See also**

`setOrthogonalization`, `BVGetOrthogonalization`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:675 <slepc4py/SLEPc/BV.pyx#L675>`

#### **getRandomContext()**

Get the `petsc4py.PETSc.Random` object associated with the BV.

Collective.

##### **Returns**

The random number generator context.

##### **Return type**

`petsc4py.PETSc.Random`

##### **See also**

`setRandomContext`, `BVGetRandomContext`

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:2133 <slepc4py/SLEPc/BV.pyx#L2133>](#)

### **getSizes()**

Get the local and global sizes, and the number of columns.

Not collective.

#### **Returns**

- **(n, N)** (**tuple** of **int**) – The local and global sizes.
- **m** (**int**) – The number of columns.

#### **Return type**

**tuple**[**LayoutSizeSpec**, **int**]

#### **See also**

[setSize](#), [setSizeFromVec](#), [BVGetSizes](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:477 <slepc4py/SLEPc/BV.pyx#L477>](#)

### **getType()**

Get the BV type of this object.

Not collective.

#### **Returns**

The basis vectors type currently being used.

#### **Return type**

**str**

#### **See also**

[setType](#), [BVGetType](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:407 <slepc4py/SLEPc/BV.pyx#L407>](#)

### **getVecType()**

Get the vector type used when creating vectors via [createVec\(\)](#).

Not collective.

#### **Returns**

The vector type.

#### **Return type**

**str**

#### **See also**

[createVec](#), [setVecType](#), [BVGetVecType](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1191 <slepc4py/SLEPc/BV.pyx#L1191>](#)

## **insertConstraints(C)**

Insert a set of vectors as constraints.

Collective.

### **Parameters**

**C** (*Vec* / *list*[*Vec*]) – Set of vectors to be inserted as constraints.

### **Returns**

Number of linearly independent constraints.

### **Return type**

*int*

## **Notes**

The constraints are relevant only during orthogonalization. Constraint vectors span a subspace that is deflated in every orthogonalization operation, so they are intended for removing those directions from the orthogonal basis computed in regular BV columns.

Constraints are not stored in regular columns, but in a special part of the storage. They can be accessed with negative indices in *getColumn()*.

This operation is DESTRUCTIVE, meaning that all data contained in the columns of the BV is lost. This is typically invoked just after creating the BV. Once a set of constraints has been set, it is not allowed to call this function again.

The vectors are copied one by one and then orthogonalized against the previous ones. If any of them is linearly dependent then it is discarded and not counted in the return value. The behavior is similar to *insertVecs()*.

### **See also**

*insertVecs*, *setNumConstraints*, *BVInsertConstraints*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1051 <slepc4py/SLEPc/BV.pyx#L1051>`

## **insertVec(j, w)**

Insert a vector into the specified column.

Logically collective.

### **Parameters**

- **j** (*int*) – The column to be overwritten.
- **w** (*Vec*) – The vector to be copied.

### **Return type**

*None*

### **See also**

*insertVecs*, *BVInsertVec*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:988 <slepc4py/SLEPc/BV.pyx#L988>`



**insertVecs**(*s*, *W*, *orth=False*)

Insert a set of vectors into the specified columns.

Collective.

**Parameters**

- **s** (*int*) – The first column to be overwritten.
- **W** (*Vec* | *list[Vec]*) – Set of vectors to be copied.
- **orth** (*bool*) – Flag indicating if the vectors must be orthogonalized.

**Returns**

Number of linearly independent vectors.

**Return type**

*int*

**Notes**

Copies the contents of vectors *W* into the BV columns *s*:*s*+*n*, where *n* is the length of *W*. If *orth* is set, then the vectors are copied one by one and then orthogonalized against the previous one. If any of them is linearly dependent then it is discarded and the not counted in the return value.

**See also**

*insertVec*, *orthogonalizeColumn*, *BVInsertVecs*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1008 <slepc4py/SLEPc/BV.pyx#L1008>](#)

**matMult**(*A*, *Y=None*)

Compute the matrix-vector product for each column,  $Y = AV$ .

Neighbor-wise collective.

**Parameters**

- **A** (*Mat*) – The matrix.
- **Y** (*BV* | *None*)

**Returns**

The result.

**Return type**

*BV*

**Notes**

Only active columns (excluding the leading ones) are processed. If *Y* is *None* a new BV is created.

It is possible to choose whether the computation is done column by column or as a dense matrix-matrix product with *setMatMultMethod()*.

**See also**

*copy*, *matMultColumn*, *matMultTranspose*, *setMatMultMethod*, *BVMatMult*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1598 <slepc4py/SLEPc/BV.pyx#L1598>](#)

**matMultColumn**(*A*, *j*)

Mat-vec product for a column, storing the result in the next column.

Neighbor-wise collective.

$$v_{j+1} = Av_j.$$

**Parameters**

- **A** (*Mat*) – The matrix.
- **j** (*int*) – Index of column.

**Return type**

*None*

**See also**

*matMult*, *BVMatMultColumn*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1743 <slepc4py/SLEPc/BV.pyx#L1743>`

**matMultHermitianTranspose**(*A*, *Y=None*)

Pre-multiplication with the conjugate transpose of a matrix.

Neighbor-wise collective.

$$Y = A^*V.$$

**Parameters**

- **A** (*Mat*) – The matrix.
- **Y** (*BV* | *None*)

**Returns**

The result.

**Return type**

*BV*

**Notes**

Only active columns (excluding the leading ones) are processed. If *Y* is *None* a new *BV* is created.

**See also**

*matMult*, *matMultHermitianTransposeColumn*, *BVMatMultHermitianTranspose*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1695 <slepc4py/SLEPc/BV.pyx#L1695>`

**matMultHermitianTransposeColumn**(*A*, *j*)

Conjugate-transpose matrix-vector product for a specified column.

Neighbor-wise collective.

Store the result in the next column:  $v_{j+1} = A^*v_j$ .

**Parameters**

- **A** (*Mat*) – The matrix.

- **j** (*int*) – Index of column.

**Return type**

None

**See also**

*matMultColumn*, *BVMatMultHermitianTransposeColumn*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1787 <slepc4py/SLEPc/BV.pyx#L1787>](#)

**matMultTranspose**(*A*, *Y=None*)

Pre-multiplication with the transpose of a matrix.

Neighbor-wise collective.

$$Y = A^T V.$$

**Parameters**

- **A** (*Mat*) – The matrix.
- **Y** (*BV* | *None*)

**Returns**

The result.

**Return type**

*BV*

**Notes**

Only active columns (excluding the leading ones) are processed. If *Y* is *None* a new *BV* is created.

**See also**

*matMult*, *matMultTransposeColumn*, *BVMatMultTranspose*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1647 <slepc4py/SLEPc/BV.pyx#L1647>](#)

**matMultTransposeColumn**(*A*, *j*)

Transpose matrix-vector product for a specified column.

Neighbor-wise collective.

Store the result in the next column:  $v_{j+1} = A^T v_j$ .

**Parameters**

- **A** (*Mat*) – The matrix.
- **j** (*int*) – Index of column.

**Return type**

None

**See also**

*matMultColumn*, *BVMatMultTransposeColumn*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1765 <slepc4py/SLEPc/BV.pyx#L1765>`

**matProject**(A, Y)

Compute the projection of a matrix onto a subspace.

Collective.

$$M = Y^*AX$$

**Parameters**

- **A** (*Mat* / *None*) – Matrix to be projected.
- **Y** (*BV*) – Left basis vectors, can be the same as self, giving  $M = X^*AX$ .

**Returns**

Projection of the matrix A onto the subspace.

**Return type**

`petsc4py.PETSc.Mat`

**Notes**

If A is None, then it is assumed that the BV already contains AX.

This operation is similar to `dot()`, with important differences. The goal is to compute the matrix resulting from the orthogonal projection of A onto the subspace spanned by the columns of the BV,  $M = X^*AX$ , or the oblique projection onto the BV along the second one Y,  $M = Y^*AX$ .

A difference with respect to `dot()` is that the standard inner product is always used, regardless of a non-standard inner product being specified with `setMatrix()`.

**See also**

`dot`, `setActiveColumns`, `setMatrix`, `BVMatProject`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1549 <slepc4py/SLEPc/BV.pyx#L1549>`

**mult**(delta, gamma, X, Q)

Compute  $Y = \gamma Y + \delta XQ$ .

Logically collective.

**Parameters**

- **delta** (*Scalar*) – Coefficient that multiplies X.
- **gamma** (*Scalar*) – Coefficient that multiplies self (Y).
- **X** (*BV*) – Input basis vectors.
- **Q** (*Mat* / *None*) – Input matrix, if not given the identity matrix is assumed.

**Return type**

*None*

**Notes**

X must be different from self (Y). The case X=Y can be addressed with `multInPlace()`.

**See also***multVec, multColumn, multInPlace, BVMult***:sources:**``Source code at slepc4py/SLEPc/BV.pyx:1809 <slepc4py/SLEPc/BV.pyx#L1809>``**multColumn**(*delta, gamma, j, q*)Compute  $y = \gamma y + \delta X q$ .

Logically collective.

Compute  $y = \gamma y + \delta X q$ , where  $y$  is the  $j$ -th column.**Parameters**

- **delta** (*Scalar*) – Coefficient that multiplies self ( $X$ ).
- **gamma** (*Scalar*) – Coefficient that multiplies  $y$ .
- **j** (*int*) – The column index.
- **q** (*Sequence*[*Scalar*]) – Input coefficients.

**Return type**

None

**See also***mult, multVec, multInPlace, BVMultColumn***:sources:**``Source code at slepc4py/SLEPc/BV.pyx:1863 <slepc4py/SLEPc/BV.pyx#L1863>``**multInPlace**(*Q, s, e*)Update a set of vectors as  $V(:, s : e - 1) = VQ(:, s : e - 1)$ .

Logically collective.

**Parameters**

- **Q** (*Mat*) – A sequential dense matrix.
- **s** (*int*) – First column to be overwritten.
- **e** (*int*) – Last column to be overwritten.

**Return type**

None

**See also***mult, multVec, BVMultInPlace***:sources:**``Source code at slepc4py/SLEPc/BV.pyx:1840 <slepc4py/SLEPc/BV.pyx#L1840>``**multVec**(*delta, gamma, y, q*)Compute  $y = \gamma y + \delta X q$ .

Logically collective.

**Parameters**

- **delta** ([Scalar](#)) – Coefficient that multiplies self (X).
- **gamma** ([Scalar](#)) – Coefficient that multiplies y.
- **y** ([Vec](#)) – Input/output vector.
- **q** ([Sequence](#)[[Scalar](#)]) – Input coefficients.

**Return type**

[None](#)

**See also**

[mult](#), [multColumn](#), [multInPlace](#), [BVMultVec](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1898 <slepc4py/SLEPc/BV.pyx#L1898>](#)

**norm**(*norm\_type=None*)

Compute the matrix norm of the BV.

Collective.

**Parameters**

**norm\_type** ([NormType](#) | [None](#)) – The norm type.

**Returns**

The norm.

**Return type**

[float](#)

**Notes**

All active columns (except the leading ones) are considered as a matrix. The allowed norms are [NORM\\_1](#), [NORM\\_FROBENIUS](#), and [NORM\\_INFINITY](#).

This operation fails if a non-standard inner product has been specified with [setMatrix\(\)](#).

**See also**

[normColumn](#), [setMatrix](#), [BVNorm](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1967 <slepc4py/SLEPc/BV.pyx#L1967>](#)

**normColumn**(*j*, *norm\_type=None*)

Compute the vector norm of a selected column.

Collective.

**Parameters**

- **j** ([int](#)) – Index of column.
- **norm\_type** ([NormType](#) | [None](#)) – The norm type.

**Returns**

The norm.

**Return type**

[float](#)

## Notes

The norm of  $v_j$  is computed (NORM\_1, NORM\_2, or NORM\_INFINITY).

If a non-standard inner product has been specified with `setMatrix()`, then the returned value is  $\sqrt{v_j^* B v_j}$ , where  $B$  is the inner product matrix (argument 'norm\_type' is ignored).

### See also

`norm`, `setMatrix`, `BVNormColumn`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1929 <slepc4py/SLEPc/BV.pyx#L1929>`

**orthogonalize**( $R=None$ ,  $**kargs$ )

Orthogonalize all columns (except leading ones) (QR decomposition).

Collective.

### Parameters

- **R** (*Mat* | *None*) – A sequential dense matrix.
- **kargs** (*Any*)

### Return type

*None*

## Notes

The output satisfies  $V_0 = VR$  (where  $V_0$  represent the input  $V$ ) and  $V^*V = I$  (or  $V^*BV = I$  if an inner product matrix  $B$  has been specified with `setMatrix()`).

### See also

`orthogonalizeColumn`, `setMatrix`, `setOrthogonalization`, `BVOrthogonalize`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2263 <slepc4py/SLEPc/BV.pyx#L2263>`

**orthogonalizeColumn**( $j$ )

Orthogonalize a column vector with respect to the previous ones.

Collective.

### Parameters

- **j** (*int*) – Index of the column to be orthogonalized.

### Returns

- **norm** (*float*) – The norm of the resulting vector.
- **lindep** (*bool*) – Flag indicating that refinement did not improve the quality of orthogonalization.

### Return type

*tuple*[*float*, *bool*]

## Notes

This function applies an orthogonal projector to project vector  $v_j$  onto the orthogonal complement of the span of the columns  $V[0..j-1]$ , where  $V[.]$  are the vectors of the BV. The columns  $V[0..j-1]$  are assumed to be mutually orthonormal.

This routine does not normalize the resulting vector.

### See also

[\*orthogonalizeVec\*](#), [\*setOrthogonalization\*](#)

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2189 <slepc4py/SLEPc/BV.pyx#L2189>`

## **orthogonalizeVec(v)**

Orthogonalize a vector with respect to all active columns.

Collective.

### Parameters

**v** (*Vec*) – Vector to be orthogonalized, modified on return.

### Returns

- **norm** (*float*) – The norm of the resulting vector.
- **lindep** (*bool*) – Flag indicating that refinement did not improve the quality of orthogonalization.

### Return type

`tuple[float, bool]`

## Notes

This function applies an orthogonal projector to project vector  $v$  onto the orthogonal complement of the span of the columns of the BV.

This routine does not normalize the resulting vector.

### See also

[\*orthogonalizeColumn\*](#), [\*setOrthogonalization\*](#)

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2153 <slepc4py/SLEPc/BV.pyx#L2153>`

## **orthonormalizeColumn(j, replace=False)**

Orthonormalize a column vector with respect to the previous ones.

Collective.

This is equivalent to a call to [\*orthogonalizeColumn\(\)\*](#) followed by a call to [\*scaleColumn\(\)\*](#) with the reciprocal of the norm.

### Parameters

- **j** (*int*) – Index of the column to be orthonormalized.
- **replace** (*bool*) – Whether it is allowed to set the vector randomly.

### Returns



- **norm** (`float`) – The norm of the resulting vector.
- **lindep** (`bool`) – Flag indicating that refinement did not improve the quality of orthogonalization.

**Return type**

`tuple[float, bool]`

**See also**

`orthogonalizeColumn`, `setOrthogonalization`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2227 <slepc4py/SLEPc/BV.pyx#L2227>`

**resize**(*m*, *copy=True*)

Change the number of columns.

Collective.

**Parameters**

- **m** (`int`) – The new number of columns.
- **copy** (`bool`) – A flag indicating whether current values should be kept.

**Return type**

`None`

**Notes**

Internal storage is reallocated. If `copy` is `True`, then the contents are copied to the leading part of the new space.

**See also**

`setSize`, `setSizeFromVec`, `BVResize`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2002 <slepc4py/SLEPc/BV.pyx#L2002>`

**restoreColumn**(*j*, *v*)

Restore a column obtained with `getColumn()`.

Logically collective.

**Parameters**

- **j** (`int`) – The index of the requested column.
- **v** (`Vec`) – The vector obtained with `getColumn()`.

**Return type**

`None`

**Notes**

The arguments must match the corresponding call to `getColumn()`.

#### See also

[getColumn](#), [BVRestoreColumn](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1422 <slepc4py/SLEPc/BV.pyx#L1422>](#)

#### **restoreMat**(*A*)

Restore the matrix obtained with [getMat\(\)](#).

Logically collective.

##### Parameters

**A** (*Mat*) – The matrix obtained with [getMat\(\)](#).

##### Return type

None

#### Notes

A call to this function must match a previous call of [getMat\(\)](#). The effect is that the contents of the matrix are copied back to the BV internal data structures.

#### See also

[getMat](#), [BVRestoreMat](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1476 <slepc4py/SLEPc/BV.pyx#L1476>](#)

#### **scale**(*alpha*)

Multiply the entries by a scalar value.

Logically collective.

##### Parameters

**alpha** (*Scalar*) – scaling factor.

##### Return type

None

#### Notes

All active columns (except the leading ones) are scaled.

#### See also

[scaleColumn](#), [setActiveColumns](#), [BVScale](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:966 <slepc4py/SLEPc/BV.pyx#L966>](#)

#### **scaleColumn**(*j*, *alpha*)

Scale a column of a BV.

Logically collective.

##### Parameters

- **j** (*int*) – column index to be scaled.

- **alpha** (*Scalar*) – scaling factor.

**Return type**

*None*

**See also**

*scale*, *BVScaleColumn*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:945 <slepc4py/SLEPc/BV.pyx#L945>`

**setActiveColumns**(*l*, *k*)

Set the columns that will be involved in operations.

Logically collective.

**Parameters**

- **l** (*int*) – The leading number of columns.
- **k** (*int*) – The active number of columns.

**Return type**

*None*

**Notes**

In operations such as *mult()* or *dot()*, only the first *k* columns are considered. This is useful when the BV is filled from left to right, so the last *m-k* columns do not have relevant information.

Also in operations such as *mult()* or *dot()*, the first *l* columns are normally not included in the computation.

In orthogonalization operations, the first *l* columns are treated differently, they participate in the orthogonalization but the computed coefficients are not stored.

Use *CURRENT* to leave any of the values unchanged. Use *DETERMINE* to set *l* to the minimum value (0) and *k* to the maximum (*m*).

**See also**

*getActiveColumns*, *setSize*, *BVSetActiveColumns*

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:887 <slepc4py/SLEPc/BV.pyx#L887>`

**setDefiniteTolerance**(*deftol*)

Set the tolerance to be used when checking a definite inner product.

Logically collective.

**Parameters**

- **deftol** (*float*) – The tolerance.

**Return type**

*None*

## Notes

When using a non-standard inner product, see `setMatrix()`, the solver needs to compute  $\sqrt{z^*Bz}$  for various vectors  $z$ . If the inner product has not been declared indefinite, the value  $z^*Bz$  must be positive, but due to rounding error a tiny value may become negative. A tolerance is used to detect this situation. Likewise, in complex arithmetic  $z^*Bz$  should be real, and we use the same tolerance to check whether a nonzero imaginary part can be considered negligible.

### See also

`setMatrix`, `getDefiniteTolerance`, `BVSetDefiniteTolerance`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:1256 <slepc4py/SLEPc/BV.pyx#L1256>`

## `setFromOptions()`

Set BV options from the options database.

Collective.

## Notes

To see all options, run your program with the `-help` option.

### See also

`setOptionsPrefix`, `BVSetFromOptions`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:656 <slepc4py/SLEPc/BV.pyx#L656>`

### Return type

`None`

## `setLeadingDimension(ld)`

Set the leading dimension.

Not collective.

### Parameters

**ld** (`int`) – The leading dimension.

### Return type

`None`

## Notes

This parameter is relevant for a BV of `BV.Type.MAT`.

### See also

`getLeadingDimension`, `BVSetLeadingDimension`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:498 <slepc4py/SLEPc/BV.pyx#L498>`

### **setMatMultMethod**(*method*)

Set the method used for the `matMult()` operation.

Logically collective.

#### **Parameters**

**method** (`MatMultType`) – The method for the `matMult()` operation.

#### **Return type**

`None`

#### **See also**

`matMult`, `getMatMultMethod`, `BVSetMatMultMethod`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:779 <slepc4py/SLEPc/BV.pyx#L779>`

### **setMatrix**(*B*, *indef=False*)

Set the bilinear form to be used for inner products.

Collective.

#### **Parameters**

- **B** (`Mat` | `None`) – The matrix of the inner product.
- **indef** (`bool`) – Whether the matrix is indefinite.

#### **Return type**

`None`

#### **Notes**

This is used to specify a non-standard inner product, whose matrix representation is given by *B*. Then, all inner products required during orthogonalization are computed as  $(x, y)_B = y^* B x$  rather than the standard form  $(x, y) = y^* x$ .

Matrix *B* must be real symmetric (or complex Hermitian). A genuine inner product requires that *B* is also positive (semi-)definite. However, we also allow for an indefinite *B* (setting `indef=True`), in which case the orthogonalization uses an indefinite inner product.

This affects operations `dot()`, `norm()`, `orthogonalize()`, and variants.

Omitting *B* has the same effect as if the identity matrix was passed.

#### **See also**

`getMatrix`, `BVSetMatrix`

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:825 <slepc4py/SLEPc/BV.pyx#L825>`

### **setNumConstraints**(*nc*)

Set the number of constraints.

Logically collective.

#### **Parameters**

**nc** (`int`) – The number of constraints.

**Return type**

None

**Notes**

This function sets the number of constraints to `nc` and marks all remaining columns as regular. Normal usage would be to call `insertConstraints()` instead.

If `nc` is smaller than the previously set value, then some of the constraints are discarded. In particular, using `nc=0` removes all constraints preserving the content of regular columns.

**See also**

`insertConstraints`, `getNumConstraints`, `BVSetNumConstraints`

**`:sources:Source code at slepc4py/SLEPc/BV.pyx:1101 <slepc4py/SLEPc/BV.pyx#L1101>`**

**`setOptionsPrefix`**(*prefix=None*)

Set the prefix used for searching for all BV options in the database.

Logically collective.

**Parameters**

**`prefix`** (*str* / *None*) – The prefix string to prepend to all BV option requests.

**Return type**

None

**Notes**

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

**See also**

`appendOptionsPrefix`, `getOptionsPrefix`, `BVGetOptionsPrefix`

**`:sources:Source code at slepc4py/SLEPc/BV.pyx:593 <slepc4py/SLEPc/BV.pyx#L593>`**

**`setOrthogonalization`**(*otype=None*, *refine=None*, *eta=None*, *block=None*)

Set the method used for the (block-)orthogonalization of vectors.

Logically collective.

Orthogonalization of vectors (classical or modified Gram-Schmidt with or without refinement), and for the block-orthogonalization (simultaneous orthogonalization of a set of vectors).

**Parameters**

- **`otype`** (*OrthogType* / *None*) – The type of orthogonalization technique.
- **`refine`** (*OrthogRefineType* / *None*) – The type of refinement.
- **`eta`** (*float* / *None*) – Parameter for selective refinement.
- **`block`** (*OrthogBlockType* / *None*) – The type of block orthogonalization.

**Return type**

None

## Notes

The default settings work well for most problems.

The parameter `eta` should be a real value between 0 and 1 (or *DETERMINE*). The value of `eta` is used only when the refinement type is *IFNEEDED*.

When using several processes, *MGS* is likely to result in bad scalability.

If the method set for block orthogonalization is *GS*, then the computation is done column by column with the vector orthogonalization.

### See also

[\*getOrthogonalization\*](#), [\*BVSetOrthogonalization\*](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:704 <slepc4py/SLEPc/BV.pyx#L704>](#)

## **setRandom()**

Set the active columns of the BV to random numbers.

Logically collective.

## Notes

All active columns (except the leading ones) are modified.

### See also

[\*setRandomContext\*](#), [\*setRandomColumn\*](#), [\*setRandomNormal\*](#), [\*BVSetRandom\*](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:2028 <slepc4py/SLEPc/BV.pyx#L2028>](#)

### Return type

None

## **setRandomColumn(j)**

Set one column of the BV to random numbers.

Logically collective.

### Parameters

**j** (*int*) – Column index to be set.

### Return type

None

### See also

[\*setRandomContext\*](#), [\*setRandom\*](#), [\*setRandomNormal\*](#), [\*BVSetRandomColumn\*](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:2076 <slepc4py/SLEPc/BV.pyx#L2076>](#)

## **setRandomCond(condn)**

Set the columns of a BV to random numbers.

Logically collective.

The generated matrix has a prescribed condition number.

**Parameters**

**condn** (*float*) – Condition number.

**Return type**

None

**See also**

[\*setRandomContext\*](#), [\*setRandomSign\*](#), [\*setRandomNormal\*](#), [\*BVSetRandomCond\*](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:2094 <slepc4py/SLEPc/BV.pyx#L2094>](#)

**setRandomContext(*rnd*)**

Set the `petsc4py.PETSc.Random` object associated with the BV.

Collective.

To be used in operations that need random numbers.

**Parameters**

**rnd** (*Random*) – The random number generator context.

**Return type**

None

**See also**

[\*getRandomContext\*](#), [\*setRandom\*](#), [\*setRandomColumn\*](#), [\*BVSetRandomContext\*](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:2114 <slepc4py/SLEPc/BV.pyx#L2114>](#)

**setRandomNormal()**

Set the active columns of the BV to normal random numbers.

Logically collective.

**Notes**

All active columns (except the leading ones) are modified.

**See also**

[\*setRandomContext\*](#), [\*setRandom\*](#), [\*setRandomSign\*](#), [\*BVSetRandomNormal\*](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:2044 <slepc4py/SLEPc/BV.pyx#L2044>](#)

**Return type**

None

**setRandomSign()**

Set the entries of a BV to values 1 or -1 with equal probability.

Logically collective.



## Notes

All active columns (except the leading ones) are modified.

### See also

[`setRandomContext`](#), [`setRandom`](#), [`setRandomNormal`](#), [`BVSetRandomSign`](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:2060](#) <[slepc4py/SLEPc/BV.pyx#L2060](#)>

### Return type

`None`

**setSize**(*sizes*, *m*)

Set the local and global sizes, and the number of columns.

Collective.

### Parameters

- **sizes** ([`LayoutSizeSpec`](#)) – The global size *N* or a two-tuple (*n*, *N*) with the local and global sizes.
- **m** (`int`) – The number of columns.

### Return type

`None`

## Notes

Either *n* or *N* (but not both) can be [`DETERMINE`](#) or `None` to have it automatically set.

### See also

[`setSizeFromVec`](#), [`getSize`](#), [`BVSetSizes`](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:426](#) <[slepc4py/SLEPc/BV.pyx#L426](#)>

**setSizeFromVec**(*w*, *m*)

Set the local and global sizes, and the number of columns.

Collective.

Local and global sizes are specified indirectly by passing a template vector.

### Parameters

- **w** ([`Vec`](#)) – The template vector.
- **m** (`int`) – The number of columns.

### Return type

`None`

### See also

[`setSize`](#), [`getSize`](#), [`BVSetSizesFromVec`](#)

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:454](#) <[slepc4py/SLEPc/BV.pyx#L454](#)>

**setType**(*bv\_type*)

Set the type for the BV object.

Logically collective.

**Parameters**

**bv\_type** (*Type* / *str*) – The basis vectors type to be used.

**Return type**

*None*

**See also**

*getType*, *BVSetType*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:388 <slepc4py/SLEPc/BV.pyx#L388>](#)

**setVecType**(*vec\_type*)

Set the vector type to be used when creating vectors via *createVec()*.

Collective.

**Parameters**

**vec\_type** (*petsc4py.PETSc.Vec.Type* / *str*) – Vector type used when creating vectors with *createVec*.

**Return type**

*None*

**Notes**

This is not needed if the BV object is set up with *setSizeFromVec()*, but may be required in the case of *setSize()* if one wants to work with non-standard vectors.

**See also**

*createVec*, *getVecType*, *setSize*, *setSizeFromVec*, *BVSetVecType*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:1166 <slepc4py/SLEPc/BV.pyx#L1166>](#)

**view**(*viewer=None*)

Print the BV data structure.

Collective.

**Parameters**

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

*None*

**See also**

*BVView*

**:sources:** [Source code at slepc4py/SLEPc/BV.pyx:192 <slepc4py/SLEPc/BV.pyx#L192>](#)

## Attributes Documentation

### `column_size`

Basis vectors column size.

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2305 <slepc4py/SLEPc/BV.pyx#L2305>`

### `local_size`

Basis vectors local size.

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2300 <slepc4py/SLEPc/BV.pyx#L2300>`

### `size`

Basis vectors global size.

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2295 <slepc4py/SLEPc/BV.pyx#L2295>`

### `sizes`

Basis vectors local and global sizes, and the number of columns.

**:sources:** `Source code at slepc4py/SLEPc/BV.pyx:2290 <slepc4py/SLEPc/BV.pyx#L2290>`

## `slepc4py.SLEPc.DS`

### `class slepc4py.SLEPc.DS`

Bases: `Object`

Direct Solver (or Dense System).

The *DS* package provides auxiliary routines that are internally used by the different slepc4py solvers. It is used to represent low-dimensional eigenproblems that must be solved within iterative solvers with direct methods. It can be seen as a structured wrapper to LAPACK functionality.

## Enumerations

<i>MatType</i>	To refer to one of the matrices stored internally in DS.
<i>ParallelType</i>	Indicates the parallel mode that the direct solver will use.
<i>StateType</i>	DS state types.
<i>Type</i>	DS type.

## `slepc4py.SLEPc.DS.MatType`

### `class slepc4py.SLEPc.DS.MatType`

Bases: `object`

To refer to one of the matrices stored internally in DS.

- *A*: first matrix of eigenproblem/singular value problem.
- *B*: second matrix of a generalized eigenproblem.
- *C*: third matrix of a quadratic eigenproblem.
- *T*: tridiagonal matrix.
- *D*: diagonal matrix.
- *Q*: orthogonal matrix of (right) Schur vectors.

- **Z**: orthogonal matrix of left Schur vectors.
- **X**: right eigenvectors.
- **Y**: left eigenvectors.
- **U**: left singular vectors.
- **V**: right singular vectors.
- **W**: workspace matrix.

#### See also

DSMatType

### Attributes Summary

<b>A</b>	Constant A of type <code>int</code>
<b>B</b>	Constant B of type <code>int</code>
<b>C</b>	Constant C of type <code>int</code>
<b>D</b>	Constant D of type <code>int</code>
<b>Q</b>	Constant Q of type <code>int</code>
<b>T</b>	Constant T of type <code>int</code>
<b>U</b>	Constant U of type <code>int</code>
<b>V</b>	Constant V of type <code>int</code>
<b>W</b>	Constant W of type <code>int</code>
<b>X</b>	Constant X of type <code>int</code>
<b>Y</b>	Constant Y of type <code>int</code>
<b>Z</b>	Constant Z of type <code>int</code>

### Attributes Documentation

**A:** `int` = **A**

Constant A of type `int`

**B:** `int` = **B**

Constant B of type `int`

**C:** `int` = **C**

Constant C of type `int`

**D:** `int` = **D**

Constant D of type `int`

**Q:** `int` = **Q**

Constant Q of type `int`

**T:** `int` = **T**

Constant T of type `int`

**U:** `int` = **U**

Constant U of type `int`

**V:** `int` = **V**

Constant V of type `int`

**W:** `int = W`  
 Constant W of type `int`

**X:** `int = X`  
 Constant X of type `int`

**Y:** `int = Y`  
 Constant Y of type `int`

**Z:** `int = Z`  
 Constant Z of type `int`

## slepc4py.SLEPc.DS.ParallelType

**class** slepc4py.SLEPc.DS.ParallelType

Bases: `object`

Indicates the parallel mode that the direct solver will use.

- *REDUNDANT*: Every process performs the computation redundantly.
- *SYNCHRONIZED*: The first process sends the result to the rest.
- *DISTRIBUTED*: Used in some cases to distribute the computation among processes.

### See also

DSParallelType

## Attributes Summary

<i>DISTRIBUTED</i>	Constant DISTRIBUTED of type <code>int</code>
<i>REDUNDANT</i>	Constant REDUNDANT of type <code>int</code>
<i>SYNCHRONIZED</i>	Constant SYNCHRONIZED of type <code>int</code>

## Attributes Documentation

**DISTRIBUTED:** `int = DISTRIBUTED`

Constant DISTRIBUTED of type `int`

**REDUNDANT:** `int = REDUNDANT`

Constant REDUNDANT of type `int`

**SYNCHRONIZED:** `int = SYNCHRONIZED`

Constant SYNCHRONIZED of type `int`

## slepc4py.SLEPc.DS.StateType

**class** slepc4py.SLEPc.DS.StateType

Bases: `object`

DS state types.

- *RAW*: Not processed yet.
- *INTERMEDIATE*: Reduced to Hessenberg or tridiagonal form (or equivalent).

- *CONDENSED*: Reduced to Schur or diagonal form (or equivalent).
- *TRUNCATED*: Condensed form truncated to a smaller size.

#### See also

DSStateType

### Attributes Summary

<i>CONDENSED</i>	Constant CONDENSED of type <code>int</code>
<i>INTERMEDIATE</i>	Constant INTERMEDIATE of type <code>int</code>
<i>RAW</i>	Constant RAW of type <code>int</code>
<i>TRUNCATED</i>	Constant TRUNCATED of type <code>int</code>

### Attributes Documentation

**CONDENSED:** `int` = **CONDENSED**

Constant CONDENSED of type `int`

**INTERMEDIATE:** `int` = **INTERMEDIATE**

Constant INTERMEDIATE of type `int`

**RAW:** `int` = **RAW**

Constant RAW of type `int`

**TRUNCATED:** `int` = **TRUNCATED**

Constant TRUNCATED of type `int`

### slepc4py.SLEPc.DS.Type

**class** `slepc4py.SLEPc.DS.Type`

Bases: `object`

DS type.

- *HEP*: Dense Hermitian Eigenvalue Problem.
- *NHEP*: Dense Non-Hermitian Eigenvalue Problem.
- *GHEP*: Dense Generalized Hermitian Eigenvalue Problem.
- *GHIEP*: Dense Generalized Hermitian Indefinite Eigenvalue Problem.
- *GNHEP*: Dense Generalized Non-Hermitian Eigenvalue Problem.
- *NHEPTS*: Dense Non-Hermitian Eigenvalue Problem (special variant intended for two-sided Krylov solvers).
- *SVD*: Dense Singular Value Decomposition.
- *HSVD*: Dense Hyperbolic Singular Value Decomposition.
- *GSVD*: Dense Generalized Singular Value Decomposition.
- *PEP*: Dense Polynomial Eigenvalue Problem.
- *NEP*: Dense Nonlinear Eigenvalue Problem.

<b>See also</b>
DSType

## Attributes Summary

<i>GHEP</i>	Object GHEP of type <code>str</code>
<i>GHIEP</i>	Object GHIEP of type <code>str</code>
<i>GNHEP</i>	Object GNHEP of type <code>str</code>
<i>GSVD</i>	Object GSVD of type <code>str</code>
<i>HEP</i>	Object HEP of type <code>str</code>
<i>HSVD</i>	Object HSVD of type <code>str</code>
<i>NEP</i>	Object NEP of type <code>str</code>
<i>NHEP</i>	Object NHEP of type <code>str</code>
<i>NHEPTS</i>	Object NHEPTS of type <code>str</code>
<i>PEP</i>	Object PEP of type <code>str</code>
<i>SVD</i>	Object SVD of type <code>str</code>

## Attributes Documentation

**GHEP:** `str` = **GHEP**

Object GHEP of type `str`

**GHIEP:** `str` = **GHIEP**

Object GHIEP of type `str`

**GNHEP:** `str` = **GNHEP**

Object GNHEP of type `str`

**GSVD:** `str` = **GSVD**

Object GSVD of type `str`

**HEP:** `str` = **HEP**

Object HEP of type `str`

**HSVD:** `str` = **HSVD**

Object HSVD of type `str`

**NEP:** `str` = **NEP**

Object NEP of type `str`

**NHEP:** `str` = **NHEP**

Object NHEP of type `str`

**NHEPTS:** `str` = **NHEPTS**

Object NHEPTS of type `str`

**PEP:** `str` = **PEP**

Object PEP of type `str`

**SVD:** `str` = **SVD**

Object SVD of type `str`

## Methods Summary

<code>allocate(ld)</code>	Allocate memory for internal storage or matrices in DS.
<code>appendOptionsPrefix([prefix])</code>	Append to the prefix used for searching for all DS options in the database.
<code>cond()</code>	Compute the inf-norm condition number of the first matrix.
<code>create([comm])</code>	Create the DS object.
<code>destroy()</code>	Destroy the DS object.
<code>duplicate()</code>	Duplicate the DS object with the same type and dimensions.
<code>getArray(matname)</code>	Return the array where the data is stored.
<code>getBlockSize()</code>	Get the block size.
<code>getCompact()</code>	Get the compact storage flag.
<code>getDimensions()</code>	Get the current dimensions.
<code>getExtraRow()</code>	Get the extra row flag.
<code>getGSVDDimensions()</code>	Get the number of columns and rows of a <i>DS</i> of type <i>GSVD</i> .
<code>getHSVDDimensions()</code>	Get the number of columns of a <i>DS</i> of type <i>HSVD</i> .
<code>getLeadingDimension()</code>	Get the leading dimension of the allocated matrices.
<code>getMat(matname)</code>	Get the requested matrix as a sequential dense Mat object.
<code>getMethod()</code>	Get the method currently used in the DS.
<code>getOptionsPrefix()</code>	Get the prefix used for searching for all DS options in the database.
<code>getPEPCoefficients()</code>	Get the polynomial basis coefficients of a <i>DS</i> of type <i>PEP</i> .
<code>getPEPDegree()</code>	Get the polynomial degree of a <i>DS</i> of type <i>PEP</i> .
<code>getParallel()</code>	Get the mode of operation in parallel runs.
<code>getRefined()</code>	Get the refined vectors flag.
<code>getSVDDimensions()</code>	Get the number of columns of a <i>DS</i> of type <i>SVD</i> .
<code>getState()</code>	Get the current state.
<code>getType()</code>	Get the DS type of this object.
<code>reset()</code>	Reset the DS object.
<code>restoreMat(matname, mat)</code>	Restore the previously seized matrix.
<code>setBlockSize(bs)</code>	Set the block size.
<code>setCompact(comp)</code>	Set the compact flag for storage of matrices.
<code>setDimensions([n, l, k])</code>	Set the matrix sizes in the DS object.
<code>setExtraRow(ext)</code>	Set a flag to indicate that the matrix has one extra row.
<code>setFromOptions()</code>	Set DS options from the options database.
<code>setGSVDDimensions(m, p)</code>	Set the number of columns and rows of a <i>DS</i> of type <i>GSVD</i> .
<code>setHSVDDimensions(m)</code>	Set the number of columns of a <i>DS</i> of type <i>HSVD</i> .
<code>setIdentity(matname)</code>	Set the identity on the active part of a matrix.
<code>setMethod(meth)</code>	Set the method to be used to solve the problem.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all DS options in the database.
<code>setPEPCoefficients(pbc)</code>	Set the polynomial basis coefficients of a <i>DS</i> of type <i>PEP</i> .
<code>setPEPDegree(deg)</code>	Set the polynomial degree of a <i>DS</i> of type <i>PEP</i> .
<code>setParallel(pmode)</code>	Set the mode of operation in parallel runs.

continues on next page



Table 19 – continued from previous page

<code>setRefined(ref)</code>	Set a flag to indicate that refined vectors must be computed.
<code>setSVDDimensions(m)</code>	Set the number of columns of a <i>DS</i> of type <i>SVD</i> .
<code>setState(state)</code>	Set the state of the DS object.
<code>setType(ds_type)</code>	Set the type for the DS object.
<code>solve()</code>	Solve the problem.
<code>truncate(n[, trim])</code>	Truncate the system represented in the DS object.
<code>updateExtraRow()</code>	Ensure that the extra row gets up-to-date after a call to <i>DS.solve()</i> .
<code>vectors([matname])</code>	Compute vectors associated to the dense system such as eigenvectors.
<code>view([viewer])</code>	Print the DS data structure.

## Attributes Summary

<code>block_size</code>	The block size.
<code>compact</code>	Compact storage of matrices.
<code>extra_row</code>	If the matrix has one extra row.
<code>method</code>	The method to be used to solve the problem.
<code>parallel</code>	The mode of operation in parallel runs.
<code>refined</code>	If refined vectors must be computed.
<code>state</code>	The state of the DS object.

## Methods Documentation

### `allocate(ld)`

Allocate memory for internal storage or matrices in DS.

Logically collective.

#### Parameters

**ld** (*int*) – Leading dimension (maximum allowed dimension for the matrices, including the extra row if present).

#### Return type

None

#### Notes

If the leading dimension is different from a previously set value, then all matrices are destroyed with `reset()`.

#### See also

`getLeadingDimension`, `setDimensions`, `setExtraRow`, `reset`, `DSAllocate`

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:338 <slepc4py/SLEPc/DS.pyx#L338>`

### `appendOptionsPrefix(prefix=None)`

Append to the prefix used for searching for all DS options in the database.

Logically collective.

**Parameters**

**prefix** (*str* / *None*) – The prefix string to prepend to all DS option requests.

**Return type**

*None*

**See also**

*setOptionsPrefix, getOptionsPrefix, DSSetOptionsPrefix*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:256 <slepc4py/SLEPc/DS.pyx#L256>](#)

**cond()**

Compute the inf-norm condition number of the first matrix.

Logically collective.

**Returns**

Condition number.

**Return type**

*float*

**See also**

*DSCond*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:905 <slepc4py/SLEPc/DS.pyx#L905>](#)

**create(comm=None)**

Create the DS object.

Collective.

**Parameters**

**comm** (*Comm* / *None*) – MPI communicator; if not provided, it defaults to all processes.

**Return type**

*Self*

**See also**

*duplicate, DSCreate*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:172 <slepc4py/SLEPc/DS.pyx#L172>](#)

**destroy()**

Destroy the DS object.

Collective.

**See also**

*DSDestroy*

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:146 <slepc4py/SLEPc/DS.pyx#L146>`

**Return type**

*Self*

**duplicate()**

Duplicate the DS object with the same type and dimensions.

Collective.

**Returns**

The new object.

**Return type**

*DS*

**Notes**

This method does not copy the matrices, and the new object does not even have internal arrays allocated. Use `allocate()` to use the new *DS*.

**See also**

`create`, `allocate`, `DSDuplicate`

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:311 <slepc4py/SLEPc/DS.pyx#L311>`

**getArray(matname)**

Return the array where the data is stored.

Not collective.

**Parameters**

**matname** (*MatType*) – The selected matrix.

**Returns**

The array.

**Return type**

*ArrayScalar*

**See also**

`DSGetArray`

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:845 <slepc4py/SLEPc/DS.pyx#L845>`

**getBlockSize()**

Get the block size.

Not collective.

**Returns**

The block size.

**Return type**

*int*

**See also**

`setBlockSize`, `DSGetBlockSize`

**:sources:** ``Source code at slepc4py/SLEPc/DS.pyx:555 <slepc4py/SLEPc/DS.pyx#L555>``

**getCompact()**

Get the compact storage flag.

Not collective.

**Returns**

The flag.

**Return type**

`bool`

**See also**

`setCompact`, `DSGetCompact`

**:sources:** ``Source code at slepc4py/SLEPc/DS.pyx:639 <slepc4py/SLEPc/DS.pyx#L639>``

**getDimensions()**

Get the current dimensions.

Not collective.

**Returns**

- **n** (`int`) – The new size.
- **l** (`int`) – Number of locked (inactive) leading columns.
- **k** (`int`) – Intermediate dimension (e.g., position of arrow).
- **t** (`int`) – Truncated length.

**Return type**

`tuple[int, int, int, int]`

**Notes**

The `t` value makes sense only if `truncate()` has been called. Otherwise it is equal to `n`.

**See also**

`setDimensions`, `truncate`, `getLeadingDimension`, `DSGetDimensions`

**:sources:** ``Source code at slepc4py/SLEPc/DS.pyx:504 <slepc4py/SLEPc/DS.pyx#L504>``

**getExtraRow()**

Get the extra row flag.

Not collective.

**Returns**

The flag.

**Return type**

`bool`

**See also**

`setExtraRow`, `DSGetExtraRow`

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:686 <slepc4py/SLEPc/DS.pyx#L686>`

**getGSVDDimensions()**

Get the number of columns and rows of a *DS* of type *GSVD*.

Not collective.

**Returns**

- **m** (`int`) – The number of columns.
- **p** (`int`) – The number of rows for the second matrix.

**Return type**

`tuple[int, int]`

**See also**

`setGSVDDimensions`, `DSGSVDGetDimensions`

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1082 <slepc4py/SLEPc/DS.pyx#L1082>`

**getHSVDDimensions()**

Get the number of columns of a *DS* of type *HSVD*.

Not collective.

**Returns**

The number of columns.

**Return type**

`int`

**See also**

`setHSVDDimensions`, `DSHSVDGetDimensions`

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1037 <slepc4py/SLEPc/DS.pyx#L1037>`

**getLeadingDimension()**

Get the leading dimension of the allocated matrices.

Not collective.

**Returns**

Leading dimension (maximum allowed dimension for the matrices).

**Return type**

`int`

**See also**

[allocate](#), [setDimensions](#), [DSGetLeadingDimension](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:362 <slepc4py/SLEPc/DS.pyx#L362>](#)

**getMat(matname)**

Get the requested matrix as a sequential dense Mat object.

Not collective.

**Parameters**

**matname** ([MatType](#)) – The requested matrix.

**Returns**

The matrix.

**Return type**

[petsc4py.PETSc.Mat](#)

**Notes**

The returned matrix has sizes equal to the current *DS* dimensions (see [setDimensions\(\)](#)), and contains the values that would be obtained with [getArray\(\)](#). If the *DS* was truncated, then the number of rows is equal to the dimension prior to truncation, see [truncate\(\)](#).

When no longer needed the user must call [restoreMat\(\)](#).

**See also**

[restoreMat](#), [setDimensions](#), [getArray](#), [truncate](#), [DSGetMat](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:789 <slepc4py/SLEPc/DS.pyx#L789>](#)

**getMethod()**

Get the method currently used in the DS.

Not collective.

**Returns**

Identifier of the method.

**Return type**

[int](#)

**See also**

[setMethod](#), [DSGetMethod](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:592 <slepc4py/SLEPc/DS.pyx#L592>](#)

**getOptionsPrefix()**

Get the prefix used for searching for all DS options in the database.

Not collective.

**Returns**

The prefix string set for this DS object.

**Return type**

*str*

**See also**

*appendOptionsPrefix*, *setOptionsPrefix*, *DSsetOptionsPrefix*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:275 <slepc4py/SLEPc/DS.pyx#L275>](#)

**getPEPCoefficients()**

Get the polynomial basis coefficients of a *DS* of type *PEP*.

Not collective.

**Returns**

Coefficients.

**Return type**

*ArrayReal*

**See also**

*setPEPCoefficients*, *DSPEPGetCoefficients*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:1176 <slepc4py/SLEPc/DS.pyx#L1176>](#)

**getPEPDegree()**

Get the polynomial degree of a *DS* of type *PEP*.

Not collective.

**Returns**

The polynomial degree.

**Return type**

*int*

**See also**

*setPEPDegree*, *DSPEPGetDegree*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:1122 <slepc4py/SLEPc/DS.pyx#L1122>](#)

**getParallel()**

Get the mode of operation in parallel runs.

Not collective.

**Returns**

The parallel mode.

**Return type**

*ParallelType*

**See also**

[`setParallel`](#), [`DSGetParallel`](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:450](#) <slepc4py/SLEPc/DS.pyx#L450>

**getRefined()**

Get the refined vectors flag.

Not collective.

**Returns**

The flag.

**Return type**

`bool`

**See also**

[`setRefined`](#), [`DSGetRefined`](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:734](#) <slepc4py/SLEPc/DS.pyx#L734>

**getSVDDimensions()**

Get the number of columns of a *DS* of type *SVD*.

Not collective.

**Returns**

The number of columns.

**Return type**

`int`

**See also**

[`setSVDDimensions`](#), [`DSSVDGetDimensions`](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:995](#) <slepc4py/SLEPc/DS.pyx#L995>

**getState()**

Get the current state.

Not collective.

**Returns**

The current state.

**Return type**

*StateType*

**See also**

[`setState`](#), [`DSGetState`](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:413](#) <slepc4py/SLEPc/DS.pyx#L413>



## **getType()**

Get the DS type of this object.

Not collective.

### **Returns**

The direct solver type currently being used.

### **Return type**

`str`

### **See also**

`setType`, `DSGetType`

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:212](#) <slepc4py/SLEPc/DS.pyx#L212>

## **reset()**

Reset the DS object.

Collective.

### **See also**

`allocate`, `DSReset`

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:160](#) <slepc4py/SLEPc/DS.pyx#L160>

### **Return type**

`None`

## **restoreMat(matname, mat)**

Restore the previously seized matrix.

Not collective.

### **Parameters**

- **matname** (`MatType`) – The selected matrix.
- **mat** (`petsc4py.PETSc.Mat`) – The matrix previously obtained with `getMat()`.

### **Return type**

`None`

### **See also**

`getMat`, `DSRestoreMat`

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:824](#) <slepc4py/SLEPc/DS.pyx#L824>

## **setBlockSize(bs)**

Set the block size.

Logically collective.

### **Parameters**

**bs** (`int`) – The block size.

**Return type**

None

**See also**`getBlockSize`, `DSSetBlockSize`**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:537` `<slepc4py/SLEPc/DS.pyx#L537>`**setCompact**(*comp*)

Set the compact flag for storage of matrices.

Logically collective.

**Parameters****comp** (*bool*) – True means compact storage.**Return type**

None

**Notes**

Compact storage is used in some *DS* types such as *DS.Type.HEP* when the matrix is tridiagonal. This flag can be used to indicate whether the user provides the matrix entries via the compact form (the tridiagonal *DS.MatType.T*) or the non-compact one (*DS.MatType.A*).

The default is False.

**See also**`getCompact`, `DSSetCompact`**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:611` `<slepc4py/SLEPc/DS.pyx#L611>`**setDimensions**(*n=None, l=None, k=None*)

Set the matrix sizes in the DS object.

Logically collective.

**Parameters**

- **n** (*int* / *None*) – The new size.
- **l** (*int* / *None*) – Number of locked (inactive) leading columns.
- **k** (*int* / *None*) – Intermediate dimension (e.g., position of arrow).

**Return type**

None

**Notes**

The internal arrays are not reallocated.

Some *DS* types have additional dimensions, e.g., the number of columns in *DS.Type.SVD*. For these, you should call a specific interface function.

#### See also

[\*getDimensions\*](#), [\*allocate\*](#), [\*DSSetDimensions\*](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:469 <slepc4py/SLEPc/DS.pyx#L469>`

#### **setExtraRow**(*ext*)

Set a flag to indicate that the matrix has one extra row.

Logically collective.

##### Parameters

**ext** (*bool*) – True if the matrix has extra row.

##### Return type

None

#### Notes

In Krylov methods it is useful that the matrix representing the direct solver has one extra row, i.e., has  $(n + 1)$  rows and  $(n + 1)$  columns. If this flag is activated, all transformations applied to the right of the matrix also affect this additional row. In that case,  $(n + 1)$  must be less or equal than the leading dimension.

The default is False.

#### See also

[\*getExtraRow\*](#), [\*solve\*](#), [\*allocate\*](#), [\*DSSetExtraRow\*](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:658 <slepc4py/SLEPc/DS.pyx#L658>`

#### **setFromOptions**()

Set DS options from the options database.

Collective.

#### Notes

To see all options, run your program with the `-help` option.

#### See also

[\*setOptionsPrefix\*](#), [\*DSSetFromOptions\*](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:294 <slepc4py/SLEPc/DS.pyx#L294>`

##### Return type

None

#### **setGSVDDimensions**(*m*, *p*)

Set the number of columns and rows of a *DS* of type *GSVD*.

Logically collective.

##### Parameters

- **m** (*int*) – The number of columns.

- **p** (*int*) – The number of rows for the second matrix.

**Return type**

None

**Notes**

This call is complementary to *setDimensions()*, to provide dimensions that are specific to this *DS.Type*.

**See also**

*setDimensions*, *getGSVDDimensions*, *DSGSVDSetDimensions*

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1056 <slepc4py/SLEPc/DS.pyx#L1056>`

**setHSVDDimensions(m)**

Set the number of columns of a *DS* of type *HSVD*.

Logically collective.

**Parameters**

**m** (*int*) – The number of columns.

**Return type**

None

**Notes**

This call is complementary to *setDimensions()*, to provide a dimension that is specific to this *DS.Type*.

**See also**

*setDimensions*, *getHSVDDimensions*, *DSHSVDSetDimensions*

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1014 <slepc4py/SLEPc/DS.pyx#L1014>`

**setIdentity(matname)**

Set the identity on the active part of a matrix.

Logically collective.

**Parameters**

**matname** (*MatType*) – The matrix to be changed.

**Return type**

None

**See also**

*DSSetIdentity*

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:885 <slepc4py/SLEPc/DS.pyx#L885>`

**setMethod(meth)**

Set the method to be used to solve the problem.

Logically collective.

**Parameters**

**meth** (*int*) – An index identifying the method.

**Return type**

None

**See also**

[\*getMethod\*](#), [\*DSSetMethod\*](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:574](#) <[slepc4py/SLEPc/DS.pyx#L574](#)>

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all DS options in the database.

Logically collective.

**Parameters**

**prefix** (*str* / *None*) – The prefix string to prepend to all DS option requests.

**Return type**

None

**Notes**

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

**See also**

[\*appendOptionsPrefix\*](#), [\*getOptionsPrefix\*](#), [\*DSSetOptionsPrefix\*](#)

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:231](#) <[slepc4py/SLEPc/DS.pyx#L231](#)>

**setPEPCoefficients**(*pbc*)

Set the polynomial basis coefficients of a *DS* of type *PEP*.

Logically collective.

**Parameters**

**pbc** (*Sequence[float]*) – Coefficients.

**Return type**

None

**Notes**

This function is required only in the case of a polynomial specified in a non-monomial basis, to provide the coefficients that will be used during the linearization, multiplying the identity blocks on the three main diagonal blocks. Depending on the polynomial basis (Chebyshev, Legendre, ...) the coefficients must be different.

There must be a total of  $3(d+1)$  coefficients, where  $d$  is the degree of the polynomial. The coefficients are arranged in three groups,  $a_i, b_i, c_i$ , according to the definition of the three-term recurrence. In the case of the monomial basis,  $a_i = 1$  and  $b_i = c_i = 0$ , in which case it is not necessary to invoke this function.

**See also**

[getPEPCoefficients](#), [DSPEPSetCoefficients](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1141 <slepc4py/SLEPc/DS.pyx#L1141>`

**setPEPDegree**(*deg*)

Set the polynomial degree of a *DS* of type *PEP*.

Logically collective.

**Parameters**

**deg** (*int*) – The polynomial degree.

**Return type**

None

**See also**

[getPEPDegree](#), [DSPEPSetDegree](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1104 <slepc4py/SLEPc/DS.pyx#L1104>`

**setParallel**(*pmode*)

Set the mode of operation in parallel runs.

Logically collective.

**Parameters**

**pmode** (*ParallelType*) – The parallel mode.

**Return type**

None

**See also**

[getParallel](#), [DSSetParallel](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:432 <slepc4py/SLEPc/DS.pyx#L432>`

**setRefined**(*ref*)

Set a flag to indicate that refined vectors must be computed.

Logically collective.

**Parameters**

**ref** (*bool*) – True if refined vectors must be used.

**Return type**

None

**Notes**

Normally the vectors returned in *DS.MatType.X* are eigenvectors of the projected matrix. With this flag activated, [vectors\(\)](#) will return the right singular vector of the smallest singular value of matrix  $\hat{A} - \eta I$ , where  $\hat{A}$  is the extended matrix (with extra row) and  $\eta$  is the Ritz value. This is used in the refined Ritz approximation.

The default is False.

#### See also

[`getRefined`](#), [`vectors`](#), [`setExtraRow`](#), [`DSSetRefined`](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:705 <slepc4py/SLEPc/DS.pyx#L705>`

#### **setSVDDimensions(*m*)**

Set the number of columns of a *DS* of type *SVD*.

Logically collective.

##### Parameters

**m** (*int*) – The number of columns.

##### Return type

None

#### Notes

This call is complementary to [`setDimensions\(\)`](#), to provide a dimension that is specific to this *DS.Type*.

#### See also

[`setDimensions`](#), [`getSVDDimensions`](#), [`DSSVDSetDimensions`](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:972 <slepc4py/SLEPc/DS.pyx#L972>`

#### **setState(*state*)**

Set the state of the DS object.

Logically collective.

##### Parameters

**state** (*StateType*) – The new state.

##### Return type

None

#### Notes

The state indicates that the dense system is in an initial state (raw), in an intermediate state (such as tridiagonal, Hessenberg or Hessenberg-triangular), in a condensed state (such as diagonal, Schur or generalized Schur), or in a truncated state.

The state is automatically changed in functions such as [`solve\(\)`](#) or [`truncate\(\)`](#). This function is normally used to return to the raw state when the condensed structure is destroyed, or to indicate that [`solve\(\)`](#) must start with a problem that already has an intermediate form.

#### See also

[`getState`](#), [`solve`](#), [`truncate`](#), [`DSSetState`](#)

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:381 <slepc4py/SLEPc/DS.pyx#L381>`

**setType**(*ds\_type*)

Set the type for the DS object.

Logically collective.

**Parameters**

**ds\_type** (*Type* / *str*) – The direct solver type to be used.

**Return type**

*None*

**See also**

*getType*, *DSSetType*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:193](#) <slepc4py/SLEPc/DS.pyx#L193>

**solve**()

Solve the problem.

Logically collective.

**Returns**

Eigenvalues or singular values.

**Return type**

*ArrayScalar*

**See also**

*DSSolve*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:924](#) <slepc4py/SLEPc/DS.pyx#L924>

**truncate**(*n*, *trim=False*)

Truncate the system represented in the DS object.

Logically collective.

**Parameters**

- **n** (*int*) – The new size.
- **trim** (*bool*) – A flag to indicate if the factorization must be trimmed.

**Return type**

*None*

**See also**

*setDimensions*, *setExtraRow*, *DSTruncate*

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:753](#) <slepc4py/SLEPc/DS.pyx#L753>

**updateExtraRow**()

Ensure that the extra row gets up-to-date after a call to *DS.solve()*.

Logically collective.



Perform all necessary operations so that the extra row gets up-to-date after a call to `DS.solve()`.

**See also**

`DSUpdateExtraRow`

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:774 <slepc4py/SLEPc/DS.pyx#L774>](#)

**Return type**

`None`

**vectors**(*matname=X*)

Compute vectors associated to the dense system such as eigenvectors.

Logically collective.

**Parameters**

**matname** – The matrix, used to indicate which vectors are required.

**Return type**

`None`

**See also**

`DSVectors`

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:952 <slepc4py/SLEPc/DS.pyx#L952>](#)

**view**(*viewer=None*)

Print the DS data structure.

Collective.

**Parameters**

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

`None`

**See also**

`DSView`

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:127 <slepc4py/SLEPc/DS.pyx#L127>](#)

## Attributes Documentation

**block\_size**

The block size.

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:1218 <slepc4py/SLEPc/DS.pyx#L1218>](#)

**compact**

Compact storage of matrices.

**:sources:** [Source code at slepc4py/SLEPc/DS.pyx:1232 <slepc4py/SLEPc/DS.pyx#L1232>](#)

## extra\_row

If the matrix has one extra row.

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1239 <slepc4py/SLEPc/DS.pyx#L1239>`

## method

The method to be used to solve the problem.

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1225 <slepc4py/SLEPc/DS.pyx#L1225>`

## parallel

The mode of operation in parallel runs.

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1211 <slepc4py/SLEPc/DS.pyx#L1211>`

## refined

If refined vectors must be computed.

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1246 <slepc4py/SLEPc/DS.pyx#L1246>`

## state

The state of the DS object.

**:sources:** `Source code at slepc4py/SLEPc/DS.pyx:1204 <slepc4py/SLEPc/DS.pyx#L1204>`

## slepc4py.SLEPc.EPS

### class slepc4py.SLEPc.EPS

Bases: `Object`

Eigenvalue Problem Solver.

The Eigenvalue Problem Solver (*EPS*) is the object provided by slepc4py for specifying a linear eigenvalue problem, either in standard or generalized form. It provides uniform and efficient access to all of the linear eigensolvers included in the package.

### Enumerations

<i>Balance</i>	EPS type of balancing used for non-Hermitian problems.
<i>CISSExtraction</i>	EPS CISS extraction technique.
<i>CISSQuadRule</i>	EPS CISS quadrature rule.
<i>Conv</i>	EPS convergence test.
<i>ConvergedReason</i>	EPS convergence reasons.
<i>ErrorType</i>	EPS error type to assess accuracy of computed solutions.
<i>Extraction</i>	EPS extraction technique.
<i>KrylovSchurBSEType</i>	EPS Krylov-Schur method for BSE problems.
<i>LanczosReorthogType</i>	EPS Lanczos reorthogonalization type.
<i>PowerShiftType</i>	EPS Power shift type.
<i>ProblemType</i>	EPS problem type.
<i>Stop</i>	EPS stopping test.
<i>Type</i>	EPS type.
<i>Which</i>	EPS desired part of spectrum.

## slepc4py.SLEPc.EPS.Balance

**class** slepc4py.SLEPc.EPS.Balance

Bases: `object`

EPS type of balancing used for non-Hermitian problems.

- *NONE*: None.
- *ONESIDE*: One-sided balancing.
- *TWOSIDE*: Two-sided balancing.
- *USER*: User-provided balancing matrices.

### See also

[EPSCBalance](#)

## Attributes Summary

<i>NONE</i>	Constant NONE of type <code>int</code>
<i>ONESIDE</i>	Constant ONESIDE of type <code>int</code>
<i>TWOSIDE</i>	Constant TWOSIDE of type <code>int</code>
<i>USER</i>	Constant USER of type <code>int</code>

## Attributes Documentation

**NONE:** `int` = **NONE**

Constant NONE of type `int`

**ONESIDE:** `int` = **ONESIDE**

Constant ONESIDE of type `int`

**TWOSIDE:** `int` = **TWOSIDE**

Constant TWOSIDE of type `int`

**USER:** `int` = **USER**

Constant USER of type `int`

## slepc4py.SLEPc.EPS.CISSExtraction

**class** slepc4py.SLEPc.EPS.CISSExtraction

Bases: `object`

EPS CISS extraction technique.

- *RITZ*: Ritz extraction.
- *HANKEL*: Extraction via Hankel eigenproblem.

### See also

[EPSCISSExtraction](#)

## Attributes Summary

<i>HANKEL</i>	Constant HANKEL of type <code>int</code>
<i>RITZ</i>	Constant RITZ of type <code>int</code>

## Attributes Documentation

**HANKEL:** `int` = HANKEL

Constant HANKEL of type `int`

**RITZ:** `int` = RITZ

Constant RITZ of type `int`

## `slepc4py.SLEPc.EPS.CISSQuadRule`

**class** `slepc4py.SLEPc.EPS.CISSQuadRule`

Bases: `object`

EPS CISS quadrature rule.

- *TRAPEZOIDAL*: Trapezoidal rule.
- *CHEBYSHEV*: Chebyshev points.

### See also

`EPSCISSQuadRule`

## Attributes Summary

<i>CHEBYSHEV</i>	Constant CHEBYSHEV of type <code>int</code>
<i>TRAPEZOIDAL</i>	Constant TRAPEZOIDAL of type <code>int</code>

## Attributes Documentation

**CHEBYSHEV:** `int` = CHEBYSHEV

Constant CHEBYSHEV of type `int`

**TRAPEZOIDAL:** `int` = TRAPEZOIDAL

Constant TRAPEZOIDAL of type `int`

## `slepc4py.SLEPc.EPS.Conv`

**class** `slepc4py.SLEPc.EPS.Conv`

Bases: `object`

EPS convergence test.

- *ABS*: Absolute convergence test.
- *REL*: Convergence test relative to the eigenvalue.
- *NORM*: Convergence test relative to the matrix norms.
- *USER*: User-defined convergence test.

<b>See also</b>
EPSCnv

## Attributes Summary

<i>ABS</i>	Constant ABS of type <code>int</code>
<i>NORM</i>	Constant NORM of type <code>int</code>
<i>REL</i>	Constant REL of type <code>int</code>
<i>USER</i>	Constant USER of type <code>int</code>

## Attributes Documentation

**ABS:** `int` = ABS

Constant ABS of type `int`

**NORM:** `int` = NORM

Constant NORM of type `int`

**REL:** `int` = REL

Constant REL of type `int`

**USER:** `int` = USER

Constant USER of type `int`

## slepc4py.SLEPc.EPS.ConvergedReason

**class** slepc4py.SLEPc.EPS.ConvergedReason

Bases: `object`

EPS convergence reasons.

- *CONVERGED\_TOL*: All eigenpairs converged to requested tolerance.
- *CONVERGED\_USER*: User-defined convergence criterion satisfied.
- *DIVERGED\_ITS*: Maximum number of iterations exceeded.
- *DIVERGED\_BREAKDOWN*: Solver failed due to breakdown.
- *DIVERGED\_SYMMETRY\_LOST*: Lanczos-type method could not preserve symmetry.
- *CONVERGED\_ITERATING*: Iteration not finished yet.

<b>See also</b>
EPSConvergedReason

## Attributes Summary

<i>CONVERGED_ITERATING</i>	Constant CONVERGED_ITERATING of type <code>int</code>
<i>CONVERGED_TOL</i>	Constant CONVERGED_TOL of type <code>int</code>
<i>CONVERGED_USER</i>	Constant CONVERGED_USER of type <code>int</code>

continues on next page

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<i>DIVERGED_BREAKDOWN</i>	Constant DIVERGED_BREAKDOWN of type <code>int</code>
<i>DIVERGED_ITS</i>	Constant DIVERGED_ITS of type <code>int</code>
<i>DIVERGED_SYMMETRY_LOST</i>	Constant DIVERGED_SYMMETRY_LOST of type <code>int</code>
<i>ITERATING</i>	Constant ITERATING of type <code>int</code>

### Attributes Documentation

**CONVERGED\_ITERATING:** `int` = **CONVERGED\_ITERATING**

Constant CONVERGED\_ITERATING of type `int`

**CONVERGED\_TOL:** `int` = **CONVERGED\_TOL**

Constant CONVERGED\_TOL of type `int`

**CONVERGED\_USER:** `int` = **CONVERGED\_USER**

Constant CONVERGED\_USER of type `int`

**DIVERGED\_BREAKDOWN:** `int` = **DIVERGED\_BREAKDOWN**

Constant DIVERGED\_BREAKDOWN of type `int`

**DIVERGED\_ITS:** `int` = **DIVERGED\_ITS**

Constant DIVERGED\_ITS of type `int`

**DIVERGED\_SYMMETRY\_LOST:** `int` = **DIVERGED\_SYMMETRY\_LOST**

Constant DIVERGED\_SYMMETRY\_LOST of type `int`

**ITERATING:** `int` = **ITERATING**

Constant ITERATING of type `int`

### `slepc4py.SLEPc.EPS.ErrorType`

**class** `slepc4py.SLEPc.EPS.ErrorType`

Bases: `object`

EPS error type to assess accuracy of computed solutions.

- *ABSOLUTE*: Absolute error.
- *RELATIVE*: Relative error.
- *BACKWARD*: Backward error.

#### See also

`EPSErrorType`

### Attributes Summary

<i>ABSOLUTE</i>	Constant ABSOLUTE of type <code>int</code>
<i>BACKWARD</i>	Constant BACKWARD of type <code>int</code>
<i>RELATIVE</i>	Constant RELATIVE of type <code>int</code>

## Attributes Documentation

**ABSOLUTE:** `int` = `ABSOLUTE`

Constant `ABSOLUTE` of type `int`

**BACKWARD:** `int` = `BACKWARD`

Constant `BACKWARD` of type `int`

**RELATIVE:** `int` = `RELATIVE`

Constant `RELATIVE` of type `int`

## slepc4py.SLEPc.EPS.Extraction

**class** `slepc4py.SLEPc.EPS.Extraction`

Bases: `object`

EPS extraction technique.

- *RITZ*: Standard Rayleigh-Ritz extraction.
- *HARMONIC*: Harmonic extraction.
- *HARMONIC\_RELATIVE*: Harmonic extraction relative to the eigenvalue.
- *HARMONIC\_RIGHT*: Harmonic extraction for rightmost eigenvalues.
- *HARMONIC\_LARGEST*: Harmonic extraction for largest magnitude (without target).
- *REFINED*: Refined extraction.
- *REFINED\_HARMONIC*: Refined harmonic extraction.

### See also

`EPSExtraction`

## Attributes Summary

<i>HARMONIC</i>	Constant <code>HARMONIC</code> of type <code>int</code>
<i>HARMONIC_LARGEST</i>	Constant <code>HARMONIC_LARGEST</code> of type <code>int</code>
<i>HARMONIC_RELATIVE</i>	Constant <code>HARMONIC_RELATIVE</code> of type <code>int</code>
<i>HARMONIC_RIGHT</i>	Constant <code>HARMONIC_RIGHT</code> of type <code>int</code>
<i>REFINED</i>	Constant <code>REFINED</code> of type <code>int</code>
<i>REFINED_HARMONIC</i>	Constant <code>REFINED_HARMONIC</code> of type <code>int</code>
<i>RITZ</i>	Constant <code>RITZ</code> of type <code>int</code>

## Attributes Documentation

**HARMONIC:** `int` = `HARMONIC`

Constant `HARMONIC` of type `int`

**HARMONIC\_LARGEST:** `int` = `HARMONIC_LARGEST`

Constant `HARMONIC_LARGEST` of type `int`

**HARMONIC\_RELATIVE:** `int` = `HARMONIC_RELATIVE`

Constant `HARMONIC_RELATIVE` of type `int`

**HARMONIC\_RIGHT:** `int` = **HARMONIC\_RIGHT**

Constant **HARMONIC\_RIGHT** of type `int`

**REFINED:** `int` = **REFINED**

Constant **REFINED** of type `int`

**REFINED\_HARMONIC:** `int` = **REFINED\_HARMONIC**

Constant **REFINED\_HARMONIC** of type `int`

**RITZ:** `int` = **RITZ**

Constant **RITZ** of type `int`

## **slepc4py.SLEPc.EPS.KrylovSchurBSEType**

**class** `slepc4py.SLEPc.EPS.KrylovSchurBSEType`

Bases: `object`

EPS Krylov-Schur method for BSE problems.

- *SHAO*: Lanczos recurrence for H square.
- *GRUNING*: Lanczos recurrence for H.
- *PROJECTEDBSE*: Lanczos where the projected problem has BSE structure.

### **See also**

`EPKrylovSchurBSEType`

## **Attributes Summary**

*GRUNING*

Constant **GRUNING** of type `int`

*PROJECTEDBSE*

Constant **PROJECTEDBSE** of type `int`

*SHAO*

Constant **SHAO** of type `int`

## **Attributes Documentation**

**GRUNING:** `int` = **GRUNING**

Constant **GRUNING** of type `int`

**PROJECTEDBSE:** `int` = **PROJECTEDBSE**

Constant **PROJECTEDBSE** of type `int`

**SHAO:** `int` = **SHAO**

Constant **SHAO** of type `int`

## **slepc4py.SLEPc.EPS.LanczosReorthogType**

**class** `slepc4py.SLEPc.EPS.LanczosReorthogType`

Bases: `object`

EPS Lanczos reorthogonalization type.

- *LOCAL*: Local reorthogonalization only.
- *FULL*: Full reorthogonalization.



- *SELECTIVE*: Selective reorthogonalization.
- *PERIODIC*: Periodic reorthogonalization.
- *PARTIAL*: Partial reorthogonalization.
- *DELAYED*: Delayed reorthogonalization.

#### See also

EPSSLanczosReorthogType

### Attributes Summary

<i>DELAYED</i>	Constant DELAYED of type <i>int</i>
<i>FULL</i>	Constant FULL of type <i>int</i>
<i>LOCAL</i>	Constant LOCAL of type <i>int</i>
<i>PARTIAL</i>	Constant PARTIAL of type <i>int</i>
<i>PERIODIC</i>	Constant PERIODIC of type <i>int</i>
<i>SELECTIVE</i>	Constant SELECTIVE of type <i>int</i>

### Attributes Documentation

**DELAYED:** *int* = DELAYED

Constant DELAYED of type *int*

**FULL:** *int* = FULL

Constant FULL of type *int*

**LOCAL:** *int* = LOCAL

Constant LOCAL of type *int*

**PARTIAL:** *int* = PARTIAL

Constant PARTIAL of type *int*

**PERIODIC:** *int* = PERIODIC

Constant PERIODIC of type *int*

**SELECTIVE:** *int* = SELECTIVE

Constant SELECTIVE of type *int*

### slepc4py.SLEPc.EPS.PowerShiftType

**class** slepc4py.SLEPc.EPS.PowerShiftType

Bases: *object*

EPS Power shift type.

- *CONSTANT*: Constant shift.
- *RAYLEIGH*: Rayleigh quotient.
- *WILKINSON*: Wilkinson shift.

<b>See also</b>
<a href="#">EPSPowerShiftType</a>

### Attributes Summary

<i>CONSTANT</i>	Constant <i>CONSTANT</i> of type <i>int</i>
<i>RAYLEIGH</i>	Constant <i>RAYLEIGH</i> of type <i>int</i>
<i>WILKINSON</i>	Constant <i>WILKINSON</i> of type <i>int</i>

### Attributes Documentation

**CONSTANT:** *int* = *CONSTANT*

Constant *CONSTANT* of type *int*

**RAYLEIGH:** *int* = *RAYLEIGH*

Constant *RAYLEIGH* of type *int*

**WILKINSON:** *int* = *WILKINSON*

Constant *WILKINSON* of type *int*

### `slepc4py.SLEPc.EPS.ProblemType`

**class** `slepc4py.SLEPc.EPS.ProblemType`

Bases: `object`

EPS problem type.

- *HEP*: Hermitian eigenproblem.
- *NHEP*: Non-Hermitian eigenproblem.
- *GHEP*: Generalized Hermitian eigenproblem.
- *GNHEP*: Generalized Non-Hermitian eigenproblem.
- *PGNHEP*: Generalized Non-Hermitian eigenproblem with positive definite *B*.
- *GHIEP*: Generalized Hermitian-indefinite eigenproblem.
- *BSE*: Structured Bethe-Salpeter eigenproblem.
- *HAMILT*: Hamiltonian eigenproblem.

<b>See also</b>
<a href="#">EPSPProblemType</a>

### Attributes Summary

<i>BSE</i>	Constant <i>BSE</i> of type <i>int</i>
<i>GHEP</i>	Constant <i>GHEP</i> of type <i>int</i>
<i>GHIEP</i>	Constant <i>GHIEP</i> of type <i>int</i>

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Table 32 – continued from previous page

<i>GNHEP</i>	Constant GNHEP of type <code>int</code>
<i>HAMILT</i>	Constant HAMILT of type <code>int</code>
<i>HEP</i>	Constant HEP of type <code>int</code>
<i>NHEP</i>	Constant NHEP of type <code>int</code>
<i>PGNHEP</i>	Constant PGNHEP of type <code>int</code>

### Attributes Documentation

**BSE:** `int` = BSE

Constant BSE of type `int`

**GHEP:** `int` = GHEP

Constant GHEP of type `int`

**GHIEP:** `int` = GHIEP

Constant GHIEP of type `int`

**GNHEP:** `int` = GNHEP

Constant GNHEP of type `int`

**HAMILT:** `int` = HAMILT

Constant HAMILT of type `int`

**HEP:** `int` = HEP

Constant HEP of type `int`

**NHEP:** `int` = NHEP

Constant NHEP of type `int`

**PGNHEP:** `int` = PGNHEP

Constant PGNHEP of type `int`

### slepc4py.SLEPc.EPS.Stop

**class** slepc4py.SLEPc.EPS.Stop

Bases: `object`

EPS stopping test.

- *BASIC*: Default stopping test.
- *USER*: User-defined stopping test.
- *THRESHOLD*: Threshold stopping test.

#### See also

EPSSStop

### Attributes Summary

<i>BASIC</i>	Constant BASIC of type <code>int</code>
<i>THRESHOLD</i>	Constant THRESHOLD of type <code>int</code>

continues on next page

Table 33 – continued from previous page

<i>USER</i>	Constant <i>USER</i> of type <i>int</i>
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### Attributes Documentation

**BASIC:** *int* = BASIC

Constant BASIC of type *int*

**THRESHOLD:** *int* = THRESHOLD

Constant THRESHOLD of type *int*

**USER:** *int* = USER

Constant USER of type *int*

### slepc4py.SLEPc.EPS.Type

**class** slepc4py.SLEPc.EPS.Type

Bases: *object*

EPS type.

Native eigenvalue solvers.

- *POWER*: Power Iteration, Inverse Iteration, RQI.
- *SUBSPACE*: Subspace Iteration.
- *ARNOLDI*: Arnoldi.
- *LANCZOS*: Lanczos.
- *KRYLOV-SCHUR*: Krylov-Schur (default).
- *GD*: Generalized Davidson.
- *JD*: Jacobi-Davidson.
- *RQCG*: Rayleigh Quotient Conjugate Gradient.
- *LOBPCG*: Locally Optimal Block Preconditioned Conjugate Gradient.
- *CISS*: Contour Integral Spectrum Slicing.
- *LYAPII*: Lyapunov inverse iteration.

Wrappers to external eigensolvers (should be enabled during installation of SLEPc).

- *LAPACK*: Sequential dense eigensolver.
- *ARPACK*: Iterative Krylov-based eigensolver.
- *BLOPEX*: Implementation of LOBPCG.
- *PRIMME*: Iterative eigensolvers of Davidson type.
- *FEAST*: Contour integral eigensolver.
- *SCALAPACK*: Parallel dense eigensolver for symmetric problems.
- *ELPA*: Parallel dense eigensolver for symmetric problems.
- *ELEMENTAL*: Parallel dense eigensolver for symmetric problems.
- *EVSL*: Iterative eigensolver based on polynomial filters.
- *CHASE*: Subspace iteration accelerated with polynomials.

<b>See also</b>
EPSType

## Attributes Summary

<i>ARNOLDI</i>	Object ARNOLDI of type <code>str</code>
<i>ARPACK</i>	Object ARPACK of type <code>str</code>
<i>BLOPEX</i>	Object BLOPEX of type <code>str</code>
<i>CHASE</i>	Object CHASE of type <code>str</code>
<i>CISS</i>	Object CISS of type <code>str</code>
<i>ELEMENTAL</i>	Object ELEMENTAL of type <code>str</code>
<i>ELPA</i>	Object ELPA of type <code>str</code>
<i>EVSL</i>	Object EVSL of type <code>str</code>
<i>FEAST</i>	Object FEAST of type <code>str</code>
<i>GD</i>	Object GD of type <code>str</code>
<i>JD</i>	Object JD of type <code>str</code>
<i>KRYLOV SCHUR</i>	Object KRYLOV SCHUR of type <code>str</code>
<i>LANCZOS</i>	Object LANCZOS of type <code>str</code>
<i>LAPACK</i>	Object LAPACK of type <code>str</code>
<i>LOBPCG</i>	Object LOBPCG of type <code>str</code>
<i>LYAPII</i>	Object LYAPII of type <code>str</code>
<i>POWER</i>	Object POWER of type <code>str</code>
<i>PRIMME</i>	Object PRIMME of type <code>str</code>
<i>RQCG</i>	Object RQCG of type <code>str</code>
<i>SCALAPACK</i>	Object SCALAPACK of type <code>str</code>
<i>SUBSPACE</i>	Object SUBSPACE of type <code>str</code>

## Attributes Documentation

**ARNOLDI:** `str` = **ARNOLDI**

Object ARNOLDI of type `str`

**ARPACK:** `str` = **ARPACK**

Object ARPACK of type `str`

**BLOPEX:** `str` = **BLOPEX**

Object BLOPEX of type `str`

**CHASE:** `str` = **CHASE**

Object CHASE of type `str`

**CISS:** `str` = **CISS**

Object CISS of type `str`

**ELEMENTAL:** `str` = **ELEMENTAL**

Object ELEMENTAL of type `str`

**ELPA:** `str` = **ELPA**

Object ELPA of type `str`

**EVSL:** `str` = **EVSL**

Object EVSL of type `str`

**FEAST:** `str` = **FEAST**

Object FEAST of type `str`

**GD:** `str` = **GD**

Object GD of type `str`

**JD:** `str` = **JD**

Object JD of type `str`

**KRYLOV SCHUR:** `str` = **KRYLOV SCHUR**

Object KRYLOV SCHUR of type `str`

**LANCZOS:** `str` = **LANCZOS**

Object LANCZOS of type `str`

**LAPACK:** `str` = **LAPACK**

Object LAPACK of type `str`

**LOBPCG:** `str` = **LOBPCG**

Object LOBPCG of type `str`

**LYAPII:** `str` = **LYAPII**

Object LYAPII of type `str`

**POWER:** `str` = **POWER**

Object POWER of type `str`

**PRIMME:** `str` = **PRIMME**

Object PRIMME of type `str`

**RQCG:** `str` = **RQCG**

Object RQCG of type `str`

**SCALAPACK:** `str` = **SCALAPACK**

Object SCALAPACK of type `str`

**SUBSPACE:** `str` = **SUBSPACE**

Object SUBSPACE of type `str`

### **slepc4py.SLEPc.EPS.Which**

**class** slepc4py.SLEPc.EPS.**Which**

Bases: `object`

EPS desired part of spectrum.

- `LARGEST_MAGNITUDE`: Largest magnitude (default).
- `SMALLEST_MAGNITUDE`: Smallest magnitude.
- `LARGEST_REAL`: Largest real parts.
- `SMALLEST_REAL`: Smallest real parts.
- `LARGEST_IMAGINARY`: Largest imaginary parts in magnitude.
- `SMALLEST_IMAGINARY`: Smallest imaginary parts in magnitude.
- `TARGET_MAGNITUDE`: Closest to target (in magnitude).
- `TARGET_REAL`: Real part closest to target.

- *TARGET\_IMAGINARY*: Imaginary part closest to target.
- *ALL*: All eigenvalues in an interval.
- *USER*: User defined selection.

#### See also

EPSWhich

### Attributes Summary

<i>ALL</i>	Constant ALL of type <code>int</code>
<i>LARGEST_IMAGINARY</i>	Constant LARGEST_IMAGINARY of type <code>int</code>
<i>LARGEST_MAGNITUDE</i>	Constant LARGEST_MAGNITUDE of type <code>int</code>
<i>LARGEST_REAL</i>	Constant LARGEST_REAL of type <code>int</code>
<i>SMALLEST_IMAGINARY</i>	Constant SMALLEST_IMAGINARY of type <code>int</code>
<i>SMALLEST_MAGNITUDE</i>	Constant SMALLEST_MAGNITUDE of type <code>int</code>
<i>SMALLEST_REAL</i>	Constant SMALLEST_REAL of type <code>int</code>
<i>TARGET_IMAGINARY</i>	Constant TARGET_IMAGINARY of type <code>int</code>
<i>TARGET_MAGNITUDE</i>	Constant TARGET_MAGNITUDE of type <code>int</code>
<i>TARGET_REAL</i>	Constant TARGET_REAL of type <code>int</code>
<i>USER</i>	Constant USER of type <code>int</code>

### Attributes Documentation

**ALL:** `int` = **ALL**

Constant ALL of type `int`

**LARGEST\_IMAGINARY:** `int` = **LARGEST\_IMAGINARY**

Constant LARGEST\_IMAGINARY of type `int`

**LARGEST\_MAGNITUDE:** `int` = **LARGEST\_MAGNITUDE**

Constant LARGEST\_MAGNITUDE of type `int`

**LARGEST\_REAL:** `int` = **LARGEST\_REAL**

Constant LARGEST\_REAL of type `int`

**SMALLEST\_IMAGINARY:** `int` = **SMALLEST\_IMAGINARY**

Constant SMALLEST\_IMAGINARY of type `int`

**SMALLEST\_MAGNITUDE:** `int` = **SMALLEST\_MAGNITUDE**

Constant SMALLEST\_MAGNITUDE of type `int`

**SMALLEST\_REAL:** `int` = **SMALLEST\_REAL**

Constant SMALLEST\_REAL of type `int`

**TARGET\_IMAGINARY:** `int` = **TARGET\_IMAGINARY**

Constant TARGET\_IMAGINARY of type `int`

**TARGET\_MAGNITUDE:** `int` = **TARGET\_MAGNITUDE**

Constant TARGET\_MAGNITUDE of type `int`

**TARGET\_REAL:** `int` = **TARGET\_REAL**

Constant TARGET\_REAL of type `int`

**USER:** `int = USER`

Constant USER of type `int`

## Methods Summary

<code>appendOptionsPrefix([prefix])</code>	Append to the prefix used for searching for all EPS options in the database.
<code>cancelMonitor()</code>	Clear all monitors for an <i>EPS</i> object.
<code>computeError(i[, etype])</code>	Compute the error associated with the i-th computed eigenpair.
<code>create([comm])</code>	Create the EPS object.
<code>destroy()</code>	Destroy the EPS object.
<code>errorView([etype, viewer])</code>	Display the errors associated with the computed solution.
<code>getArbitrarySelection()</code>	Get the arbitrary selection function.
<code>getArnoldiDelayed()</code>	Get the type of reorthogonalization used during the Arnoldi iteration.
<code>getBV()</code>	Get the basis vectors object associated to the eigensolver.
<code>getBalance()</code>	Get the balancing type used by the EPS, and the associated parameters.
<code>getCISSExtraction()</code>	Get the extraction technique used in the CISS solver.
<code>getCISSKSPs()</code>	Get the array of linear solver objects associated with the CISS solver.
<code>getCISSQuadRule()</code>	Get the quadrature rule used in the CISS solver.
<code>getCISSRefinement()</code>	Get the values of various refinement parameters in the CISS solver.
<code>getCISSSizes()</code>	Get the values of various size parameters in the CISS solver.
<code>getCISSThreshold()</code>	Get the values of various threshold parameters in the CISS solver.
<code>getCISSUseST()</code>	Get the flag indicating the use of the <i>ST</i> object in the CISS solver.
<code>getConverged()</code>	Get the number of converged eigenpairs.
<code>getConvergedReason()</code>	Get the reason why the <code>solve()</code> iteration was stopped.
<code>getConvergenceTest()</code>	Get how to compute the error estimate used in the convergence test.
<code>getDS()</code>	Get the direct solver associated to the eigensolver.
<code>getDimensions()</code>	Get number of eigenvalues to compute and the dimension of the subspace.
<code>getEigenpair(i[, Vr, Vi])</code>	Get the i-th solution of the eigenproblem as computed by <code>solve()</code> .
<code>getEigenvalue(i)</code>	Get the i-th eigenvalue as computed by <code>solve()</code> .
<code>getEigenvalueComparison()</code>	Get the eigenvalue comparison function.
<code>getEigenvector(i[, Vr, Vi])</code>	Get the i-th right eigenvector as computed by <code>solve()</code> .
<code>getErrorEstimate(i)</code>	Get the error estimate associated to the i-th computed eigenpair.
<code>getExtraction()</code>	Get the extraction type used by the EPS object.
<code>getGDBOrth()</code>	Get the orthogonalization used in the search subspace.

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<code>getGDBlockSize()</code>	Get the number of vectors to be added to the searching space.
<code>getGDDoubleExpansion()</code>	Get a flag indicating whether the double expansion variant is active.
<code>getGDInitialSize()</code>	Get the initial size of the searching space.
<code>getGDKrylovStart()</code>	Get a flag indicating if the search subspace is started with a Krylov basis.
<code>getGDRestart()</code>	Get the number of vectors of the search space after restart.
<code>getInterval()</code>	Get the computational interval for spectrum slicing.
<code>getInvariantSubspace()</code>	Get an orthonormal basis of the computed invariant subspace.
<code>getIterationNumber()</code>	Get the current iteration number.
<code>getJDBOrth()</code>	Get the orthogonalization used in the search subspace.
<code>getJDBlockSize()</code>	Get the number of vectors to be added to the searching space.
<code>getJDConstCorrectionTol()</code>	Get the flag indicating if the dynamic stopping is being used.
<code>getJDFix()</code>	Get the threshold for changing the target in the correction equation.
<code>getJDInitialSize()</code>	Get the initial size of the searching space.
<code>getJDKrylovStart()</code>	Get a flag indicating if the search subspace is started with a Krylov basis.
<code>getJDRestart()</code>	Get the number of vectors of the search space after restart.
<code>getKrylovSchurBSEType()</code>	Get the method used for BSE structured eigenproblems (Krylov-Schur).
<code>getKrylovSchurDetectZeros()</code>	Get the flag that enforces zero detection in spectrum slicing.
<code>getKrylovSchurDimensions()</code>	Get the dimensions used for each subsolve step (spectrum slicing).
<code>getKrylovSchurInertias()</code>	Get the values of the shifts and their corresponding inertias.
<code>getKrylovSchurKSP()</code>	Get the linear solver object associated with the internal <a href="#">EPS</a> object.
<code>getKrylovSchurLocking()</code>	Get the locking flag used in the Krylov-Schur method.
<code>getKrylovSchurPartitions()</code>	Get the number of partitions of the communicator (spectrum slicing).
<code>getKrylovSchurRestart()</code>	Get the restart parameter used in the Krylov-Schur method.
<code>getKrylovSchurSubcommInfo()</code>	Get information related to the case of doing spectrum slicing.
<code>getKrylovSchurSubcommMats()</code>	Get the eigenproblem matrices stored in the subcommunicator.
<code>getKrylovSchurSubcommPairs(i[, v])</code>	Get the i-th eigenpair stored in the multi-communicator of the process.
<code>getKrylovSchurSubintervals()</code>	Get the points that delimit the subintervals.
<code>getLOBPCGBlockSize()</code>	Get the block size used in the LOBPCG method.
<code>getLOBPCGLocking()</code>	Get the locking flag used in the LOBPCG method.
<code>getLOBPCGRestart()</code>	Get the restart parameter used in the LOBPCG method.

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<code>getLanczosReorthogType()</code>	Get the type of reorthogonalization used during the Lanczos iteration.
<code>getLeftEigenvector(i[, Wr, Wi])</code>	Get the $i$ -th left eigenvector as computed by <code>solve()</code> .
<code>getLyapIIRanks()</code>	Get the rank values used for the Lyapunov step.
<code>getMonitor()</code>	Get the list of monitor functions.
<code>getOperators()</code>	Get the matrices associated with the eigenvalue problem.
<code>getOptionsPrefix()</code>	Get the prefix used for searching for all EPS options in the database.
<code>getPowerShiftType()</code>	Get the type of shifts used during the power iteration.
<code>getProblemType()</code>	Get the problem type from the EPS object.
<code>getPurify()</code>	Get the flag indicating whether purification is activated or not.
<code>getRG()</code>	Get the region object associated to the eigensolver.
<code>getRQCGReset()</code>	Get the reset parameter used in the RQCG method.
<code>getST()</code>	Get the spectral transformation object associated to the eigensolver.
<code>getStoppingTest()</code>	Get the stopping test function.
<code>getTarget()</code>	Get the value of the target.
<code>getThreshold()</code>	Get the threshold used in the threshold stopping test.
<code>getTolerances()</code>	Get the tolerance and max.
<code>getTrackAll()</code>	Get the flag indicating if all residual norms must be computed or not.
<code>getTrueResidual()</code>	Get the flag indicating if true residual must be computed explicitly.
<code>getTwoSided()</code>	Get the flag indicating if a two-sided variant of the algorithm is being used.
<code>getType()</code>	Get the EPS type of this object.
<code>getWhichEigenpairs()</code>	Get which portion of the spectrum is to be sought.
<code>isGeneralized()</code>	Tell if the EPS object corresponds to a generalized eigenproblem.
<code>isHermitian()</code>	Tell if the EPS object corresponds to a Hermitian eigenproblem.
<code>isPositive()</code>	Eigenproblem requiring a positive (semi-) definite matrix $B$ .
<code>isStructured()</code>	Tell if the EPS object corresponds to a structured eigenvalue problem.
<code>reset()</code>	Reset the EPS object.
<code>setArbitrarySelection(arbitrary[, args, kargs])</code>	Set an arbitrary selection criterion function.
<code>setArnoldiDelayed(delayed)</code>	Set (toggle) delayed reorthogonalization in the Arnoldi iteration.
<code>setBV(bv)</code>	Set a basis vectors object associated to the eigensolver.
<code>setBalance([balance, iterations, cutoff])</code>	Set the balancing technique to be used by the eigensolver.
<code>setCISSExtraction(extraction)</code>	Set the extraction technique used in the CISS solver.
<code>setCISSQuadRule(quad)</code>	Set the quadrature rule used in the CISS solver.
<code>setCISSRefinement([inner, blsize])</code>	Set the values of various refinement parameters in the CISS solver.
<code>setCISSSizes([ip, bs, ms, npart, bsmax, ...])</code>	Set the values of various size parameters in the CISS solver.

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Table 36 – continued from previous page

<code>setCISSThreshold</code> ([delta, spur])	Set the values of various threshold parameters in the CISS solver.
<code>setCISSUseST</code> (usest)	Set a flag indicating that the CISS solver will use the <i>ST</i> object.
<code>setConvergenceTest</code> (conv)	Set how to compute the error estimate used in the convergence test.
<code>setDS</code> (ds)	Set a direct solver object associated to the eigensolver.
<code>setDeflationSpace</code> (space)	Set vectors to form a basis of the deflation space.
<code>setDimensions</code> ([nev, ncv, mpd])	Set number of eigenvalues to compute and the dimension of the subspace.
<code>setEigenvalueComparison</code> (comparison[, args, ...])	Set an eigenvalue comparison function.
<code>setExtraction</code> (extraction)	Set the extraction type used by the eigensolver.
<code>setFromOptions</code> ()	Set EPS options from the options database.
<code>setGDBOrth</code> (borth)	Set the orthogonalization that will be used in the search subspace.
<code>setGDBlockSize</code> (bs)	Set the number of vectors to be added to the searching space.
<code>setGDDoubleExpansion</code> (doubleexp)	Set that the search subspace is expanded with double expansion.
<code>setGDInitialSize</code> (initialsize)	Set the initial size of the searching space.
<code>setGDKrylovStart</code> ([krylovstart])	Set (toggle) starting the search subspace with a Krylov basis.
<code>setGDRestart</code> ([minv, plusk])	Set the number of vectors of the search space after restart.
<code>setInitialSpace</code> (space)	Set the initial space from which the eigensolver starts to iterate.
<code>setInterval</code> (inta, intb)	Set the computational interval for spectrum slicing.
<code>setJDBOrth</code> (borth)	Set the orthogonalization that will be used in the search subspace.
<code>setJDBlockSize</code> (bs)	Set the number of vectors to be added to the searching space.
<code>setJDConstCorrectionTol</code> (constant)	Deactivate the dynamic stopping criterion.
<code>setJDFix</code> (fix)	Set the threshold for changing the target in the correction equation.
<code>setJDInitialSize</code> (initialsize)	Set the initial size of the searching space.
<code>setJDKrylovStart</code> ([krylovstart])	Set (toggle) starting the search subspace with a Krylov basis.
<code>setJDRestart</code> ([minv, plusk])	Set the number of vectors of the search space after restart.
<code>setKrylovSchurBSEType</code> (bse)	Set the Krylov-Schur variant used for BSE structured eigenproblems.
<code>setKrylovSchurDetectZeros</code> (detect)	Set the flag that enforces zero detection in spectrum slicing.
<code>setKrylovSchurDimensions</code> ([nev, ncv, mpd])	Set the dimensions used for each subsolve step (spectrum slicing).
<code>setKrylovSchurLocking</code> (lock)	Set (toggle) locking/non-locking variants of the Krylov-Schur method.
<code>setKrylovSchurPartitions</code> (npart)	Set the number of partitions of the communicator (spectrum slicing).
<code>setKrylovSchurRestart</code> (keep)	Set the restart parameter for the Krylov-Schur method.

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<code>setKrylovSchurSubintervals(subint)</code>	Set the subinterval boundaries.
<code>setLOBPCGBlockSize(bs)</code>	Set the block size of the LOBPCG method.
<code>setLOBPCGLocking(lock)</code>	Toggle between locking and non-locking (LOBPCG method).
<code>setLOBPCGRestart(restart)</code>	Set the restart parameter for the LOBPCG method.
<code>setLanczosReorthogType(reorthog)</code>	Set the type of reorthogonalization used during the Lanczos iteration.
<code>setLeftInitialSpace(space)</code>	Set a left initial space from which the eigensolver starts to iterate.
<code>setLyapIIRanks([rkc, rkl])</code>	Set the ranks used in the solution of the Lyapunov equation.
<code>setMonitor(monitor[, args, kargs])</code>	Append a monitor function to the list of monitors.
<code>setOperators(A[, B])</code>	Set the matrices associated with the eigenvalue problem.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all EPS options in the database.
<code>setPowerShiftType(shift)</code>	Set the type of shifts used during the power iteration.
<code>setProblemType(problem_type)</code>	Set the type of the eigenvalue problem.
<code>setPurify([purify])</code>	Set (toggle) eigenvector purification.
<code>setRG(rg)</code>	Set a region object associated to the eigensolver.
<code>setRQCGReset(nrest)</code>	Set the reset parameter of the RQCG iteration.
<code>setST(st)</code>	Set a spectral transformation object associated to the eigensolver.
<code>setStoppingTest(stopping[, args, kargs])</code>	Set a function to decide when to stop the outer iteration of the eigensolver.
<code>setTarget(target)</code>	Set the value of the target.
<code>setThreshold(thres[, rel])</code>	Set the threshold used in the threshold stopping test.
<code>setTolerances([tol, max_it])</code>	Set the tolerance and max.
<code>setTrackAll(trackall)</code>	Set if the solver must compute the residual of all approximate eigenpairs.
<code>setTrueResidual(trueres)</code>	Set if the solver must compute the true residual explicitly or not.
<code>setTwoSided(twosided)</code>	Set to use a two-sided variant that also computes left eigenvectors.
<code>setType(eps_type)</code>	Set the particular solver to be used in the EPS object.
<code>setUp()</code>	Set up all the internal data structures.
<code>setWhichEigenpairs(which)</code>	Set which portion of the spectrum is to be sought.
<code>solve()</code>	Solve the eigensystem.
<code>updateKrylovSchurSubcommMats([s, a, Au, t, ...])</code>	Update the eigenproblem matrices stored internally in the communicator.
<code>valuesView([viewer])</code>	Display the computed eigenvalues in a viewer.
<code>vectorsView([viewer])</code>	Output computed eigenvectors to a viewer.
<code>view([viewer])</code>	Print the EPS data structure.

### Attributes Summary

<code>bv</code>	The basis vectors ( <i>BV</i> ) object associated.
<code>ds</code>	The direct solver ( <i>DS</i> ) object associated.
<code>extraction</code>	The type of extraction technique to be employed.
<code>max_it</code>	The maximum iteration count.
<code>problem_type</code>	The type of the eigenvalue problem.

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<i>purify</i>	Eigenvector purification.
<i>rg</i>	The region ( <i>RG</i> ) object associated.
<i>st</i>	The spectral transformation ( <i>ST</i> ) object associated.
<i>target</i>	The value of the target.
<i>tol</i>	The tolerance.
<i>track_all</i>	Compute the residual norm of all approximate eigenpairs.
<i>true_residual</i>	Compute the true residual explicitly.
<i>two_sided</i>	Two-sided that also computes left eigenvectors.
<i>which</i>	The portion of the spectrum to be sought.

## Methods Documentation

**appendOptionsPrefix**(*prefix=None*)

Append to the prefix used for searching for all EPS options in the database.

Logically collective.

### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all EPS option requests.

### Return type

*None*

### See also

*setOptionsPrefix*, *getOptionsPrefix*, *EPSAppendOptionsPrefix*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:513 <slepc4py/SLEPc/EPS.pyx#L513>`

**cancelMonitor**()

Clear all monitors for an *EPS* object.

Logically collective.

### See also

*EPSMonitorCancel*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:1860 <slepc4py/SLEPc/EPS.pyx#L1860>`

### Return type

*None*

**computeError**(*i*, *etype=None*)

Compute the error associated with the *i*-th computed eigenpair.

Collective.

Compute the error (based on the residual norm) associated with the *i*-th computed eigenpair.

### Parameters

- **i** (*int*) – Index of the solution to be considered.
- **etype** (*ErrorType* / *None*) – The error type to compute.

**Returns**

The error bound, computed in various ways from the residual norm  $\|Ax - \lambda Bx\|_2$  where  $\lambda$  is the eigenvalue and  $x$  is the eigenvector.

**Return type**

`float`

**Notes**

The index `i` should be a value between 0 and `nconv-1` (see `getConverged()`).

If the computation of left eigenvectors was enabled with `setTwoSided()`, then the error will be computed using the maximum of the value above and the left residual norm  $\|y^*A - \lambda y^*B\|_2$ , where  $y$  is the approximate left eigenvector.

**See also**

`getErrorEstimate`, `setTwoSided`, `EPSComputeError`

**`:sources:`** [Source code at slepc4py/SLEPc/EPS.pyx:2213](#) <slepc4py/SLEPc/EPS.pyx#L2213>

**`create(comm=None)`**

Create the EPS object.

Collective.

**Parameters**

**`comm`** (`Comm` / `None`) – MPI communicator; if not provided, it defaults to all processes.

**Return type**

*Self*

**See also**

`EPSCreate`

**`:sources:`** [Source code at slepc4py/SLEPc/EPS.pyx:396](#) <slepc4py/SLEPc/EPS.pyx#L396>

**`destroy()`**

Destroy the EPS object.

Collective.

**See also**

`EPSTDestroy`

**`:sources:`** [Source code at slepc4py/SLEPc/EPS.pyx:370](#) <slepc4py/SLEPc/EPS.pyx#L370>

**Return type**

*Self*

**`errorView(etype=None, viewer=None)`**

Display the errors associated with the computed solution.

Collective.

Display the errors and the eigenvalues.

#### Parameters

- **etype** (`ErrorType` / `None`) – The error type to compute.
- **viewer** (`petsc4py.PETSc.Viewer` / `None`) – Visualization context; if not provided, the standard output is used.

#### Return type

`None`

#### Notes

By default, this function checks the error of all eigenpairs and prints the eigenvalues if all of them are below the requested tolerance. If the viewer has format `ASCII_INFO_DETAIL` then a table with eigenvalues and corresponding errors is printed.

#### See also

`solve`, `valuesView`, `vectorsView`, `EPSErrorView`

**`:sources:`** [Source code at slepc4py/SLEPc/EPs.pyx:2256 <slepc4py/SLEPc/EPs.pyx#L2256>](#)

#### `getArbitrarySelection()`

Get the arbitrary selection function.

Not collective.

#### Returns

The arbitrary selection function.

#### Return type

`EPsArbitraryFunction`

#### See also

`setArbitrarySelection`

**`:sources:`** [Source code at slepc4py/SLEPc/EPs.pyx:1754 <slepc4py/SLEPc/EPs.pyx#L1754>](#)

#### `getArnoldiDelayed()`

Get the type of reorthogonalization used during the Arnoldi iteration.

Not collective.

#### Returns

True if delayed reorthogonalization is to be used.

#### Return type

`bool`

#### See also

`setArnoldiDelayed`, `EPsArnoldiGetDelayed`

**`:sources:`** [Source code at slepc4py/SLEPc/EPs.pyx:2411 <slepc4py/SLEPc/EPs.pyx#L2411>](#)

### getBV()

Get the basis vectors object associated to the eigensolver.

Not collective.

#### Returns

The basis vectors context.

#### Return type

*BV*

#### See also

*setBV*, *EPSGetBV*

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:1407 <slepc4py/SLEPc/EPS.pyx#L1407>](#)

### getBalance()

Get the balancing type used by the EPS, and the associated parameters.

Not collective.

#### Returns

- **balance** (*Balance*) – The balancing method.
- **iterations** (*int*) – Number of iterations of the balancing algorithm.
- **cutoff** (*float*) – Cutoff value.

#### Return type

*tuple*[*Balance*, *int*, *float*]

#### See also

*setBalance*, *EPSGetBalance*

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:693 <slepc4py/SLEPc/EPS.pyx#L693>](#)

### getCISSExtraction()

Get the extraction technique used in the CISS solver.

Not collective.

#### Returns

The extraction technique.

#### Return type

*CISSExtraction*

#### See also

*setCISSExtraction*, *EPSCISSGetExtraction*

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:3957 <slepc4py/SLEPc/EPS.pyx#L3957>](#)



### **getCISSKSPs()**

Get the array of linear solver objects associated with the CISS solver.

Not collective.

#### **Returns**

The linear solver objects.

#### **Return type**

`list of petsc4py.PETSc.KSP`

### **Notes**

The number of `petsc4py.PETSc.KSP` solvers is equal to the number of integration points divided by the number of partitions. This value is halved in the case of real matrices with a region centered at the real axis.

#### **See also**

`setCISSSizes`, `EPSCISSGetKSPs`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:4238 <slepc4py/SLEPc/EPs.pyx#L4238>`

### **getCISSQuadRule()**

Get the quadrature rule used in the CISS solver.

Not collective.

#### **Returns**

The quadrature rule.

#### **Return type**

`CISSQuadRule`

#### **See also**

`setCISSQuadRule`, `EPSCISSGetQuadRule`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:3994 <slepc4py/SLEPc/EPs.pyx#L3994>`

### **getCISSRefinement()**

Get the values of various refinement parameters in the CISS solver.

Not collective.

#### **Returns**

- **inner** (`int`) – Number of iterative refinement iterations (inner loop).
- **blsize** (`int`) – Number of iterative refinement iterations (blocksize loop).

#### **Return type**

`tuple[int, int]`

#### **See also**

`setCISSRefinement`, `EPSCISSGetRefinement`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:4169 <slepc4py/SLEPc/EPs.pyx#L4169>`

### **getCISSSizes()**

Get the values of various size parameters in the CISS solver.

Not collective.

#### **Returns**

- **ip** (`int`) – Number of integration points.
- **bs** (`int`) – Block size.
- **ms** (`int`) – Moment size.
- **npart** (`int`) – Number of partitions when splitting the communicator.
- **bsmax** (`int`) – Maximum block size.
- **realmats** (`bool`) – True if A and B are real.

#### **Return type**

`tuple[int, int, int, int, int, bool]`

#### **See also**

`setCISSSizes`, `EPSCISSGetSizes`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:4067 <slepc4py/SLEPc/EPs.pyx#L4067>`

### **getCISSThreshold()**

Get the values of various threshold parameters in the CISS solver.

Not collective.

#### **Returns**

- **delta** (`float`) – Threshold for numerical rank.
- **spur** (`float`) – Spurious threshold (to discard spurious eigenpairs).

#### **Return type**

`tuple[float, float]`

#### **See also**

`setCISSThreshold`, `EPSCISSGetThreshold`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:4124 <slepc4py/SLEPc/EPs.pyx#L4124>`

### **getCISSUseST()**

Get the flag indicating the use of the `ST` object in the CISS solver.

Not collective.

#### **Returns**

Whether to use the `ST` object or not.

#### **Return type**

`bool`

**See also**

[setCISSUseST](#), [EPSCISSGetUseST](#)

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:4219](#) <[slepc4py/SLEPc/EPs.pyx#L4219](#)>

**getConverged()**

Get the number of converged eigenpairs.

Not collective.

**Returns**

**nconv** – Number of converged eigenpairs.

**Return type**

`int`

**Notes**

This function should be called after [solve\(\)](#) has finished.

The value `nconv` may be different from the number of requested solutions `nev`, but not larger than `ncv`, see [setDimensions\(\)](#).

**See also**

[setDimensions](#), [solve](#), [getEigenpair](#), [EPSGetConverged](#)

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:1960](#) <[slepc4py/SLEPc/EPs.pyx#L1960](#)>

**getConvergedReason()**

Get the reason why the [solve\(\)](#) iteration was stopped.

Not collective.

**Returns**

Negative value indicates diverged, positive value converged.

**Return type**

[ConvergedReason](#)

**See also**

[setTolerances](#), [solve](#), [EPSGetConvergedReason](#)

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:1941](#) <[slepc4py/SLEPc/EPs.pyx#L1941](#)>

**getConvergenceTest()**

Get how to compute the error estimate used in the convergence test.

Not collective.

**Returns**

The method used to compute the error estimate used in the convergence test.

**Return type**

[Conv](#)

**See also**

[`setConvergenceTest`](#), [`EPSGetConvergenceTest`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:1177 <slepc4py/SLEPc/EPS.pyx#L1177>`

**getDS()**

Get the direct solver associated to the eigensolver.

Not collective.

**Returns**

The direct solver context.

**Return type**

*DS*

**See also**

[`setDS`](#), [`EPSGetDS`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:1444 <slepc4py/SLEPc/EPS.pyx#L1444>`

**getDimensions()**

Get number of eigenvalues to compute and the dimension of the subspace.

Not collective.

**Returns**

- **nev** (*int*) – Number of eigenvalues to compute.
- **ncv** (*int*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int*) – Maximum dimension allowed for the projected problem.

**Return type**

`tuple[int, int, int]`

**See also**

[`setDimensions`](#), [`EPSGetDimensions`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:1290 <slepc4py/SLEPc/EPS.pyx#L1290>`

**getEigenpair(*i*, *Vr*=None, *Vi*=None)**

Get the *i*-th solution of the eigenproblem as computed by [`solve\(\)`](#).

Collective.

The solution consists of both the eigenvalue and the eigenvector.

**Parameters**

- **i** (*int*) – Index of the solution to be obtained.
- **Vr** (*Vec* / *None*) – Placeholder for the returned eigenvector (real part).
- **Vi** (*Vec* / *None*) – Placeholder for the returned eigenvector (imaginary part).

**Returns**

**e** – The computed eigenvalue. It will be a real variable in case of a Hermitian or generalized Hermitian eigenproblem. Otherwise it will be a complex variable (possibly with zero imaginary part).

**Return type**

*Scalar*

**Notes**

The index **i** should be a value between 0 and `nconv-1` (see `getConverged()`). Eigenpairs are indexed according to the ordering criterion established with `setWhichEigenpairs()`.

The 2-norm of the eigenvector is one unless the problem is generalized Hermitian. In this case the eigenvector is normalized with respect to the norm defined by the **B** matrix.

**See also**

`solve`, `getConverged`, `setWhichEigenpairs`, `EPSGetEigenpair`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2090 <slepc4py/SLEPc/EPs.pyx#L2090>`

**getEigenvalue(i)**

Get the *i*-th eigenvalue as computed by `solve()`.

Not collective.

**Parameters**

**i** (*int*) – Index of the solution to be obtained.

**Returns**

The computed eigenvalue. It will be a real variable in case of a Hermitian or generalized Hermitian eigenproblem. Otherwise it will be a complex variable (possibly with zero imaginary part).

**Return type**

*Scalar*

**Notes**

The index **i** should be a value between 0 and `nconv-1` (see `getConverged()`). Eigenpairs are indexed according to the ordering criterion established with `setWhichEigenpairs()`.

**See also**

`getConverged`, `setWhichEigenpairs`, `getEigenpair`, `EPSGetEigenvalue`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1986 <slepc4py/SLEPc/EPs.pyx#L1986>`

**getEigenvalueComparison()**

Get the eigenvalue comparison function.

Not collective.

**Returns**

The eigenvalue comparison function.

**Return type***EPSEigenvalueComparison***See also***setEigenvalueComparison***:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1801 <slepc4py/SLEPc/EPs.pyx#L1801>`**getEigenvector**(*i*, *Vr*=None, *Vi*=None)Get the *i*-th right eigenvector as computed by *solve()*.

Collective.

**Parameters**

- **i** (*int*) – Index of the solution to be obtained.
- **Vr** (*Vec* / *None*) – Placeholder for the returned eigenvector (real part).
- **Vi** (*Vec* / *None*) – Placeholder for the returned eigenvector (imaginary part).

**Return type***None***Notes**

The index *i* should be a value between 0 and *nconv*-1 (see *getConverged()*). Eigenpairs are indexed according to the ordering criterion established with *setWhichEigenpairs()*.

The 2-norm of the eigenvector is one unless the problem is generalized Hermitian. In this case the eigenvector is normalized with respect to the norm defined by the B matrix.

**See also***getConverged*, *setWhichEigenpairs*, *getEigenpair*, *EPsGetEigenvector***:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2024 <slepc4py/SLEPc/EPs.pyx#L2024>`**getErrorEstimate**(*i*)Get the error estimate associated to the *i*-th computed eigenpair.

Not collective.

**Parameters**

- **i** (*int*) – Index of the solution to be considered.

**Returns**

Error estimate.

**Return type***float***Notes**

This is the error estimate used internally by the eigensolver. The actual error bound can be computed with *computeError()*.

**See also**

[\*computeError\*](#), [\*EPSGetErrorEstimate\*](#)

**`:sources: `Source code at slepc4py/SLEPc/EPS.pyx:2184 <slepc4py/SLEPc/EPS.pyx#L2184>``**

**getExtraction()**

Get the extraction type used by the EPS object.

Not collective.

**Returns**

The method of extraction.

**Return type**

[\*Extraction\*](#)

**See also**

[\*setExtraction\*](#), [\*EPSGetExtraction\*](#)

**`:sources: `Source code at slepc4py/SLEPc/EPS.pyx:768 <slepc4py/SLEPc/EPS.pyx#L768>``**

**getGDBOrth()**

Get the orthogonalization used in the search subspace.

Not collective.

Get the orthogonalization used in the search subspace in case of generalized Hermitian problems.

**Returns**

Whether to B-orthogonalize the search subspace.

**Return type**

[\*bool\*](#)

**See also**

[\*setGDBOrth\*](#), [\*EPSGDGetBOrth\*](#)

**`:sources: `Source code at slepc4py/SLEPc/EPS.pyx:3334 <slepc4py/SLEPc/EPS.pyx#L3334>``**

**getGDBlockSize()**

Get the number of vectors to be added to the searching space.

Not collective.

Get the number of vectors to be added to the searching space in every iteration.

**Returns**

The number of vectors added to the search space in every iteration.

**Return type**

[\*int\*](#)

**See also**

[\*setGDBlockSize\*](#), [\*EPSCGGetBlockSize\*](#)

**:sources:** `Source code at slepc4py/SLEPc/EPSC.pyx:3193 <slepc4py/SLEPc/EPSC.pyx#L3193>`

**getGDDoubleExpansion()**

Get a flag indicating whether the double expansion variant is active.

Not collective.

Get a flag indicating whether the double expansion variant has been activated or not.

**Returns**

True if using double expansion.

**Return type**

`bool`

**See also**

[\*setGDDoubleExpansion\*](#), [\*EPSCGGetDoubleExpansion\*](#)

**:sources:** `Source code at slepc4py/SLEPc/EPSC.pyx:3382 <slepc4py/SLEPc/EPSC.pyx#L3382>`

**getGDInitialSize()**

Get the initial size of the searching space.

Not collective.

**Returns**

The number of vectors of the initial searching subspace.

**Return type**

`int`

**See also**

[\*setGDInitialSize\*](#), [\*EPSCGGetInitialSize\*](#)

**:sources:** `Source code at slepc4py/SLEPc/EPSC.pyx:3294 <slepc4py/SLEPc/EPSC.pyx#L3294>`

**getGDKrylovStart()**

Get a flag indicating if the search subspace is started with a Krylov basis.

Not collective.

**Returns**

True if starting the search subspace with a Krylov basis.

**Return type**

`bool`

**See also**

[\*setGDKrylovStart\*](#), [\*EPSCGGetKrylovStart\*](#)



**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3153 <slepc4py/SLEPc/EPS.pyx#L3153>`

### **getGDRestart()**

Get the number of vectors of the search space after restart.

Not collective.

Get the number of vectors of the search space after restart and the number of vectors saved from the previous iteration.

#### **Returns**

- **minv** (`int`) – The number of vectors of the search subspace after restart.
- **plusk** (`int`) – The number of vectors saved from the previous iteration.

#### **Return type**

`tuple[int, int]`

#### **See also**

`setGDRestart`, `EPSPGGetRestart`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3241 <slepc4py/SLEPc/EPS.pyx#L3241>`

### **getInterval()**

Get the computational interval for spectrum slicing.

Not collective.

#### **Returns**

- **inta** (`float`) – The left end of the interval.
- **intb** (`float`) – The right end of the interval.

#### **Return type**

`tuple[float, float]`

#### **Notes**

If the interval was not set by the user, then zeros are returned.

#### **See also**

`setInterval`, `EPSGetInterval`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:972 <slepc4py/SLEPc/EPS.pyx#L972>`

### **getInvariantSubspace()**

Get an orthonormal basis of the computed invariant subspace.

Collective.

#### **Returns**

Basis of the invariant subspace.

#### **Return type**

`list of petsc4py.PETSc.Vec`

## Notes

This function should be called after `solve()` has finished.

The returned vectors span an invariant subspace associated with the computed eigenvalues. An invariant subspace  $X$  of  $A$  satisfies  $Ax \in X$ , for all  $x \in X$  (a similar definition applies for generalized eigenproblems).

### See also

`getEigenpair`, `getConverged`, `solve`, `EPSGetInvariantSubspace`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:2140 <slepc4py/SLEPc/EPS.pyx#L2140>`

## `getIterationNumber()`

Get the current iteration number.

Not collective.

If the call to `solve()` is complete, then it returns the number of iterations carried out by the solution method.

### Returns

Iteration number.

### Return type

`int`

### See also

`getConvergedReason`, `setTolerances`, `EPSGetIterationNumber`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:1919 <slepc4py/SLEPc/EPS.pyx#L1919>`

## `getJDBOrth()`

Get the orthogonalization used in the search subspace.

Not collective.

Get the orthogonalization used in the search subspace in case of generalized Hermitian problems.

### Returns

Whether to B-orthogonalize the search subspace.

### Return type

`bool`

### See also

`setJDBOrth`, `EPSJDGetBOrth`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3693 <slepc4py/SLEPc/EPS.pyx#L3693>`

## `getJDBlockSize()`

Get the number of vectors to be added to the searching space.

Not collective.

Get the number of vectors to be added to the searching space in every iteration.

**Returns**

The number of vectors added to the search space in every iteration.

**Return type**

`int`

**See also**

`setJDBlockSize`, `EPSJDGetBlockSize`

**:sources:** ``Source code at slepc4py/SLEPc/EPS.pyx:3464 <slepc4py/SLEPc/EPS.pyx#L3464>``

**getJDConstCorrectionTol()**

Get the flag indicating if the dynamic stopping is being used.

Not collective.

**Returns**

True if the dynamic stopping criterion is not being used.

**Return type**

`bool`

**See also**

`setJDConstCorrectionTol`, `EPSJDGetConstCorrectionTol`

**:sources:** ``Source code at slepc4py/SLEPc/EPS.pyx:3653 <slepc4py/SLEPc/EPS.pyx#L3653>``

**getJDFix()**

Get the threshold for changing the target in the correction equation.

Not collective.

**Returns**

The threshold for changing the target.

**Return type**

`float`

**See also**

`setJDFix`, `EPSJDGetFix`

**:sources:** ``Source code at slepc4py/SLEPc/EPS.pyx:3608 <slepc4py/SLEPc/EPS.pyx#L3608>``

**getJDInitialSize()**

Get the initial size of the searching space.

Not collective.

**Returns**

The number of vectors of the initial searching subspace.

**Return type**

`int`

**See also**

[`setJDInitialSize`](#), [`EPSJDGetInitialSize`](#)

**:sources:** [`Source code at slepc4py/SLEPc/EPS.pyx:3565 <slepc4py/SLEPc/EPS.pyx#L3565>`](#)

**getJDKrylovStart()**

Get a flag indicating if the search subspace is started with a Krylov basis.

Not collective.

**Returns**

True if starting the search subspace with a Krylov basis.

**Return type**

`bool`

**See also**

[`setJDKrylovStart`](#), [`EPSJDGetKrylovStart`](#)

**:sources:** [`Source code at slepc4py/SLEPc/EPS.pyx:3424 <slepc4py/SLEPc/EPS.pyx#L3424>`](#)

**getJDRestart()**

Get the number of vectors of the search space after restart.

Not collective.

Get the number of vectors of the search space after restart and the number of vectors saved from the previous iteration.

**Returns**

- **minv** (`int`) – The number of vectors of the search subspace after restart.
- **plusk** (`int`) – The number of vectors saved from the previous iteration.

**Return type**

`tuple[int, int]`

**See also**

[`setJDRestart`](#), [`EPSJDGetRestart`](#)

**:sources:** [`Source code at slepc4py/SLEPc/EPS.pyx:3512 <slepc4py/SLEPc/EPS.pyx#L3512>`](#)

**getKrylovSchurBSEType()**

Get the method used for BSE structured eigenproblems (Krylov-Schur).

Not collective.

**Returns**

The BSE method.

**Return type**

`KrylovSchurBSEType`

**See also**

[\*setKrylovSchurBSEType\*](#), [\*EPSKrylovSchurGetBSEType\*](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2499 <slepc4py/SLEPc/EPs.pyx#L2499>`

**getKrylovSchurDetectZeros()**

Get the flag that enforces zero detection in spectrum slicing.

Not collective.

**Returns**

The zero detection flag.

**Return type**

`bool`

**See also**

[\*setKrylovSchurDetectZeros\*](#), [\*EPSKrylovSchurGetDetectZeros\*](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2693 <slepc4py/SLEPc/EPs.pyx#L2693>`

**getKrylovSchurDimensions()**

Get the dimensions used for each subsolve step (spectrum slicing).

Not collective.

**Returns**

- **nev** (`int`) – Number of eigenvalues to compute.
- **ncv** (`int`) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (`int`) – Maximum dimension allowed for the projected problem.

**Return type**

`tuple[int, int, int]`

**See also**

[\*setKrylovSchurDimensions\*](#), [\*EPSKrylovSchurGetDimensions\*](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2752 <slepc4py/SLEPc/EPs.pyx#L2752>`

**getKrylovSchurInertias()**

Get the values of the shifts and their corresponding inertias.

Not collective.

Get the values of the shifts and their corresponding inertias in case of doing spectrum slicing for a computational interval.

**Returns**

- **shifts** (`ArrayReal`) – The values of the shifts used internally in the solver.
- **inertias** (`ArrayInt`) – The values of the inertia in each shift.

**Return type**

tuple[ArrayReal, ArrayInt]

**Notes**

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with *setInterval()* and *SINVERT* is set.

If called after *solve()*, all shifts used internally by the solver are returned (including both endpoints and any intermediate ones). If called before *solve()* and after *setUp()* then only the information of the endpoints of subintervals is available.

**See also**

*setInterval*, *setKrylovSchurSubintervals*, *EPSKrylovSchurGetInertias*

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:3050 <slepc4py/SLEPc/EPs.pyx#L3050>`

**getKrylovSchurKSP()**

Get the linear solver object associated with the internal *EPS* object.

Collective.

Get the linear solver object associated with the internal *EPS* object in case of doing spectrum slicing for a computational interval.

**Returns**

The linear solver object.

**Return type**

`petsc4py.PETSc.KSP`

**Notes**

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with *setInterval()* and *SINVERT* is set.

When invoked to compute all eigenvalues in an interval with spectrum slicing, *KRYLOV SCHUR* creates another *EPS* object internally that is used to compute eigenvalues by chunks near selected shifts. This function allows access to the KSP object associated to this internal *EPS* object.

In case of having more than one partition, the returned KSP will be different in MPI processes belonging to different partitions. Hence, if required, *setKrylovSchurPartitions()* must be called BEFORE this function.

**See also**

*setInterval*, *setKrylovSchurPartitions*, *EPSKrylovSchurGetKSP*

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:3094 <slepc4py/SLEPc/EPs.pyx#L3094>`

**getKrylovSchurLocking()**

Get the locking flag used in the Krylov-Schur method.

Not collective.

**Returns**

The locking flag.

**Return type**

`bool`

**See also**

`setKrylovSchurLocking`, `EPKrylovSchurGetLocking`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:2586 <slepc4py/SLEPc/EPS.pyx#L2586>`

**getKrylovSchurPartitions()**

Get the number of partitions of the communicator (spectrum slicing).

Not collective.

**Returns**

The number of partitions.

**Return type**

`int`

**See also**

`setKrylovSchurPartitions`, `EPKrylovSchurGetPartitions`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:2640 <slepc4py/SLEPc/EPS.pyx#L2640>`

**getKrylovSchurRestart()**

Get the restart parameter used in the Krylov-Schur method.

Not collective.

**Returns**

The number of vectors to be kept at restart.

**Return type**

`float`

**See also**

`setKrylovSchurRestart`, `EPKrylovSchurGetRestart`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:2542 <slepc4py/SLEPc/EPS.pyx#L2542>`

**getKrylovSchurSubcommInfo()**

Get information related to the case of doing spectrum slicing.

Collective on the subcommunicator.

Get information related to the case of doing spectrum slicing for a computational interval with multiple communicators.

**Returns**

- **k** (`int`) – Index of the subinterval for the calling process.
- **n** (`int`) – Number of eigenvalues found in the k-th subinterval.

- `v` (`petsc4py.PETSc.Vec`) – A vector owned by processes in the subcommunicator with dimensions compatible for locally computed eigenvectors.

#### Return type

`tuple[int, int, Vec]`

#### Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with `setInterval()` and `SINVERT` is set.

#### See also

`getKrylovSchurSubcommPairs`, `EPKrylovSchurGetSubcommInfo`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2777 <slepc4py/SLEPc/EPs.pyx#L2777>`

#### `getKrylovSchurSubcommMats()`

Get the eigenproblem matrices stored in the subcommunicator.

Collective on the subcommunicator.

Get the eigenproblem matrices stored internally in the subcommunicator to which the calling process belongs.

#### Returns

- `A` (`petsc4py.PETSc.Mat`) – The matrix associated with the eigensystem.
- `B` (`petsc4py.PETSc.Mat`) – The second matrix in the case of generalized eigenproblems.

#### Return type

`tuple[Mat, Mat] | tuple[Mat, None]`

#### Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with `setInterval()` and `SINVERT` is set. And is relevant only when the number of partitions (`setKrylovSchurPartitions()`) is larger than one.

This is the analog of `getOperators()`, but returns the matrices distributed differently (in the subcommunicator rather than in the parent communicator).

These matrices should not be modified by the user.

#### See also

`setInterval`, `setKrylovSchurPartitions`, `EPKrylovSchurGetSubcommMats`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2855 <slepc4py/SLEPc/EPs.pyx#L2855>`

#### `getKrylovSchurSubcommPairs(i, v=None)`

Get the *i*-th eigenpair stored in the multi-communicator of the process.

Collective on the subcommunicator (if *v* is given).

Get the *i*-th eigenpair stored internally in the multi-communicator to which the calling process belongs.

#### Parameters



- **i** (*int*) – Index of the solution to be obtained.
- **v** (*Vec* | *None*) – Placeholder for the returned eigenvector.

#### Returns

The computed eigenvalue.

#### Return type

*Scalar*

#### Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with *setInterval()* and *SINVERT* is set. And is relevant only when the number of partitions (*setKrylovSchurPartitions()*) is larger than one.

Argument *v* must be a valid *Vec* object, created by calling *getKrylovSchurSubcommInfo()*.

The index *i* should be a value between 0 and *n*-1, where *n* is the number of vectors in the local subinterval, see *getKrylovSchurSubcommInfo()*.

#### See also

*getKrylovSchurSubcommMats*, *EPKrylovSchurGetSubcommPairs*

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2811 <slepc4py/SLEPc/EPs.pyx#L2811>`

#### **getKrylovSchurSubintervals()**

Get the points that delimit the subintervals.

Not collective.

Get the points that delimit the subintervals used in spectrum slicing with several partitions.

#### Returns

Real values specifying subintervals.

#### Return type

*ArrayReal*

#### Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with *setInterval()* and *SINVERT* is set.

If the user passed values with *setKrylovSchurSubintervals()*, then the same values are returned here. Otherwise, the values computed internally are obtained.

#### See also

*setKrylovSchurSubintervals*, *EPKrylovSchurGetSubintervals*

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:3012 <slepc4py/SLEPc/EPs.pyx#L3012>`

#### **getLOBPCGBlockSize()**

Get the block size used in the LOBPCG method.

Not collective.

**Returns**

The block size.

**Return type**

`int`

**See also**

`setLOBPCGBlockSize`, `EPSLOBPCGGetBlockSize`

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:3777](#) <slepc4py/SLEPc/EPS.pyx#L3777>

**getLOBPCGLocking()**

Get the locking flag used in the LOBPCG method.

Not collective.

**Returns**

The locking flag.

**Return type**

`bool`

**See also**

`setLOBPCGLocking`, `EPSLOBPCGGetLocking`

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:3864](#) <slepc4py/SLEPc/EPS.pyx#L3864>

**getLOBPCGRestart()**

Get the restart parameter used in the LOBPCG method.

Not collective.

**Returns**

The restart parameter.

**Return type**

`float`

**See also**

`setLOBPCGRestart`, `EPSLOBPCGGetRestart`

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:3821](#) <slepc4py/SLEPc/EPS.pyx#L3821>

**getLanczosReorthogType()**

Get the type of reorthogonalization used during the Lanczos iteration.

Not collective.

**Returns**

The type of reorthogonalization.

**Return type**

`LanczosReorthogType`

#### See also

[setLanczosReorthogType](#), [EPSLanczosGetReorthog](#)

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:2453 <slepc4py/SLEPc/EPS.pyx#L2453>`

**getLeftEigenvector**(*i*, *Wr*=None, *Wi*=None)

Get the *i*-th left eigenvector as computed by [solve\(\)](#).

Collective.

#### Parameters

- **i** ([int](#)) – Index of the solution to be obtained.
- **Wr** ([Vec](#) / [None](#)) – Placeholder for the returned left eigenvector (real part).
- **Wi** ([Vec](#) / [None](#)) – Placeholder for the returned left eigenvector (imaginary part).

#### Return type

[None](#)

#### Notes

The index *i* should be a value between 0 and `nconv-1` (see [getConverged\(\)](#)). Eigensolutions are indexed according to the ordering criterion established with [setWhichEigenpairs\(\)](#).

Left eigenvectors are available only if the twosided flag was set with [setTwoSided\(\)](#).

#### See also

[getConverged](#), [setWhichEigenpairs](#), [getEigenpair](#), [EPSGetLeftEigenvector](#)

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:2058 <slepc4py/SLEPc/EPS.pyx#L2058>`

**getLyapIIRanks()**

Get the rank values used for the Lyapunov step.

Not collective.

#### Returns

- **rkc** ([int](#)) – The compressed rank.
- **rkl** ([int](#)) – The Lyapunov rank.

#### Return type

[tuple](#)[[int](#), [int](#)]

#### See also

[setLyapIIRanks](#), [EPSLyapIIGetRanks](#)

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3915 <slepc4py/SLEPc/EPS.pyx#L3915>`

**getMonitor()**

Get the list of monitor functions.

Not collective.

**Returns**

The list of monitor functions.

**Return type**

*EPSPMonitorFunction*

**See also**

*setMonitor*

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1843 <slepc4py/SLEPc/EPs.pyx#L1843>`

**getOperators()**

Get the matrices associated with the eigenvalue problem.

Collective.

**Returns**

- **A** (`petsc4py.PETSc.Mat`) – The matrix associated with the eigensystem.
- **B** (`petsc4py.PETSc.Mat`) – The second matrix in the case of generalized eigenproblems.

**Return type**

`tuple[Mat, Mat] | tuple[Mat, None]`

**See also**

*setOperators*, *EPsGetOperators*

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1518 <slepc4py/SLEPc/EPs.pyx#L1518>`

**getOptionsPrefix()**

Get the prefix used for searching for all EPS options in the database.

Not collective.

**Returns**

The prefix string set for this EPS object.

**Return type**

`str`

**See also**

*setOptionsPrefix*, *appendOptionsPrefix*, *EPsGetOptionsPrefix*

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:463 <slepc4py/SLEPc/EPs.pyx#L463>`

**getPowerShiftType()**

Get the type of shifts used during the power iteration.

Not collective.

**Returns**

The type of shift.

**Return type**  
*PowerShiftType*

**See also**

*setPowerShiftType*, *EPSPowerGetShiftType*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:2364 <slepc4py/SLEPc/EPS.pyx#L2364>`

**getProblemType()**

Get the problem type from the EPS object.

Not collective.

**Returns**  
The problem type that was previously set.

**Return type**  
*ProblemType*

**See also**

*setProblemType*, *EPSGetProblemType*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:553 <slepc4py/SLEPc/EPS.pyx#L553>`

**getPurify()**

Get the flag indicating whether purification is activated or not.

Not collective.

**Returns**  
Whether purification is activated or not.

**Return type**  
*bool*

**See also**

*setPurify*, *EPSGetPurify*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:1132 <slepc4py/SLEPc/EPS.pyx#L1132>`

**getRG()**

Get the region object associated to the eigensolver.

Not collective.

**Returns**  
The region context.

**Return type**  
*RG*

**See also**

`setRG`, `EPSGetRG`

**:sources:** ``Source code at slepc4py/SLEPc/EPs.pyx:1481 <slepc4py/SLEPc/EPs.pyx#L1481>``

**getRQCGReset()**

Get the reset parameter used in the RQCG method.

Not collective.

**Returns**

The number of iterations between resets.

**Return type**

`int`

**See also**

`setRQCGReset`, `EPsRQCGGetReset`

**:sources:** ``Source code at slepc4py/SLEPc/EPs.pyx:3740 <slepc4py/SLEPc/EPs.pyx#L3740>``

**getST()**

Get the spectral transformation object associated to the eigensolver.

Not collective.

**Returns**

The spectral transformation.

**Return type**

`ST`

**See also**

`setST`, `EPSGetST`

**:sources:** ``Source code at slepc4py/SLEPc/EPs.pyx:1370 <slepc4py/SLEPc/EPs.pyx#L1370>``

**getStoppingTest()**

Get the stopping test function.

Not collective.

**Returns**

The stopping test function.

**Return type**

`EPsStoppingFunction`

**See also**

`setStoppingTest`

**:sources:** ``Source code at slepc4py/SLEPc/EPs.pyx:1708 <slepc4py/SLEPc/EPs.pyx#L1708>``

### **getTarget()**

Get the value of the target.

Not collective.

#### **Returns**

The value of the target.

#### **Return type**

*Scalar*

### **Notes**

If the target was not set by the user, then zero is returned.

#### **See also**

*setTarget*, *EPSGetTarget*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:922 <slepc4py/SLEPc/EPS.pyx#L922>`

### **getThreshold()**

Get the threshold used in the threshold stopping test.

Not collective.

#### **Returns**

- **thres** (*float*) – The threshold.
- **rel** (*bool*) – Whether the threshold is relative or not.

#### **Return type**

*tuple*[*float*, *bool*]

#### **See also**

*setThreshold*, *EPSGetThreshold*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:871 <slepc4py/SLEPc/EPS.pyx#L871>`

### **getTolerances()**

Get the tolerance and max. iter. count used for convergence tests.

Not collective.

Get the tolerance and iteration limit used by the default EPS convergence tests.

#### **Returns**

- **tol** (*float*) – The convergence tolerance.
- **max\_it** (*int*) – The maximum number of iterations.

#### **Return type**

*tuple*[*float*, *int*]

**See also**

[\*setTolerances\*](#), [\*EPSGetTolerances\*](#)

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:1032 <slepc4py/SLEPc/EPs.pyx#L1032>](#)

**getTrackAll()**

Get the flag indicating if all residual norms must be computed or not.

Not collective.

**Returns**

Whether the solver computes all residuals or not.

**Return type**

`bool`

**See also**

[\*setTrackAll\*](#), [\*EPSGetTrackAll\*](#)

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:1253 <slepc4py/SLEPc/EPs.pyx#L1253>](#)

**getTrueResidual()**

Get the flag indicating if true residual must be computed explicitly.

Not collective.

**Returns**

Whether the solver computes true residuals or not.

**Return type**

`bool`

**See also**

[\*setTrueResidual\*](#), [\*EPSGetTrueResidual\*](#)

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:1216 <slepc4py/SLEPc/EPs.pyx#L1216>](#)

**getTwoSided()**

Get the flag indicating if a two-sided variant of the algorithm is being used.

Not collective.

**Returns**

Whether the two-sided variant is to be used or not.

**Return type**

`bool`

**See also**

[\*setTwoSided\*](#), [\*EPSGetTwoSided\*](#)



**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:1085 <slepc4py/SLEPc/EPS.pyx#L1085>`

#### **getType()**

Get the EPS type of this object.

Not collective.

##### **Returns**

The solver currently being used.

##### **Return type**

`str`

##### **See also**

`setType`, `EPSGetType`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:444 <slepc4py/SLEPc/EPS.pyx#L444>`

#### **getWhichEigenpairs()**

Get which portion of the spectrum is to be sought.

Not collective.

##### **Returns**

The portion of the spectrum to be sought by the solver.

##### **Return type**

`Which`

##### **See also**

`setWhichEigenpairs`, `EPSGetWhichEigenpairs`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:815 <slepc4py/SLEPc/EPS.pyx#L815>`

#### **isGeneralized()**

Tell if the EPS object corresponds to a generalized eigenproblem.

Not collective.

##### **Returns**

True if the problem is generalized.

##### **Return type**

`bool`

##### **See also**

`isHermitian`, `isPositive`, `isStructured`, `EPSIsGeneralized`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:608 <slepc4py/SLEPc/EPS.pyx#L608>`

#### **isHermitian()**

Tell if the EPS object corresponds to a Hermitian eigenproblem.

Not collective.

**Returns**

True if the problem is Hermitian.

**Return type**

bool

**See also**

*isGeneralized*, *isPositive*, *isStructured*, *EPSIsHermitian*

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:627](#) <slepc4py/SLEPc/EPS.pyx#L627>

**isPositive()**

Eigenproblem requiring a positive (semi-) definite matrix  $B$ .

Not collective.

Tell if the EPS corresponds to an eigenproblem requiring a positive (semi-) definite matrix  $B$ .

**Returns**

True if the problem is positive (semi-) definite.

**Return type**

bool

**See also**

*isGeneralized*, *isHermitian*, *isStructured*, *EPSIsPositive*

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:646](#) <slepc4py/SLEPc/EPS.pyx#L646>

**isStructured()**

Tell if the EPS object corresponds to a structured eigenvalue problem.

Not collective.

**Returns**

True if the problem is structured.

**Return type**

bool

**Notes**

The result will be True if the problem type has been set to some structured type such as [BSE](#). This is independent of whether the input matrix has been built with a certain structure with a helper function.

**See also**

*isGeneralized*, *isHermitian*, *isPositive*, *EPSIsStructured*

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:668](#) <slepc4py/SLEPc/EPS.pyx#L668>

**reset()**

Reset the EPS object.

Collective.

#### See also

[EPSReset](#)

**`:sources:`** `Source code at slepc4py/SLEPc/EPS.pyx:384 <slepc4py/SLEPc/EPS.pyx#L384>`

#### Return type

`None`

**`setArbitrarySelection`**(*arbitrary*, *args*=`None`, *kargs*=`None`)

Set an arbitrary selection criterion function.

Logically collective.

Set a function to look for eigenvalues according to an arbitrary selection criterion. This criterion can be based on a computation involving the current eigenvector approximation.

#### See also

[getArbitrarySelection](#), [EPSSetArbitrarySelection](#)

**`:sources:`** `Source code at slepc4py/SLEPc/EPS.pyx:1725 <slepc4py/SLEPc/EPS.pyx#L1725>`

#### Parameters

- **`arbitrary`** ([EPSSetArbitraryFunction](#) | `None`)
- **`args`** (`tuple`[`Any`, ...] | `None`)
- **`kargs`** (`dict`[`str`, `Any`] | `None`)

#### Return type

`None`

**`setArnoldiDelayed`**(*delayed*)

Set (toggle) delayed reorthogonalization in the Arnoldi iteration.

Logically collective.

#### Parameters

**`delayed`** (`bool`) – True if delayed reorthogonalization is to be used.

#### Return type

`None`

#### Notes

This call is only relevant if the type was set to [EPS.Type.ARNOLDI](#) with [setType\(\)](#).

Delayed reorthogonalization is an aggressive optimization for the Arnoldi eigensolver than may provide better scalability, but sometimes makes the solver converge more slowly compared to the default algorithm.

#### See also

[getArnoldiDelayed](#), [EPSArnoldiSetDelayed](#)

**`:sources:`** `Source code at slepc4py/SLEPc/EPS.pyx:2383 <slepc4py/SLEPc/EPS.pyx#L2383>`

**setBV**(*bv*)

Set a basis vectors object associated to the eigensolver.

Collective.

**Parameters**

**bv** (*BV*) – The basis vectors context.

**Return type**

*None*

**See also**

*getBV*, *EPSSetBV*

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:1427 <slepc4py/SLEPc/EPs.pyx#L1427>](#)

**setBalance**(*balance=None*, *iterations=None*, *cutoff=None*)

Set the balancing technique to be used by the eigensolver.

Logically collective.

**Parameters**

- **balance** (*Balance* / *None*) – The balancing method.
- **iterations** (*int* / *None*) – Number of iterations of the balancing algorithm.
- **cutoff** (*float* / *None*) – Cutoff value.

**Return type**

*None*

**Notes**

When balancing is enabled, the solver works implicitly with matrix  $DAD^{-1}$ , where  $D$  is an appropriate diagonal matrix. This improves the accuracy of the computed results in some cases.

Balancing makes sense only for non-Hermitian problems when the required precision is high (i.e., with a small tolerance).

By default, balancing is disabled. The two-sided method is much more effective than the one-sided counterpart, but it requires the system matrices to have the `Mat.multTranspose()` operation defined.

The parameter `iterations` is the number of iterations performed by the method. The `cutoff` value is used only in the two-side variant.

**See also**

*setBalance*, *EPSGetBalance*

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:718 <slepc4py/SLEPc/EPs.pyx#L718>](#)

**setCISSExtraction**(*extraction*)

Set the extraction technique used in the CISS solver.

Logically collective.

**Parameters**

**extraction** (*CISSExtraction*) – The extraction technique.

**Return type**

None

**See also**[getCISSExtraction](#), [EPSCISSSetExtraction](#)**:sources:**``Source code at slepc4py/SLEPc/EPs.pyx:3939 <slepc4py/SLEPc/EPs.pyx#L3939>``**setCISSQuadRule**(*quad*)

Set the quadrature rule used in the CISS solver.

Logically collective.

**Parameters****quad** ([CISSQuadRule](#)) – The quadrature rule.**Return type**

None

**See also**[getCISSQuadRule](#), [EPSCISSSetQuadRule](#)**:sources:**``Source code at slepc4py/SLEPc/EPs.pyx:3976 <slepc4py/SLEPc/EPs.pyx#L3976>``**setCISSRefinement**(*inner=None*, *blsize=None*)

Set the values of various refinement parameters in the CISS solver.

Logically collective.

**Parameters**

- **inner** (*int* / *None*) – Number of iterative refinement iterations (inner loop).
- **blsize** (*int* / *None*) – Number of iterative refinement iterations (blocksize loop).

**Return type**

None

**See also**[getCISSRefinement](#), [EPSCISSSetRefinement](#)**:sources:**``Source code at slepc4py/SLEPc/EPs.pyx:4146 <slepc4py/SLEPc/EPs.pyx#L4146>``**setCISSSizes**(*ip=None*, *bs=None*, *ms=None*, *npart=None*, *bsmax=None*, *realmats=False*)

Set the values of various size parameters in the CISS solver.

Logically collective.

**Parameters**

- **ip** (*int* / *None*) – Number of integration points.
- **bs** (*int* / *None*) – Block size.
- **ms** (*int* / *None*) – Moment size.

- **npart** (*int* / *None*) – Number of partitions when splitting the communicator.
- **bsmax** (*int* / *None*) – Maximum block size.
- **realmats** (*bool*) – True if A and B are real.

**Return type**

*None*

**Notes**

The default number of partitions is 1. This means the internal `petsc4py.PETSc.KSP` object is shared among all processes of the *EPS* communicator. Otherwise, the communicator is split into `npart` communicators, so that `npart` `petsc4py.PETSc.KSP` solves proceed simultaneously.

**See also**

`getCISSSizes`, `setCISSThreshold`, `setCISSRefinement`, `EPSCISSSetSizes`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4013 <slepc4py/SLEPc/EPS.pyx#L4013>`

**setCISSThreshold**(*delta=None*, *spur=None*)

Set the values of various threshold parameters in the CISS solver.

Logically collective.

**Parameters**

- **delta** (*float* / *None*) – Threshold for numerical rank.
- **spur** (*float* / *None*) – Spurious threshold (to discard spurious eigenpairs).

**Return type**

*None*

**See also**

`getCISSThreshold`, `EPSCISSSetThreshold`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4101 <slepc4py/SLEPc/EPS.pyx#L4101>`

**setCISSUseST**(*usest*)

Set a flag indicating that the CISS solver will use the *ST* object.

Logically collective.

**Parameters**

**usest** (*bool*) – Whether to use the *ST* object or not.

**Return type**

*None*

**Notes**

When this option is set, the linear solves can be configured by setting options for the `petsc4py.PETSc.KSP` object obtained with `ST.getKSP()`. Otherwise, several `petsc4py.PETSc.KSP` objects are created, which can be accessed with `getCISSKSPs()`.

The default is to use the *ST*, unless several partitions have been specified, see `setCISSSizes()`.

**See also**

[`getCISSTUseST`](#), [`getCISSTKSPs`](#), [`setCISSTSizes`](#), [`EPSCISSTSetUseST`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:4191 <slepc4py/SLEPc/EPs.pyx#L4191>`

**setConvergenceTest**(*conv*)

Set how to compute the error estimate used in the convergence test.

Logically collective.

**Parameters**

**conv** (*Conv*) – The method used to compute the error estimate used in the convergence test.

**Return type**

None

**See also**

[`getConvergenceTest`](#), [`EPSetConvergenceTest`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1197 <slepc4py/SLEPc/EPs.pyx#L1197>`

**setDS**(*ds*)

Set a direct solver object associated to the eigensolver.

Collective.

**Parameters**

**ds** (*DS*) – The direct solver context.

**Return type**

None

**See also**

[`getDS`](#), [`EPSetDS`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1464 <slepc4py/SLEPc/EPs.pyx#L1464>`

**setDeflationSpace**(*space*)

Set vectors to form a basis of the deflation space.

Collective.

**Parameters**

**space** (*Vec* | *list*[*Vec*]) – Set of basis vectors of the deflation space.

**Return type**

None

**Notes**

When a deflation space is given, the eigensolver seeks the eigensolution in the restriction of the problem to the orthogonal complement of this space. This can be used for instance in the case that an invariant subspace is known beforehand (such as the nullspace of the matrix).

These vectors do not persist from one `solve()` call to the other, so the deflation space should be set every time.

The vectors do not need to be mutually orthonormal, since they are explicitly orthonormalized internally.

#### See also

`setInitialSpace`, `EPSSetDeflationSpace`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1576 <slepc4py/SLEPc/EPs.pyx#L1576>`

**setDimensions**(*nev=None, ncv=None, mpd=None*)

Set number of eigenvalues to compute and the dimension of the subspace.

Logically collective.

#### Parameters

- **nev** (*int* / *None*) – Number of eigenvalues to compute.
- **ncv** (*int* / *None*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int* / *None*) – Maximum dimension allowed for the projected problem.

#### Return type

*None*

#### Notes

Use *DETERMINE* for *ncv* and *mpd* to assign a reasonably good value, which is dependent on the solution method.

The parameters *ncv* and *mpd* are intimately related, so that the user is advised to set one of them at most. Normal usage is the following:

- In cases where *nev* is small, the user sets *ncv* (a reasonable default is  $2 * nev$ ).
- In cases where *nev* is large, the user sets *mpd*.

The value of *ncv* should always be between *nev* and (*nev* + *mpd*), typically  $ncv = nev + mpd$ . If *nev* is not too large,  $mpd = nev$  is a reasonable choice, otherwise a smaller value should be used.

When computing all eigenvalues in an interval, see `setInterval()`, these parameters lose relevance, and tuning must be done with `setKrylovSchurDimensions()`.

#### See also

`getDimensions`, `setKrylovSchurDimensions`, `EPSSetDimensions`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1315 <slepc4py/SLEPc/EPs.pyx#L1315>`

**setEigenvalueComparison**(*comparison, args=None, kargs=None*)

Set an eigenvalue comparison function.

Logically collective.



## Notes

This eigenvalue comparison function is used when `setWhichEigenpairs()` is set to `EPS.Which.USER`.

### See also

`getEigenvalueComparison`, `EPSSetEigenvalueComparison`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1771 <slepc4py/SLEPc/EPs.pyx#L1771>`

### Parameters

- **comparison** (`EPSEigenvalueComparison` | `None`)
- **args** (`tuple[Any, ...]` | `None`)
- **kargs** (`dict[str, Any]` | `None`)

### Return type

`None`

## **setExtraction(extraction)**

Set the extraction type used by the eigensolver.

Logically collective.

### Parameters

**extraction** (`Extraction`) – The extraction method to be used by the solver.

### Return type

`None`

## Notes

Not all eigensolvers support all types of extraction.

By default, a standard Rayleigh-Ritz extraction is used. Other extractions may be useful when computing interior eigenvalues.

Harmonic-type extractions are used in combination with a *target*, see `setTarget()`.

### See also

`getExtraction`, `setTarget`, `EPSSetExtraction`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:787 <slepc4py/SLEPc/EPs.pyx#L787>`

## **setFromOptions()**

Set EPS options from the options database.

Collective.

## Notes

To see all options, run your program with the `-help` option.

This routine must be called before `setUp()` if the user is to be allowed to set the solver type.

**See also**

[setOptionsPrefix](#), [EPSSetFromOptions](#)

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:532](#) <[slepc4py/SLEPc/EPS.pyx#L532](#)>

**Return type**

None

**setGDBOrth**(*borth*)

Set the orthogonalization that will be used in the search subspace.

Logically collective.

Set the orthogonalization that will be used in the search subspace in case of generalized Hermitian problems.

**Parameters**

**borth** (*bool*) – Whether to B-orthogonalize the search subspace.

**Return type**

None

**See also**

[getGDBOrth](#), [EPSCGSetBOrth](#)

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:3313](#) <[slepc4py/SLEPc/EPS.pyx#L3313](#)>

**setGDBBlockSize**(*bs*)

Set the number of vectors to be added to the searching space.

Logically collective.

Set the number of vectors to be added to the searching space in every iteration.

**Parameters**

**bs** (*int*) – The number of vectors added to the search space in every iteration.

**Return type**

None

**See also**

[getGDBBlockSize](#), [EPSCGSetBlockSize](#)

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:3172](#) <[slepc4py/SLEPc/EPS.pyx#L3172](#)>

**setGDDoubleExpansion**(*doubleexp*)

Set that the search subspace is expanded with double expansion.

Logically collective.

**Parameters**

**doubleexp** (*bool*) – True if using double expansion.

**Return type**

None

## Notes

In the double expansion variant the search subspace is expanded with  $K[Ax, Bx]$  (double expansion) instead of the classic  $Kr$ , where  $K$  is the preconditioner,  $x$  the selected approximate eigenvector and  $r$  its associated residual vector.

### See also

[`getGDDoubleExpansion`](#), [`EPSCGSetDoubleExpansion`](#)

**:sources:** [Source code at slepc4py/SLEPc/EPSCG.pyx:3356 <slepc4py/SLEPc/EPSCG.pyx#L3356>](#)

**setGDInitialSize**(*initialsize*)

Set the initial size of the searching space.

Logically collective.

### Parameters

**initialsize** (*int*) – The number of vectors of the initial searching subspace.

### Return type

None

## Notes

If the flag in [`setGDKrylovStart\(\)`](#) is set to `False` and the user provides vectors with [`setInitialSpace\(\)`](#), up to *initialsize* vectors will be used; and if the provided vectors are not enough, the solver completes the subspace with random vectors. In case the [`setGDKrylovStart\(\)`](#) flag is `True`, the solver gets the first vector provided by the user or, if not available, a random vector, and expands the Krylov basis up to *initialsize* vectors.

### See also

[`setGDKrylovStart`](#), [`getGDInitialSize`](#), [`EPSCGSetInitialSize`](#)

**:sources:** [Source code at slepc4py/SLEPc/EPSCG.pyx:3266 <slepc4py/SLEPc/EPSCG.pyx#L3266>](#)

**setGDKrylovStart**(*krylovstart=True*)

Set (toggle) starting the search subspace with a Krylov basis.

Logically collective.

### Parameters

**krylovstart** (*bool*) – True if starting the search subspace with a Krylov basis.

### Return type

None

### See also

[`setGDInitialSize`](#), [`setGDKrylovStart`](#), [`EPSCGSetKrylovStart`](#)

**:sources:** [Source code at slepc4py/SLEPc/EPSCG.pyx:3135 <slepc4py/SLEPc/EPSCG.pyx#L3135>](#)

**setGDRestart**(*minv=None, plusk=None*)

Set the number of vectors of the search space after restart.

Logically collective.

Set the number of vectors of the search space after restart and the number of vectors saved from the previous iteration.

**Parameters**

- **minv** (*int*) – The number of vectors of the search subspace after restart.
- **plusk** (*int*) – The number of vectors saved from the previous iteration.

**Return type**

*None*

**See also**

*getGDRestart*, *EPSetRestart*

**:sources:** [Source code at slepc4py/SLEPc/EP.pyx:3215 <slepc4py/SLEPc/EP.pyx#L3215>](#)

**setInitialSpace**(*space*)

Set the initial space from which the eigensolver starts to iterate.

Collective.

**Parameters**

**space** (*Vec* | *list[Vec]*) – Set of basis vectors of the initial space.

**Return type**

*None*

**Notes**

Some solvers start to iterate on a single vector (initial vector). In that case, only the first vector is taken into account and the other vectors are ignored. But other solvers such as *SUBSPACE* are able to make use of the whole initial subspace as an initial guess.

These vectors do not persist from one *solve()* call to the other, so the initial space should be set every time.

The vectors do not need to be mutually orthonormal, since they are explicitly orthonormalized internally.

Common usage of this function is when the user can provide a rough approximation of the wanted eigenspace. Then, convergence may be faster.

**See also**

*setDeflationSpace*, *setLeftInitialSpace*, *EPSetInitialSpace*

**:sources:** [Source code at slepc4py/SLEPc/EP.pyx:1614 <slepc4py/SLEPc/EP.pyx#L1614>](#)

**setInterval**(*inta, intb*)

Set the computational interval for spectrum slicing.

Logically collective.

**Parameters**

- **inta** (*float*) – The left end of the interval.
- **intb** (*float*) – The right end of the interval.

**Return type**

None

**Notes**

Spectrum slicing is a technique employed for computing all eigenvalues of symmetric eigenproblems in a given interval. This function provides the interval to be considered. It must be used in combination with *EPS.Which.ALL*, see *setWhichEigenpairs()*.

A computational interval is also needed when using polynomial filters, see *STFILTER*.

**See also**

*getInterval*, *setWhichEigenpairs*, *EPSSetInterval*, *STFILTER*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:998 <slepc4py/SLEPc/EPS.pyx#L998>`

**setJDBOrth**(*borth*)

Set the orthogonalization that will be used in the search subspace.

Logically collective.

Set the orthogonalization that will be used in the search subspace in case of generalized Hermitian problems.

**Parameters**

**borth** (*bool*) – Whether to B-orthogonalize the search subspace.

**Return type**

None

**See also**

*getJDBOrth*, *EPSJDSetBOrth*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3672 <slepc4py/SLEPc/EPS.pyx#L3672>`

**setJDBlockSize**(*bs*)

Set the number of vectors to be added to the searching space.

Logically collective.

Set the number of vectors to be added to the searching space in every iteration.

**Parameters**

**bs** (*int*) – The number of vectors added to the search space in every iteration.

**Return type**

None

**See also**

*getJDBlockSize*, *EPSJDSetBlockSize*

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3443 <slepc4py/SLEPc/EPS.pyx#L3443>`

**setJDConstCorrectionTol**(*constant*)

Deactivate the dynamic stopping criterion.

Logically collective.

**Parameters**

**constant** (*bool*) – If False, the `petsc4py.PETSc.KSP` relative tolerance is set to  $0.5^{**i}$ .

**Return type**

None

**Notes**

If this flag is set to False, then the `petsc4py.PETSc.KSP` relative tolerance is dynamically set to  $0.5^{**i}$ , where *i* is the number of *EPS* iterations since the last converged value. By the default, a constant tolerance is used.

**See also**

`getJDConstCorrectionTol`, `EPSJDSetConstCorrectionTol`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3627 <slepc4py/SLEPc/EPS.pyx#L3627>`

**setJDFix**(*fix*)

Set the threshold for changing the target in the correction equation.

Logically collective.

**Parameters**

**fix** (*float*) – The threshold for changing the target.

**Return type**

None

**Notes**

The target in the correction equation is fixed at the first iterations. When the norm of the residual vector is lower than the *fix* value, the target is set to the corresponding eigenvalue.

**See also**

`getJDFix`, `EPSJDSetFix`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3584 <slepc4py/SLEPc/EPS.pyx#L3584>`

**setJDInitialSize**(*initialsize*)

Set the initial size of the searching space.

Logically collective.

**Parameters**

**initialsize** (*int*) – The number of vectors of the initial searching subspace.

**Return type**

None

## Notes

If the flag in `setJDKrylovStart()` is set to `False` and the user provides vectors with `setInitialSpace()`, up to `initialsize` vectors will be used; and if the provided vectors are not enough, the solver completes the subspace with random vectors. In case the `setJDKrylovStart()` flag is `True`, the solver gets the first vector provided by the user or, if not available, a random vector, and expands the Krylov basis up to `initialsize` vectors.

### See also

`setJDKrylovStart`, `getJDInitialSize`, `EPSJDSetInitialSize`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3537 <slepc4py/SLEPc/EPS.pyx#L3537>`

**setJDKrylovStart** (*krylovstart=True*)

Set (toggle) starting the search subspace with a Krylov basis.

Logically collective.

### Parameters

**krylovstart** (*bool*) – True if starting the search subspace with a Krylov basis.

### Return type

`None`

### See also

`setJDInitialSize`, `setJDKrylovStart`, `EPSJDSetKrylovStart`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3406 <slepc4py/SLEPc/EPS.pyx#L3406>`

**setJDRestart** (*minv=None, plusk=None*)

Set the number of vectors of the search space after restart.

Logically collective.

Set the number of vectors of the search space after restart and the number of vectors saved from the previous iteration.

### Parameters

- **minv** (*int* / *None*) – The number of vectors of the search subspace after restart.
- **plusk** (*int* / *None*) – The number of vectors saved from the previous iteration.

### Return type

`None`

### See also

`getJDRestart`, `EPSJDSetRestart`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3486 <slepc4py/SLEPc/EPS.pyx#L3486>`

**setKrylovSchurBSEType** (*bse*)

Set the Krylov-Schur variant used for BSE structured eigenproblems.

Logically collective.

#### Parameters

**bse** (*KrylovSchurBSEType*) – The BSE method.

#### Return type

None

#### Notes

This call is only relevant if the type was set to *EPS.Type.KRYLOV SCHUR* with *setType()* and the problem type to *EPS.ProblemType.BSE* with *setProblemType()*.

#### See also

*createMatBSE*, *getKrylovSchurBSEType*, *EP SKrylovSchurSetBSEType*

**:sources:** [Source code at slepc4py/SLEPc/EP S.pyx:2475 <slepc4py/SLEPc/EP S.pyx#L2475>](#)

#### **setKrylovSchurDetectZeros**(*detect*)

Set the flag that enforces zero detection in spectrum slicing.

Logically collective.

Set a flag to enforce the detection of zeros during the factorizations throughout the spectrum slicing computation.

#### Parameters

**detect** (*bool*) – True if zeros must checked for.

#### Return type

None

#### Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with *setInterval()* and *SINVERT* is set.

A zero in the factorization indicates that a shift coincides with an eigenvalue.

This flag is turned off by default, and may be necessary in some cases, especially when several partitions are being used. This feature currently requires an external package for factorizations with support for zero detection, e.g., MUMPS.

#### See also

*setInterval*, *getKrylovSchurDetectZeros*, *EP SKrylovSchurSetDetectZeros*

**:sources:** [Source code at slepc4py/SLEPc/EP S.pyx:2659 <slepc4py/SLEPc/EP S.pyx#L2659>](#)

#### **setKrylovSchurDimensions**(*nev=None*, *ncv=None*, *mpd=None*)

Set the dimensions used for each subsolve step (spectrum slicing).

Logically collective.

#### Parameters

- **nev** (*int* / *None*) – Number of eigenvalues to compute.
- **ncv** (*int* / *None*) – Maximum dimension of the subspace to be used by the solver.



- **mpd** (*int* / *None*) – Maximum dimension allowed for the projected problem.

**Return type**

*None*

## Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with *setInterval()* and *SINVERT* is set.

The meaning of the parameters is the same as in *setDimensions()*, but the ones here apply to every subsolve done by the child *EPS* object.

### See also

*setInterval*, *getKrylovSchurDimensions*, *EPSKrylovSchurSetDimensions*

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:2712 <slepc4py/SLEPc/EPs.pyx#L2712>](#)

**setKrylovSchurLocking**(*lock*)

Set (toggle) locking/non-locking variants of the Krylov-Schur method.

Logically collective.

### Parameters

**lock** (*bool*) – True if the locking variant must be selected.

**Return type**

*None*

## Notes

The default is to lock converged eigenpairs when the method restarts. This behavior can be changed so that all directions are kept in the working subspace even if already converged to working accuracy (the non-locking variant).

### See also

*getKrylovSchurLocking*, *EPSKrylovSchurSetLocking*

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:2561 <slepc4py/SLEPc/EPs.pyx#L2561>](#)

**setKrylovSchurPartitions**(*npart*)

Set the number of partitions of the communicator (spectrum slicing).

Logically collective.

Set the number of partitions for the case of doing spectrum slicing for a computational interval with the communicator split in several sub-communicators.

### Parameters

**npart** (*int*) – The number of partitions.

**Return type**

*None*

## Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with `setInterval()` and `SINVERT` is set.

By default, `npart=1` so all processes in the communicator participate in the processing of the whole interval. If `npart>1` then the interval is divided into `npart` subintervals, each of them being processed by a subset of processes.

The interval is split proportionally unless the separation points are specified with `setKrylovSchurSubintervals()`.

### See also

`setInterval`, `getKrylovSchurPartitions`, `EPKrylovSchurSetPartitions`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2605 <slepc4py/SLEPc/EPs.pyx#L2605>`

## `setKrylovSchurRestart(keep)`

Set the restart parameter for the Krylov-Schur method.

Logically collective.

It is the proportion of basis vectors that must be kept after restart.

### Parameters

**keep** (*float*) – The number of vectors to be kept at restart.

### Return type

`None`

## Notes

Allowed values are in the range `[0.1,0.9]`. The default is `0.5`.

### See also

`getKrylovSchurRestart`, `EPKrylovSchurSetRestart`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2518 <slepc4py/SLEPc/EPs.pyx#L2518>`

## `setKrylovSchurSubintervals(subint)`

Set the subinterval boundaries.

Logically collective.

Set the subinterval boundaries for spectrum slicing with a computational interval with several partitions.

### Parameters

**subint** (*Sequence[`float`]*) – Real values specifying subintervals.

### Return type

`None`

## Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with `setInterval()` and `SINVERT` is set.

This function must be called after `setKrylovSchurPartitions()`. For `npart` partitions, the argument `subint` must contain `npart+1` real values sorted in ascending order: `subint_0`, `subint_1`, ..., `subint_npart`, where the first and last values must coincide with the interval endpoints set with `setInterval()`. The subintervals are then defined by two consecutive points: `[subint_0, subint_1]`, `[subint_1, subint_2]`, and so on.

### See also

`setInterval`, `setKrylovSchurPartitions`, `EPSKrylovSchurSetSubintervals`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:2968 <slepc4py/SLEPc/EPS.pyx#L2968>`

## `setLOBPCGBlockSize(bs)`

Set the block size of the LOBPCG method.

Logically collective.

### Parameters

**bs** (*int*) – The block size.

### Return type

None

### See also

`getLOBPCGBlockSize`, `EPSLOBPCGSetBlockSize`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3759 <slepc4py/SLEPc/EPS.pyx#L3759>`

## `setLOBPCGLocking(lock)`

Toggle between locking and non-locking (LOBPCG method).

Logically collective.

### Parameters

**lock** (*bool*) – True if the locking variant must be selected.

### Return type

None

## Notes

This flag refers to soft locking (converged vectors within the current block iterate), since hard locking is always used (when `nev` is larger than the block size).

### See also

`getLOBPCGLocking`, `EPSLOBPCGSetLocking`

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:3840 <slepc4py/SLEPc/EPS.pyx#L3840>`

### **setLOBPCGRestart**(*restart*)

Set the restart parameter for the LOBPCG method.

Logically collective.

#### **Parameters**

**restart** (*float*) – The percentage of the block of vectors to force a restart.

#### **Return type**

*None*

### **Notes**

The meaning of this parameter is the proportion of vectors within the current block iterate that must have converged in order to force a restart with hard locking. Allowed values are in the range [0.1,1.0]. The default is 0.9.

#### **See also**

[\*getLOBPCGRestart\*](#), [\*EPSLOBPCGSetRestart\*](#)

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:3796 <slepc4py/SLEPc/EPS.pyx#L3796>](#)

### **setLanczosReorthogType**(*reorthog*)

Set the type of reorthogonalization used during the Lanczos iteration.

Logically collective.

#### **Parameters**

**reorthog** ([\*LanczosReorthogType\*](#)) – The type of reorthogonalization.

#### **Return type**

*None*

### **Notes**

This call is only relevant if the type was set to [\*EPS.Type.LANCZOS\*](#) with [\*setType\(\)\*](#).

#### **See also**

[\*getLanczosReorthogType\*](#), [\*EPSLanczosSetReorthog\*](#)

**:sources:** [Source code at slepc4py/SLEPc/EPS.pyx:2430 <slepc4py/SLEPc/EPS.pyx#L2430>](#)

### **setLeftInitialSpace**(*space*)

Set a left initial space from which the eigensolver starts to iterate.

Collective.

#### **Parameters**

**space** ([\*Vec\*](#) | [\*list\[Vec\]\*](#)) – Set of basis vectors of the left initial space.

#### **Return type**

*None*

## Notes

Left initial vectors are used to initiate the left search space in two-sided eigensolvers. Users should pass here an approximation of the left eigenspace, if available.

The same comments in `setInitialSpace()` are applicable here.

### See also

`setInitialSpace`, `setTwoSided`, `EPSSetLeftInitialSpace`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1652 <slepc4py/SLEPc/EPs.pyx#L1652>`

**setLyapIIRanks**(*rkc=None*, *rkl=None*)

Set the ranks used in the solution of the Lyapunov equation.

Logically collective.

### Parameters

- **rkc** (*int* / *None*) – The compressed rank.
- **rkl** (*int* / *None*) – The Lyapunov rank.

### Return type

*None*

## Notes

Lyapunov inverse iteration needs to solve a large-scale Lyapunov equation at each iteration of the eigensolver. For this, an iterative solver (*LME*) is used, which requires to prescribe the rank of the solution matrix *X*. This is the meaning of parameter *rkl*. Later, this matrix is compressed into another matrix of rank *rkc*. If not provided, *rkl* is a small multiple of *rkc*.

### See also

`getLyapIIRanks`, `EPsLyapIISetRanks`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:3883 <slepc4py/SLEPc/EPs.pyx#L3883>`

**setMonitor**(*monitor*, *args=None*, *kargs=None*)

Append a monitor function to the list of monitors.

Logically collective.

### See also

`getMonitor`, `cancelMonitor`, `EPsMonitorSet`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1818 <slepc4py/SLEPc/EPs.pyx#L1818>`

### Parameters

- **monitor** (*EPsMonitorFunction* / *None*)
- **args** (*tuple*[*Any*, ...] / *None*)
- **kargs** (*dict*[*str*, *Any*] / *None*)

**Return type**

None

**setOperators**(*A*, *B=None*)

Set the matrices associated with the eigenvalue problem.

Collective.

**Parameters**

- **A** (*Mat*) – The matrix associated with the eigensystem.
- **B** (*Mat* / *None*) – The second matrix in the case of generalized eigenproblems; if not provided, a standard eigenproblem is assumed.

**Return type**

None

**Notes**

It must be called before `setUp()`. If it is called again after `setUp()` and the matrix sizes have changed then the *EPS* object is reset.

For structured eigenproblem types such as *BSE*, see `setProblemType()`, the provided matrices must have been created with the corresponding helper function, i.e., `createMatBSE()`.

**See also**

`getOperators`, `solve`, `setUp`, `reset`, `setProblemType`, `EPSSetOperators`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1545 <slepc4py/SLEPc/EPs.pyx#L1545>`

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all EPS options in the database.

Logically collective.

**Parameters**

**prefix** (*str* / *None*) – The prefix string to prepend to all EPS option requests.

**Return type**

None

**Notes**

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

For example, to distinguish between the runtime options for two different EPS contexts, one could call:

```
E1.setOptionsPrefix("eig1_")
E2.setOptionsPrefix("eig2_")
```

**See also**

`appendOptionsPrefix`, `getOptionsPrefix`, `EPSGetOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:482 <slepc4py/SLEPc/EPs.pyx#L482>`

### **setPowerShiftType**(*shift*)

Set the type of shifts used during the power iteration.

Logically collective.

This can be used to emulate the Rayleigh Quotient Iteration (RQI) method.

#### **Parameters**

**shift** (`PowerShiftType`) – The type of shift.

#### **Return type**

None

### **Notes**

This call is only relevant if the type was set to `EPS.Type.POWER` with `setType()`.

By default, shifts are constant (`EPS.PowerShiftType.CONSTANT`) and the iteration is the simple power method (or inverse iteration if a shift-and-invert transformation is being used).

A variable shift can be specified (`EPS.PowerShiftType.RAYLEIGH` or `EPS.PowerShiftType.WILKINSON`). In this case, the iteration behaves rather like a cubic converging method as RQI.

#### **See also**

`getPowerShiftType`, `EPSPowerSetShiftType`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2328 <slepc4py/SLEPc/EPs.pyx#L2328>`

### **setProblemType**(*problem\_type*)

Set the type of the eigenvalue problem.

Logically collective.

#### **Parameters**

**problem\_type** (`ProblemType`) – The problem type to be set.

#### **Return type**

None

### **Notes**

This function must be used to instruct SLEPc to exploit symmetry or other kind of structure. If no problem type is specified, by default a non-Hermitian problem is assumed (either standard or generalized). If the user knows that the problem is Hermitian (i.e.,  $A = A^*$ ) or generalized Hermitian (i.e.,  $A = A^*$ ,  $B = B^*$ , and  $B$  positive definite) then it is recommended to set the problem type so that eigensolver can exploit these properties.

If the user does not call this function, the solver will use a reasonable guess.

For structured problem types such as `BSE`, the matrices passed in via `setOperators()` must have been created with the corresponding helper function, i.e., `createMatBSE()`.

#### **See also**

`setOperators`, `createMatBSE`, `getProblemType`, `EPSSetProblemType`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:572 <slepc4py/SLEPc/EPs.pyx#L572>`

**setPurify**(*purify=True*)

Set (toggle) eigenvector purification.

Logically collective.

**Parameters**

**purify** (*bool*) – True to activate purification (default).

**Return type**

None

**Notes**

By default, eigenvectors of generalized symmetric eigenproblems are purified in order to purge directions in the nullspace of matrix  $B$ . If the user knows that  $B$  is non-singular, then purification can be safely deactivated and some computational cost is avoided (this is particularly important in interval computations).

**See also**

[getPurify](#), [setInterval](#), [EPSSetPurify](#)

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:1151 <slepc4py/SLEPc/EPs.pyx#L1151>](#)

**setRG**(*rg*)

Set a region object associated to the eigensolver.

Collective.

**Parameters**

**rg** (*RG*) – The region context.

**Return type**

None

**See also**

[getRG](#), [EPSSetRG](#)

**:sources:** [Source code at slepc4py/SLEPc/EPs.pyx:1501 <slepc4py/SLEPc/EPs.pyx#L1501>](#)

**setRQCGReset**(*nrest*)

Set the reset parameter of the RQCG iteration.

Logically collective.

**Parameters**

**nrest** (*int*) – The number of iterations between resets.

**Return type**

None

**Notes**

Every *nrest* iterations the solver performs a Rayleigh-Ritz projection step.



**See also**

[`getRQCGReset`](#), [`EPSRQCSetReset`](#)

**:sources:** ``Source code at slepc4py/SLEPc/EPS.pyx:3717 <slepc4py/SLEPc/EPS.pyx#L3717>``

**setST**(*st*)

Set a spectral transformation object associated to the eigensolver.

Collective.

**Parameters**

**st** (*ST*) – The spectral transformation.

**Return type**

`None`

**See also**

[`getST`](#), [`EPSSetST`](#)

**:sources:** ``Source code at slepc4py/SLEPc/EPS.pyx:1390 <slepc4py/SLEPc/EPS.pyx#L1390>``

**setStoppingTest**(*stopping*, *args=None*, *kargs=None*)

Set a function to decide when to stop the outer iteration of the eigensolver.

Logically collective.

**See also**

[`getStoppingTest`](#), [`EPSSetStoppingTestFunction`](#)

**:sources:** ``Source code at slepc4py/SLEPc/EPS.pyx:1684 <slepc4py/SLEPc/EPS.pyx#L1684>``

**Parameters**

- **stopping** ([`EPSStoppingFunction`](#) | `None`)
- **args** (`tuple`[*Any*, ...] | `None`)
- **kargs** (`dict`[*str*, *Any*] | `None`)

**Return type**

`None`

**setTarget**(*target*)

Set the value of the target.

Logically collective.

**Parameters**

**target** ([`Scalar`](#)) – The value of the target.

**Return type**

`None`

## Notes

The target is a scalar value used to determine the portion of the spectrum of interest. It is used in combination with `setWhichEigenpairs()`.

When PETSc is built with real scalars, it is not possible to specify a complex target.

### See also

`getTarget`, `EPSSetTarget`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:945 <slepc4py/SLEPc/EPs.pyx#L945>`

**setThreshold**(*thres*, *rel*=*False*)

Set the threshold used in the threshold stopping test.

Logically collective.

### Parameters

- **thres** (*float*) – The threshold.
- **rel** (*bool*) – Whether the threshold is relative or not.

### Return type

*None*

## Notes

This function internally sets a special stopping test based on the threshold, where eigenvalues are computed in sequence until one of the computed eigenvalues is below/above the threshold (depending on whether largest or smallest eigenvalues are computed). The details are given in `EPSSetThreshold`.

### See also

`setStoppingTest`, `getThreshold`, `EPSSetThreshold`

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:893 <slepc4py/SLEPc/EPs.pyx#L893>`

**setTolerances**(*tol*=*None*, *max\_it*=*None*)

Set the tolerance and max. iter. used by the default EPS convergence tests.

Logically collective.

### Parameters

- **tol** (*float* / *None*) – The convergence tolerance.
- **max\_it** (*int* / *None*) – The maximum number of iterations.

### Return type

*None*

## Notes

Use `DETERMINE` for *max\_it* to assign a reasonably good value, which is dependent on the solution method.

**See also**

[`getTolerances`](#), [`EPSSetTolerances`](#)

**`:sources:`**``Source code at slepc4py/SLEPc/EPs.pyx:1057 <slepc4py/SLEPc/EPs.pyx#L1057>``

**`setTrackAll`**(*trackall*)

Set if the solver must compute the residual of all approximate eigenpairs.

Logically collective.

**Parameters**

**`trackall`** (*bool*) – Whether to compute all residuals or not.

**Return type**

`None`

**See also**

[`getTrackAll`](#), [`EPSSetTrackAll`](#)

**`:sources:`**``Source code at slepc4py/SLEPc/EPs.pyx:1272 <slepc4py/SLEPc/EPs.pyx#L1272>``

**`setTrueResidual`**(*trueeres*)

Set if the solver must compute the true residual explicitly or not.

Logically collective.

**Parameters**

**`trueeres`** (*bool*) – Whether the solver computes true residuals or not.

**Return type**

`None`

**See also**

[`getTrueResidual`](#), [`EPSSetTrueResidual`](#)

**`:sources:`**``Source code at slepc4py/SLEPc/EPs.pyx:1235 <slepc4py/SLEPc/EPs.pyx#L1235>``

**`setTwoSided`**(*twosided*)

Set to use a two-sided variant that also computes left eigenvectors.

Logically collective.

**Parameters**

**`twosided`** (*bool*) – Whether the two-sided variant is to be used or not.

**Return type**

`None`

**Notes**

If the user sets `twosided` to `True` then the solver uses a variant of the algorithm that computes both right and left eigenvectors. This is usually much more costly. This option is not available in all solvers.

When using two-sided solvers, the problem matrices must have both the `Mat.mult` and `Mat.multTranspose` operations defined.

#### See also

[`getTwoSided`](#), [`getLeftEigenvector`](#), [`EPSSetTwoSided`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1104 <slepc4py/SLEPc/EPs.pyx#L1104>`

**setType**(*eps\_type*)

Set the particular solver to be used in the EPS object.

Logically collective.

#### Parameters

**eps\_type** (*Type* / *str*) – The solver to be used.

#### Return type

None

#### Notes

The default is *KRYLOV SCHUR*. Normally, it is best to use [`setFromOptions\(\)`](#) and then set the EPS type from the options database rather than by using this routine. Using the options database provides the user with maximum flexibility in evaluating the different available methods.

#### See also

[`getType`](#), [`EPSSetType`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:417 <slepc4py/SLEPc/EPs.pyx#L417>`

**setUp()**

Set up all the internal data structures.

Collective.

#### Notes

Sets up all the internal data structures necessary for the execution of the eigensolver. This includes the setup of the internal *ST* object.

This function need not be called explicitly in most cases, since [`solve\(\)`](#) calls it. It can be useful when one wants to measure the set-up time separately from the solve time.

#### See also

[`solve`](#), [`setInitialSpace`](#), [`setDeflationSpace`](#), [`EPSSetUp`](#)

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:1875 <slepc4py/SLEPc/EPs.pyx#L1875>`

#### Return type

None

### **setWhichEigenpairs(*which*)**

Set which portion of the spectrum is to be sought.

Logically collective.

#### **Parameters**

**which** ([Which](#)) – The portion of the spectrum to be sought by the solver.

#### **Return type**

None

### **Notes**

Not all eigensolvers implemented in EPS account for all the possible values. Also, some values make sense only for certain types of problems. If SLEPc is compiled for real numbers [EPS.Which.LARGEST\\_IMAGINARY](#) and [EPS.Which.SMALLEST\\_IMAGINARY](#) use the absolute value of the imaginary part for eigenvalue selection.

The target is a scalar value provided with [setTarget\(\)](#).

The criterion [EPS.Which.TARGET\\_IMAGINARY](#) is available only in case PETSc and SLEPc have been built with complex scalars.

[EPS.Which.ALL](#) is intended for use in combination with an interval (see [setInterval\(\)](#)), when all eigenvalues within the interval are requested, or in the context of the [EPS.Type.CISS](#) solver for computing all eigenvalues in a region.

#### **See also**

[setTarget](#), [setInterval](#), [getWhichEigenpairs](#), [EPSSetWhichEigenpairs](#)

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:834 <slepc4py/SLEPc/EPS.pyx#L834>`

### **solve()**

Solve the eigensystem.

Collective.

### **Notes**

The problem matrices are specified with [setOperators\(\)](#).

[solve\(\)](#) will return without generating an error regardless of whether all requested solutions were computed or not. Call [getConverged\(\)](#) to get the actual number of computed solutions, and [getConvergedReason\(\)](#) to determine if the solver converged or failed and why.

#### **See also**

[setUp](#), [setOperators](#), [getConverged](#), [getConvergedReason](#), [EPSSolve](#)

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:1897 <slepc4py/SLEPc/EPS.pyx#L1897>`

#### **Return type**

None

**updateKrylovSchurSubcommMats**(*s=1.0*, *a=1.0*, *Au=None*, *t=1.0*, *b=1.0*, *Bu=None*, *structure=None*, *globalup=False*)

Update the eigenproblem matrices stored internally in the communicator.

Collective.

Update the eigenproblem matrices stored internally in the subcommunicator to which the calling process belongs.

#### Parameters

- **s** (*Scalar*) – Scalar that multiplies the existing A matrix.
- **a** (*Scalar*) – Scalar used in the axpy operation on A.
- **Au** (*petsc4py.PETSc.Mat* / *None*) – The matrix used in the axpy operation on A.
- **t** (*Scalar*) – Scalar that multiplies the existing B matrix.
- **b** (*Scalar*) – Scalar used in the axpy operation on B.
- **Bu** (*petsc4py.PETSc.Mat* / *None*) – The matrix used in the axpy operation on B.
- **structure** (*petsc4py.PETSc.Mat.Structure* / *None*) – Either same, different, or a subset of the non-zero sparsity pattern.
- **globalup** (*bool*) – Whether global matrices must be updated or not.

#### Return type

*None*

#### Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with *setInterval()* and *SINVERT* is set. And is relevant only when the number of partitions (*setKrylovSchurPartitions()*) is larger than one.

This function modifies the eigenproblem matrices at subcommunicator level, and optionally updates the global matrices in the parent communicator. The updates are expressed as  $A \leftarrow sA + aAu$ ,  $B \leftarrow tB + bBu$ .

It is possible to update one of the matrices, or both.

The matrices Au and Bu must be equal in all subcommunicators.

The structure flag is passed to the *petsc4py.PETSc.Mat.axpy* operations to perform the updates.

If *globalup* is True, communication is carried out to reconstruct the updated matrices in the parent communicator.

#### See also

*setInterval*, *setKrylovSchurPartitions*, *EPSCrylovSchurUpdateSubcommMats*

**:sources:** `Source code at slepc4py/SLEPc/EPs.pyx:2897 <slepc4py/SLEPc/EPs.pyx#L2897>`

**valuesView**(*viewer=None*)

Display the computed eigenvalues in a viewer.

Collective.

#### Parameters

- **viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

None

**See also**`solve`, `vectorsView`, `errorView`, `EPSValuesView`**`:sources:`**``Source code at slepc4py/SLEPc/EPS.pyx:2288 <slepc4py/SLEPc/EPS.pyx#L2288>``**`vectorsView`**(*viewer=None*)

Output computed eigenvectors to a viewer.

Collective.

**Parameters****`viewer`** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.**Return type**

None

**See also**`solve`, `valuesView`, `errorView`, `EPSVectorsView`**`:sources:`**``Source code at slepc4py/SLEPc/EPS.pyx:2307 <slepc4py/SLEPc/EPS.pyx#L2307>``**`view`**(*viewer=None*)

Print the EPS data structure.

Collective.

**Parameters****`viewer`** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.**Return type**

None

**See also**`EPSView`**`:sources:`**``Source code at slepc4py/SLEPc/EPS.pyx:351 <slepc4py/SLEPc/EPS.pyx#L351>``**Attributes Documentation****`bv`**The basis vectors (*BV*) object associated.**`:sources:`**``Source code at slepc4py/SLEPc/EPS.pyx:4343 <slepc4py/SLEPc/EPS.pyx#L4343>``**`ds`**The direct solver (*DS*) object associated.**`:sources:`**``Source code at slepc4py/SLEPc/EPS.pyx:4357 <slepc4py/SLEPc/EPS.pyx#L4357>``

**extraction**

The type of extraction technique to be employed.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4273 <slepc4py/SLEPc/EPS.pyx#L4273>`

**max\_it**

The maximum iteration count.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4301 <slepc4py/SLEPc/EPS.pyx#L4301>`

**problem\_type**

The type of the eigenvalue problem.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4266 <slepc4py/SLEPc/EPS.pyx#L4266>`

**purify**

Eigenvector purification.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4322 <slepc4py/SLEPc/EPS.pyx#L4322>`

**rg**

The region (*RG*) object associated.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4350 <slepc4py/SLEPc/EPS.pyx#L4350>`

**st**

The spectral transformation (*ST*) object associated.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4336 <slepc4py/SLEPc/EPS.pyx#L4336>`

**target**

The value of the target.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4287 <slepc4py/SLEPc/EPS.pyx#L4287>`

**tol**

The tolerance.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4294 <slepc4py/SLEPc/EPS.pyx#L4294>`

**track\_all**

Compute the residual norm of all approximate eigenpairs.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4329 <slepc4py/SLEPc/EPS.pyx#L4329>`

**true\_residual**

Compute the true residual explicitly.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4315 <slepc4py/SLEPc/EPS.pyx#L4315>`

**two\_sided**

Two-sided that also computes left eigenvectors.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4308 <slepc4py/SLEPc/EPS.pyx#L4308>`

**which**

The portion of the spectrum to be sought.

**:sources:** `Source code at slepc4py/SLEPc/EPS.pyx:4280 <slepc4py/SLEPc/EPS.pyx#L4280>`



## slepc4py.SLEPc.FN

**class** slepc4py.SLEPc.FN

Bases: `Object`

Mathematical Function.

The *FN* package provides the functionality to represent a simple mathematical function such as an exponential, a polynomial or a rational function. This is used as a building block for defining the function associated to the nonlinear eigenproblem, as well as for specifying which function to use when computing the action of a matrix function on a vector.

### Enumerations

<i>CombineType</i>	FN type of combination of child functions.
<i>ParallelType</i>	FN parallel types.
<i>Type</i>	FN type.

## slepc4py.SLEPc.FN.CombineType

**class** slepc4py.SLEPc.FN.CombineType

Bases: `object`

FN type of combination of child functions.

- *ADD*: Addition  $f(x) = f_1(x) + f_2(x)$
- *MULTIPLY*: Multiplication  $f(x) = f_1(x)f_2(x)$
- *DIVIDE*: Division  $f(x) = f_1(x)/f_2(x)$
- *COMPOSE*: Composition  $f(x) = f_2(f_1(x))$

#### See also

FNCombineType

### Attributes Summary

<i>ADD</i>	Constant ADD of type <code>int</code>
<i>COMPOSE</i>	Constant COMPOSE of type <code>int</code>
<i>DIVIDE</i>	Constant DIVIDE of type <code>int</code>
<i>MULTIPLY</i>	Constant MULTIPLY of type <code>int</code>

### Attributes Documentation

**ADD:** `int` = *ADD*

Constant ADD of type `int`

**COMPOSE:** `int` = *COMPOSE*

Constant COMPOSE of type `int`

**DIVIDE:** `int` = *DIVIDE*

Constant DIVIDE of type `int`

**MULTIPLY:** `int` = **MULTIPLY**  
Constant MULTIPLY of type `int`

### **slepc4py.SLEPc.FN.ParallelType**

**class** `slepc4py.SLEPc.FN.ParallelType`

Bases: `object`

FN parallel types.

- *REDUNDANT*: Every process performs the computation redundantly.
- *SYNCHRONIZED*: The first process sends the result to the rest.

#### **See also**

`FNParallelType`

### **Attributes Summary**

<i>REDUNDANT</i>	Constant REDUNDANT of type <code>int</code>
<i>SYNCHRONIZED</i>	Constant SYNCHRONIZED of type <code>int</code>

### **Attributes Documentation**

**REDUNDANT:** `int` = **REDUNDANT**  
Constant REDUNDANT of type `int`

**SYNCHRONIZED:** `int` = **SYNCHRONIZED**  
Constant SYNCHRONIZED of type `int`

### **slepc4py.SLEPc.FN.Type**

**class** `slepc4py.SLEPc.FN.Type`

Bases: `object`

FN type.

- *COMBINE*: A math function defined by combining two functions.
- *RATIONAL*: A rational function  $f(x) = p(x)/q(x)$ .
- *EXP*: The exponential function  $f(x) = e^x$ .
- *LOG*: The logarithm function  $f(x) = \log x$ .
- *PHI*: One of the Phi\_k functions with index k.
- *SQRT*: The square root function  $f(x) = \sqrt{x}$ .
- *INVSQRT*: The inverse square root function.

#### **See also**

`FNType`

## Attributes Summary

<i>COMBINE</i>	Object COMBINE of type <i>str</i>
<i>EXP</i>	Object EXP of type <i>str</i>
<i>INVSQRT</i>	Object INVSQRT of type <i>str</i>
<i>LOG</i>	Object LOG of type <i>str</i>
<i>PHI</i>	Object PHI of type <i>str</i>
<i>RATIONAL</i>	Object RATIONAL of type <i>str</i>
<i>SQRT</i>	Object SQRT of type <i>str</i>

## Attributes Documentation

**COMBINE:** *str* = COMBINE

Object COMBINE of type *str*

**EXP:** *str* = EXP

Object EXP of type *str*

**INVSQRT:** *str* = INVSQRT

Object INVSQRT of type *str*

**LOG:** *str* = LOG

Object LOG of type *str*

**PHI:** *str* = PHI

Object PHI of type *str*

**RATIONAL:** *str* = RATIONAL

Object RATIONAL of type *str*

**SQRT:** *str* = SQRT

Object SQRT of type *str*

## Methods Summary

<i>appendOptionsPrefix</i> ([prefix])	Append to the prefix used for searching for all FN options in the database.
<i>create</i> ([comm])	Create the FN object.
<i>destroy</i> ()	Destroy the FN object.
<i>duplicate</i> ([comm])	Duplicate the FN object copying all parameters.
<i>evaluateDerivative</i> (x)	Compute the value of the derivative $f'(x)$ for a given x.
<i>evaluateFunction</i> (x)	Compute the value of the function $f(x)$ for a given x.
<i>evaluateFunctionMat</i> (A[, B])	Compute the value of the function $f(A)$ for a given matrix A.
<i>evaluateFunctionMatVec</i> (A[, v])	Compute the first column of the matrix $f(A)$ .
<i>getCombineChildren</i> ()	Get the two child functions that constitute this combined function.
<i>getMethod</i> ()	Get the method currently used for matrix functions.
<i>getOptionsPrefix</i> ()	Get the prefix used for searching for all FN options in the database.
<i>getParallel</i> ()	Get the mode of operation in parallel runs.

continues on next page

Table 42 – continued from previous page

<code>getPhiIndex()</code>	Get the index of the phi-function.
<code>getRationalDenominator()</code>	Get the coefficients of the denominator of the rational function.
<code>getRationalNumerator()</code>	Get the coefficients of the numerator of the rational function.
<code>getScale()</code>	Get the scaling parameters that define the mathematical function.
<code>getType()</code>	Get the FN type of this object.
<code>setCombineChildren(comb, f1, f2)</code>	Set the two child functions that constitute this combined function.
<code>setFromOptions()</code>	Set FN options from the options database.
<code>setMethod(meth)</code>	Set the method to be used to evaluate functions of matrices.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all FN options in the database.
<code>setParallel(pmode)</code>	Set the mode of operation in parallel runs.
<code>setPhiIndex(k)</code>	Set the index of the phi-function.
<code>setRationalDenominator(alpha)</code>	Set the coefficients of the denominator of the rational function.
<code>setRationalNumerator(alpha)</code>	Set the coefficients of the numerator of the rational function.
<code>setScale([alpha, beta])</code>	Set the scaling parameters that define the mathematical function.
<code>setType(fn_type)</code>	Set the type for the FN object.
<code>view([viewer])</code>	Print the FN data structure.

### Attributes Summary

<code>method</code>	The method to be used to evaluate functions of matrices.
<code>parallel</code>	The mode of operation in parallel runs.

### Methods Documentation

#### `appendOptionsPrefix(prefix=None)`

Append to the prefix used for searching for all FN options in the database.

Logically collective.

##### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all FN option requests.

##### Return type

*None*

#### See also

`setOptionsPrefix`, `getOptionsPrefix`, `FNAppendOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:266 <slepc4py/SLEPc/FN.pyx#L266>`

**create**(*comm=None*)

Create the FN object.

Collective.

**Parameters**

**comm** (*Comm* / *None*) – MPI communicator; if not provided, it defaults to all processes.

**Return type**

*Self*

**See also**

[FNCreate](#)

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:182 <slepc4py/SLEPc/FN.pyx#L182>](#)

**destroy**()

Destroy the FN object.

Collective.

**See also**

[FNDestroy](#)

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:168 <slepc4py/SLEPc/FN.pyx#L168>](#)

**Return type**

*Self*

**duplicate**(*comm=None*)

Duplicate the FN object copying all parameters.

Collective.

Duplicate the FN object copying all parameters, possibly with a different communicator.

**Parameters**

**comm** (*Comm* / *None*) – MPI communicator; if not provided, it defaults to the object's communicator.

**Returns**

The new object.

**Return type**

*FN*

**See also**

[create](#), [FNDuplicate](#)

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:321 <slepc4py/SLEPc/FN.pyx#L321>](#)

**evaluateDerivative**(*x*)

Compute the value of the derivative  $f'(x)$  for a given *x*.

Not collective.

**Parameters**

**x** ([Scalar](#)) – Value where the derivative must be evaluated.

**Returns**

The result of  $f'(x)$ .

**Return type**

[Scalar](#)

**Notes**

Scaling factors are taken into account, so the actual derivative evaluation will return  $abf'(ax)$ .

**See also**

[evaluateFunction](#), [setScale](#), [FNEvaluateDerivative](#)

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:382 <slepc4py/SLEPc/FN.pyx#L382>](#)

**evaluateFunction(x)**

Compute the value of the function  $f(x)$  for a given x.

Not collective.

**Parameters**

**x** ([Scalar](#)) – Value where the function must be evaluated.

**Returns**

The result of  $f(x)$ .

**Return type**

[Scalar](#)

**Notes**

Scaling factors are taken into account, so the actual function evaluation will return  $bf(ax)$ .

**See also**

[evaluateDerivative](#), [evaluateFunctionMat](#), [setScale](#), [FNEvaluateFunction](#)

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:352 <slepc4py/SLEPc/FN.pyx#L352>](#)

**evaluateFunctionMat(A, B=None)**

Compute the value of the function  $f(A)$  for a given matrix A.

Logically collective.

**Parameters**

- **A** ([Mat](#)) – Matrix on which the function must be evaluated.
- **B** ([Mat](#) / [None](#)) – Placeholder for the result.

**Returns**

The result of  $f(A)$ .

**Return type**

[petsc4py.PETSc.Mat](#)

## Notes

Scaling factors are taken into account, so the actual function evaluation will return  $bf(aA)$ .

### See also

`evaluateFunction`, `evaluateFunctionMatVec`, `FNEvaluateFunctionMat`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:412 <slepc4py/SLEPc/FN.pyx#L412>`

**evaluateFunctionMatVec**(*A*, *v=None*)

Compute the first column of the matrix  $f(A)$ .

Logically collective.

### Parameters

- **A** (*Mat*) – Matrix on which the function must be evaluated.
- **v** (*Vec* | *None*)

### Returns

The first column of the result  $f(A)$ .

### Return type

`petsc4py.PETSc.Vec`

## Notes

This operation is similar to `evaluateFunctionMat()` but returns only the first column of  $f(A)$ , hence saving computations in most cases.

### See also

`evaluateFunctionMat`, `FNEvaluateFunctionMatVec`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:443 <slepc4py/SLEPc/FN.pyx#L443>`

**getCombineChildren()**

Get the two child functions that constitute this combined function.

Not collective.

Get the two child functions that constitute this combined function, and the way they must be combined.

### Returns

- **comb** (*CombineType*) – How to combine the functions (addition, multiplication, division, composition).
- **f1** (*FN*) – First function.
- **f2** (*FN*) – Second function.

### Return type

`tuple[CombineType, FN, FN]`

**See also**

[\*setCombineChildren\*](#), [\*FNCombineGetChildren\*](#)

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:725](#) <[slepc4py/SLEPc/FN.pyx#L725](#)>

**getMethod()**

Get the method currently used for matrix functions.

Not collective.

**Returns**

An index identifying the method.

**Return type**

`int`

**See also**

[\*setMethod\*](#), [\*FNGetMethod\*](#)

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:546](#) <[slepc4py/SLEPc/FN.pyx#L546](#)>

**getOptionsPrefix()**

Get the prefix used for searching for all FN options in the database.

Not collective.

**Returns**

The prefix string set for this FN object.

**Return type**

`str`

**See also**

[\*setOptionsPrefix\*](#), [\*appendOptionsPrefix\*](#), [\*FNGetOptionsPrefix\*](#)

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:285](#) <[slepc4py/SLEPc/FN.pyx#L285](#)>

**getParallel()**

Get the mode of operation in parallel runs.

Not collective.

**Returns**

The parallel mode.

**Return type**

[\*ParallelType\*](#)

**See also**

[\*setParallel\*](#), [\*FNGetParallel\*](#)



**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:588 <slepc4py/SLEPc/FN.pyx#L588>`

#### **getPhiIndex()**

Get the index of the phi-function.

Not collective.

##### **Returns**

The index.

##### **Return type**

`int`

##### **See also**

`setPhiIndex`, `FNPhiGetIndex`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:778 <slepc4py/SLEPc/FN.pyx#L778>`

#### **getRationalDenominator()**

Get the coefficients of the denominator of the rational function.

Not collective.

##### **Returns**

Coefficients.

##### **Return type**

`ArrayScalar`

##### **See also**

`setRationalDenominator`, `FN RationalGetDenominator`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:674 <slepc4py/SLEPc/FN.pyx#L674>`

#### **getRationalNumerator()**

Get the coefficients of the numerator of the rational function.

Not collective.

##### **Returns**

Coefficients.

##### **Return type**

`ArrayScalar`

##### **See also**

`setRationalNumerator`, `FN RationalGetNumerator`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:629 <slepc4py/SLEPc/FN.pyx#L629>`

#### **getScale()**

Get the scaling parameters that define the mathematical function.

Not collective.

### Returns

- **alpha** (*Scalar*) – Inner scaling (argument).
- **beta** (*Scalar*) – Outer scaling (result).

### Return type

`tuple[Scalar, Scalar]`

### See also

`setScale`, `FNGetScale`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:496 <slepc4py/SLEPc/FN.pyx#L496>`

### `getType()`

Get the FN type of this object.

Not collective.

### Returns

The math function type currently being used.

### Return type

`str`

### See also

`setType`, `FNGetType`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:222 <slepc4py/SLEPc/FN.pyx#L222>`

### `setCombineChildren(comb, f1, f2)`

Set the two child functions that constitute this combined function.

Logically collective.

Set the two child functions that constitute this combined function, and the way they must be combined.

### Parameters

- **comb** (*CombineType*) – How to combine the functions (addition, multiplication, division, composition).
- **f1** (*FN*) – First function.
- **f2** (*FN*) – Second function.

### Return type

`None`

### See also

`getCombineChildren`, `FNCombineSetChildren`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:699 <slepc4py/SLEPc/FN.pyx#L699>`

### **setFromOptions()**

Set FN options from the options database.

Collective.

#### **Notes**

To see all options, run your program with the `-help` option.

#### **See also**

`setOptionsPrefix`, `FNSetFromOptions`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:304 <slepc4py/SLEPc/FN.pyx#L304>`

#### **Return type**

`None`

### **setMethod(meth)**

Set the method to be used to evaluate functions of matrices.

Logically collective.

#### **Parameters**

**meth** (`int`) – An index identifying the method.

#### **Return type**

`None`

#### **Notes**

In some *FN* types there are more than one algorithms available for computing matrix functions. In that case, this function allows choosing the wanted method.

If `meth` is currently set to 0 and the input argument of `FN.evaluateFunctionMat()` is a symmetric/Hermitian matrix, then the computation is done via the eigendecomposition, rather than with the general algorithm.

#### **See also**

`getMethod`, `FNSetMethod`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:517 <slepc4py/SLEPc/FN.pyx#L517>`

### **setOptionsPrefix(prefix=None)**

Set the prefix used for searching for all FN options in the database.

Logically collective.

#### **Parameters**

**prefix** (`str` / `None`) – The prefix string to prepend to all FN option requests.

#### **Return type**

`None`

## Notes

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

### See also

`appendOptionsPrefix`, `getOptionsPrefix`, `FNGetOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:241 <slepc4py/SLEPc/FN.pyx#L241>`

## `setParallel(pmode)`

Set the mode of operation in parallel runs.

Logically collective.

### Parameters

**pmode** (`ParallelType`) – The parallel mode.

### Return type

`None`

## Notes

This is relevant only when the function is evaluated on a matrix, with either `evaluateFunctionMat()` or `evaluateFunctionMatVec()`.

### See also

`evaluateFunctionMat`, `getParallel`, `FNSetParallel`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:565 <slepc4py/SLEPc/FN.pyx#L565>`

## `setPhiIndex(k)`

Set the index of the phi-function.

Logically collective.

### Parameters

**k** (`int`) – The index.

### Return type

`None`

## Notes

If not set, the default index is 1.

### See also

`getPhiIndex`, `FNPhiSetIndex`

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:756 <slepc4py/SLEPc/FN.pyx#L756>`

### **setRationalDenominator**(*alpha*)

Set the coefficients of the denominator of the rational function.

Logically collective.

#### **Parameters**

**alpha** (*Sequence*[*Scalar*]) – Coefficients.

#### **Return type**

*None*

#### **See also**

*setRationalNumerator*, *FN*RationalSetDenominator

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:654 <slepc4py/SLEPc/FN.pyx#L654>](#)

### **setRationalNumerator**(*alpha*)

Set the coefficients of the numerator of the rational function.

Logically collective.

#### **Parameters**

**alpha** (*Sequence*[*Scalar*]) – Coefficients.

#### **Return type**

*None*

#### **See also**

*setRationalDenominator*, *FN*RationalSetNumerator

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:609 <slepc4py/SLEPc/FN.pyx#L609>](#)

### **setScale**(*alpha=None*, *beta=None*)

Set the scaling parameters that define the mathematical function.

Logically collective.

#### **Parameters**

- **alpha** (*Scalar* / *None*) – Inner scaling (argument), default is 1.0.
- **beta** (*Scalar* / *None*) – Outer scaling (result), default is 1.0.

#### **Return type**

*None*

#### **See also**

*getScale*, *evaluateFunction*, *FN*SetScale

**:sources:** [Source code at slepc4py/SLEPc/FN.pyx:473 <slepc4py/SLEPc/FN.pyx#L473>](#)

### **setType**(*fn\_type*)

Set the type for the FN object.

Logically collective.

#### Parameters

**fn\_type** (*Type* / *str*) – The math function type to be used.

#### Return type

*None*

#### See also

*getType*, *FNSetType*

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:203 <slepc4py/SLEPc/FN.pyx#L203>`

**view**(*viewer=None*)

Print the FN data structure.

Collective.

#### Parameters

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

#### Return type

*None*

#### See also

*FNView*

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:149 <slepc4py/SLEPc/FN.pyx#L149>`

### Attributes Documentation

#### method

The method to be used to evaluate functions of matrices.

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:799 <slepc4py/SLEPc/FN.pyx#L799>`

#### parallel

The mode of operation in parallel runs.

**:sources:** `Source code at slepc4py/SLEPc/FN.pyx:806 <slepc4py/SLEPc/FN.pyx#L806>`

### slepc4py.SLEPc.LME

**class** `slepc4py.SLEPc.LME`

Bases: *Object*

Linear Matrix Equation.

Linear Matrix Equation (*LME*) is the object provided by slepc4py for solving linear matrix equations such as Lyapunov or Sylvester where the solution has low rank.

## Enumerations

<i>ConvergedReason</i>	LME convergence reasons.
<i>ProblemType</i>	LME problem type.
<i>Type</i>	LME type.

### `slepc4py.SLEPc.LME.ConvergedReason`

**class** `slepc4py.SLEPc.LME.ConvergedReason`

Bases: `object`

LME convergence reasons.

- *CONVERGED\_TOL*: All eigenpairs converged to requested tolerance.
- *DIVERGED\_ITS*: Maximum number of iterations exceeded.
- *DIVERGED\_BREAKDOWN*: Solver failed due to breakdown.
- *CONVERGED\_ITERATING*: Iteration not finished yet.

#### See also

`LMEConvergedReason`

### Attributes Summary

<i>CONVERGED_ITERATING</i>	Constant <i>CONVERGED_ITERATING</i> of type <code>int</code>
<i>CONVERGED_TOL</i>	Constant <i>CONVERGED_TOL</i> of type <code>int</code>
<i>DIVERGED_BREAKDOWN</i>	Constant <i>DIVERGED_BREAKDOWN</i> of type <code>int</code>
<i>DIVERGED_ITS</i>	Constant <i>DIVERGED_ITS</i> of type <code>int</code>
<i>ITERATING</i>	Constant <i>ITERATING</i> of type <code>int</code>

### Attributes Documentation

**CONVERGED\_ITERATING:** `int` = *CONVERGED\_ITERATING*

Constant *CONVERGED\_ITERATING* of type `int`

**CONVERGED\_TOL:** `int` = *CONVERGED\_TOL*

Constant *CONVERGED\_TOL* of type `int`

**DIVERGED\_BREAKDOWN:** `int` = *DIVERGED\_BREAKDOWN*

Constant *DIVERGED\_BREAKDOWN* of type `int`

**DIVERGED\_ITS:** `int` = *DIVERGED\_ITS*

Constant *DIVERGED\_ITS* of type `int`

**ITERATING:** `int` = *ITERATING*

Constant *ITERATING* of type `int`

## slepc4py.SLEPc.LME.ProblemType

**class** slepc4py.SLEPc.LME.ProblemType

Bases: `object`

LME problem type.

- *LYAPUNOV*: Continuous-time Lyapunov.
- *SYLVESTER*: Continuous-time Sylvester.
- *GEN\_LYAPUNOV*: Generalized Lyapunov.
- *GEN\_SYLVESTER*: Generalized Sylvester.
- *DT\_LYAPUNOV*: Discrete-time Lyapunov.
- *STEIN*: Stein.

### See also

LMEProblemType

## Attributes Summary

<i>DT_LYAPUNOV</i>	Constant DT_LYAPUNOV of type <code>int</code>
<i>GEN_LYAPUNOV</i>	Constant GEN_LYAPUNOV of type <code>int</code>
<i>GEN_SYLVESTER</i>	Constant GEN_SYLVESTER of type <code>int</code>
<i>LYAPUNOV</i>	Constant LYAPUNOV of type <code>int</code>
<i>STEIN</i>	Constant STEIN of type <code>int</code>
<i>SYLVESTER</i>	Constant SYLVESTER of type <code>int</code>

## Attributes Documentation

**DT\_LYAPUNOV:** `int` = DT\_LYAPUNOV

Constant DT\_LYAPUNOV of type `int`

**GEN\_LYAPUNOV:** `int` = GEN\_LYAPUNOV

Constant GEN\_LYAPUNOV of type `int`

**GEN\_SYLVESTER:** `int` = GEN\_SYLVESTER

Constant GEN\_SYLVESTER of type `int`

**LYAPUNOV:** `int` = LYAPUNOV

Constant LYAPUNOV of type `int`

**STEIN:** `int` = STEIN

Constant STEIN of type `int`

**SYLVESTER:** `int` = SYLVESTER

Constant SYLVESTER of type `int`



## slepc4py.SLEPc.LME.Type

**class** slepc4py.SLEPc.LME.Type

Bases: object

LME type.

- *KRYLOV*: Restarted Krylov solver.

### See also

LMEType

## Attributes Summary

*KRYLOV*

Object KRYLOV of type *str*

## Attributes Documentation

**KRYLOV:** *str* = *KRYLOV*

Object KRYLOV of type *str*

## Methods Summary

<i>appendOptionsPrefix</i> ([prefix])	Append to the prefix used for searching in the database.
<i>cancelMonitor</i> ()	Clear all monitors for an <i>LME</i> object.
<i>computeError</i> ()	Compute the error associated with the last equation solved.
<i>create</i> ([comm])	Create the LME object.
<i>destroy</i> ()	Destroy the LME object.
<i>getBV</i> ()	Get the basis vector object associated to the LME object.
<i>getCoefficients</i> ()	Get the coefficient matrices of the matrix equation.
<i>getConvergedReason</i> ()	Get the reason why the <i>solve</i> () iteration was stopped.
<i>getDimensions</i> ()	Get the dimension of the subspace used by the solver.
<i>getErrorEstimate</i> ()	Get the error estimate obtained during the solve.
<i>getErrorIfNotConverged</i> ()	Get if <i>solve</i> () generates an error if the solver does not converge.
<i>getIterationNumber</i> ()	Get the current iteration number.
<i>getMonitor</i> ()	Get the list of monitor functions.
<i>getOptionsPrefix</i> ()	Get the prefix used for searching for all LME options in the database.
<i>getProblemType</i> ()	Get the LME problem type of this object.
<i>getRHS</i> ()	Get the right-hand side of the matrix equation.
<i>getSolution</i> ()	Get the solution of the matrix equation.
<i>getTolerances</i> ()	Get the tolerance and maximum iteration count.
<i>getType</i> ()	Get the LME type of this object.
<i>reset</i> ()	Reset the LME object.
<i>setBV</i> (bv)	Set a basis vector object to the LME object.

continues on next page

Table 48 – continued from previous page

<code>setCoefficients(A[, B, D, E])</code>	Set the coefficient matrices.
<code>setDimensions(ncv)</code>	Set the dimension of the subspace to be used by the solver.
<code>setErrorIfNotConverged([flg])</code>	Set <code>solve()</code> to generate an error if the solver has not converged.
<code>setFromOptions()</code>	Set LME options from the options database.
<code>setMonitor(monitor[, args, kargs])</code>	Append a monitor function to the list of monitors.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all LME options in the database.
<code>setProblemType(lme_problem_type)</code>	Set the LME problem type of this object.
<code>setRHS(C)</code>	Set the right-hand side of the matrix equation.
<code>setSolution([X])</code>	Set the placeholder for the solution of the matrix equation.
<code>setTolerances([tol, max_it])</code>	Set the tolerance and maximum iteration count.
<code>setType(lme_type)</code>	Set the particular solver to be used in the LME object.
<code>setUp()</code>	Set up all the internal necessary data structures.
<code>solve()</code>	Solve the linear matrix equation.
<code>view([viewer])</code>	Print the LME data structure.

## Attributes Summary

<code>bv</code>	The basis vectors ( <i>BV</i> ) object associated to the LME object.
<code>fn</code>	The math function ( <i>FN</i> ) object associated to the LME object.
<code>max_it</code>	The maximum iteration count used by the LME convergence tests.
<code>tol</code>	The tolerance value used by the LME convergence tests.

## Methods Documentation

### `appendOptionsPrefix(prefix=None)`

Append to the prefix used for searching in the database.

Logically collective.

Append to the prefix used for searching for all LME options in the database.

#### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all LME option requests.

#### Return type

*None*

#### See also

`setOptionsPrefix`, `getOptionsPrefix`, `LMEAppendOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:512 <slepc4py/SLEPc/LME.pyx#L512>`

### **cancelMonitor()**

Clear all monitors for an *LME* object.

Logically collective.

#### **See also**

[LMEMonitorCancel](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:713 <slepc4py/SLEPc/LME.pyx#L713>](#)

#### **Return type**

*None*

### **computeError()**

Compute the error associated with the last equation solved.

Collective.

#### **Returns**

The error.

#### **Return type**

*float*

### **Notes**

The error is based on the residual norm.

This function is not scalable (in terms of memory or parallel communication), so it should not be called except in the case of small problem size. For large equations, use [getErrorEstimate\(\)](#).

#### **See also**

[getErrorEstimate](#), [LMEComputeError](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:435 <slepc4py/SLEPc/LME.pyx#L435>](#)

### **create(comm=None)**

Create the LME object.

Collective.

#### **Parameters**

**comm** (*Comm* | *None*) – MPI communicator. If not provided, it defaults to all processes.

#### **Return type**

*Self*

#### **See also**

[LMECreat](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:121 <slepc4py/SLEPc/LME.pyx#L121>](#)

## **destroy()**

Destroy the LME object.

Collective.

### **See also**

[LMEDestroy](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:95](#) <[slepc4py/SLEPc/LME.pyx#L95](#)>

### **Return type**

*Self*

## **getBV()**

Get the basis vector object associated to the LME object.

Not collective.

### **Returns**

The basis vectors context.

### **Return type**

*BV*

### **See also**

[setBV](#), [LMEGetBV](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:638](#) <[slepc4py/SLEPc/LME.pyx#L638](#)>

## **getCoefficients()**

Get the coefficient matrices of the matrix equation.

Collective.

### **Returns**

- **A** ([petsc4py.PETSc.Mat](#)) – First coefficient matrix.
- **B** ([petsc4py.PETSc.Mat](#)) – Second coefficient matrix, if available.
- **D** ([petsc4py.PETSc.Mat](#)) – Third coefficient matrix, if available.
- **E** ([petsc4py.PETSc.Mat](#)) – Fourth coefficient matrix, if available.

### **Return type**

*tuple*[*Mat*, *Mat* | *None*, *Mat* | *None*, *Mat* | *None*]

### **See also**

[setCoefficients](#), [LMEGetCoefficients](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:264](#) <[slepc4py/SLEPc/LME.pyx#L264](#)>

### **getConvergedReason()**

Get the reason why the *solve()* iteration was stopped.

Not collective.

#### **Returns**

Negative value indicates diverged, positive value converged.

#### **Return type**

*ConvergedReason*

#### **See also**

*setTolerances*, *solve*, *setErrorIfNotConverged*, *LMEGetConvergedReason*

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:780 <slepc4py/SLEPc/LME.pyx#L780>](#)

### **getDimensions()**

Get the dimension of the subspace used by the solver.

Not collective.

#### **Returns**

Maximum dimension of the subspace to be used by the solver.

#### **Return type**

*int*

#### **See also**

*setDimensions*, *LMEGetDimensions*

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:601 <slepc4py/SLEPc/LME.pyx#L601>](#)

### **getErrorEstimate()**

Get the error estimate obtained during the solve.

Not collective.

#### **Returns**

The error estimate.

#### **Return type**

*float*

### **Notes**

This is the error estimated internally by the solver. The actual error bound can be computed with *computeError()*. Note that some solvers may not be able to provide an error estimate.

#### **See also**

*computeError*, *LMEGetErrorEstimate*

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:410 <slepc4py/SLEPc/LME.pyx#L410>](#)

### **getErrorIfNotConverged()**

Get if *solve()* generates an error if the solver does not converge.

Not collective.

Get a flag indicating whether *solve()* will generate an error if the solver does not converge.

#### **Returns**

True indicates you want the error generated.

#### **Return type**

bool

#### **See also**

*setErrorIfNotConverged*, *LMEGetErrorIfNotConverged*

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:823 <slepc4py/SLEPc/LME.pyx#L823>](#)

### **getIterationNumber()**

Get the current iteration number.

Not collective.

If the call to *solve()* is complete, then it returns the number of iterations carried out by the solution method.

#### **Returns**

Iteration number.

#### **Return type**

int

#### **See also**

*getConvergedReason*, *LMEGetIterationNumber*

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:758 <slepc4py/SLEPc/LME.pyx#L758>](#)

### **getMonitor()**

Get the list of monitor functions.

Not collective.

#### **Returns**

The list of monitor functions.

#### **Return type**

*LMEMonitorFunction*

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:700 <slepc4py/SLEPc/LME.pyx#L700>](#)

### **getOptionsPrefix()**

Get the prefix used for searching for all LME options in the database.

Not collective.

#### **Returns**

The prefix string set for this LME object.

**Return type**

`str`

**See also**

`setOptionsPrefix`, `appendOptionsPrefix`, `LMEGetOptionsPrefix`

**:sources:** ``Source code at slepc4py/SLEPc/LME.pyx:462 <slepc4py/SLEPc/LME.pyx#L462>``

**getProblemType()**

Get the LME problem type of this object.

Not collective.

**Returns**

The problem type currently being used.

**Return type**

`ProblemType`

**See also**

`setProblemType`, `LMEGetProblemType`

**:sources:** ``Source code at slepc4py/SLEPc/LME.pyx:206 <slepc4py/SLEPc/LME.pyx#L206>``

**getRHS()**

Get the right-hand side of the matrix equation.

Collective.

**Returns**

**C** – The low-rank matrix.

**Return type**

`petsc4py.PETSc.Mat`

**See also**

`setRHS`, `LMEGetRHS`

**:sources:** ``Source code at slepc4py/SLEPc/LME.pyx:327 <slepc4py/SLEPc/LME.pyx#L327>``

**getSolution()**

Get the solution of the matrix equation.

Collective.

**Returns**

**X** – The low-rank matrix.

**Return type**

`petsc4py.PETSc.Mat`

## Notes

If called after `solve()`, `X` will contain the solution of the equation.

The matrix `X` may have been passed by the user via `setSolution()`, although this is not required.

### See also

`solve`, `setSolution`, `LMEGetSolution`

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:382 <slepc4py/SLEPc/LME.pyx#L382>`

## `getTolerances()`

Get the tolerance and maximum iteration count.

Not collective.

### Returns

- `tol` (`float`) – The convergence tolerance.
- `max_it` (`int`) – The maximum number of iterations.

### Return type

`tuple[float, int]`

### See also

`setTolerances`, `LMEGetTolerances`

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:553 <slepc4py/SLEPc/LME.pyx#L553>`

## `getType()`

Get the LME type of this object.

Not collective.

### Returns

The solver currently being used.

### Return type

`str`

### See also

`setType`, `LMEGetType`

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:169 <slepc4py/SLEPc/LME.pyx#L169>`

## `reset()`

Reset the LME object.

Collective.

### See also

`LMEReset`



**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:109 <slepc4py/SLEPc/LME.pyx#L109>`

**Return type**

None

**setBV(bv)**

Set a basis vector object to the LME object.

Collective.

**Parameters**

**bv** (*BV*) – The basis vectors context.

**Return type**

None

**See also**

[getBV](#), [LMEsetBV](#)

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:658 <slepc4py/SLEPc/LME.pyx#L658>`

**setCoefficients(A, B=None, D=None, E=None)**

Set the coefficient matrices.

Collective.

Set the coefficient matrices that define the linear matrix equation to be solved.

**Parameters**

- **A** (*Mat*) – First coefficient matrix
- **B** (*Mat* | *None*) – Second coefficient matrix, optional
- **D** (*Mat* | *None*) – Third coefficient matrix, optional
- **E** (*Mat* | *None*) – Fourth coefficient matrix, optional

**Return type**

None

**Notes**

The matrix equation takes the general form  $AXE + DXB = C$ , where matrix  $C$  is not provided here but with [setRHS\(\)](#). Not all four matrices must be passed.

This must be called before [setUp\(\)](#). If called again after [setUp\(\)](#) then the *LME* object is reset.

**See also**

[getCoefficients](#), [solve](#), [setRHS](#), [setUp](#), [LMEsetCoefficients](#)

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:225 <slepc4py/SLEPc/LME.pyx#L225>`

**setDimensions(ncv)**

Set the dimension of the subspace to be used by the solver.

Logically collective.

**Parameters**

**ncv** (*int*) – Maximum dimension of the subspace to be used by the solver.

**Return type**

None

**See also**

[\*getDimensions\*](#), [\*LMESetDimensions\*](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:620 <slepc4py/SLEPc/LME.pyx#L620>](#)

**setErrorIfNotConverged**(*flag=True*)

Set [\*solve\(\)\*](#) to generate an error if the solver has not converged.

Logically collective.

**Parameters**

**flag** (*bool*) – True indicates you want the error generated.

**Return type**

None

**Notes**

Normally SLEPc continues if the solver fails to converge, you can call [\*getConvergedReason\(\)\*](#) after a [\*solve\(\)\*](#) to determine if it has converged.

**See also**

[\*getConvergedReason\*](#), [\*solve\*](#), [\*LMESetErrorIfNotConverged\*](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:799 <slepc4py/SLEPc/LME.pyx#L799>](#)

**setFromOptions**()

Set LME options from the options database.

Collective.

**Notes**

To see all options, run your program with the `-help` option.

This routine must be called before [\*setUp\(\)\*](#) if the user is to be allowed to set the solver type.

**See also**

[\*setOptionsPrefix\*](#), [\*LMESetFromOptions\*](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:534 <slepc4py/SLEPc/LME.pyx#L534>](#)

**Return type**

None

**setMonitor**(*monitor*, *args=None*, *kargs=None*)

Append a monitor function to the list of monitors.

Logically collective.

**See also**

[\*getMonitor\*](#), [\*cancelMonitor\*](#), [\*LMEMonitorSet\*](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:675 <slepc4py/SLEPc/LME.pyx#L675>](#)

**Parameters**

- **monitor** ([\*LMEMonitorFunction\*](#) / *None*)
- **args** ([\*tuple\*](#)[*Any*, ...] / *None*)
- **kargs** ([\*dict\*](#)[*str*, *Any*] / *None*)

**Return type**

[\*None\*](#)

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all LME options in the database.

Logically collective.

**Parameters**

**prefix** ([\*str\*](#) / *None*) – The prefix string to prepend to all LME option requests.

**Return type**

[\*None\*](#)

**Notes**

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

For example, to distinguish between the runtime options for two different LME contexts, one could call:

```
L1.setOptionsPrefix("lme1_")
L2.setOptionsPrefix("lme2_")
```

**See also**

[\*appendOptionsPrefix\*](#), [\*getOptionsPrefix\*](#), [\*LMEGetOptionsPrefix\*](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:481 <slepc4py/SLEPc/LME.pyx#L481>](#)

**setProblemType**(*lme\_problem\_type*)

Set the LME problem type of this object.

Logically collective.

**Parameters**

**lme\_problem\_type** ([\*ProblemType\*](#) / *str*) – The problem type to be used.

**Return type**

[\*None\*](#)

#### See also

[`getProblemType`](#), [`LMESetProblemType`](#)

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:188 <slepc4py/SLEPc/LME.pyx#L188>`

#### **setRHS(*C*)**

Set the right-hand side of the matrix equation.

Collective.

##### **Parameters**

**C** (*Mat*) – The right-hand side matrix

##### **Return type**

None

#### **Notes**

The matrix equation takes the general form  $AXE + DXB = C$ , where matrix  $C$  is given with this function. It must be a low-rank matrix of type `petsc4py.PETSc.Mat.Type.LRC`, that is,  $C = UDV^*$  where  $D$  is diagonal of order  $k$ , and  $U, V$  are dense tall-skinny matrices with  $k$  columns. No sparse matrix must be provided when creating the LRC matrix.

In equation types that require  $C$  to be symmetric, such as Lyapunov,  $C$  must be created with  $V = U$ .

#### See also

[`getRHS`](#), [`setSolution`](#), [`LMESetRHS`](#)

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:298 <slepc4py/SLEPc/LME.pyx#L298>`

#### **setSolution(*X=None*)**

Set the placeholder for the solution of the matrix equation.

Collective.

##### **Parameters**

**X** (*Mat* / *None*) – The solution matrix

##### **Return type**

None

#### **Notes**

The matrix equation takes the general form  $AXE + DXB = C$ , where the solution matrix is of low rank and is written in factored form  $X = UDV^*$ . This function provides a matrix object of type `petsc4py.PETSc.Mat.Type.LRC` that stores  $U, V$  and (optionally)  $D$ . These factors will be computed during [`solve\(\)`](#).

In equation types whose solution  $X$  is symmetric, such as Lyapunov,  $X$  must be created with  $V = U$ .

If the user provides  $X$  with this function, then the solver will return a solution with rank at most the number of columns of  $U$ . Alternatively, it is possible to let the solver choose the rank of the solution, by passing `None` and then calling [`getSolution\(\)`](#) after [`solve\(\)`](#).

#### See also

[solve](#), [setRHS](#), [getSolution](#), [LMESetSolution](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:347](#) <[slepc4py/SLEPc/LME.pyx#L347](#)>

**setTolerances**(*tol=None, max\_it=None*)

Set the tolerance and maximum iteration count.

Logically collective.

Set the tolerance and maximum iteration count used by the default LME convergence tests.

#### Parameters

- **tol** (*float* / *None*) – The convergence tolerance.
- **max\_it** (*int* / *None*) – The maximum number of iterations.

#### Return type

*None*

#### See also

[getTolerances](#), [LMESetTolerances](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:575](#) <[slepc4py/SLEPc/LME.pyx#L575](#)>

**setType**(*lme\_type*)

Set the particular solver to be used in the LME object.

Logically collective.

#### Parameters

- **lme\_type** (*Type* / *str*) – The solver to be used.

#### Return type

*None*

#### Notes

The default is KRYLOV. Normally, it is best to use [setFromOptions\(\)](#) and then set the LME type from the options database rather than by using this routine. Using the options database provides the user with maximum flexibility in evaluating the different available methods.

#### See also

[getType](#), [LMESetType](#)

**:sources:** [Source code at slepc4py/SLEPc/LME.pyx:142](#) <[slepc4py/SLEPc/LME.pyx#L142](#)>

**setUp()**

Set up all the internal necessary data structures.

Collective.

Set up all the internal data structures necessary for the execution of the eigensolver.

#### See also

`solve`, `LMESetUp`

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:726 <slepc4py/SLEPc/LME.pyx#L726>`

#### Return type

`None`

#### `solve()`

Solve the linear matrix equation.

Collective.

#### Notes

The matrix coefficients are specified with `setCoefficients()`. The right-hand side is specified with `setRHS()`. The placeholder for the solution is specified with `setSolution()`.

#### See also

`setCoefficients`, `setRHS`, `setSolution`, `LMESolve`

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:741 <slepc4py/SLEPc/LME.pyx#L741>`

#### Return type

`None`

#### `view(viewer=None)`

Print the LME data structure.

Collective.

#### Parameters

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

#### Return type

`None`

#### See also

`LMEView`

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:76 <slepc4py/SLEPc/LME.pyx#L76>`

#### Attributes Documentation

##### `bv`

The basis vectors (*BV*) object associated to the LME object.

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:868 <slepc4py/SLEPc/LME.pyx#L868>`

##### `fn`

The math function (*FN*) object associated to the LME object.

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:861 <slepc4py/SLEPc/LME.pyx#L861>`

**max\_it**

The maximum iteration count used by the LME convergence tests.

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:854 <slepc4py/SLEPc/LME.pyx#L854>`

**tol**

The tolerance value used by the LME convergence tests.

**:sources:** `Source code at slepc4py/SLEPc/LME.pyx:847 <slepc4py/SLEPc/LME.pyx#L847>`

## slepc4py.SLEPc.MFN

**class** slepc4py.SLEPc.MFN

Bases: `Object`

Matrix Function.

Matrix Function (*MFN*) is the object provided by slepc4py for computing the action of a matrix function on a vector. Given a matrix  $A$  and a vector  $b$ , the call `mfn.solve(b, x)` computes  $x = f(A)b$ , where  $f$  is a function such as the exponential.

## Enumerations

<i>ConvergedReason</i> Type	MFN convergence reasons. MFN type.
--------------------------------	---------------------------------------

## slepc4py.SLEPc.MFN.ConvergedReason

**class** slepc4py.SLEPc.MFN.ConvergedReason

Bases: `object`

MFN convergence reasons.

- *CONVERGED\_TOL*: All eigenpairs converged to requested tolerance.
- *CONVERGED\_ITS*: Solver completed the requested number of steps.
- *DIVERGED\_ITS*: Maximum number of iterations exceeded.
- *DIVERGED\_BREAKDOWN*: Generic breakdown in method.

### See also

`MFNConvergedReason`

## Attributes Summary

<i>CONVERGED_ITERATING</i>	Constant <i>CONVERGED_ITERATING</i> of type <code>int</code>
<i>CONVERGED_ITS</i>	Constant <i>CONVERGED_ITS</i> of type <code>int</code>
<i>CONVERGED_TOL</i>	Constant <i>CONVERGED_TOL</i> of type <code>int</code>
<i>DIVERGED_BREAKDOWN</i>	Constant <i>DIVERGED_BREAKDOWN</i> of type <code>int</code>
<i>DIVERGED_ITS</i>	Constant <i>DIVERGED_ITS</i> of type <code>int</code>
<i>ITERATING</i>	Constant <i>ITERATING</i> of type <code>int</code>

## Attributes Documentation

**CONVERGED\_ITERATING:** `int` = `CONVERGED_ITERATING`

Constant `CONVERGED_ITERATING` of type `int`

**CONVERGED\_ITS:** `int` = `CONVERGED_ITS`

Constant `CONVERGED_ITS` of type `int`

**CONVERGED\_TOL:** `int` = `CONVERGED_TOL`

Constant `CONVERGED_TOL` of type `int`

**DIVERGED\_BREAKDOWN:** `int` = `DIVERGED_BREAKDOWN`

Constant `DIVERGED_BREAKDOWN` of type `int`

**DIVERGED\_ITS:** `int` = `DIVERGED_ITS`

Constant `DIVERGED_ITS` of type `int`

**ITERATING:** `int` = `ITERATING`

Constant `ITERATING` of type `int`

## `slepc4py.SLEPc.MFN.Type`

**class** `slepc4py.SLEPc.MFN.Type`

Bases: `object`

MFN type.

- *KRYLOV*: Restarted Krylov solver.
- *EXPOKIT*: Implementation of the method in Expokit.

### See also

`MFNType`

## Attributes Summary

<i>EXPOKIT</i>	Object <code>EXPOKIT</code> of type <code>str</code>
<i>KRYLOV</i>	Object <code>KRYLOV</code> of type <code>str</code>

## Attributes Documentation

**EXPOKIT:** `str` = `EXPOKIT`

Object `EXPOKIT` of type `str`

**KRYLOV:** `str` = `KRYLOV`

Object `KRYLOV` of type `str`

## Methods Summary

<i>appendOptionsPrefix</i> ([prefix])	Append to the prefix used for searching for all MFN options in the database.
<i>cancelMonitor</i> ()	Clear all monitors for an <i>MFN</i> object.

continues on next page



Table 53 – continued from previous page

<code>create([comm])</code>	Create the MFN object.
<code>destroy()</code>	Destroy the MFN object.
<code>getBV()</code>	Get the basis vector object associated to the MFN object.
<code>getConvergedReason()</code>	Get the reason why the <code>solve()</code> iteration was stopped.
<code>getDimensions()</code>	Get the dimension of the subspace used by the solver.
<code>getErrorIfNotConverged()</code>	Get if <code>solve()</code> generates an error if the solver does not converge.
<code>getFN()</code>	Get the math function object associated to the MFN object.
<code>getIterationNumber()</code>	Get the current iteration number.
<code>getMonitor()</code>	Get the list of monitor functions.
<code>getOperator()</code>	Get the matrix associated with the MFN object.
<code>getOptionsPrefix()</code>	Get the prefix used for searching for all MFN options in the database.
<code>getTolerances()</code>	Get the tolerance and maximum iteration count.
<code>getType()</code>	Get the MFN type of this object.
<code>reset()</code>	Reset the MFN object.
<code>setBV(bv)</code>	Set a basis vector object associated to the MFN object.
<code>setDimensions(ncv)</code>	Set the dimension of the subspace to be used by the solver.
<code>setErrorIfNotConverged([flg])</code>	Set <code>solve()</code> to generate an error if the solver does not converge.
<code>setFN(fn)</code>	Set a math function object associated to the MFN object.
<code>setFromOptions()</code>	Set MFN options from the options database.
<code>setMonitor(monitor[, args, kargs])</code>	Append a monitor function to the list of monitors.
<code>setOperator(A)</code>	Set the matrix associated with the MFN object.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all MFN options in the database.
<code>setTolerances([tol, max_it])</code>	Set the tolerance and maximum iteration count.
<code>setType(mfn_type)</code>	Set the particular solver to be used in the MFN object.
<code>setUp()</code>	Set up all the necessary internal data structures.
<code>solve(b, x)</code>	Solve the matrix function problem.
<code>solveTranspose(b, x)</code>	Solve the transpose matrix function problem.
<code>view([viewer])</code>	Print the MFN data structure.

### Attributes Summary

<code>bv</code>	The basis vectors ( <i>BV</i> ) object associated to the MFN object.
<code>fn</code>	The math function ( <i>FN</i> ) object associated to the MFN object.
<code>max_it</code>	The maximum iteration count used by the MFN convergence tests.
<code>tol</code>	The tolerance count used by the MFN convergence tests.

## Methods Documentation

### **appendOptionsPrefix**(*prefix=None*)

Append to the prefix used for searching for all MFN options in the database.

Logically collective.

#### **Parameters**

**prefix** (*str* / *None*) – The prefix string to prepend to all MFN option requests.

#### **Return type**

*None*

#### **See also**

*setOptionsPrefix*, *getOptionsPrefix*, *MFNAppendOptionsPrefix*

**:sources:** [Source code at slepc4py/SLEPc/MFN.pyx:219 <slepc4py/SLEPc/MFN.pyx#L219>](#)

### **cancelMonitor**()

Clear all monitors for an *MFN* object.

Logically collective.

#### **See also**

*MFNMonitorCancel*

**:sources:** [Source code at slepc4py/SLEPc/MFN.pyx:498 <slepc4py/SLEPc/MFN.pyx#L498>](#)

#### **Return type**

*None*

### **create**(*comm=None*)

Create the MFN object.

Collective.

#### **Parameters**

**comm** (*Comm* / *None*) – MPI communicator. If not provided, it defaults to all processes.

#### **Return type**

*Self*

#### **See also**

*MFNCreate*

**:sources:** [Source code at slepc4py/SLEPc/MFN.pyx:102 <slepc4py/SLEPc/MFN.pyx#L102>](#)

### **destroy**()

Destroy the MFN object.

Logically collective.

<b>See also</b>
<a href="#">MFNDestroy</a>

**:sources:** [Source code at slepc4py/SLEPc/MFN.pyx:76 <slepc4py/SLEPc/MFN.pyx#L76>](#)

**Return type**

*Self*

**getBV()**

Get the basis vector object associated to the MFN object.

Not collective.

**Returns**

The basis vectors context.

**Return type**

*BV*

<b>See also</b>
<a href="#">setBV</a> , <a href="#">MFNGetBV</a>

**:sources:** [Source code at slepc4py/SLEPc/MFN.pyx:379 <slepc4py/SLEPc/MFN.pyx#L379>](#)

**getConvergedReason()**

Get the reason why the *solve()* iteration was stopped.

Not collective.

**Returns**

Negative value indicates diverged, positive value converged.

**Return type**

*ConvergedReason*

<b>See also</b>
<a href="#">setTolerances</a> , <a href="#">solve</a> , <a href="#">setErrorIfNotConverged</a> , <a href="#">MFNGetConvergedReason</a>

**:sources:** [Source code at slepc4py/SLEPc/MFN.pyx:607 <slepc4py/SLEPc/MFN.pyx#L607>](#)

**getDimensions()**

Get the dimension of the subspace used by the solver.

Not collective.

**Returns**

Maximum dimension of the subspace to be used by the solver.

**Return type**

*int*

**See also**

[`setDimensions`](#), [`MFNGetDimensions`](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:305 <slepc4py/SLEPc/MFN.pyx#L305>`

**getErrorIfNotConverged()**

Get if `solve()` generates an error if the solver does not converge.

Not collective.

Get a flag indicating whether `solve()` will generate an error if the solver does not converge.

**Returns**

True indicates you want the error generated.

**Return type**

`bool`

**See also**

[`setErrorIfNotConverged`](#), [`MFNGetErrorIfNotConverged`](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:650 <slepc4py/SLEPc/MFN.pyx#L650>`

**getFN()**

Get the math function object associated to the MFN object.

Not collective.

**Returns**

The math function context.

**Return type**

`FN`

**See also**

[`setFN`](#), [`MFNGetFN`](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:342 <slepc4py/SLEPc/MFN.pyx#L342>`

**getIterationNumber()**

Get the current iteration number.

Not collective.

Get the current iteration number. If the call to `solve()` is complete, then it returns the number of iterations carried out by the solution method.

**Returns**

Iteration number.

**Return type**

`int`

**See also**

[\*getConvergedReason\*](#), [\*MFNGetIterationNumber\*](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:584 <slepc4py/SLEPc/MFN.pyx#L584>`

**getMonitor()**

Get the list of monitor functions.

Not collective.

**Returns**

The list of monitor functions.

**Return type**

[\*MFNMonitorFunction\*](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:485 <slepc4py/SLEPc/MFN.pyx#L485>`

**getOperator()**

Get the matrix associated with the MFN object.

Collective.

**Returns**

The matrix for which the matrix function is to be computed.

**Return type**

[\*petsc4py.PETSc.Mat\*](#)

**See also**

[\*setOperator\*](#), [\*MFNGetOperator\*](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:416 <slepc4py/SLEPc/MFN.pyx#L416>`

**getOptionsPrefix()**

Get the prefix used for searching for all MFN options in the database.

Not collective.

**Returns**

The prefix string set for this MFN object.

**Return type**

[\*str\*](#)

**See also**

[\*setOptionsPrefix\*](#), [\*appendOptionsPrefix\*](#), [\*MFNGetOptionsPrefix\*](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:169 <slepc4py/SLEPc/MFN.pyx#L169>`

**getTolerances()**

Get the tolerance and maximum iteration count.

Not collective.

### Returns

- `tol` (`float`) – The convergence tolerance.
- `max_it` (`int`) – The maximum number of iterations.

### Return type

`tuple[float, int]`

### See also

`setTolerances`, `MFNGetTolerances`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:257 <slepc4py/SLEPc/MFN.pyx#L257>`

### `getType()`

Get the MFN type of this object.

Not collective.

### Returns

The solver currently being used.

### Return type

`str`

### See also

`setType`, `MFNGetType`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:150 <slepc4py/SLEPc/MFN.pyx#L150>`

### `reset()`

Reset the MFN object.

Collective.

### See also

`MFNReset`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:90 <slepc4py/SLEPc/MFN.pyx#L90>`

### Return type

`None`

### `setBV(bv)`

Set a basis vector object associated to the MFN object.

Collective.

### Parameters

**`bv`** (`BV`) – The basis vectors context.

### Return type

`None`

#### See also

[getBV](#), [MFNSetBV](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:399 <slepc4py/SLEPc/MFN.pyx#L399>`

#### **setDimensions**(*ncv*)

Set the dimension of the subspace to be used by the solver.

Logically collective.

##### Parameters

**ncv** (*int*) – Maximum dimension of the subspace to be used by the solver.

##### Return type

None

#### See also

[getDimensions](#), [MFNSetDimensions](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:324 <slepc4py/SLEPc/MFN.pyx#L324>`

#### **setErrorIfNotConverged**(*flag=True*)

Set [solve\(\)](#) to generate an error if the solver does not converge.

Logically collective.

##### Parameters

**flag** (*bool*) – True indicates you want the error generated.

##### Return type

None

#### Notes

Normally SLEPc continues if the solver fails to converge, you can call [getConvergedReason\(\)](#) after a [solve\(\)](#) to determine if it has converged.

#### See also

[getConvergedReason](#), [solve](#), [MFNSetErrorIfNotConverged](#)

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:626 <slepc4py/SLEPc/MFN.pyx#L626>`

#### **setFN**(*fn*)

Set a math function object associated to the MFN object.

Collective.

##### Parameters

**fn** (*FN*) – The math function context.

##### Return type

None

#### See also

[getFN](#), [MFNSetFN](#)

**:sources:** ``Source code at slepc4py/SLEPc/MFN.pyx:362 <slepc4py/SLEPc/MFN.pyx#L362>``

#### **setFromOptions()**

Set MFN options from the options database.

Collective.

#### Notes

To see all options, run your program with the `-help` option.

This routine must be called before [setUp\(\)](#) if the user is to be allowed to set the solver type.

#### See also

[setOptionsPrefix](#), [MFNSetFromOptions](#)

**:sources:** ``Source code at slepc4py/SLEPc/MFN.pyx:238 <slepc4py/SLEPc/MFN.pyx#L238>``

#### Return type

`None`

#### **setMonitor**(*monitor*, *args*=*None*, *kargs*=*None*)

Append a monitor function to the list of monitors.

Logically collective.

#### See also

[getMonitor](#), [cancelMonitor](#), [MFNMonitorSet](#)

**:sources:** ``Source code at slepc4py/SLEPc/MFN.pyx:460 <slepc4py/SLEPc/MFN.pyx#L460>``

#### Parameters

- **monitor** ([MFNMonitorFunction](#) | `None`)
- **args** ([tuple](#)[*Any*, ...] | `None`)
- **kargs** ([dict](#)[*str*, *Any*] | `None`)

#### Return type

`None`

#### **setOperator**(*A*)

Set the matrix associated with the MFN object.

Collective.

#### Parameters

**A** ([Mat](#)) – The problem matrix.

#### Return type

`None`



## Notes

This must be called before `setUp()`. If called again after `setUp()` then the *MFN* object is reset.

### See also

`getOperator`, `MFNSetOperator`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:436 <slepc4py/SLEPc/MFN.pyx#L436>`

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all MFN options in the database.

Logically collective.

### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all MFN option requests.

### Return type

*None*

## Notes

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

For example, to distinguish between the runtime options for two different MFN contexts, one could call:

```
M1.setOptionsPrefix("mfn1_")
M2.setOptionsPrefix("mfn2_")
```

### See also

`appendOptionsPrefix`, `getOptionsPrefix`, `MFNGetOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:188 <slepc4py/SLEPc/MFN.pyx#L188>`

**setTolerances**(*tol=None*, *max\_it=None*)

Set the tolerance and maximum iteration count.

Logically collective.

Set the tolerance and maximum iteration count used by the default MFN convergence tests.

### Parameters

- **tol** (*float* / *None*) – The convergence tolerance.
- **max\_it** (*int* / *None*) – The maximum number of iterations.

### Return type

*None*

### See also

`getTolerances`, `MFNSetTolerances`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:279 <slepc4py/SLEPc/MFN.pyx#L279>`

**setType**(*mfn\_type*)

Set the particular solver to be used in the MFN object.

Logically collective.

**Parameters**

**mfn\_type** (*Type* / *str*) – The solver to be used.

**Return type**

*None*

**Notes**

The default is KRYLOV. Normally, it is best to use `setFromOptions()` and then set the MFN type from the options database rather than by using this routine. Using the options database provides the user with maximum flexibility in evaluating the different available methods.

**See also**

`getType`, `MFNSetType`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:123 <slepc4py/SLEPc/MFN.pyx#L123>`

**setUp()**

Set up all the necessary internal data structures.

Collective.

Set up all the internal data structures necessary for the execution of the eigensolver.

**See also**

`solve`, `MFNSetUp`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:513 <slepc4py/SLEPc/MFN.pyx#L513>`

**Return type**

*None*

**solve**(*b*, *x*)

Solve the matrix function problem.

Collective.

Given a vector *b*, the vector  $x = f(A)b$  is returned.

**Parameters**

- **b** (*Vec*) – The right hand side vector.
- **x** (*Vec*) – The solution.

**Return type**

*None*

## Notes

The matrix  $A$  is specified with `setOperator()`. The function  $f$  is specified via the `FN` object obtained with `getFN()` or set with `setFN()`.

### See also

`setOperator`, `getFN`, `solveTranspose`, `MFNSolve`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:528 <slepc4py/SLEPc/MFN.pyx#L528>`

## `solveTranspose(b, x)`

Solve the transpose matrix function problem.

Collective.

Given a vector  $b$ , the vector  $x = f(A^T)b$  is returned.

### Parameters

- **b** (`Vec`) – The right hand side vector.
- **x** (`Vec`) – The solution.

### Return type

`None`

## Notes

The matrix  $A$  is specified with `setOperator()`. The function  $f$  is specified via the `FN` object obtained with `getFN()` or set with `setFN()`.

### See also

`setOperator`, `getFN`, `solve`, `MFNSolveTranspose`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:556 <slepc4py/SLEPc/MFN.pyx#L556>`

## `view(viewer=None)`

Print the MFN data structure.

Collective.

### Parameters

**viewer** (`Viewer` / `None`) – Visualization context; if not provided, the standard output is used.

### Return type

`None`

### See also

`MFNView`

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:57 <slepc4py/SLEPc/MFN.pyx#L57>`

## Attributes Documentation

**bv**

The basis vectors (*BV*) object associated to the MFN object.

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:695 <slepc4py/SLEPc/MFN.pyx#L695>`

**fn**

The math function (*FN*) object associated to the MFN object.

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:688 <slepc4py/SLEPc/MFN.pyx#L688>`

**max\_it**

The maximum iteration count used by the MFN convergence tests.

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:681 <slepc4py/SLEPc/MFN.pyx#L681>`

**tol**

The tolerance count used by the MFN convergence tests.

**:sources:** `Source code at slepc4py/SLEPc/MFN.pyx:674 <slepc4py/SLEPc/MFN.pyx#L674>`

## slepc4py.SLEPc.NEP

**class** slepc4py.SLEPc.NEP

Bases: `Object`

Nonlinear Eigenvalue Problem Solver.

The Nonlinear Eigenvalue Problem (*NEP*) solver is the object provided by slepc4py for specifying an eigenvalue problem that is nonlinear with respect to the eigenvalue (not the eigenvector). This is intended for general non-linear problems (rather than polynomial eigenproblems) described as  $T(\lambda)x = 0$ .

## Enumerations

<i>CISSExtraction</i>	NEP CISS extraction technique.
<i>Conv</i>	NEP convergence test.
<i>ConvergedReason</i>	NEP convergence reasons.
<i>ErrorType</i>	NEP error type to assess accuracy of computed solutions.
<i>ProblemType</i>	NEP problem type.
<i>Refine</i>	NEP refinement strategy.
<i>RefineScheme</i>	NEP scheme for solving linear systems during iterative refinement.
<i>Stop</i>	NEP stopping test.
<i>Type</i>	NEP type.
<i>Which</i>	NEP desired part of spectrum.

## slepc4py.SLEPc.NEP.CISSExtraction

**class** slepc4py.SLEPc.NEP.CISSExtraction

Bases: `object`

NEP CISS extraction technique.

- *RITZ*: Ritz extraction.
- *HANKEL*: Extraction via Hankel eigenproblem.

- [CAA](#): Communication-avoiding Arnoldi.

#### See also

[NEPCISSExtraction](#)

### Attributes Summary

<a href="#">CAA</a>	Constant CAA of type <a href="#">int</a>
<a href="#">HANKEL</a>	Constant HANKEL of type <a href="#">int</a>
<a href="#">RITZ</a>	Constant RITZ of type <a href="#">int</a>

### Attributes Documentation

**CAA:** [int](#) = CAA

Constant CAA of type [int](#)

**HANKEL:** [int](#) = HANKEL

Constant HANKEL of type [int](#)

**RITZ:** [int](#) = RITZ

Constant RITZ of type [int](#)

### slepc4py.SLEPc.NEP.Conv

**class** `slepc4py.SLEPc.NEP.Conv`

Bases: [object](#)

NEP convergence test.

- [ABS](#): Absolute convergence test.
- [REL](#): Convergence test relative to the eigenvalue.
- [NORM](#): Convergence test relative to the matrix norms.
- [USER](#): User-defined convergence test.

#### See also

[NEPConv](#)

### Attributes Summary

<a href="#">ABS</a>	Constant ABS of type <a href="#">int</a>
<a href="#">NORM</a>	Constant NORM of type <a href="#">int</a>
<a href="#">REL</a>	Constant REL of type <a href="#">int</a>
<a href="#">USER</a>	Constant USER of type <a href="#">int</a>

## Attributes Documentation

**ABS:** `int` = `ABS`

Constant ABS of type `int`

**NORM:** `int` = `NORM`

Constant NORM of type `int`

**REL:** `int` = `REL`

Constant REL of type `int`

**USER:** `int` = `USER`

Constant USER of type `int`

## slepc4py.SLEPc.NEP.ConvergedReason

**class** slepc4py.SLEPc.NEP.ConvergedReason

Bases: `object`

NEP convergence reasons.

- `CONVERGED_TOL`: All eigenpairs converged to requested tolerance.
- `CONVERGED_USER`: User-defined convergence criterion satisfied.
- `DIVERGED_ITS`: Maximum number of iterations exceeded.
- `DIVERGED_BREAKDOWN`: Solver failed due to breakdown.
- `DIVERGED_LINEAR_SOLVE`: Inner linear solve failed.
- `DIVERGED_SUBSPACE_EXHAUSTED`: Run out of space for the basis in an unrestarted solver.
- `CONVERGED_ITERATING`: Iteration not finished yet.

### See also

NEPConvergedReason

## Attributes Summary

<code>CONVERGED_ITERATING</code>	Constant <code>CONVERGED_ITERATING</code> of type <code>int</code>
<code>CONVERGED_TOL</code>	Constant <code>CONVERGED_TOL</code> of type <code>int</code>
<code>CONVERGED_USER</code>	Constant <code>CONVERGED_USER</code> of type <code>int</code>
<code>DIVERGED_BREAKDOWN</code>	Constant <code>DIVERGED_BREAKDOWN</code> of type <code>int</code>
<code>DIVERGED_ITS</code>	Constant <code>DIVERGED_ITS</code> of type <code>int</code>
<code>DIVERGED_LINEAR_SOLVE</code>	Constant <code>DIVERGED_LINEAR_SOLVE</code> of type <code>int</code>
<code>DIVERGED_SUBSPACE_EXHAUSTED</code>	Constant <code>DIVERGED_SUBSPACE_EXHAUSTED</code> of type <code>int</code>
<code>ITERATING</code>	Constant <code>ITERATING</code> of type <code>int</code>

## Attributes Documentation

**CONVERGED\_ITERATING:** `int` = `CONVERGED_ITERATING`

Constant `CONVERGED_ITERATING` of type `int`

**CONVERGED\_TOL:** `int` = **CONVERGED\_TOL**

Constant **CONVERGED\_TOL** of type `int`

**CONVERGED\_USER:** `int` = **CONVERGED\_USER**

Constant **CONVERGED\_USER** of type `int`

**DIVERGED\_BREAKDOWN:** `int` = **DIVERGED\_BREAKDOWN**

Constant **DIVERGED\_BREAKDOWN** of type `int`

**DIVERGED\_ITS:** `int` = **DIVERGED\_ITS**

Constant **DIVERGED\_ITS** of type `int`

**DIVERGED\_LINEAR\_SOLVE:** `int` = **DIVERGED\_LINEAR\_SOLVE**

Constant **DIVERGED\_LINEAR\_SOLVE** of type `int`

**DIVERGED\_SUBSPACE\_EXHAUSTED:** `int` = **DIVERGED\_SUBSPACE\_EXHAUSTED**

Constant **DIVERGED\_SUBSPACE\_EXHAUSTED** of type `int`

**ITERATING:** `int` = **ITERATING**

Constant **ITERATING** of type `int`

### **slepc4py.SLEPc.NEP.ErrorType**

**class** `slepc4py.SLEPc.NEP.ErrorType`

Bases: `object`

NEP error type to assess accuracy of computed solutions.

- *ABSOLUTE*: Absolute error.
- *RELATIVE*: Relative error.
- *BACKWARD*: Backward error.

#### **See also**

`NEPErrorType`

### **Attributes Summary**

<i>ABSOLUTE</i>	Constant <b>ABSOLUTE</b> of type <code>int</code>
<i>BACKWARD</i>	Constant <b>BACKWARD</b> of type <code>int</code>
<i>RELATIVE</i>	Constant <b>RELATIVE</b> of type <code>int</code>

### **Attributes Documentation**

**ABSOLUTE:** `int` = **ABSOLUTE**

Constant **ABSOLUTE** of type `int`

**BACKWARD:** `int` = **BACKWARD**

Constant **BACKWARD** of type `int`

**RELATIVE:** `int` = **RELATIVE**

Constant **RELATIVE** of type `int`

## slepc4py.SLEPc.NEP.ProblemType

**class** slepc4py.SLEPc.NEP.ProblemType

Bases: `object`

NEP problem type.

- *GENERAL*: General nonlinear eigenproblem.
- *RATIONAL*: NEP defined in split form with all  $f_i$  rational.

### See also

NEPProblemType

## Attributes Summary

<i>GENERAL</i>	Constant GENERAL of type <code>int</code>
<i>RATIONAL</i>	Constant RATIONAL of type <code>int</code>

## Attributes Documentation

**GENERAL:** `int` = GENERAL

Constant GENERAL of type `int`

**RATIONAL:** `int` = RATIONAL

Constant RATIONAL of type `int`

## slepc4py.SLEPc.NEP.Refine

**class** slepc4py.SLEPc.NEP.Refine

Bases: `object`

NEP refinement strategy.

- *NONE*: No refinement.
- *SIMPLE*: Refine eigenpairs one by one.
- *MULTIPLE*: Refine all eigenpairs simultaneously (invariant pair).

### See also

NEPRefine

## Attributes Summary

<i>MULTIPLE</i>	Constant MULTIPLE of type <code>int</code>
<i>NONE</i>	Constant NONE of type <code>int</code>
<i>SIMPLE</i>	Constant SIMPLE of type <code>int</code>



## Attributes Documentation

**MULTIPLE:** `int` = `MULTIPLE`

Constant `MULTIPLE` of type `int`

**NONE:** `int` = `NONE`

Constant `NONE` of type `int`

**SIMPLE:** `int` = `SIMPLE`

Constant `SIMPLE` of type `int`

## `slepc4py.SLEPc.NEP.RefineScheme`

**class** `slepc4py.SLEPc.NEP.RefineScheme`

Bases: `object`

NEP scheme for solving linear systems during iterative refinement.

- *SCHUR*: Schur complement.
- *MBE*: Mixed block elimination.
- *EXPLICIT*: Build the explicit matrix.

### See also

`NEPRefineScheme`

## Attributes Summary

<i>EXPLICIT</i>	Constant <code>EXPLICIT</code> of type <code>int</code>
<i>MBE</i>	Constant <code>MBE</code> of type <code>int</code>
<i>SCHUR</i>	Constant <code>SCHUR</code> of type <code>int</code>

## Attributes Documentation

**EXPLICIT:** `int` = `EXPLICIT`

Constant `EXPLICIT` of type `int`

**MBE:** `int` = `MBE`

Constant `MBE` of type `int`

**SCHUR:** `int` = `SCHUR`

Constant `SCHUR` of type `int`

## `slepc4py.SLEPc.NEP.Stop`

**class** `slepc4py.SLEPc.NEP.Stop`

Bases: `object`

NEP stopping test.

- *BASIC*: Default stopping test.
- *USER*: User-defined stopping test.

<b>See also</b>
NEPStop

## Attributes Summary

<i>BASIC</i>	Constant BASIC of type <code>int</code>
<i>USER</i>	Constant USER of type <code>int</code>

## Attributes Documentation

**BASIC:** `int` = BASIC

Constant BASIC of type `int`

**USER:** `int` = USER

Constant USER of type `int`

## slepc4py.SLEPc.NEP.Type

**class** slepc4py.SLEPc.NEP.Type

Bases: `object`

NEP type.

- *RII*: Residual inverse iteration.
- *SLP*: Successive linear problems.
- *NARNOLDI*: Nonlinear Arnoldi.
- *NLEIGS*: Fully rational Krylov method for nonlinear eigenproblems.
- *CISS*: Contour integral spectrum slice.
- *INTERPOL*: Polynomial interpolation.

<b>See also</b>
NEPType

## Attributes Summary

<i>CISS</i>	Object CISS of type <code>str</code>
<i>INTERPOL</i>	Object INTERPOL of type <code>str</code>
<i>NARNOLDI</i>	Object NARNOLDI of type <code>str</code>
<i>NLEIGS</i>	Object NLEIGS of type <code>str</code>
<i>RII</i>	Object RII of type <code>str</code>
<i>SLP</i>	Object SLP of type <code>str</code>

## Attributes Documentation

**CISS:** `str` = CISS

Object CISS of type `str`

**INTERPOL:** `str` = INTERPOL

Object INTERPOL of type `str`

**NARNOLDI:** `str` = NARNOLDI

Object NARNOLDI of type `str`

**NLEIGS:** `str` = NLEIGS

Object NLEIGS of type `str`

**RII:** `str` = RII

Object RII of type `str`

**SLP:** `str` = SLP

Object SLP of type `str`

## slepc4py.SLEPc.NEP.Which

**class** slepc4py.SLEPc.NEP.**Which**

Bases: `object`

NEP desired part of spectrum.

- *LARGEST\_MAGNITUDE*: Largest magnitude (default).
- *SMALLEST\_MAGNITUDE*: Smallest magnitude.
- *LARGEST\_REAL*: Largest real parts.
- *SMALLEST\_REAL*: Smallest real parts.
- *LARGEST\_IMAGINARY*: Largest imaginary parts in magnitude.
- *SMALLEST\_IMAGINARY*: Smallest imaginary parts in magnitude.
- *TARGET\_MAGNITUDE*: Closest to target (in magnitude).
- *TARGET\_REAL*: Real part closest to target.
- *TARGET\_IMAGINARY*: Imaginary part closest to target.
- *ALL*: All eigenvalues in a region.
- *USER*: User defined selection.

### See also

NEPWhich

## Attributes Summary

<i>ALL</i>	Constant ALL of type <code>int</code>
<i>LARGEST_IMAGINARY</i>	Constant LARGEST_IMAGINARY of type <code>int</code>
<i>LARGEST_MAGNITUDE</i>	Constant LARGEST_MAGNITUDE of type <code>int</code>
<i>LARGEST_REAL</i>	Constant LARGEST_REAL of type <code>int</code>

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<i>SMALLEST_IMAGINARY</i>	Constant <i>SMALLEST_IMAGINARY</i> of type <i>int</i>
<i>SMALLEST_MAGNITUDE</i>	Constant <i>SMALLEST_MAGNITUDE</i> of type <i>int</i>
<i>SMALLEST_REAL</i>	Constant <i>SMALLEST_REAL</i> of type <i>int</i>
<i>TARGET_IMAGINARY</i>	Constant <i>TARGET_IMAGINARY</i> of type <i>int</i>
<i>TARGET_MAGNITUDE</i>	Constant <i>TARGET_MAGNITUDE</i> of type <i>int</i>
<i>TARGET_REAL</i>	Constant <i>TARGET_REAL</i> of type <i>int</i>
<i>USER</i>	Constant <i>USER</i> of type <i>int</i>

### Attributes Documentation

**ALL:** *int* = *ALL*

Constant *ALL* of type *int*

**LARGEST\_IMAGINARY:** *int* = *LARGEST\_IMAGINARY*

Constant *LARGEST\_IMAGINARY* of type *int*

**LARGEST\_MAGNITUDE:** *int* = *LARGEST\_MAGNITUDE*

Constant *LARGEST\_MAGNITUDE* of type *int*

**LARGEST\_REAL:** *int* = *LARGEST\_REAL*

Constant *LARGEST\_REAL* of type *int*

**SMALLEST\_IMAGINARY:** *int* = *SMALLEST\_IMAGINARY*

Constant *SMALLEST\_IMAGINARY* of type *int*

**SMALLEST\_MAGNITUDE:** *int* = *SMALLEST\_MAGNITUDE*

Constant *SMALLEST\_MAGNITUDE* of type *int*

**SMALLEST\_REAL:** *int* = *SMALLEST\_REAL*

Constant *SMALLEST\_REAL* of type *int*

**TARGET\_IMAGINARY:** *int* = *TARGET\_IMAGINARY*

Constant *TARGET\_IMAGINARY* of type *int*

**TARGET\_MAGNITUDE:** *int* = *TARGET\_MAGNITUDE*

Constant *TARGET\_MAGNITUDE* of type *int*

**TARGET\_REAL:** *int* = *TARGET\_REAL*

Constant *TARGET\_REAL* of type *int*

**USER:** *int* = *USER*

Constant *USER* of type *int*

### Methods Summary

<i>appendOptionsPrefix</i> ([prefix])	Append to the prefix used for searching for all NEP options in the database.
<i>applyResolvent</i> (omega, v, r[, rg])	Apply the resolvent $T^{-1}(z)$ to a given vector.
<i>cancelMonitor</i> ()	Clear all monitors for a <i>NEP</i> object.
<i>computeError</i> (i[, etype])	Compute the error associated with the i-th computed eigenpair.
<i>create</i> ([comm])	Create the NEP object.
<i>destroy</i> ()	Destroy the NEP object.

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<code>errorView([etype, viewer])</code>	Display the errors associated with the computed solution.
<code>getBV()</code>	Get the basis vectors object associated to the eigensolver.
<code>getCISSExtraction()</code>	Get the extraction technique used in the CISS solver.
<code>getCISSKSPs()</code>	Get the list of linear solver objects associated with the CISS solver.
<code>getCISSRefinement()</code>	Get the values of various refinement parameters in the CISS solver.
<code>getCISSSizes()</code>	Get the values of various size parameters in the CISS solver.
<code>getCISSThreshold()</code>	Get the values of various threshold parameters in the CISS solver.
<code>getConverged()</code>	Get the number of converged eigenpairs.
<code>getConvergedReason()</code>	Get the reason why the <code>solve()</code> iteration was stopped.
<code>getConvergenceTest()</code>	Get the method used to compute the error estimate used in the convergence test.
<code>getDS()</code>	Get the direct solver associated to the eigensolver.
<code>getDimensions()</code>	Get the number of eigenvalues to compute.
<code>getEigenpair(i[, Vr, Vi])</code>	Get the i-th solution of the eigenproblem as computed by <code>solve()</code> .
<code>getEigenvalueComparison()</code>	Get the eigenvalue comparison function.
<code>getErrorEstimate(i)</code>	Get the error estimate associated to the i-th computed eigenpair.
<code>getFunction()</code>	Get the function to compute the nonlinear Function $T(\lambda)$ .
<code>getInterpolInterpolation()</code>	Get the tolerance and maximum degree for the interpolation polynomial.
<code>getInterpolPEP()</code>	Get the associated polynomial eigensolver object.
<code>getIterationNumber()</code>	Get the current iteration number.
<code>getJacobian()</code>	Get the function to compute the Jacobian $T'(\lambda)$ and $J$ .
<code>getLeftEigenvector(i, Wr[, Wi])</code>	Get the i-th left eigenvector as computed by <code>solve()</code> .
<code>getMonitor()</code>	Get the list of monitor functions.
<code>getNArnoldiKSP()</code>	Get the linear solver object associated with the nonlinear eigensolver.
<code>getNArnoldiLagPreconditioner()</code>	Get how often the preconditioner is rebuilt.
<code>getNLEIGSEPS()</code>	Get the linear eigensolver object associated with the nonlinear eigensolver.
<code>getNLEIGSFullBasis()</code>	Get the flag that indicates if NLEIGS is using the full-basis variant.
<code>getNLEIGSInterpolation()</code>	Get the tolerance and maximum degree for the interpolation polynomial.
<code>getNLEIGSKSPs()</code>	Get the list of linear solver objects associated with the NLEIGS solver.
<code>getNLEIGSLocking()</code>	Get the locking flag used in the NLEIGS method.
<code>getNLEIGSRKShifts()</code>	Get the list of shifts used in the Rational Krylov method.
<code>getNLEIGSRestart()</code>	Get the restart parameter used in the NLEIGS method.

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<code>getOptionsPrefix()</code>	Get the prefix used for searching for all NEP options in the database.
<code>getProblemType()</code>	Get the problem type from the <i>NEP</i> object.
<code>getRG()</code>	Get the region object associated to the eigensolver.
<code>getRIIConstCorrectionTol()</code>	Get the constant tolerance flag.
<code>getRIIDeflationThreshold()</code>	Get the threshold value that controls deflation.
<code>getRIIHermitian()</code>	Get if the Hermitian version must be used by the solver.
<code>getRIIKSP()</code>	Get the linear solver object associated with the non-linear eigensolver.
<code>getRIILagPreconditioner()</code>	Get how often the preconditioner is rebuilt.
<code>getRIIMaximumIterations()</code>	Get the maximum number of inner iterations of RII.
<code>getRefine()</code>	Get the refinement strategy used by the NEP object.
<code>getRefineKSP()</code>	Get the KSP object used by the eigensolver in the refinement phase.
<code>getSLPDeflationThreshold()</code>	Get the threshold value that controls deflation.
<code>getSLPEPS()</code>	Get the linear eigensolver object associated with the nonlinear eigensolver.
<code>getSLPEPSLeft()</code>	Get the left eigensolver.
<code>getSLPKSP()</code>	Get the linear solver object associated with the non-linear eigensolver.
<code>getSplitOperator()</code>	Get the operator of the nonlinear eigenvalue problem in split form.
<code>getSplitPreconditioner()</code>	Get the operator of the split preconditioner.
<code>getStoppingTest()</code>	Get the stopping test function.
<code>getTarget()</code>	Get the value of the target.
<code>getTolerances()</code>	Get the tolerance and maximum iteration count.
<code>getTrackAll()</code>	Get the flag indicating whether all residual norms must be computed.
<code>getTwoSided()</code>	Get the flag indicating if a two-sided variant is being used.
<code>getType()</code>	Get the NEP type of this object.
<code>getWhichEigenpairs()</code>	Get which portion of the spectrum is to be sought.
<code>reset()</code>	Reset the NEP object.
<code>setBV(bv)</code>	Set the basis vectors object associated to the eigensolver.
<code>setCISSExtraction(extraction)</code>	Set the extraction technique used in the CISS solver.
<code>setCISSRefinement([inner, bsize])</code>	Set the values of various refinement parameters in the CISS solver.
<code>setCISSSizes([ip, bs, ms, npart, bsmx, ...])</code>	Set the values of various size parameters in the CISS solver.
<code>setCISSThreshold([delta, spur])</code>	Set the values of various threshold parameters in the CISS solver.
<code>setConvergenceTest(conv)</code>	Set how to compute the error estimate used in the convergence test.
<code>setDS(ds)</code>	Set a direct solver object associated to the eigensolver.
<code>setDimensions([nev, ncv, mpd])</code>	Set the number of eigenvalues to compute.
<code>setEigenvalueComparison(comparison[, args, ...])</code>	Set an eigenvalue comparison function.
<code>setFromOptions()</code>	Set NEP options from the options database.
<code>setFunction(function[, F, P, args, kargs])</code>	Set the function to compute the nonlinear Function $T(\lambda)$ .

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<i>setInitialSpace</i> (space)	Set the initial space from which the eigensolver starts to iterate.
<i>setInterpolInterpolation</i> ([tol, deg])	Set the tolerance and maximum degree for the interpolation polynomial.
<i>setInterpolPEP</i> (pep)	Set a polynomial eigensolver object associated to the nonlinear eigensolver.
<i>setJacobian</i> (jacobian[, J, args, kargs])	Set the function to compute the Jacobian $T'(\lambda)$ .
<i>setMonitor</i> (monitor[, args, kargs])	Append a monitor function to the list of monitors.
<i>setArnoldiKSP</i> (ksp)	Set a linear solver object associated to the nonlinear eigensolver.
<i>setArnoldiLagPreconditioner</i> (lag)	Set when the preconditioner is rebuilt in the nonlinear solve.
<i>setNLEIGSEPS</i> (eps)	Set a linear eigensolver object associated to the nonlinear eigensolver.
<i>setNLEIGSFullBasis</i> ([fullbasis])	Set TOAR-basis (default) or full-basis variants of the NLEIGS method.
<i>setNLEIGSInterpolation</i> ([tol, deg])	Set the tolerance and maximum degree for the interpolation polynomial.
<i>setNLEIGSLocking</i> (lock)	Toggle between locking and non-locking variants of the NLEIGS method.
<i>setNLEIGSRKShifts</i> (shifts)	Set a list of shifts to be used in the Rational Krylov method.
<i>setNLEIGSRestart</i> (keep)	Set the restart parameter for the NLEIGS method.
<i>setOptionsPrefix</i> ([prefix])	Set the prefix used for searching for all NEP options in the database.
<i>setProblemType</i> (problem_type)	Set the type of the eigenvalue problem.
<i>setRG</i> (rg)	Set a region object associated to the eigensolver.
<i>setRIIConstCorrectionTol</i> (cct)	Set a flag to keep the tolerance used in the linear solver constant.
<i>setRIIDeflationThreshold</i> (deftol)	Set the threshold used to switch between deflated and non-deflated.
<i>setRIIHermitian</i> (herm)	Set a flag to use the Hermitian version of the solver.
<i>setRIIKSP</i> (ksp)	Set a linear solver object associated to the nonlinear eigensolver.
<i>setRIILagPreconditioner</i> (lag)	Set when the preconditioner is rebuilt in the nonlinear solve.
<i>setRIIMaximumIterations</i> (its)	Set the max.
<i>setRefine</i> (ref[, npart, tol, its, scheme])	Set the refinement strategy used by the NEP object.
<i>setSLPDeflationThreshold</i> (deftol)	Set the threshold used to switch between deflated and non-deflated.
<i>setSLPEPS</i> (eps)	Set a linear eigensolver object associated to the nonlinear eigensolver.
<i>setSLPEPSLeft</i> (eps)	Set a linear eigensolver object associated to the nonlinear eigensolver.
<i>setSLPKSP</i> (ksp)	Set a linear solver object associated to the nonlinear eigensolver.
<i>setSplitOperator</i> (A, f[, structure])	Set the operator of the nonlinear eigenvalue problem in split form.
<i>setSplitPreconditioner</i> (P[, structure])	Set the operator in split form.
<i>setStoppingTest</i> (stopping[, args, kargs])	Set a function to decide when to stop the outer iteration of the eigensolver.
<i>setTarget</i> (target)	Set the value of the target.

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<code>setTolerances([tol, maxit])</code>	Set the tolerance and max.
<code>setTrackAll(trackall)</code>	Set if the solver must compute the residual of all approximate eigenpairs.
<code>setTwoSided(twosided)</code>	Set the solver to use a two-sided variant.
<code>setType(nep_type)</code>	Set the particular solver to be used in the NEP object.
<code>setUp()</code>	Set up all the internal data structures.
<code>setWhichEigenpairs(which)</code>	Set which portion of the spectrum is to be sought.
<code>solve()</code>	Solve the nonlinear eigenproblem.
<code>valuesView([viewer])</code>	Display the computed eigenvalues in a viewer.
<code>vectorsView([viewer])</code>	Output computed eigenvectors to a viewer.
<code>view([viewer])</code>	Print the NEP data structure.

## Attributes Summary

<code>bv</code>	The basis vectors ( <i>BV</i> ) object associated.
<code>ds</code>	The direct solver ( <i>DS</i> ) object associated.
<code>max_it</code>	The maximum iteration count used by the NEP convergence tests.
<code>problem_type</code>	The problem type from the NEP object.
<code>rg</code>	The region ( <i>RG</i> ) object associated.
<code>target</code>	The value of the target.
<code>tol</code>	The tolerance used by the NEP convergence tests.
<code>track_all</code>	Compute the residual of all approximate eigenpairs.
<code>which</code>	The portion of the spectrum to be sought.

## Methods Documentation

### `appendOptionsPrefix(prefix=None)`

Append to the prefix used for searching for all NEP options in the database.

Logically collective.

#### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all NEP option requests.

#### Return type

*None*

#### See also

`setOptionsPrefix`, `getOptionsPrefix`, `NEPAppendOptionsPrefix`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:385 <slepc4py/SLEPc/NEP.pyx#L385>``

### `applyResolvent(omega, v, r, rg=None)`

Apply the resolvent  $T^{-1}(z)$  to a given vector.

Collective.

#### Parameters

- **omega** (*Scalar*) – Value where the resolvent must be evaluated.
- **v** (*Vec*) – Input vector.



- **r** ([Vec](#)) – Placeholder for the result vector.
- **rg** ([RG](#) / [None](#)) – Region.

**Return type**

[None](#)

**Notes**

The resolvent  $T^{-1}(z) = \sum_i (z - \lambda_i)^{-1} x_i y_i^*$  is evaluated at  $z = \omega$  and the matrix-vector product  $r = T^{-1}(\omega)v$  is computed. Vectors  $x_i, y_i$  are right and left eigenvectors, respectively, normalized so that  $y_i^* T'(\lambda_i) x_i = 1$ . The sum contains only eigenvectors that have been previously computed with [solve\(\)](#), and if a region **rg** is given then only those corresponding to eigenvalues inside the region are considered.

**See also**

[solve](#), [getLeftEigenvector](#), [NEPApplyResolvent](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1834](#) <[slepc4py/SLEPc/NEP.pyx#L1834](#)>

**cancelMonitor()**

Clear all monitors for a [NEP](#) object.

Logically collective.

**See also**

[NEPMonitorCancel](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1159](#) <[slepc4py/SLEPc/NEP.pyx#L1159](#)>

**Return type**

[None](#)

**computeError(i, etype=None)**

Compute the error associated with the i-th computed eigenpair.

Collective.

Compute the error (based on the residual norm) associated with the i-th computed eigenpair.

**Parameters**

- **i** ([int](#)) – Index of the solution to be considered.
- **etype** ([ErrorType](#) / [None](#)) – The error type to compute.

**Returns**

The error bound, computed in various ways from the residual norm  $\|T(\lambda)x\|_2$  where  $\lambda$  is the eigenvalue and  $x$  is the eigenvector.

**Return type**

[float](#)

## Notes

The index `i` should be a value between 0 and `nconv-1` (see `getConverged()`).

If the computation of left eigenvectors was enabled with `setTwoSided()`, then the error will be computed using the maximum of the value above and the left residual norm  $\|y^*T(\lambda)\|_2$ , where  $y$  is the approximate left eigenvector.

### See also

`getErrorEstimate`, `setTwoSided`, `NEPComputeError`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1385 <slepc4py/SLEPc/NEP.pyx#L1385>`

**create**(*comm=None*)

Create the NEP object.

Collective.

#### Parameters

**comm** (*Comm* / *None*) – MPI communicator. If not provided, it defaults to all processes.

#### Return type

*Self*

### See also

`NEPCreate`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:268 <slepc4py/SLEPc/NEP.pyx#L268>`

**destroy**()

Destroy the NEP object.

Collective.

### See also

`NEPDestroy`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:242 <slepc4py/SLEPc/NEP.pyx#L242>`

#### Return type

*Self*

**errorView**(*etype=None*, *viewer=None*)

Display the errors associated with the computed solution.

Collective.

Display the errors and the eigenvalues.

#### Parameters

- **etype** (*ErrorType* / *None*) – The error type to compute.
- **viewer** (*petsc4py.PETSc.Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

None

**Notes**

By default, this function checks the error of all eigenpairs and prints the eigenvalues if all of them are below the requested tolerance. If the viewer has format ASCII\_INFO\_DETAIL then a table with eigenvalues and corresponding errors is printed.

**See also**

*solve*, *valuesView*, *vectorsView*, *NEPErrorView*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1428 <slepc4py/SLEPc/NEP.pyx#L1428>](#)

**getBV()**

Get the basis vectors object associated to the eigensolver.

Not collective.

**Returns**

The basis vectors context.

**Return type**

*BV*

**See also**

*setBV*, *NEPGetBV*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:877 <slepc4py/SLEPc/NEP.pyx#L877>](#)

**getCISSExtraction()**

Get the extraction technique used in the CISS solver.

Not collective.

**Returns**

The extraction technique.

**Return type**

*CISSExtraction*

**See also**

*setCISSExtraction*, *NEPCISSGetExtraction*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2792 <slepc4py/SLEPc/NEP.pyx#L2792>](#)

**getCISSKSPs()**

Get the list of linear solver objects associated with the CISS solver.

Collective.

**Returns**

The linear solver objects.

**Return type**

list of `petsc4py.PETSc.KSP`

**Notes**

The number of `petsc4py.PETSc.KSP` solvers is equal to the number of integration points divided by the number of partitions. This value is halved in the case of real matrices with a region centered at the real axis.

**See also**

`setCISSSizes`, `NEPCISSGetKSPs`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2989 <slepc4py/SLEPc/NEP.pyx#L2989>`

**getCISSRefinement()**

Get the values of various refinement parameters in the CISS solver.

Not collective.

**Returns**

- **inner** (`int`) – Number of iterative refinement iterations (inner loop).
- **bsize** (`int`) – Number of iterative refinement iterations (blocksize loop).

**Return type**

tuple[int, int]

**See also**

`setCISSRefinement`, `NEPCISSGetRefinement`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2967 <slepc4py/SLEPc/NEP.pyx#L2967>`

**getCISSSizes()**

Get the values of various size parameters in the CISS solver.

Not collective.

**Returns**

- **ip** (`int`) – Number of integration points.
- **bs** (`int`) – Block size.
- **ms** (`int`) – Moment size.
- **npart** (`int`) – Number of partitions when splitting the communicator.
- **bsmax** (`int`) – Maximum block size.
- **realmats** (`bool`) – True if A and B are real.

**Return type**

tuple[int, int, int, int, int, bool]

**See also**

[`setCISSizes`](#), [`NEPCISSGetSizes`](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2865](#) <[slepc4py/SLEPc/NEP.pyx#L2865](#)>

**getCISSThreshold()**

Get the values of various threshold parameters in the CISS solver.

Not collective.

**Returns**

- **delta** ([`float`](#)) – Threshold for numerical rank.
- **spur** ([`float`](#)) – Spurious threshold (to discard spurious eigenpairs).

**Return type**

[`tuple`](#)[[`float`](#), [`float`](#)]

**See also**

[`setCISSThreshold`](#), [`NEPCISSGetThreshold`](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2922](#) <[slepc4py/SLEPc/NEP.pyx#L2922](#)>

**getConverged()**

Get the number of converged eigenpairs.

Not collective.

**Returns**

**nconv** – Number of converged eigenpairs.

**Return type**

[`int`](#)

**Notes**

This function should be called after [`solve\(\)`](#) has finished.

The value **nconv** may be different from the number of requested solutions **nev**, but not larger than **ncv**, see [`setDimensions\(\)`](#).

**See also**

[`setDimensions`](#), [`solve`](#), [`getEigenpair`](#), [`NEPGetConverged`](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1256](#) <[slepc4py/SLEPc/NEP.pyx#L1256](#)>

**getConvergedReason()**

Get the reason why the [`solve\(\)`](#) iteration was stopped.

Not collective.

**Returns**

Negative value indicates diverged, positive value converged.

**Return type**  
*ConvergedReason*

**See also**

*setTolerances*, *solve*, *NEPGetConvergedReason*

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1237 <slepc4py/SLEPc/NEP.pyx#L1237>`

**getConvergenceTest()**

Get the method used to compute the error estimate used in the convergence test.

Not collective.

**Returns**

The method used to compute the error estimate used in the convergence test.

**Return type**

*Conv*

**See also**

*setConvergenceTest*, *NEPGetConvergenceTest*

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:624 <slepc4py/SLEPc/NEP.pyx#L624>`

**getDS()**

Get the direct solver associated to the eigensolver.

Not collective.

**Returns**

The direct solver context.

**Return type**

*DS*

**See also**

*setDS*, *NEPGetDS*

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:951 <slepc4py/SLEPc/NEP.pyx#L951>`

**getDimensions()**

Get the number of eigenvalues to compute.

Not collective.

Get the number of eigenvalues to compute, and the dimension of the subspace.

**Returns**

- **nev** (*int*) – Number of eigenvalues to compute.
- **ncv** (*int*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int*) – Maximum dimension allowed for the projected problem.

**Return type**  
tuple[int, int, int]

**See also**

[setDimensions](#), [NEPGetDimensions](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:795 <slepc4py/SLEPc/NEP.pyx#L795>`

**getEigenpair**(*i*, *Vr=None*, *Vi=None*)

Get the *i*-th solution of the eigenproblem as computed by [solve\(\)](#).

Collective.

The solution consists of both the eigenvalue and the eigenvector.

**Parameters**

- **i** (*int*) – Index of the solution to be obtained.
- **Vr** (*Vec* / *None*) – Placeholder for the returned eigenvector (real part).
- **Vi** (*Vec* / *None*) – Placeholder for the returned eigenvector (imaginary part).

**Returns**

The computed eigenvalue.

**Return type**

complex

**Notes**

The index *i* should be a value between 0 and `nconv-1` (see [getConverged\(\)](#)). Eigenpairs are indexed according to the ordering criterion established with [setWhichEigenpairs\(\)](#).

The eigenvector is normalized to have unit norm.

**See also**

[solve](#), [getConverged](#), [setWhichEigenpairs](#), [NEPGetEigenpair](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1282 <slepc4py/SLEPc/NEP.pyx#L1282>`

**getEigenvalueComparison**()

Get the eigenvalue comparison function.

Not collective.

**Returns**

The eigenvalue comparison function.

**Return type**

[NEPEigenvalueComparison](#)

**See also**

[setEigenvalueComparison](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1100 <slepc4py/SLEPc/NEP.pyx#L1100>`

### **getErrorEstimate(*i*)**

Get the error estimate associated to the *i*-th computed eigenpair.

Not collective.

#### **Parameters**

**i** (*int*) – Index of the solution to be considered.

#### **Returns**

Error estimate.

#### **Return type**

*float*

### **Notes**

This is the error estimate used internally by the eigensolver. The actual error bound can be computed with *computeError()*.

#### **See also**

*computeError*, *NEPGetErrorEstimate*

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1356 <slepc4py/SLEPc/NEP.pyx#L1356>`

### **getFunction()**

Get the function to compute the nonlinear Function  $T(\lambda)$ .

Collective.

Get the function to compute the nonlinear Function  $T(\lambda)$  and the matrix.

#### **Returns**

- **F** (*petsc4py.PETSc.Mat*) – Function matrix.
- **P** (*petsc4py.PETSc.Mat*) – Preconditioner matrix (usually the same as the F).
- **function** (*NEPFunction*) – Function evaluation routine.

#### **Return type**

*tuple*[*petsc4py.PETSc.Mat*, *petsc4py.PETSc.Mat*, *NEPFunction*]

#### **See also**

*setFunction*, *NEPGetFunction*

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1540 <slepc4py/SLEPc/NEP.pyx#L1540>`

### **getInterpolInterpolation()**

Get the tolerance and maximum degree for the interpolation polynomial.

Not collective.

#### **Returns**

- **tol** (*float*) – The tolerance to stop computing polynomial coefficients.
- **deg** (*int*) – The maximum degree of interpolation.



**Return type**  
`tuple[float, int]`

**See also**

`setInterpolInterpolation`, `NEPInterpolGetInterpolation`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2444 <slepc4py/SLEPc/NEP.pyx#L2444>`

**getInterpolPEP()**

Get the associated polynomial eigensolver object.

Collective.

**Returns**  
The polynomial eigensolver.

**Return type**  
`PEP`

**See also**

`setInterpolPEP`, `NEPInterpolGetPEP`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2401 <slepc4py/SLEPc/NEP.pyx#L2401>`

**getIterationNumber()**

Get the current iteration number.

Not collective.

If the call to `solve()` is complete, then it returns the number of iterations carried out by the solution method.

**Returns**  
Iteration number.

**Return type**  
`int`

**See also**

`getConvergedReason`, `setTolerances`, `NEPGetIterationNumber`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1215 <slepc4py/SLEPc/NEP.pyx#L1215>`

**getJacobian()**

Get the function to compute the Jacobian  $T'(\lambda)$  and  $J$ .

Collective.

Get the function to compute the Jacobian  $T'(\lambda)$  and the matrix.

**Returns**

- `J` (`petsc4py.PETSc.Mat`) – Jacobian matrix.
- `jacobian` (`NEPJacobian`) – Jacobian evaluation routine.

**Return type**

tuple[petsc4py.PETSc.Mat, *NEPJacobian*]

**See also**

*setJacobian*, *NEPGetJacobian*

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1606 <slepc4py/SLEPc/NEP.pyx#L1606>`

**getLeftEigenvector**(*i*, *Wr*, *Wi=None*)

Get the *i*-th left eigenvector as computed by *solve()*.

Collective.

**Parameters**

- **i** (*int*) – Index of the solution to be obtained.
- **Wr** (*Vec*) – Placeholder for the returned eigenvector (real part).
- **Wi** (*Vec* / *None*) – Placeholder for the returned eigenvector (imaginary part).

**Return type**

*None*

**Notes**

The index *i* should be a value between 0 and *nconv*-1 (see *getConverged()*). Eigensolutions are indexed according to the ordering criterion established with *setWhichEigenpairs()*.

Left eigenvectors are available only if the *twosided* flag was set with *setTwoSided()*.

**See also**

*getEigenpair*, *getConverged*, *setTwoSided*, *NEPGetLeftEigenvector*

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1323 <slepc4py/SLEPc/NEP.pyx#L1323>`

**getMonitor()**

Get the list of monitor functions.

Not collective.

**Returns**

The list of monitor functions.

**Return type**

*NEPMonitorFunction*

**See also**

*setMonitor*

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1142 <slepc4py/SLEPc/NEP.pyx#L1142>`

### **getNArnoldiKSP()**

Get the linear solver object associated with the nonlinear eigensolver.

Collective.

#### **Returns**

The linear solver object.

#### **Return type**

`petsc4py.PETSc.KSP`

#### **See also**

`setNArnoldiKSP`, `NEPNArnoldiGetKSP`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2318 <slepc4py/SLEPc/NEP.pyx#L2318>`

### **getNArnoldiLagPreconditioner()**

Get how often the preconditioner is rebuilt.

Not collective.

#### **Returns**

The lag parameter.

#### **Return type**

`int`

#### **See also**

`setNArnoldiLagPreconditioner`, `NEPNArnoldiGetLagPreconditioner`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2363 <slepc4py/SLEPc/NEP.pyx#L2363>`

### **getNLEIGSEPS()**

Get the linear eigensolver object associated with the nonlinear eigensolver.

Collective.

#### **Returns**

The linear eigensolver.

#### **Return type**

`EPS`

#### **See also**

`setNLEIGSEPS`, `NEPNLEIGSGetEPS`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2675 <slepc4py/SLEPc/NEP.pyx#L2675>`

### **getNLEIGSFULLBASIS()**

Get the flag that indicates if NLEIGS is using the full-basis variant.

Not collective.

**Returns**

True if the full-basis variant is selected.

**Return type**

bool

**See also**

[\*setNLEIGSFULLBasis\*](#), [\*NEPNLEIGSGetFullBasis\*](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2639](#) <[slepc4py/SLEPc/NEP.pyx#L2639](#)>

**getNLEIGSInterpolation()**

Get the tolerance and maximum degree for the interpolation polynomial.

Not collective.

Get the tolerance and maximum degree when building the interpolation via divided differences.

**Returns**

- **tol** (float) – The tolerance to stop computing divided differences.
- **deg** (int) – The maximum degree of interpolation.

**Return type**

tuple[float, int]

**See also**

[\*setNLEIGSInterpolation\*](#), [\*NEPNLEIGSGetInterpolation\*](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2581](#) <[slepc4py/SLEPc/NEP.pyx#L2581](#)>

**getNLEIGSKSPs()**

Get the list of linear solver objects associated with the NLEIGS solver.

Collective.

**Returns**

The linear solver objects.

**Return type**

list of `petsc4py.PETSc.KSP`

**Notes**

The number of `petsc4py.PETSc.KSP` solvers is equal to the number of shifts provided by the user, or 1 if the user did not provide shifts.

**See also**

[\*setNLEIGSRKShifts\*](#), [\*NEPNLEIGSGetKSPs\*](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2747](#) <[slepc4py/SLEPc/NEP.pyx#L2747](#)>

### **getNLEIGSLocking()**

Get the locking flag used in the NLEIGS method.

Not collective.

#### **Returns**

The locking flag.

#### **Return type**

*bool*

#### **See also**

*setNLEIGSLocking*, *NEPNLEIGSGetLocking*

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:2536 <slepc4py/SLEPc/NEP.pyx#L2536>``

### **getNLEIGSRKShifts()**

Get the list of shifts used in the Rational Krylov method.

Not collective.

#### **Returns**

The shift values.

#### **Return type**

*ArrayScalar*

#### **See also**

*setNLEIGSRKShifts*, *NEPNLEIGSGetRKShifts*

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:2722 <slepc4py/SLEPc/NEP.pyx#L2722>``

### **getNLEIGSRestart()**

Get the restart parameter used in the NLEIGS method.

Not collective.

#### **Returns**

The number of vectors to be kept at restart.

#### **Return type**

*float*

#### **See also**

*setNLEIGSRestart*, *NEPNLEIGSGetRestart*

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:2492 <slepc4py/SLEPc/NEP.pyx#L2492>``

### **getOptionsPrefix()**

Get the prefix used for searching for all NEP options in the database.

Not collective.

**Returns**

The prefix string set for this NEP object.

**Return type**

`str`

**See also**

`setOptionsPrefix`, `appendOptionsPrefix`, `NEPGetOptionsPrefix`

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:335 <slepc4py/SLEPc/NEP.pyx#L335>](#)

**getProblemType()**

Get the problem type from the *NEP* object.

Not collective.

**Returns**

The problem type that was previously set.

**Return type**

*ProblemType*

**See also**

`setProblemType`, `NEPGetProblemType`

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:423 <slepc4py/SLEPc/NEP.pyx#L423>](#)

**getRG()**

Get the region object associated to the eigensolver.

Not collective.

**Returns**

The region context.

**Return type**

*RG*

**See also**

`setRG`, `NEPGetRG`

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:914 <slepc4py/SLEPc/NEP.pyx#L914>](#)

**getRIIConstCorrectionTol()**

Get the constant tolerance flag.

Not collective.

**Returns**

If `True`, the `petsc4py.PETSc.KSP` relative tolerance is constant.

**Return type**

`bool`

**See also**

`setRIIConstCorrectionTol`, `NEPRIIGetConstCorrectionTol`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:1942 <slepc4py/SLEPc/NEP.pyx#L1942>``

**getRIIDeflationThreshold()**

Get the threshold value that controls deflation.

Not collective.

**Returns**

The threshold value.

**Return type**

`float`

**See also**

`setRIIDeflationThreshold`, `NEPRIIGetDeflationThreshold`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:2081 <slepc4py/SLEPc/NEP.pyx#L2081>``

**getRIIHermitian()**

Get if the Hermitian version must be used by the solver.

Not collective.

**Returns**

If True, the Hermitian version is used.

**Return type**

`bool`

**See also**

`setRIIHermitian`, `NEPRIIGetHermitian`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:2031 <slepc4py/SLEPc/NEP.pyx#L2031>``

**getRIIKSP()**

Get the linear solver object associated with the nonlinear eigensolver.

Collective.

**Returns**

The linear solver object.

**Return type**

`petsc4py.PETSc.KSP`

**See also**

`setRIIKSP`, `NEPRIIGetKSP`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:2117 <slepc4py/SLEPc/NEP.pyx#L2117>``

### **getRIILagPreconditioner()**

Get how often the preconditioner is rebuilt.

Not collective.

#### **Returns**

The lag parameter.

#### **Return type**

`int`

#### **See also**

`setRIILagPreconditioner`, `NEPRIIGetLagPreconditioner`

**`:sources:`** `Source code at slepc4py/SLEPc/NEP.pyx:1898 <slepc4py/SLEPc/NEP.pyx#L1898>`

### **getRIIMaximumIterations()**

Get the maximum number of inner iterations of RII.

Not collective.

#### **Returns**

Maximum inner iterations.

#### **Return type**

`int`

#### **See also**

`setRIIMaximumIterations`, `NEPRIIGetMaximumIterations`

**`:sources:`** `Source code at slepc4py/SLEPc/NEP.pyx:1983 <slepc4py/SLEPc/NEP.pyx#L1983>`

### **getRefine()**

Get the refinement strategy used by the NEP object.

Not collective.

#### **Returns**

- **ref** (*Refine*) – The refinement type.
- **npart** (`int`) – The number of partitions of the communicator.
- **tol** (`float`) – The convergence tolerance.
- **its** (`int`) – The maximum number of refinement iterations.
- **scheme** (*RefineScheme*) – Scheme for solving linear systems.

#### **Return type**

`tuple[Refine, int, float, int, RefineScheme]`

#### **See also**

`setRefine`, `NEPGetRefine`

**`:sources:`** `Source code at slepc4py/SLEPc/NEP.pyx:663 <slepc4py/SLEPc/NEP.pyx#L663>`



### **getRefineKSP()**

Get the KSP object used by the eigensolver in the refinement phase.

Collective.

#### **Returns**

The linear solver object.

#### **Return type**

`petsc4py.PETSc.KSP`

#### **See also**

`setRefine`, `NEPRefineGetKSP`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:738 <slepc4py/SLEPc/NEP.pyx#L738>`

### **getSLPDeflationThreshold()**

Get the threshold value that controls deflation.

Not collective.

#### **Returns**

The threshold value.

#### **Return type**

`float`

#### **See also**

`setSLPDeflationThreshold`, `NEPSLPGetDeflationThreshold`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2167 <slepc4py/SLEPc/NEP.pyx#L2167>`

### **getSLPEPS()**

Get the linear eigensolver object associated with the nonlinear eigensolver.

Collective.

#### **Returns**

The linear eigensolver.

#### **Return type**

`EPS`

#### **See also**

`setSLPEPS`, `NEPSLPGetEPS`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2203 <slepc4py/SLEPc/NEP.pyx#L2203>`

### **getSLPEPSLeft()**

Get the left eigensolver.

Collective.

**Returns**

The linear eigensolver.

**Return type**

*EPS*

**See also**

*setSLPEPSLeft*, *NEPSLPGetEPSLeft*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2242](#) <[slepc4py/SLEPc/NEP.pyx#L2242](#)>

**getSLPKSP()**

Get the linear solver object associated with the nonlinear eigensolver.

Collective.

**Returns**

The linear solver object.

**Return type**

*petsc4py.PETSc.KSP*

**See also**

*setSLPKSP*, *NEPSLPGetKSP*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2279](#) <[slepc4py/SLEPc/NEP.pyx#L2279](#)>

**getSplitOperator()**

Get the operator of the nonlinear eigenvalue problem in split form.

Collective.

**Returns**

- **A** (list of *petsc4py.PETSc.Mat*) – Coefficient matrices of the split form.
- **f** (list of *FN*) – Scalar functions of the split form.
- **structure** (*petsc4py.PETSc.Mat.Structure*) – Structure flag for matrices.

**Return type**

*tuple*[list[*petsc4py.PETSc.Mat*], list[*FN*], *petsc4py.PETSc.Mat.Structure*]

**See also**

*setSplitOperator*, *NEPGetSplitOperatorInfo*, *NEPGetSplitOperatorTerm*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1683](#) <[slepc4py/SLEPc/NEP.pyx#L1683](#)>

**getSplitPreconditioner()**

Get the operator of the split preconditioner.

Not collective.

**Returns**

- **P** (list of `petsc4py.PETSc.Mat`) – Coefficient matrices of the split preconditioner.
- **structure** (`petsc4py.PETSc.Mat.Structure`) – Structure flag for matrices.

**Return type**

`tuple[list[petsc4py.PETSc.Mat], petsc4py.PETSc.Mat.Structure]`

**See also**

`setSplitPreconditioner`, `NEPGetSplitPreconditionerTerm`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:1752 <slepc4py/SLEPc/NEP.pyx#L1752>``

**getStoppingTest()**

Get the stopping test function.

Not collective.

**Returns**

The stopping test function.

**Return type**

*NEPStoppingFunction*

**See also**

`setStoppingTest`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:1053 <slepc4py/SLEPc/NEP.pyx#L1053>``

**getTarget()**

Get the value of the target.

Not collective.

**Returns**

The value of the target.

**Return type**

*Scalar*

**Notes**

If the target was not set by the user, then zero is returned.

**See also**

`setTarget`, `NEPGetTarget`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:521 <slepc4py/SLEPc/NEP.pyx#L521>``

**getTolerances()**

Get the tolerance and maximum iteration count.

Not collective.

Get the tolerance and maximum iteration count used by the default NEP convergence tests.

### Returns

- **tol** (`float`) – The convergence tolerance.
- **maxit** (`int`) – The maximum number of iterations.

### Return type

`tuple[float, int]`

### See also

`setTolerances`, `NEPGetTolerances`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:571 <slepc4py/SLEPc/NEP.pyx#L571>`

### `getTrackAll()`

Get the flag indicating whether all residual norms must be computed.

Not collective.

### Returns

Whether the solver computes all residuals or not.

### Return type

`bool`

### See also

`setTrackAll`, `NEPGetTrackAll`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:758 <slepc4py/SLEPc/NEP.pyx#L758>`

### `getTwoSided()`

Get the flag indicating if a two-sided variant is being used.

Not collective.

Get the flag indicating whether a two-sided variant of the algorithm is being used or not.

### Returns

Whether the two-sided variant is to be used or not.

### Return type

`bool`

### See also

`setTwoSided`, `NEPGetTwoSided`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1781 <slepc4py/SLEPc/NEP.pyx#L1781>`

### `getType()`

Get the NEP type of this object.

Not collective.

### Returns

The solver currently being used.

**Return type**

`str`

**See also**

`setType`, `NEPGetType`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:316 <slepc4py/SLEPc/NEP.pyx#L316>``

**getWhichEigenpairs()**

Get which portion of the spectrum is to be sought.

Not collective.

**Returns**

The portion of the spectrum to be sought by the solver.

**Return type**

`Which`

**See also**

`setWhichEigenpairs`, `NEPGetWhichEigenpairs`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:467 <slepc4py/SLEPc/NEP.pyx#L467>``

**reset()**

Reset the NEP object.

Collective.

**See also**

`NEPReset`

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:256 <slepc4py/SLEPc/NEP.pyx#L256>``

**Return type**

`None`

**setBV(bv)**

Set the basis vectors object associated to the eigensolver.

Collective.

**Parameters**

**bv** (`BV`) – The basis vectors context.

**Return type**

`None`

**See also**

`getBV`, `NEPSetBV`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:897 <slepc4py/SLEPc/NEP.pyx#L897>`

**setCISSExtraction**(*extraction*)

Set the extraction technique used in the CISS solver.

Logically collective.

**Parameters**

**extraction** (`CISSExtraction`) – The extraction technique.

**Return type**

`None`

**See also**

`getCISSExtraction`, `NEPCISSSetExtraction`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2774 <slepc4py/SLEPc/NEP.pyx#L2774>`

**setCISSRefinement**(*inner=None*, *blsize=None*)

Set the values of various refinement parameters in the CISS solver.

Logically collective.

**Parameters**

- **inner** (`int` / `None`) – Number of iterative refinement iterations (inner loop).
- **blsize** (`int` / `None`) – Number of iterative refinement iterations (blocksize loop).

**Return type**

`None`

**See also**

`getCISSRefinement`, `NEPCISSSetRefinement`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2944 <slepc4py/SLEPc/NEP.pyx#L2944>`

**setCISSSizes**(*ip=None*, *bs=None*, *ms=None*, *npart=None*, *bsmax=None*, *realmats=False*)

Set the values of various size parameters in the CISS solver.

Logically collective.

**Parameters**

- **ip** (`int` / `None`) – Number of integration points.
- **bs** (`int` / `None`) – Block size.
- **ms** (`int` / `None`) – Moment size.
- **npart** (`int` / `None`) – Number of partitions when splitting the communicator.
- **bsmax** (`int` / `None`) – Maximum block size.
- **realmats** (`bool`) – True if A and B are real.

**Return type**

`None`

## Notes

The default number of partitions is 1. This means the internal `petsc4py.PETSc.KSP` object is shared among all processes of the `NEP` communicator. Otherwise, the communicator is split into `npart` communicators, so that `npart` `petsc4py.PETSc.KSP` solves proceed simultaneously.

### See also

`getCISSSizes`, `setCISSThreshold`, `setCISSRefinement`, `NEPCISSSetSizes`

**`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:2811 <slepc4py/SLEPc/NEP.pyx#L2811>``**

**`setCISSThreshold(delta=None, spur=None)`**

Set the values of various threshold parameters in the CISS solver.

Logically collective.

### Parameters

- **`delta`** (`float` / `None`) – Threshold for numerical rank.
- **`spur`** (`float` / `None`) – Spurious threshold (to discard spurious eigenpairs).

### Return type

`None`

### See also

`getCISSThreshold`, `NEPCISSSetThreshold`

**`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:2899 <slepc4py/SLEPc/NEP.pyx#L2899>``**

**`setConvergenceTest(conv)`**

Set how to compute the error estimate used in the convergence test.

Logically collective.

### Parameters

- **`conv`** (`Conv`) – The method used to compute the error estimate used in the convergence test.

### Return type

`None`

### See also

`getConvergenceTest`, `NEPSetConvergenceTest`

**`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:644 <slepc4py/SLEPc/NEP.pyx#L644>``**

**`setDS(ds)`**

Set a direct solver object associated to the eigensolver.

Collective.

### Parameters

- **`ds`** (`DS`) – The direct solver context.

**Return type**

None

**See also**[getDS](#), [NEPSetDS](#)**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:971 <slepc4py/SLEPc/NEP.pyx#L971>`**setDimensions**(*nev=None, ncv=None, mpd=None*)

Set the number of eigenvalues to compute.

Logically collective.

Set the number of eigenvalues to compute and the dimension of the subspace.

**Parameters**

- **nev** (*int* / *None*) – Number of eigenvalues to compute.
- **ncv** (*int* / *None*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int* / *None*) – Maximum dimension allowed for the projected problem.

**Return type**

None

**Notes**

Use [DETERMINE](#) for *ncv* and *mpd* to assign a reasonably good value, which is dependent on the solution method.

The parameters *ncv* and *mpd* are intimately related, so that the user is advised to set one of them at most. Normal usage is the following:

- In cases where *nev* is small, the user sets *ncv* (a reasonable default is  $2 * nev$ ).
- In cases where *nev* is large, the user sets *mpd*.

The value of *ncv* should always be between *nev* and (*nev* + *mpd*), typically  $ncv = nev + mpd$ . If *nev* is not too large,  $mpd = nev$  is a reasonable choice, otherwise a smaller value should be used.

**See also**[getDimensions](#), [NEPSetDimensions](#)**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:823 <slepc4py/SLEPc/NEP.pyx#L823>`**setEigenvalueComparison**(*comparison, args=None, kargs=None*)

Set an eigenvalue comparison function.

Logically collective.

**Notes**

This eigenvalue comparison function is used when [setWhichEigenpairs\(\)](#) is set to [NEP.Which.USER](#).



#### See also

[getEigenvalueComparison](#), [NEPSetEigenvalueComparison](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1070 <slepc4py/SLEPc/NEP.pyx#L1070>`

#### Parameters

- **comparison** ([NEPEigenvalueComparison](#) | *None*)
- **args** (*tuple*[*Any*, ...] | *None*)
- **kargs** (*dict*[*str*, *Any*] | *None*)

#### Return type

*None*

#### **setFromOptions()**

Set NEP options from the options database.

Collective.

#### Notes

To see all options, run your program with the `-help` option.

This routine must be called before [setUp\(\)](#) if the user is to be allowed to set the solver type.

#### See also

[setOptionsPrefix](#), [NEPSetFromOptions](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:404 <slepc4py/SLEPc/NEP.pyx#L404>`

#### Return type

*None*

#### **setFunction(function, F=None, P=None, args=None, kargs=None)**

Set the function to compute the nonlinear Function  $T(\lambda)$ .

Collective.

Set the function to compute the nonlinear Function  $T(\lambda)$  as well as the location to store the matrix.

#### Parameters

- **function** ([NEPFunction](#)) – Function evaluation routine.
- **F** ([petsc4py.PETSc.Mat](#) | *None*) – Function matrix.
- **P** ([petsc4py.PETSc.Mat](#) | *None*) – Preconditioner matrix (usually the same as F).
- **args** (*tuple*[*Any*, ...] | *None*)
- **kargs** (*dict*[*str*, *Any*] | *None*)

#### Return type

*None*

**See also**

[`setJacobian`](#), [`getFunction`](#), [`NEPSetFunction`](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1500](#) <[slepc4py/SLEPc/NEP.pyx#L1500](#)>

**setInitialSpace**(*space*)

Set the initial space from which the eigensolver starts to iterate.

Collective.

**Parameters**

**space** (*Vec*) – The initial space.

**Return type**

*None*

**Notes**

Some solvers start to iterate on a single vector (initial vector). In that case, only the first vector is taken into account and the other vectors are ignored.

These vectors do not persist from one [`solve\(\)`](#) call to the other, so the initial space should be set every time.

The vectors do not need to be mutually orthonormal, since they are explicitly orthonormalized internally.

Common usage of this function is when the user can provide a rough approximation of the wanted eigenspace. Then, convergence may be faster.

**See also**

[`setUp`](#), [`NEPSetInitialSpace`](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:990](#) <[slepc4py/SLEPc/NEP.pyx#L990](#)>

**setInterpolInterpolation**(*tol=None, deg=None*)

Set the tolerance and maximum degree for the interpolation polynomial.

Collective.

**Parameters**

- **tol** (*float* / *None*) – The tolerance to stop computing polynomial coefficients.
- **deg** (*int* / *None*) – The maximum degree of interpolation.

**Return type**

*None*

**See also**

[`getInterpolInterpolation`](#), [`NEPInterpolSetInterpolation`](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2421](#) <[slepc4py/SLEPc/NEP.pyx#L2421](#)>

### **setInterpolPEP**(*pep*)

Set a polynomial eigensolver object associated to the nonlinear eigensolver.

Collective.

#### **Parameters**

**pep** ([PEP](#)) – The polynomial eigensolver.

#### **Return type**

`None`

#### **See also**

[getInterpolPEP](#), [NEPInterpolSetPEP](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2384 <slepc4py/SLEPc/NEP.pyx#L2384>](#)

### **setJacobian**(*jacobian*, *J=None*, *args=None*, *kargs=None*)

Set the function to compute the Jacobian  $T'(\lambda)$ .

Collective.

Set the function to compute the Jacobian  $T'(\lambda)$  as well as the location to store the matrix.

#### **Parameters**

- **jacobian** ([NEPJacobian](#)) – Jacobian evaluation routine.
- **J** ([petsc4py.PETSc.Mat](#) | `None`) – Jacobian matrix.
- **args** ([tuple](#)[`Any`, ...] | `None`)
- **kargs** ([dict](#)[`str`, `Any`] | `None`)

#### **Return type**

`None`

#### **See also**

[setFunction](#), [getJacobian](#), [NEPSetJacobian](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1570 <slepc4py/SLEPc/NEP.pyx#L1570>](#)

### **setMonitor**(*monitor*, *args=None*, *kargs=None*)

Append a monitor function to the list of monitors.

Logically collective.

#### **See also**

[getMonitor](#), [cancelMonitor](#), [NEPMonitorSet](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1117 <slepc4py/SLEPc/NEP.pyx#L1117>](#)

#### **Parameters**

- **monitor** ([NEPMonitorFunction](#) | `None`)
- **args** ([tuple](#)[`Any`, ...] | `None`)

- **kargs** (*dict[str, Any] | None*)

**Return type**

*None*

**setNArnoldiKSP(*ksp*)**

Set a linear solver object associated to the nonlinear eigensolver.

Collective.

**Parameters**

**ksp** (*KSP*) – The linear solver object.

**Return type**

*None*

**See also**

*getNArnoldiKSP*, *NEPNArnoldiSetKSP*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2301 <slepc4py/SLEPc/NEP.pyx#L2301>](#)

**setNArnoldiLagPreconditioner(*lag*)**

Set when the preconditioner is rebuilt in the nonlinear solve.

Logically collective.

**Parameters**

**lag** (*int*) – 0 indicates NEVER rebuild, 1 means rebuild every time the Jacobian is computed within the nonlinear iteration, 2 means every second time the Jacobian is built, etc.

**Return type**

*None*

**Notes**

The default is 1. The preconditioner is ALWAYS built in the first iteration of a nonlinear solve.

**See also**

*getNArnoldiLagPreconditioner*, *NEPNArnoldiSetLagPreconditioner*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2338 <slepc4py/SLEPc/NEP.pyx#L2338>](#)

**setNLEIGSEPS(*eps*)**

Set a linear eigensolver object associated to the nonlinear eigensolver.

Collective.

**Parameters**

**eps** (*EPS*) – The linear eigensolver.

**Return type**

*None*

#### See also

[getNLEIGSEPS](#), [NEPNLEIGSSetEPS](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2658 <slepc4py/SLEPc/NEP.pyx#L2658>`

**setNLEIGSFULLBASIS**(*fullbasis=True*)

Set TOAR-basis (default) or full-basis variants of the NLEIGS method.

Logically collective.

Toggle between TOAR-basis (default) and full-basis variants of the NLEIGS method.

#### Parameters

**fullbasis** (*bool*) – True if the full-basis variant must be selected.

#### Return type

*None*

#### Notes

The default is to use a compact representation of the Krylov basis, that is,  $V = (I \otimes U)S$ , with a *BV* of type *TENSOR*. This behavior can be changed so that the full basis  $V$  is explicitly stored and operated with. This variant is more expensive in terms of memory and computation, but is necessary in some cases, particularly for two-sided computations, see [setTwoSided\(\)](#).

In the full-basis variant, the NLEIGS solver uses an *EPS* object to explicitly solve the linearized eigenproblem, see [getNLEIGSEPS\(\)](#).

#### See also

[setTwoSided](#), [getNLEIGSFULLBASIS](#), [getNLEIGSEPS](#), [NEPNLEIGSSetFULLBASIS](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2606 <slepc4py/SLEPc/NEP.pyx#L2606>`

**setNLEIGSINTERPOLATION**(*tol=None, deg=None*)

Set the tolerance and maximum degree for the interpolation polynomial.

Collective.

Set the tolerance and maximum degree when building the interpolation via divided differences.

#### Parameters

- **tol** (*float* / *None*) – The tolerance to stop computing divided differences.
- **deg** (*int* / *None*) – The maximum degree of interpolation.

#### Return type

*None*

#### See also

[getNLEIGSINTERPOLATION](#), [NEPNLEIGSSetINTERPOLATION](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2555 <slepc4py/SLEPc/NEP.pyx#L2555>`

### **setNLEIGSLocking**(*lock*)

Toggle between locking and non-locking variants of the NLEIGS method.

Logically collective.

#### **Parameters**

**lock** (*bool*) – True if the locking variant must be selected.

#### **Return type**

*None*

### **Notes**

The default is to lock converged eigenpairs when the method restarts. This behavior can be changed so that all directions are kept in the working subspace even if already converged to working accuracy (the non-locking variant).

#### **See also**

[\*getNLEIGSLocking\*](#), [\*NEPNLEIGSSetLocking\*](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2511 <slepc4py/SLEPc/NEP.pyx#L2511>](#)

### **setNLEIGSRKShifts**(*shifts*)

Set a list of shifts to be used in the Rational Krylov method.

Collective.

#### **Parameters**

**shifts** (*Sequence[Scalar]*) – Values specifying the shifts.

#### **Return type**

*None*

### **Notes**

If only one shift is provided, the built subspace is equivalent to shift-and-invert Krylov-Schur (provided that the absolute convergence criterion is used). Otherwise, the rational Krylov variant is run.

#### **See also**

[\*getNLEIGSRKShifts\*](#), [\*getNLEIGSKSPs\*](#), [\*NEPNLEIGSSetRKShifts\*](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:2695 <slepc4py/SLEPc/NEP.pyx#L2695>](#)

### **setNLEIGSRestart**(*keep*)

Set the restart parameter for the NLEIGS method.

Logically collective.

The proportion of basis vectors that must be kept after restart.

#### **Parameters**

**keep** (*float*) – The number of vectors to be kept at restart.

#### **Return type**

*None*

## Notes

Allowed values are in the range [0.1,0.9]. The default is 0.5.

### See also

[getNLEIGSRestart](#), [NEPNLEIGSSetRestart](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2468 <slepc4py/SLEPc/NEP.pyx#L2468>`

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all NEP options in the database.

Logically collective.

### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all NEP option requests.

### Return type

*None*

## Notes

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

For example, to distinguish between the runtime options for two different NEP contexts, one could call:

```
N1.setOptionsPrefix("nep1_")
N2.setOptionsPrefix("nep2_")
```

### See also

[appendOptionsPrefix](#), [getOptionsPrefix](#), [NEPGetOptionsPrefix](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:354 <slepc4py/SLEPc/NEP.pyx#L354>`

**setProblemType**(*problem\_type*)

Set the type of the eigenvalue problem.

Logically collective.

### Parameters

**problem\_type** (*ProblemType*) – The problem type to be set.

### Return type

*None*

## Notes

This function is used to provide a hint to the *NEP* solver to exploit certain properties of the nonlinear eigenproblem. This hint may be used or not, depending on the solver. By default, no particular structure is assumed.

**See also**

`getProblemType`, `NEPSetProblemType`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:442 <slepc4py/SLEPc/NEP.pyx#L442>`

**setRG**(*rg*)

Set a region object associated to the eigensolver.

Collective.

**Parameters**

**rg** (*RG*) – The region context.

**Return type**

None

**See also**

`getRG`, `NEPSetRG`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:934 <slepc4py/SLEPc/NEP.pyx#L934>`

**setRIIConstCorrectionTol**(*cct*)

Set a flag to keep the tolerance used in the linear solver constant.

Logically collective.

**Parameters**

**cct** (*bool*) – If True, the `petsc4py.PETSc.KSP` relative tolerance is constant.

**Return type**

None

**Notes**

By default, an exponentially decreasing tolerance is set in the KSP used within the nonlinear iteration, so that each Newton iteration requests better accuracy than the previous one. The constant correction tolerance flag stops this behavior.

**See also**

`getRIIConstCorrectionTol`, `NEPRIISetConstCorrectionTol`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1917 <slepc4py/SLEPc/NEP.pyx#L1917>`

**setRIIDeflationThreshold**(*deftol*)

Set the threshold used to switch between deflated and non-deflated.

Logically collective.

Set the threshold value used to switch between deflated and non-deflated iteration.

**Parameters**

**deftol** (*float*) – The threshold value.



**Return type**

None

**Notes**

Normally, the solver iterates on the extended problem in order to deflate previously converged eigenpairs. If this threshold is set to a nonzero value, then once the residual error is below this threshold the solver will continue the iteration without deflation. The intention is to be able to improve the current eigenpair further, despite having previous eigenpairs with somewhat bad precision.

**See also**

`getRIIDeflationThreshold`, `NEPRIISetDeflationThreshold`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2050 <slepc4py/SLEPc/NEP.pyx#L2050>`

**setRIIHermitian**(*herm*)

Set a flag to use the Hermitian version of the solver.

Logically collective.

Set a flag to indicate if the Hermitian version of the scalar nonlinear equation must be used by the solver.

**Parameters**

**herm** (*bool*) – If True, the Hermitian version is used.

**Return type**

None

**Notes**

By default, the scalar nonlinear equation  $x^*T(\sigma)^{-1}T(z)x = 0$  is solved at each step of the nonlinear iteration. When this flag is set the simpler form  $x^*T(z)x = 0$  is used, which is supposed to be valid only for Hermitian problems.

**See also**

`getRIIHermitian`, `NEPRIISetHermitian`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2002 <slepc4py/SLEPc/NEP.pyx#L2002>`

**setRIIKSP**(*ksp*)

Set a linear solver object associated to the nonlinear eigensolver.

Collective.

**Parameters**

**ksp** (*KSP*) – The linear solver object.

**Return type**

None

**See also**

`getRIIKSP`, `NEPRIISetKSP`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2100 <slepc4py/SLEPc/NEP.pyx#L2100>`

**setRIILagPreconditioner**(*lag*)

Set when the preconditioner is rebuilt in the nonlinear solve.

Logically collective.

**Parameters**

**lag** (*int*) – 0 indicates NEVER rebuild, 1 means rebuild every time the Jacobian is computed within the nonlinear iteration, 2 means every second time the Jacobian is built, etc.

**Return type**

None

**See also**

`getRIILagPreconditioner`, `NEPRIISetLagPreconditioner`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1878 <slepc4py/SLEPc/NEP.pyx#L1878>`

**setRIIMaximumIterations**(*its*)

Set the max. number of inner iterations to be used in the RII solver.

Logically collective.

These are the Newton iterations related to the computation of the nonlinear Rayleigh functional.

**Parameters**

**its** (*int*) – Maximum inner iterations.

**Return type**

None

**See also**

`getRIIMaximumIterations`, `NEPRIISetMaximumIterations`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1962 <slepc4py/SLEPc/NEP.pyx#L1962>`

**setRefine**(*ref*, *npart*=None, *tol*=None, *its*=None, *scheme*=None)

Set the refinement strategy used by the NEP object.

Logically collective.

Set the refinement strategy used by the NEP object and the associated parameters.

**Parameters**

- **ref** (*Refine*) – The refinement type.
- **npart** (*int* / *None*) – The number of partitions of the communicator.
- **tol** (*float* / *None*) – The convergence tolerance.
- **its** (*int* / *None*) – The maximum number of refinement iterations.
- **scheme** (*RefineScheme* / *None*) – Scheme for solving linear systems.

**Return type**

None

#### See also

[`getRefine`](#), [`NEPSetRefine`](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:694 <slepc4py/SLEPc/NEP.pyx#L694>`

**setSLPDeflationThreshold**(*deftol*)

Set the threshold used to switch between deflated and non-deflated.

Logically collective.

#### Parameters

**deftol** (*float*) – The threshold value.

#### Return type

None

#### Notes

Normally, the solver iterates on the extended problem in order to deflate previously converged eigenpairs. If this threshold is set to a nonzero value, then once the residual error is below this threshold the solver will continue the iteration without deflation. The intention is to be able to improve the current eigenpair further, despite having previous eigenpairs with somewhat bad precision.

#### See also

[`getSLPDeflationThreshold`](#), [`NEPSLPSetDeflationThreshold`](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2139 <slepc4py/SLEPc/NEP.pyx#L2139>`

**setSLPEPS**(*eps*)

Set a linear eigensolver object associated to the nonlinear eigensolver.

Collective.

#### Parameters

**eps** (*EPS*) – The linear eigensolver.

#### Return type

None

#### See also

[`getSLPEPS`](#), [`NEPSLPSetEPS`](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2186 <slepc4py/SLEPc/NEP.pyx#L2186>`

**setSLPEPSLeft**(*eps*)

Set a linear eigensolver object associated to the nonlinear eigensolver.

Collective.

Used to compute left eigenvectors in the two-sided variant of SLP.

#### Parameters

**eps** (*EPS*) – The linear eigensolver.

**Return type**

None

**See also**`setTwoSided`, `setSLPEPS`, `getSLPEPSLeft`, `NEPSLPSetEPSLeft`**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2223` `<slepc4py/SLEPc/NEP.pyx#L2223>`**setSLPKSP**(*ksp*)

Set a linear solver object associated to the nonlinear eigensolver.

Collective.

**Parameters****ksp** (*KSP*) – The linear solver object.**Return type**

None

**See also**`getSLPKSP`, `NEPSLPSetKSP`**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:2262` `<slepc4py/SLEPc/NEP.pyx#L2262>`**setSplitOperator**(*A*, *f*, *structure=None*)

Set the operator of the nonlinear eigenvalue problem in split form.

Collective.

**Parameters**

- **A** (*petsc4py.PETSc.Mat* / *list*[*petsc4py.PETSc.Mat*]) – Coefficient matrices of the split form.
- **f** (*FN* / *list*[*FN*]) – Scalar functions of the split form.
- **structure** (*petsc4py.PETSc.Mat.Structure* / *None*) – Structure flag for matrices.

**Return type**

None

**Notes**

The nonlinear operator is written as  $T(\lambda) = \sum_i A_i f_i(\lambda)$ , for  $i = 1, \dots, n$ . The derivative  $T'(\lambda)$  can be obtained using the derivatives of  $f_i$ .

The structure flag provides information about  $A_i$ 's nonzero pattern.

This function must be called before `setUp()`. If it is called again after `setUp()` then the *NEP* object is reset.

**See also**`getSplitOperator`, `NEPSetSplitOperator`**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1632` `<slepc4py/SLEPc/NEP.pyx#L1632>`

**setSplitPreconditioner**(*P*, *structure=None*)

Set the operator in split form.

Collective.

Set the operator in split form from which to build the preconditioner to be used when solving the nonlinear eigenvalue problem in split form.

**Parameters**

- **P** (*petsc4py.PETSc.Mat* / *list[petsc4py.PETSc.Mat]*) – Coefficient matrices of the split preconditioner.
- **structure** (*petsc4py.PETSc.Mat.Structure* / *None*) – Structure flag for matrices.

**Return type**

*None*

**See also**

*getSplitPreconditioner*, *NEPSetSplitPreconditioner*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1719](#) <slepc4py/SLEPc/NEP.pyx#L1719>

**setStoppingTest**(*stopping*, *args=None*, *kargs=None*)

Set a function to decide when to stop the outer iteration of the eigensolver.

Logically collective.

**See also**

*getStoppingTest*, *NEPSetStoppingTestFunction*

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1029](#) <slepc4py/SLEPc/NEP.pyx#L1029>

**Parameters**

- **stopping** (*NEPStoppingFunction* / *None*)
- **args** (*tuple[Any, ...]* / *None*)
- **kargs** (*dict[str, Any]* / *None*)

**Return type**

*None*

**setTarget**(*target*)

Set the value of the target.

Logically collective.

**Parameters**

**target** (*Scalar*) – The value of the target.

**Return type**

*None*

## Notes

The target is a scalar value used to determine the portion of the spectrum of interest. It is used in combination with `setWhichEigenpairs()`.

When PETSc is built with real scalars, it is not possible to specify a complex target.

### See also

`getTarget`, `setWhichEigenpairs`, `NEPSetTarget`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:544 <slepc4py/SLEPc/NEP.pyx#L544>`

**setTolerances**(*tol=None, maxit=None*)

Set the tolerance and max. iteration count used in convergence tests.

Logically collective.

### Parameters

- **tol** (*float* / *None*) – The convergence tolerance.
- **maxit** (*int* / *None*) – The maximum number of iterations.

### Return type

*None*

## Notes

Use `DETERMINE` for `max_it` to assign a reasonably good value, which is dependent on the solution method.

### See also

`getTolerances`, `NEPSetTolerances`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:596 <slepc4py/SLEPc/NEP.pyx#L596>`

**setTrackAll**(*trackall*)

Set if the solver must compute the residual of all approximate eigenpairs.

Logically collective.

### Parameters

- **trackall** (*bool*) – Whether to compute all residuals or not.

### Return type

*None*

### See also

`getTrackAll`, `NEPSetTrackAll`

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:777 <slepc4py/SLEPc/NEP.pyx#L777>`

**setTwoSided**(*twosided*)

Set the solver to use a two-sided variant.

Logically collective.

Set the solver to use a two-sided variant so that left eigenvectors are also computed.

**Parameters**

**twosided** (*bool*) – Whether the two-sided variant is to be used or not.

**Return type**

*None*

**Notes**

If the user sets `twosided` to `True` then the solver uses a variant of the algorithm that computes both right and left eigenvectors. This is usually much more costly. This option is not available in all solvers.

When using two-sided solvers, the problem matrices must have both the `Mat.mult` and `Mat.multTranspose` operations defined.

**See also**

[`getTwoSided`](#), [`getLeftEigenvector`](#), [`NEPSetTwoSided`](#)

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:1803 <slepc4py/SLEPc/NEP.pyx#L1803>``

**setType**(*nep\_type*)

Set the particular solver to be used in the NEP object.

Logically collective.

**Parameters**

**nep\_type** (*Type* / *str*) – The solver to be used.

**Return type**

*None*

**Notes**

The default is *RII*. Normally, it is best to use [`setFromOptions\(\)`](#) and then set the NEP type from the options database rather than by using this routine. Using the options database provides the user with maximum flexibility in evaluating the different available methods.

**See also**

[`getType`](#), [`NEPSetType`](#)

**:sources:** ``Source code at slepc4py/SLEPc/NEP.pyx:289 <slepc4py/SLEPc/NEP.pyx#L289>``

**setUp()**

Set up all the internal data structures.

Collective.

**Notes**

Sets up all the internal data structures necessary for the execution of the eigensolver.

This function need not be called explicitly in most cases, since [`solve\(\)`](#) calls it. It can be useful when one wants to measure the set-up time separately from the solve time.

#### See also

[solve](#), [NEPSetUp](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:1174 <slepc4py/SLEPc/NEP.pyx#L1174>`

#### Return type

None

#### **setWhichEigenpairs**(*which*)

Set which portion of the spectrum is to be sought.

Logically collective.

#### Parameters

**which** ([Which](#)) – The portion of the spectrum to be sought by the solver.

#### Return type

None

#### Notes

Not all eigensolvers implemented in NEP account for all the possible values. Also, some values make sense only for certain types of problems. If SLEPc is compiled for real numbers [NEP.Which.LARGEST\\_IMAGINARY](#) and [NEP.Which.SMALLEST\\_IMAGINARY](#) use the absolute value of the imaginary part for eigenvalue selection.

The target is a scalar value provided with [setTarget\(\)](#).

The criterion [NEP.Which.TARGET\\_IMAGINARY](#) is available only in case PETSc and SLEPc have been built with complex scalars.

[NEP.Which.ALL](#) is intended for use in the context of the [PEP.Type.CISS](#) solver for computing all eigenvalues in a region.

#### See also

[getWhichEigenpairs](#), [setTarget](#), [PEPSetWhichEigenpairs](#)

**:sources:** `Source code at slepc4py/SLEPc/NEP.pyx:486 <slepc4py/SLEPc/NEP.pyx#L486>`

#### **solve()**

Solve the nonlinear eigenproblem.

Collective.

#### Notes

[solve\(\)](#) will return without generating an error regardless of whether all requested solutions were computed or not. Call [getConverged\(\)](#) to get the actual number of computed solutions, and [getConvergedReason\(\)](#) to determine if the solver converged or failed and why.

#### See also

[setUp](#), [getConverged](#), [getConvergedReason](#), [NEPSolve](#)



**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1195](#) <[slepc4py/SLEPc/NEP.pyx#L1195](#)>

**Return type**

None

**valuesView**(viewer=None)

Display the computed eigenvalues in a viewer.

Collective.

**Parameters**

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

None

**See also**

[solve](#), [vectorsView](#), [errorView](#), [NEPValuesView](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1460](#) <[slepc4py/SLEPc/NEP.pyx#L1460](#)>

**vectorsView**(viewer=None)

Output computed eigenvectors to a viewer.

Collective.

**Parameters**

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

None

**See also**

[solve](#), [valuesView](#), [errorView](#), [NEPVectorsView](#)

**:sources:** [Source code at slepc4py/SLEPc/NEP.pyx:1479](#) <[slepc4py/SLEPc/NEP.pyx#L1479](#)>

**view**(viewer=None)

Print the NEP data structure.

Collective.

**Parameters**

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

None

**See also**

[NEPView](#)

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:223 <slepc4py/SLEPc/NEP.pyx#L223>``

## Attributes Documentation

### **bv**

The basis vectors (*BV*) object associated.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3058 <slepc4py/SLEPc/NEP.pyx#L3058>``

### **ds**

The direct solver (*DS*) object associated.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3072 <slepc4py/SLEPc/NEP.pyx#L3072>``

### **max\_it**

The maximum iteration count used by the NEP convergence tests.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3044 <slepc4py/SLEPc/NEP.pyx#L3044>``

### **problem\_type**

The problem type from the NEP object.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3016 <slepc4py/SLEPc/NEP.pyx#L3016>``

### **rg**

The region (*RG*) object associated.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3065 <slepc4py/SLEPc/NEP.pyx#L3065>``

### **target**

The value of the target.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3030 <slepc4py/SLEPc/NEP.pyx#L3030>``

### **tol**

The tolerance used by the NEP convergence tests.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3037 <slepc4py/SLEPc/NEP.pyx#L3037>``

### **track\_all**

Compute the residual of all approximate eigenpairs.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3051 <slepc4py/SLEPc/NEP.pyx#L3051>``

### **which**

The portion of the spectrum to be sought.

`:sources:`Source code at slepc4py/SLEPc/NEP.pyx:3023 <slepc4py/SLEPc/NEP.pyx#L3023>``

## slepc4py.SLEPc.PEP

### **class** slepc4py.SLEPc.PEP

Bases: `Object`

Polynomial Eigenvalue Problem Solver.

The Polynomial Eigenvalue Problem (*PEP*) solver is the object provided by slepc4py for specifying a polynomial eigenvalue problem. Apart from the specific solvers for this type of problems, there is an *EPS*-based solver, i.e., it uses a solver from *EPS* to solve a generalized eigenproblem obtained after linearization.

## Enumerations

<i>Basis</i>	PEP basis type for the representation of the polynomial.
<i>CISSExtraction</i>	PEP CISS extraction technique.
<i>Conv</i>	PEP convergence test.
<i>ConvergedReason</i>	PEP convergence reasons.
<i>ErrorType</i>	PEP error type to assess accuracy of computed solutions.
<i>Extract</i>	PEP extraction strategy used.
<i>JDProjection</i>	PEP type of projection to be used in the Jacobi-Davidson solver.
<i>ProblemType</i>	PEP problem type.
<i>Refine</i>	PEP refinement strategy.
<i>RefineScheme</i>	PEP scheme for solving linear systems during iterative refinement.
<i>Scale</i>	PEP scaling strategy.
<i>Stop</i>	PEP stopping test.
<i>Type</i>	PEP type.
<i>Which</i>	PEP desired part of spectrum.

### slepc4py.SLEPc.PEP.Basis

**class** slepc4py.SLEPc.PEP.Basis

Bases: `object`

PEP basis type for the representation of the polynomial.

- *MONOMIAL*: Monomials (default).
- *CHEBYSHEV1*: Chebyshev polynomials of the 1st kind.
- *CHEBYSHEV2*: Chebyshev polynomials of the 2nd kind.
- *LEGENDRE*: Legendre polynomials.
- *LAGUERRE*: Laguerre polynomials.
- *HERMITE*: Hermite polynomials.

#### See also

PEPBasis

### Attributes Summary

<i>CHEBYSHEV1</i>	Constant CHEBYSHEV1 of type <code>int</code>
<i>CHEBYSHEV2</i>	Constant CHEBYSHEV2 of type <code>int</code>
<i>HERMITE</i>	Constant HERMITE of type <code>int</code>
<i>LAGUERRE</i>	Constant LAGUERRE of type <code>int</code>
<i>LEGENDRE</i>	Constant LEGENDRE of type <code>int</code>
<i>MONOMIAL</i>	Constant MONOMIAL of type <code>int</code>

Attributes Documentation

**CHEBYSHEV1:** `int` = **CHEBYSHEV1**  
Constant CHEBYSHEV1 of type `int`

**CHEBYSHEV2:** `int` = **CHEBYSHEV2**  
Constant CHEBYSHEV2 of type `int`

**HERMITE:** `int` = **HERMITE**  
Constant HERMITE of type `int`

**LAGUERRE:** `int` = **LAGUERRE**  
Constant LAGUERRE of type `int`

**LEGENDRE:** `int` = **LEGENDRE**  
Constant LEGENDRE of type `int`

**MONOMIAL:** `int` = **MONOMIAL**  
Constant MONOMIAL of type `int`

slepc4py.SLEPc.PEP.CISSExtraction

**class** slepc4py.SLEPc.PEP.CISSExtraction  
Bases: `object`  
PEP CISS extraction technique.

- *RITZ*: Ritz extraction.
- *HANKEL*: Extraction via Hankel eigenproblem.
- *CAA*: Communication-avoiding Arnoldi.

See also
PEPCISSExtraction

Attributes Summary

<i>CAA</i>	Constant CAA of type <code>int</code>
<i>HANKEL</i>	Constant HANKEL of type <code>int</code>
<i>RITZ</i>	Constant RITZ of type <code>int</code>

Attributes Documentation

**CAA:** `int` = **CAA**  
Constant CAA of type `int`

**HANKEL:** `int` = **HANKEL**  
Constant HANKEL of type `int`

**RITZ:** `int` = **RITZ**  
Constant RITZ of type `int`

## slepc4py.SLEPc.PEP.Conv

**class** slepc4py.SLEPc.PEP.Conv

Bases: `object`

PEP convergence test.

- *ABS*: Absolute convergence test.
- *REL*: Convergence test relative to the eigenvalue.
- *NORM*: Convergence test relative to the matrix norms.
- *USER*: User-defined convergence test.

### See also

PEPConv

## Attributes Summary

<i>ABS</i>	Constant ABS of type <code>int</code>
<i>NORM</i>	Constant NORM of type <code>int</code>
<i>REL</i>	Constant REL of type <code>int</code>
<i>USER</i>	Constant USER of type <code>int</code>

## Attributes Documentation

**ABS:** `int` = **ABS**

Constant ABS of type `int`

**NORM:** `int` = **NORM**

Constant NORM of type `int`

**REL:** `int` = **REL**

Constant REL of type `int`

**USER:** `int` = **USER**

Constant USER of type `int`

## slepc4py.SLEPc.PEP.ConvergedReason

**class** slepc4py.SLEPc.PEP.ConvergedReason

Bases: `object`

PEP convergence reasons.

- *CONVERGED\_TOL*: All eigenpairs converged to requested tolerance.
- *CONVERGED\_USER*: User-defined convergence criterion satisfied.
- *DIVERGED\_ITS*: Maximum number of iterations exceeded.
- *DIVERGED\_BREAKDOWN*: Solver failed due to breakdown.
- *DIVERGED\_SYMMETRY\_LOST*: Lanczos-type method could not preserve symmetry.
- *CONVERGED\_ITERATING*: Iteration not finished yet.

<b>See also</b>
PEPConvergedReason

## Attributes Summary

<i>CONVERGED_ITERATING</i>	Constant CONVERGED_ITERATING of type <code>int</code>
<i>CONVERGED_TOL</i>	Constant CONVERGED_TOL of type <code>int</code>
<i>CONVERGED_USER</i>	Constant CONVERGED_USER of type <code>int</code>
<i>DIVERGED_BREAKDOWN</i>	Constant DIVERGED_BREAKDOWN of type <code>int</code>
<i>DIVERGED_ITS</i>	Constant DIVERGED_ITS of type <code>int</code>
<i>DIVERGED_SYMMETRY_LOST</i>	Constant DIVERGED_SYMMETRY_LOST of type <code>int</code>
<i>ITERATING</i>	Constant ITERATING of type <code>int</code>

## Attributes Documentation

**CONVERGED\_ITERATING:** `int` = CONVERGED\_ITERATING

Constant CONVERGED\_ITERATING of type `int`

**CONVERGED\_TOL:** `int` = CONVERGED\_TOL

Constant CONVERGED\_TOL of type `int`

**CONVERGED\_USER:** `int` = CONVERGED\_USER

Constant CONVERGED\_USER of type `int`

**DIVERGED\_BREAKDOWN:** `int` = DIVERGED\_BREAKDOWN

Constant DIVERGED\_BREAKDOWN of type `int`

**DIVERGED\_ITS:** `int` = DIVERGED\_ITS

Constant DIVERGED\_ITS of type `int`

**DIVERGED\_SYMMETRY\_LOST:** `int` = DIVERGED\_SYMMETRY\_LOST

Constant DIVERGED\_SYMMETRY\_LOST of type `int`

**ITERATING:** `int` = ITERATING

Constant ITERATING of type `int`

## slepc4py.SLEPc.PEP.ErrorType

**class** slepc4py.SLEPc.PEP.**ErrorType**

Bases: `object`

PEP error type to assess accuracy of computed solutions.

- *ABSOLUTE*: Absolute error.
- *RELATIVE*: Relative error.
- *BACKWARD*: Backward error.

<b>See also</b>
PEPErrorType

## Attributes Summary

<i>ABSOLUTE</i>	Constant ABSOLUTE of type <code>int</code>
<i>BACKWARD</i>	Constant BACKWARD of type <code>int</code>
<i>RELATIVE</i>	Constant RELATIVE of type <code>int</code>

## Attributes Documentation

**ABSOLUTE:** `int` = **ABSOLUTE**

Constant ABSOLUTE of type `int`

**BACKWARD:** `int` = **BACKWARD**

Constant BACKWARD of type `int`

**RELATIVE:** `int` = **RELATIVE**

Constant RELATIVE of type `int`

## `slepc4py.SLEPc.PEP.Extract`

**class** `slepc4py.SLEPc.PEP.Extract`

Bases: `object`

PEP extraction strategy used.

PEP extraction strategy used to obtain eigenvectors of the PEP from the eigenvectors of the linearization.

- *NONE*: Use the first block.
- *NORM*: Use the first or last block depending on norm of H.
- *RESIDUAL*: Use the block with smallest residual.
- *STRUCTURED*: Combine all blocks in a certain way.

### See also

`PEPExtract`

## Attributes Summary

<i>NONE</i>	Constant NONE of type <code>int</code>
<i>NORM</i>	Constant NORM of type <code>int</code>
<i>RESIDUAL</i>	Constant RESIDUAL of type <code>int</code>
<i>STRUCTURED</i>	Constant STRUCTURED of type <code>int</code>

## Attributes Documentation

**NONE:** `int` = **NONE**

Constant NONE of type `int`

**NORM:** `int` = **NORM**

Constant NORM of type `int`

**RESIDUAL:** `int` = **RESIDUAL**

Constant RESIDUAL of type `int`

**STRUCTURED:** `int` = **STRUCTURED**

Constant STRUCTURED of type `int`

### **slepc4py.SLEPc.PEP.JDProjection**

**class** `slepc4py.SLEPc.PEP.JDProjection`

Bases: `object`

PEP type of projection to be used in the Jacobi-Davidson solver.

- *HARMONIC*: Harmonic projection.
- *ORTHOGONAL*: Orthogonal projection.

#### **See also**

`PEPJDDProjection`

### **Attributes Summary**

*HARMONIC*

Constant HARMONIC of type `int`

*ORTHOGONAL*

Constant ORTHOGONAL of type `int`

### **Attributes Documentation**

**HARMONIC:** `int` = **HARMONIC**

Constant HARMONIC of type `int`

**ORTHOGONAL:** `int` = **ORTHOGONAL**

Constant ORTHOGONAL of type `int`

### **slepc4py.SLEPc.PEP.ProblemType**

**class** `slepc4py.SLEPc.PEP.ProblemType`

Bases: `object`

PEP problem type.

- *GENERAL*: No structure.
- *HERMITIAN*: Hermitian structure.
- *HYPERBOLIC*: QEP with Hermitian matrices,  $M > 0$ ,  $(x^T C x)^2 > 4(x^T M x)(x^T K x)$ .
- *GYROSCOPIC*: QEP with  $M$ ,  $K$  Hermitian,  $M > 0$ ,  $C$  skew-Hermitian.

#### **See also**

`PEPProblemType`



## Attributes Summary

<i>GENERAL</i>	Constant GENERAL of type <code>int</code>
<i>GYROSCOPIC</i>	Constant GYROSCOPIC of type <code>int</code>
<i>HERMITIAN</i>	Constant HERMITIAN of type <code>int</code>
<i>HYPERBOLIC</i>	Constant HYPERBOLIC of type <code>int</code>

## Attributes Documentation

**GENERAL:** `int` = GENERAL

Constant GENERAL of type `int`

**GYROSCOPIC:** `int` = GYROSCOPIC

Constant GYROSCOPIC of type `int`

**HERMITIAN:** `int` = HERMITIAN

Constant HERMITIAN of type `int`

**HYPERBOLIC:** `int` = HYPERBOLIC

Constant HYPERBOLIC of type `int`

## slepc4py.SLEPc.PEP.Refine

**class** slepc4py.SLEPc.PEP.Refine

Bases: `object`

PEP refinement strategy.

- *NONE*: No refinement.
- *SIMPLE*: Refine eigenpairs one by one.
- *MULTIPLE*: Refine all eigenpairs simultaneously (invariant pair).

### See also

PEPRefine

## Attributes Summary

<i>MULTIPLE</i>	Constant MULTIPLE of type <code>int</code>
<i>NONE</i>	Constant NONE of type <code>int</code>
<i>SIMPLE</i>	Constant SIMPLE of type <code>int</code>

## Attributes Documentation

**MULTIPLE:** `int` = MULTIPLE

Constant MULTIPLE of type `int`

**NONE:** `int` = NONE

Constant NONE of type `int`

**SIMPLE:** `int` = SIMPLE

Constant SIMPLE of type `int`

## slepc4py.SLEPc.PEP.RefineScheme

**class** slepc4py.SLEPc.PEP.RefineScheme

Bases: `object`

PEP scheme for solving linear systems during iterative refinement.

- *SCHUR*: Schur complement.
- *MBE*: Mixed block elimination.
- *EXPLICIT*: Build the explicit matrix.

### See also

`PEPRefineScheme`

## Attributes Summary

<i>EXPLICIT</i>	Constant EXPLICIT of type <code>int</code>
<i>MBE</i>	Constant MBE of type <code>int</code>
<i>SCHUR</i>	Constant SCHUR of type <code>int</code>

## Attributes Documentation

**EXPLICIT:** `int` = **EXPLICIT**

Constant EXPLICIT of type `int`

**MBE:** `int` = **MBE**

Constant MBE of type `int`

**SCHUR:** `int` = **SCHUR**

Constant SCHUR of type `int`

## slepc4py.SLEPc.PEP.Scale

**class** slepc4py.SLEPc.PEP.Scale

Bases: `object`

PEP scaling strategy.

- *NONE*: No scaling.
- *SCALAR*: Parameter scaling.
- *DIAGONAL*: Diagonal scaling.
- *BOTH*: Both parameter and diagonal scaling.

### See also

`PEPScale`

## Attributes Summary

<i>BOTH</i>	Constant BOTH of type <code>int</code>
<i>DIAGONAL</i>	Constant DIAGONAL of type <code>int</code>
<i>NONE</i>	Constant NONE of type <code>int</code>
<i>SCALAR</i>	Constant SCALAR of type <code>int</code>

## Attributes Documentation

**BOTH:** `int` = **BOTH**

Constant BOTH of type `int`

**DIAGONAL:** `int` = **DIAGONAL**

Constant DIAGONAL of type `int`

**NONE:** `int` = **NONE**

Constant NONE of type `int`

**SCALAR:** `int` = **SCALAR**

Constant SCALAR of type `int`

## `slepc4py.SLEPc.PEP.Stop`

**class** `slepc4py.SLEPc.PEP.Stop`

Bases: `object`

PEP stopping test.

- *BASIC*: Default stopping test.
- *USER*: User-defined stopping test.

### See also

`PEPStop`

## Attributes Summary

<i>BASIC</i>	Constant BASIC of type <code>int</code>
<i>USER</i>	Constant USER of type <code>int</code>

## Attributes Documentation

**BASIC:** `int` = **BASIC**

Constant BASIC of type `int`

**USER:** `int` = **USER**

Constant USER of type `int`

## slepc4py.SLEPc.PEP.Type

**class** slepc4py.SLEPc.PEP.Type

Bases: `object`

PEP type.

- *TOAR*: Two-level orthogonal Arnoldi.
- *STOAR*: Symmetric TOAR.
- *QARNOLDI*: Q-Arnoldi for quadratic problems.
- *LINEAR*: Linearization via EPS.
- *JD*: Polynomial Jacobi-Davidson.
- *CISS*: Contour integral spectrum slice.

### See also

PEPType

## Attributes Summary

<i>CISS</i>	Object CISS of type <code>str</code>
<i>JD</i>	Object JD of type <code>str</code>
<i>LINEAR</i>	Object LINEAR of type <code>str</code>
<i>QARNOLDI</i>	Object QARNOLDI of type <code>str</code>
<i>STOAR</i>	Object STOAR of type <code>str</code>
<i>TOAR</i>	Object TOAR of type <code>str</code>

## Attributes Documentation

**CISS:** `str` = CISS

Object CISS of type `str`

**JD:** `str` = JD

Object JD of type `str`

**LINEAR:** `str` = LINEAR

Object LINEAR of type `str`

**QARNOLDI:** `str` = QARNOLDI

Object QARNOLDI of type `str`

**STOAR:** `str` = STOAR

Object STOAR of type `str`

**TOAR:** `str` = TOAR

Object TOAR of type `str`

## slepc4py.SLEPc.PEP.Which

**class** slepc4py.SLEPc.PEP.Which

Bases: `object`

PEP desired part of spectrum.

- *LARGEST\_MAGNITUDE*: Largest magnitude (default).
- *SMALLEST\_MAGNITUDE*: Smallest magnitude.
- *LARGEST\_REAL*: Largest real parts.
- *SMALLEST\_REAL*: Smallest real parts.
- *LARGEST\_IMAGINARY*: Largest imaginary parts in magnitude.
- *SMALLEST\_IMAGINARY*: Smallest imaginary parts in magnitude.
- *TARGET\_MAGNITUDE*: Closest to target (in magnitude).
- *TARGET\_REAL*: Real part closest to target.
- *TARGET\_IMAGINARY*: Imaginary part closest to target.
- *ALL*: All eigenvalues in an interval.
- *USER*: User-defined criterion.

### See also

PEPWhich

## Attributes Summary

<i>ALL</i>	Constant ALL of type <code>int</code>
<i>LARGEST_IMAGINARY</i>	Constant LARGEST_IMAGINARY of type <code>int</code>
<i>LARGEST_MAGNITUDE</i>	Constant LARGEST_MAGNITUDE of type <code>int</code>
<i>LARGEST_REAL</i>	Constant LARGEST_REAL of type <code>int</code>
<i>SMALLEST_IMAGINARY</i>	Constant SMALLEST_IMAGINARY of type <code>int</code>
<i>SMALLEST_MAGNITUDE</i>	Constant SMALLEST_MAGNITUDE of type <code>int</code>
<i>SMALLEST_REAL</i>	Constant SMALLEST_REAL of type <code>int</code>
<i>TARGET_IMAGINARY</i>	Constant TARGET_IMAGINARY of type <code>int</code>
<i>TARGET_MAGNITUDE</i>	Constant TARGET_MAGNITUDE of type <code>int</code>
<i>TARGET_REAL</i>	Constant TARGET_REAL of type <code>int</code>
<i>USER</i>	Constant USER of type <code>int</code>

## Attributes Documentation

**ALL:** `int` = ALL

Constant ALL of type `int`

**LARGEST\_IMAGINARY:** `int` = LARGEST\_IMAGINARY

Constant LARGEST\_IMAGINARY of type `int`

**LARGEST\_MAGNITUDE:** `int` = LARGEST\_MAGNITUDE

Constant LARGEST\_MAGNITUDE of type `int`

**LARGEST\_REAL: int = LARGEST\_REAL**  
Constant LARGEST\_REAL of type `int`

**SMALLEST\_IMAGINARY: int = SMALLEST\_IMAGINARY**  
Constant SMALLEST\_IMAGINARY of type `int`

**SMALLEST\_MAGNITUDE: int = SMALLEST\_MAGNITUDE**  
Constant SMALLEST\_MAGNITUDE of type `int`

**SMALLEST\_REAL: int = SMALLEST\_REAL**  
Constant SMALLEST\_REAL of type `int`

**TARGET\_IMAGINARY: int = TARGET\_IMAGINARY**  
Constant TARGET\_IMAGINARY of type `int`

**TARGET\_MAGNITUDE: int = TARGET\_MAGNITUDE**  
Constant TARGET\_MAGNITUDE of type `int`

**TARGET\_REAL: int = TARGET\_REAL**  
Constant TARGET\_REAL of type `int`

**USER: int = USER**  
Constant USER of type `int`

## Methods Summary

<code>appendOptionsPrefix([prefix])</code>	Append to the prefix used for searching for all PEP options in the database.
<code>cancelMonitor()</code>	Clear all monitors for a <a href="#">PEP</a> object.
<code>computeError(i[, etype])</code>	Compute the error associated with the i-th computed eigenpair.
<code>create([comm])</code>	Create the PEP object.
<code>destroy()</code>	Destroy the PEP object.
<code>errorView([etype, viewer])</code>	Display the errors associated with the computed solution.
<code>getBV()</code>	Get the basis vectors object associated to the eigensolver.
<code>getBasis()</code>	Get the type of polynomial basis used.
<code>getCISSExtraction()</code>	Get the extraction technique used in the CISS solver.
<code>getCISSKSPs()</code>	Get the array of linear solver objects associated with the CISS solver.
<code>getCISSRefinement()</code>	Get the values of various refinement parameters in the CISS solver.
<code>getCISSSizes()</code>	Get the values of various size parameters in the CISS solver.
<code>getCISSThreshold()</code>	Get the values of various threshold parameters in the CISS solver.
<code>getConverged()</code>	Get the number of converged eigenpairs.
<code>getConvergedReason()</code>	Get the reason why the <code>solve()</code> iteration was stopped.
<code>getConvergenceTest()</code>	Get the method used to compute the error estimate used in the convergence test.
<code>getDS()</code>	Get the direct solver associated to the eigensolver.

continues on next page

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<code>getDimensions()</code>	Get the number of eigenvalues to compute and the dimension of the subspace.
<code>getEigenpair(i[, Vr, Vi])</code>	Get the i-th solution of the eigenproblem as computed by <code>solve()</code> .
<code>getEigenvalueComparison()</code>	Get the eigenvalue comparison function.
<code>getErrorEstimate(i)</code>	Get the error estimate associated to the i-th computed eigenpair.
<code>getExtract()</code>	Get the extraction technique used by the <i>PEP</i> object.
<code>getInterval()</code>	Get the computational interval for spectrum slicing.
<code>getIterationNumber()</code>	Get the current iteration number.
<code>getJDFix()</code>	Get threshold for changing the target in the correction equation.
<code>getJDMINimalityIndex()</code>	Get the maximum allowed value of the minimality index.
<code>getJDProjection()</code>	Get the type of projection to be used in the Jacobi-Davidson solver.
<code>getJDRestart()</code>	Get the restart parameter used in the Jacobi-Davidson method.
<code>getJDReusePreconditioner()</code>	Get the flag for reusing the preconditioner.
<code>getLinearEPS()</code>	Get the eigensolver object associated to the polynomial eigenvalue solver.
<code>getLinearExplicitMatrix()</code>	Get if the matrices for the linearization are built explicitly.
<code>getLinearLinearization()</code>	Get the coeffs.
<code>getMonitor()</code>	Get the list of monitor functions.
<code>getOperators()</code>	Get the matrices associated with the eigenvalue problem.
<code>getOptionsPrefix()</code>	Get the prefix used for searching for all PEP options in the database.
<code>getProblemType()</code>	Get the problem type from the PEP object.
<code>getQArnoldiLocking()</code>	Get the locking flag used in the Q-Arnoldi method.
<code>getQArnoldiRestart()</code>	Get the restart parameter used in the Q-Arnoldi method.
<code>getRG()</code>	Get the region object associated to the eigensolver.
<code>getRefine()</code>	Get the refinement strategy used by the PEP object.
<code>getRefineKSP()</code>	Get the KSP object used by the eigensolver in the refinement phase.
<code>getST()</code>	Get the spectral transformation object associated to the eigensolver.
<code>getSTOARCheckEigenvalueType()</code>	Get the flag for the eigenvalue type check in spectrum slicing.
<code>getSTOARDetectZeros()</code>	Get the flag that enforces zero detection in spectrum slicing.
<code>getSTOARDimensions()</code>	Get the dimensions used for each subsolve step.
<code>getSTOARInertias()</code>	Get the values of the shifts and their corresponding inertias.
<code>getSTOARLinearization()</code>	Get the coefficients that define the linearization of a quadratic eigenproblem.
<code>getSTOARLocking()</code>	Get the locking flag used in the STOAR method.
<code>getScale([DI, Dr])</code>	Get the strategy used for scaling the polynomial eigenproblem.
<code>getStoppingTest()</code>	Get the stopping test function.

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Table 83 – continued from previous page

<i>getTOARLocking()</i>	Get the locking flag used in the TOAR method.
<i>getTOARRestart()</i>	Get the restart parameter used in the TOAR method.
<i>getTarget()</i>	Get the value of the target.
<i>getTolerances()</i>	Get the tolerance and maximum iteration count.
<i>getTrackAll()</i>	Get the flag indicating whether all residual norms must be computed.
<i>getType()</i>	Get the PEP type of this object.
<i>getWhichEigenpairs()</i>	Get which portion of the spectrum is to be sought.
<i>reset()</i>	Reset the PEP object.
<i>setBV(bv)</i>	Set a basis vectors object associated to the eigensolver.
<i>setBasis(basis)</i>	Set the type of polynomial basis used.
<i>setCISSExtraction(extraction)</i>	Set the extraction technique used in the CISS solver.
<i>setCISSRefinement([inner, blsize])</i>	Set the values of various refinement parameters in the CISS solver.
<i>setCISSSizes([ip, bs, ms, npart, bsmax, ...])</i>	Set the values of various size parameters in the CISS solver.
<i>setCISSThreshold([delta, spur])</i>	Set the values of various threshold parameters in the CISS solver.
<i>setConvergenceTest(conv)</i>	Set how to compute the error estimate used in the convergence test.
<i>setDS(ds)</i>	Set a direct solver object associated to the eigensolver.
<i>setDimensions([nev, ncv, mpd])</i>	Set the number of eigenvalues to compute and the dimension of the subspace.
<i>setEigenvalueComparison(comparison[, args, ...])</i>	Set an eigenvalue comparison function.
<i>setExtract(extract)</i>	Set the extraction strategy to be used.
<i>setFromOptions()</i>	Set PEP options from the options database.
<i>setInitialSpace(space)</i>	Set the initial space from which the eigensolver starts to iterate.
<i>setInterval(inta, intb)</i>	Set the computational interval for spectrum slicing.
<i>setJDFix(fix)</i>	Set the threshold for changing the target in the correction equation.
<i>setJDMinimalityIndex(flag)</i>	Set the maximum allowed value for the minimality index.
<i>setJDProjection(proj)</i>	Set the type of projection to be used in the Jacobi-Davidson solver.
<i>setJDRestart(keep)</i>	Set the restart parameter for the Jacobi-Davidson method.
<i>setJDReusePreconditioner(flag)</i>	Set a flag indicating whether the preconditioner must be reused or not.
<i>setLinearEPS(eps)</i>	Set an eigensolver object associated to the polynomial eigenvalue solver.
<i>setLinearExplicitMatrix(flag)</i>	Set flag to explicitly build the matrices for the linearization.
<i>setLinearLinearization([alpha, beta])</i>	Set the coefficients that define the linearization of a quadratic eigenproblem.
<i>setMonitor(monitor[, args, kargs])</i>	Append a monitor function to the list of monitors.
<i>setOperators(operators)</i>	Set the matrices associated with the eigenvalue problem.
<i>setOptionsPrefix([prefix])</i>	Set the prefix used for searching for all PEP options in the database.

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Table 83 – continued from previous page

<code>setProblemType(problem_type)</code>	Set the type of the polynomial eigenvalue problem.
<code>setQArnoldiLocking(lock)</code>	Toggle between locking and non-locking variants of the Q-Arnoldi method.
<code>setQArnoldiRestart(keep)</code>	Set the restart parameter for the Q-Arnoldi method.
<code>setRG(rg)</code>	Set a region object associated to the eigensolver.
<code>setRefine(ref[, npart, tol, its, scheme])</code>	Set the refinement strategy used by the PEP object.
<code>setST(st)</code>	Set a spectral transformation object associated to the eigensolver.
<code>setSTOARCheckEigenvalueType(flag)</code>	Set flag to check if all eigenvalues have the same definite type.
<code>setSTOARDetectZeros(detect)</code>	Set flag to enforce detection of zeros during the factorizations.
<code>setSTOARDimensions([nev, ncv, mpd])</code>	Set the dimensions used for each subsolve step.
<code>setSTOARLinearization([alpha, beta])</code>	Set the coefficients that define the linearization of a quadratic eigenproblem.
<code>setSTOARLocking(lock)</code>	Toggle between locking and non-locking variants of the STOAR method.
<code>setScale(scale[, alpha, DI, Dr, its, lbda])</code>	Set the scaling strategy to be used.
<code>setStoppingTest(stopping[, args, kargs])</code>	Set a function to decide when to stop the outer iteration of the eigensolver.
<code>setTOARLocking(lock)</code>	Toggle between locking and non-locking variants of the TOAR method.
<code>setTOARRestart(keep)</code>	Set the restart parameter for the TOAR method.
<code>setTarget(target)</code>	Set the value of the target.
<code>setTolerances([tol, max_it])</code>	Set the tolerance and maximum iteration count.
<code>setTrackAll(trackall)</code>	Set flag to compute the residual of all approximate eigenpairs.
<code>setType(pep_type)</code>	Set the particular solver to be used in the PEP object.
<code>setUp()</code>	Set up all the internal data structures.
<code>setWhichEigenpairs(which)</code>	Set which portion of the spectrum is to be sought.
<code>solve()</code>	Solve the polynomial eigenproblem.
<code>valuesView([viewer])</code>	Display the computed eigenvalues in a viewer.
<code>vectorsView([viewer])</code>	Output computed eigenvectors to a viewer.
<code>view([viewer])</code>	Print the PEP data structure.

### Attributes Summary

<code>bv</code>	The basis vectors ( <i>BV</i> ) object associated.
<code>ds</code>	The direct solver ( <i>DS</i> ) object associated.
<code>extract</code>	The type of extraction technique to be employed.
<code>max_it</code>	The maximum iteration count.
<code>problem_type</code>	The type of the eigenvalue problem.
<code>rg</code>	The region ( <i>RG</i> ) object associated.
<code>st</code>	The spectral transformation ( <i>ST</i> ) object associated.
<code>target</code>	The value of the target.
<code>tol</code>	The tolerance.
<code>track_all</code>	Compute the residual norm of all approximate eigenpairs.
<code>which</code>	The portion of the spectrum to be sought.

## Methods Documentation

**appendOptionsPrefix**(*prefix=None*)

Append to the prefix used for searching for all PEP options in the database.

Logically collective.

**Parameters**

**prefix** (*str* / *None*) – The prefix string to prepend to all PEP option requests.

**Return type**

*None*

**See also**

*setOptionsPrefix*, *getOptionsPrefix*, *PEPAppendOptionsPrefix*

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:467](#) <slepc4py/SLEPc/PEP.pyx#L467>

**cancelMonitor**()

Clear all monitors for a *PEP* object.

Logically collective.

**See also**

*PEPMonitorCancel*

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1589](#) <slepc4py/SLEPc/PEP.pyx#L1589>

**Return type**

*None*

**computeError**(*i*, *etype=None*)

Compute the error associated with the *i*-th computed eigenpair.

Collective.

Compute the error (based on the residual norm) associated with the *i*-th computed eigenpair.

**Parameters**

- **i** (*int*) – Index of the solution to be considered.
- **etype** (*ErrorType* / *None*) – The error type to compute.

**Returns**

The error bound, computed in various ways from the residual norm  $\|P(\lambda)x\|_2$  where  $\lambda$  is the eigenvalue and  $x$  is the eigenvector.

**Return type**

*float*

## Notes

The index `i` should be a value between 0 and `nconv-1` (see `getConverged()`).

### See also

`getErrorEstimate`, `PEPComputeError`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1785 <slepc4py/SLEPc/PEP.pyx#L1785>`

**create**(*comm=None*)

Create the PEP object.

Collective.

#### Parameters

**comm** (*Comm* / *None*) – MPI communicator. If not provided, it defaults to all processes.

#### Return type

*Self*

### See also

`PEPCreate`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:350 <slepc4py/SLEPc/PEP.pyx#L350>`

**destroy**()

Destroy the PEP object.

Collective.

### See also

`PEPDestroy`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:324 <slepc4py/SLEPc/PEP.pyx#L324>`

#### Return type

*Self*

**errorView**(*etype=None*, *viewer=None*)

Display the errors associated with the computed solution.

Collective.

Display the errors and the eigenvalues.

#### Parameters

- **etype** (*ErrorType* / *None*) – The error type to compute.
- **viewer** (*petsc4py.PETSc.Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

#### Return type

*None*

## Notes

By default, this function checks the error of all eigenpairs and prints the eigenvalues if all of them are below the requested tolerance. If the viewer has format ASCII\_INFO\_DETAIL then a table with eigenvalues and corresponding errors is printed.

### See also

[\*solve\*](#), [\*valuesView\*](#), [\*vectorsView\*](#), [\*PEPErrorView\*](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1823](#) <[slepc4py/SLEPc/PEP.pyx#L1823](#)>

## getBV()

Get the basis vectors object associated to the eigensolver.

Not collective.

### Returns

The basis vectors context.

### Return type

*BV*

### See also

[\*setBV\*](#), [\*PEPGetBV\*](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1250](#) <[slepc4py/SLEPc/PEP.pyx#L1250](#)>

## getBasis()

Get the type of polynomial basis used.

Not collective.

### Returns

The basis that was previously set.

### Return type

*Basis*

### See also

[\*setBasis\*](#), [\*PEPGetBasis\*](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:505](#) <[slepc4py/SLEPc/PEP.pyx#L505](#)>

## getCISSExtraction()

Get the extraction technique used in the CISS solver.

Not collective.

### Returns

The extraction technique.

### Return type

*CISSExtraction*

**See also**

`setCISSExtraction`, `PEPCISSGetExtraction`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2729 <slepc4py/SLEPc/PEP.pyx#L2729>`

**getCISSKSPs()**

Get the array of linear solver objects associated with the CISS solver.

Collective.

**Returns**

The linear solver objects.

**Return type**

list of `petsc4py.PETSc.KSP`

**Notes**

The number of `petsc4py.PETSc.KSP` solvers is equal to the number of integration points divided by the number of partitions. This value is halved in the case of real matrices with a region centered at the real axis.

**See also**

`setCISSSizes`, `PEPCISSGetKSPs`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2926 <slepc4py/SLEPc/PEP.pyx#L2926>`

**getCISSRefinement()**

Get the values of various refinement parameters in the CISS solver.

Not collective.

**Returns**

- **inner** (`int`) – Number of iterative refinement iterations (inner loop).
- **blsize** (`int`) – Number of iterative refinement iterations (blocksize loop).

**Return type**

tuple[int, int]

**See also**

`setCISSRefinement`, `PEPCISSGetRefinement`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2904 <slepc4py/SLEPc/PEP.pyx#L2904>`

**getCISSSizes()**

Get the values of various size parameters in the CISS solver.

Not collective.

**Returns**

- **ip** (`int`) – Number of integration points.

- **bs** (`int`) – Block size.
- **ms** (`int`) – Moment size.
- **npart** (`int`) – Number of partitions when splitting the communicator.
- **bsmax** (`int`) – Maximum block size.
- **realmats** (`bool`) – True if A and B are real.

**Return type**

`tuple[int, int, int, int, int, bool]`

**See also**

`setCISSSizes`, `PEPCISSGetSizes`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2802 <slepc4py/SLEPc/PEP.pyx#L2802>`

**getCISSThreshold()**

Get the values of various threshold parameters in the CISS solver.

Not collective.

**Returns**

- **delta** (`float`) – Threshold for numerical rank.
- **spur** (`float`) – Spurious threshold (to discard spurious eigenpairs).

**Return type**

`tuple[float, float]`

**See also**

`setCISSThreshold`, `PEPCISSGetThreshold`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2859 <slepc4py/SLEPc/PEP.pyx#L2859>`

**getConverged()**

Get the number of converged eigenpairs.

Not collective.

**Returns**

**nconv** – Number of converged eigenpairs.

**Return type**

`int`

**Notes**

This function should be called after `solve()` has finished.

The value `nconv` may be different from the number of requested solutions `nev`, but not larger than `ncv`, see `setDimensions()`.

**See also**

*setDimensions*, *solve*, *getEigenpair*, *PEPGetConverged*

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1689 <slepc4py/SLEPc/PEP.pyx#L1689>](#)

**getConvergedReason()**

Get the reason why the *solve()* iteration was stopped.

Not collective.

**Returns**

Negative value indicates diverged, positive value converged.

**Return type**

*ConvergedReason*

**See also**

*setTolerances*, *solve*, *PEPGetConvergedReason*

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1670 <slepc4py/SLEPc/PEP.pyx#L1670>](#)

**getConvergenceTest()**

Get the method used to compute the error estimate used in the convergence test.

Not collective.

**Returns**

The method used to compute the error estimate used in the convergence test.

**Return type**

*Conv*

**See also**

*setConvergenceTest*, *PEPGetConvergenceTest*

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:820 <slepc4py/SLEPc/PEP.pyx#L820>](#)

**getDS()**

Get the direct solver associated to the eigensolver.

Not collective.

**Returns**

The direct solver context.

**Return type**

*DS*

**See also**

*setDS*, *PEPGetDS*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1324 <slepc4py/SLEPc/PEP.pyx#L1324>`

### **getDimensions()**

Get the number of eigenvalues to compute and the dimension of the subspace.

Not collective.

#### **Returns**

- **nev** (*int*) – Number of eigenvalues to compute.
- **ncv** (*int*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int*) – Maximum dimension allowed for the projected problem.

#### **Return type**

`tuple[int, int, int]`

#### **See also**

`setDimensions`, `PEPGetDimensions`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1038 <slepc4py/SLEPc/PEP.pyx#L1038>`

### **getEigenpair(*i*, *Vr*=None, *Vi*=None)**

Get the *i*-th solution of the eigenproblem as computed by `solve()`.

Collective.

The solution consists of both the eigenvalue and the eigenvector.

#### **Parameters**

- **i** (*int*) – Index of the solution to be obtained.
- **Vr** (*Vec* | *None*) – Placeholder for the returned eigenvector (real part).
- **Vi** (*Vec* | *None*) – Placeholder for the returned eigenvector (imaginary part).

#### **Returns**

The computed eigenvalue.

#### **Return type**

`complex`

### **Notes**

The index *i* should be a value between 0 and `nconv-1` (see `getConverged()`). Eigenpairs are indexed according to the ordering criterion established with `setWhichEigenpairs()`.

The eigenvector is normalized to have unit norm.

#### **See also**

`solve`, `getConverged`, `setWhichEigenpairs`, `PEPGetEigenpair`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1715 <slepc4py/SLEPc/PEP.pyx#L1715>`



### **getEigenvalueComparison()**

Get the eigenvalue comparison function.

Not collective.

#### **Returns**

The eigenvalue comparison function.

#### **Return type**

*PEPEigenvalueComparison*

#### **See also**

*setEigenvalueComparison*

**`:sources:`Source code at slepc4py/SLEPc/PEP.pyx:1530 <slepc4py/SLEPc/PEP.pyx#L1530>``**

### **getErrorEstimate(*i*)**

Get the error estimate associated to the *i*-th computed eigenpair.

Not collective.

#### **Parameters**

***i*** (*int*) – Index of the solution to be considered.

#### **Returns**

Error estimate.

#### **Return type**

*float*

#### **Notes**

This is the error estimate used internally by the eigensolver. The actual error bound can be computed with *computeError()*.

#### **See also**

*computeError*, *PEPGetErrorEstimate*

**`:sources:`Source code at slepc4py/SLEPc/PEP.pyx:1756 <slepc4py/SLEPc/PEP.pyx#L1756>``**

### **getExtract()**

Get the extraction technique used by the *PEP* object.

Not collective.

#### **Returns**

The extraction strategy.

#### **Return type**

*Extract*

#### **See also**

*setExtract*, *PEPGetExtract*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:979 <slepc4py/SLEPc/PEP.pyx#L979>`

### **getInterval()**

Get the computational interval for spectrum slicing.

Not collective.

#### **Returns**

- **inta** (`float`) – The left end of the interval.
- **intb** (`float`) – The right end of the interval.

#### **Return type**

`tuple[float, float]`

### **Notes**

If the interval was not set by the user, then zeros are returned.

#### **See also**

`setInterval`, `PEPGetInterval`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:764 <slepc4py/SLEPc/PEP.pyx#L764>`

### **getIterationNumber()**

Get the current iteration number.

Not collective.

If the call to `solve()` is complete, then it returns the number of iterations carried out by the solution method.

#### **Returns**

Iteration number.

#### **Return type**

`int`

#### **See also**

`getConvergedReason`, `setTolerances`, `PEPGetIterationNumber`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1648 <slepc4py/SLEPc/PEP.pyx#L1648>`

### **getJDFix()**

Get threshold for changing the target in the correction equation.

Not collective.

#### **Returns**

The threshold for changing the target.

#### **Return type**

`float`

**See also**

*setJDFix*, *PEPJGetFix*

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:2566 <slepc4py/SLEPc/PEP.pyx#L2566>``

**getJDMinimalityIndex()**

Get the maximum allowed value of the minimality index.

Not collective.

**Returns**

The maximum minimality index.

**Return type**

*int*

**See also**

*setJDMinimalityIndex*, *PEPJGetMinimalityIndex*

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:2653 <slepc4py/SLEPc/PEP.pyx#L2653>``

**getJDProjection()**

Get the type of projection to be used in the Jacobi-Davidson solver.

Not collective.

**Returns**

The type of projection.

**Return type**

*JDProjection*

**See also**

*setJDProjection*, *PEPJGetProjection*

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:2690 <slepc4py/SLEPc/PEP.pyx#L2690>``

**getJDRestart()**

Get the restart parameter used in the Jacobi-Davidson method.

Not collective.

**Returns**

The number of vectors to be kept at restart.

**Return type**

*float*

**See also**

*setJDRestart*, *PEPJGetRestart*

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:2523 <slepc4py/SLEPc/PEP.pyx#L2523>``

### **getJDReusePreconditioner()**

Get the flag for reusing the preconditioner.

Not collective.

#### **Returns**

The reuse flag.

#### **Return type**

`bool`

#### **See also**

`setJDReusePreconditioner`, `PEPJDGetReusePreconditioner`

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2610 <slepc4py/SLEPc/PEP.pyx#L2610>](#)

### **getLinearEPS()**

Get the eigensolver object associated to the polynomial eigenvalue solver.

Collective.

#### **Returns**

The linear eigensolver.

#### **Return type**

`EPS`

#### **See also**

`setLinearEPS`, `PEPLinearGetEPS`

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1912 <slepc4py/SLEPc/PEP.pyx#L1912>](#)

### **getLinearExplicitMatrix()**

Get if the matrices for the linearization are built explicitly.

Not collective.

#### **Returns**

Boolean flag indicating if the matrices are built explicitly.

#### **Return type**

`bool`

#### **See also**

`getLinearExplicitMatrix`, `PEPLinearSetExplicitMatrix`

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1993 <slepc4py/SLEPc/PEP.pyx#L1993>](#)

### **getLinearLinearization()**

Get the coeffs. defining the linearization of a quadratic eigenproblem.

Not collective.

#### **Returns**

- **alpha** (`float`) – First parameter of the linearization.
- **beta** (`float`) – Second parameter of the linearization.

**Return type**

`tuple[float, float]`

**See also**

`setLinearLinearization`, `PEPLinearGetLinearization`

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:1953 <slepc4py/SLEPc/PEP.pyx#L1953>``

**getMonitor()**

Get the list of monitor functions.

Not collective.

**Returns**

The list of monitor functions.

**Return type**

`PEPMonitorFunction`

**See also**

`setMonitor`

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:1572 <slepc4py/SLEPc/PEP.pyx#L1572>``

**getOperators()**

Get the matrices associated with the eigenvalue problem.

Collective.

**Returns**

The matrices associated with the eigensystem.

**Return type**

`list of petsc4py.PETSc.Mat`

**See also**

`setOperators`, `PEPGetOperators`

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:1361 <slepc4py/SLEPc/PEP.pyx#L1361>``

**getOptionsPrefix()**

Get the prefix used for searching for all PEP options in the database.

Not collective.

**Returns**

The prefix string set for this PEP object.

**Return type**

`str`

**See also**

[\*setOptionsPrefix\*](#), [\*appendOptionsPrefix\*](#), [\*PEPGetOptionsPrefix\*](#)

**:sources:** [`Source code at slepc4py/SLEPc/PEP.pyx:417 <slepc4py/SLEPc/PEP.pyx#L417>`](#)

**getProblemType()**

Get the problem type from the PEP object.

Not collective.

**Returns**

The problem type that was previously set.

**Return type**

*ProblemType*

**See also**

[\*setProblemType\*](#), [\*PEPGetProblemType\*](#)

**:sources:** [`Source code at slepc4py/SLEPc/PEP.pyx:553 <slepc4py/SLEPc/PEP.pyx#L553>`](#)

**getQArnoldiLocking()**

Get the locking flag used in the Q-Arnoldi method.

Not collective.

**Returns**

The locking flag.

**Return type**

*bool*

**See also**

[\*setQArnoldiLocking\*](#), [\*PEPQArnoldiGetLocking\*](#)

**:sources:** [`Source code at slepc4py/SLEPc/PEP.pyx:2084 <slepc4py/SLEPc/PEP.pyx#L2084>`](#)

**getQArnoldiRestart()**

Get the restart parameter used in the Q-Arnoldi method.

Not collective.

**Returns**

The number of vectors to be kept at restart.

**Return type**

*float*

**See also**

[\*setQArnoldiRestart\*](#), [\*PEPQArnoldiGetRestart\*](#)

**:sources:** [`Source code at slepc4py/SLEPc/PEP.pyx:2040 <slepc4py/SLEPc/PEP.pyx#L2040>`](#)

## getRG()

Get the region object associated to the eigensolver.

Not collective.

### Returns

The region context.

### Return type

*RG*

#### See also

[setRG](#), [PEPGetRG](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1287 <slepc4py/SLEPc/PEP.pyx#L1287>](#)

## getRefine()

Get the refinement strategy used by the PEP object.

Not collective.

### Returns

- **ref** (*Refine*) – The refinement type.
- **npart** (*int*) – The number of partitions of the communicator.
- **tol** (*float*) – The convergence tolerance.
- **its** (*int*) – The maximum number of refinement iterations.
- **scheme** (*RefineScheme*) – Scheme for solving linear systems.

### Return type

*tuple[Refine, int, float, int, RefineScheme]*

#### See also

[setRefine](#), [PEPGetRefine](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:859 <slepc4py/SLEPc/PEP.pyx#L859>](#)

## getRefineKSP()

Get the KSP object used by the eigensolver in the refinement phase.

Collective.

### Returns

The linear solver object.

### Return type

*petsc4py.PETSc.KSP*

#### See also

[setRefine](#), [PEPRefineGetKSP](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:934 <slepc4py/SLEPc/PEP.pyx#L934>](#)

## getST()

Get the spectral transformation object associated to the eigensolver.

Not collective.

### Returns

The spectral transformation.

### Return type

*ST*

#### See also

*setST*, *PEPGetST*

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1114 <slepc4py/SLEPc/PEP.pyx#L1114>](#)

## getSTOARCheckEigenvalueType()

Get the flag for the eigenvalue type check in spectrum slicing.

Not collective.

### Returns

Whether the eigenvalue type is checked or not.

### Return type

*bool*

#### See also

*setSTOARCheckEigenvalueType*, *PEPSTOARGetCheckEigenvalueType*

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2476 <slepc4py/SLEPc/PEP.pyx#L2476>](#)

## getSTOARDetectZeros()

Get the flag that enforces zero detection in spectrum slicing.

Not collective.

### Returns

The zero detection flag.

### Return type

*bool*

#### See also

*setSTOARDetectZeros*, *PEPSTOARGetDetectZeros*

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2316 <slepc4py/SLEPc/PEP.pyx#L2316>](#)

## getSTOARDimensions()

Get the dimensions used for each subsolve step.

Not collective.

### Returns



- **nev** (*int*) – Number of eigenvalues to compute.
- **ncv** (*int*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int*) – Maximum dimension allowed for the projected problem.

**Return type**

`tuple[int, int, int]`

**See also**

`setSTOARDimensions`, `PEPSTOARGetDimensions`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2375 <slepc4py/SLEPc/PEP.pyx#L2375>`

**getSTOARInertias()**

Get the values of the shifts and their corresponding inertias.

Not collective.

Get the values of the shifts and their corresponding inertias in case of doing spectrum slicing for a computational interval.

**Returns**

- **shifts** (*ArrayReal*) – The values of the shifts used internally in the solver.
- **inertias** (*ArrayInt*) – The values of the inertia in each shift.

**Return type**

`tuple[ArrayReal, ArrayInt]`

**Notes**

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with `setInterval()` and `SINVERT` is set.

If called after `solve()`, all shifts used internally by the solver are returned (including both endpoints and any intermediate ones). If called before `solve()` and after `setUp()` then only the information of the endpoints of subintervals is available.

**See also**

`setInterval`, `PEPSTOARGetInertias`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2400 <slepc4py/SLEPc/PEP.pyx#L2400>`

**getSTOARLinearization()**

Get the coefficients that define the linearization of a quadratic eigenproblem.

Not collective.

**Returns**

- **alpha** (*float*) – First parameter of the linearization.
- **beta** (*float*) – Second parameter of the linearization.

**Return type**

`tuple[float, float]`

**See also**

[setSTOARLinearization](#), [PEPSTOARGetLinearization](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2217](#) <[slepc4py/SLEPc/PEP.pyx#L2217](#)>

**getSTOARLocking()**

Get the locking flag used in the STOAR method.

Not collective.

**Returns**

The locking flag.

**Return type**

`bool`

**See also**

[setSTOARLocking](#), [PEPSTOARGetLocking](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2264](#) <[slepc4py/SLEPc/PEP.pyx#L2264](#)>

**getScale(Dl=None, Dr=None)**

Get the strategy used for scaling the polynomial eigenproblem.

Not collective.

**Parameters**

- **Dl** (`petsc4py.PETSc.Vec` / `None`) – Placeholder for the returned left diagonal matrix.
- **Dr** (`petsc4py.PETSc.Vec` / `None`) – Placeholder for the returned right diagonal matrix.

**Returns**

- **scale** (`Scale`) – The scaling strategy.
- **alpha** (`float`) – The scaling factor.
- **its** (`int`) – The number of iterations of diagonal scaling.
- **lbda** (`float`) – Approximation of the wanted eigenvalues (modulus).

**Return type**

`tuple[Scale, float, int, float]`

**See also**

[setScale](#), [PEPGetScale](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1151](#) <[slepc4py/SLEPc/PEP.pyx#L1151](#)>

**getStoppingTest()**

Get the stopping test function.

Not collective.

**Returns**

The stopping test function.

**Return type***PEPStoppingFunction***See also***setStoppingTest*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1483 <slepc4py/SLEPc/PEP.pyx#L1483>`

**getTOARLocking()**

Get the locking flag used in the TOAR method.

Not collective.

**Returns**

The locking flag.

**Return type***bool***See also***setTOARLocking*, *PEPTOARGetLocking*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2175 <slepc4py/SLEPc/PEP.pyx#L2175>`

**getTOARRestart()**

Get the restart parameter used in the TOAR method.

Not collective.

**Returns**

The number of vectors to be kept at restart.

**Return type***float***See also***setTOARRestart*, *PEPTOARGetRestart*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2131 <slepc4py/SLEPc/PEP.pyx#L2131>`

**getTarget()**

Get the value of the target.

Not collective.

**Returns**

The value of the target.

**Return type***Scalar*

## Notes

If the target was not set by the user, then zero is returned.

### See also

`setTarget`, `PEPGetTarget`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:658 <slepc4py/SLEPc/PEP.pyx#L658>`

## `getTolerances()`

Get the tolerance and maximum iteration count.

Not collective.

Get the tolerance and maximum iteration count used by the default PEP convergence tests.

### Returns

- `tol` (`float`) – The convergence tolerance.
- `max_it` (`int`) – The maximum number of iterations.

### Return type

`tuple[float, int]`

### See also

`setTolerances`, `PEPGetTolerances`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:708 <slepc4py/SLEPc/PEP.pyx#L708>`

## `getTrackAll()`

Get the flag indicating whether all residual norms must be computed.

Not collective.

### Returns

Whether the solver computes all residuals or not.

### Return type

`bool`

### See also

`setTrackAll`, `PEPGetTrackAll`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:998 <slepc4py/SLEPc/PEP.pyx#L998>`

## `getType()`

Get the PEP type of this object.

Not collective.

### Returns

The solver currently being used.

### Return type

`str`

**See also**

`setType`, `PEPGetType`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:398` `<slepc4py/SLEPc/PEP.pyx#L398>`

**getWhichEigenpairs()**

Get which portion of the spectrum is to be sought.

Not collective.

**Returns**

The portion of the spectrum to be sought by the solver.

**Return type**

*Which*

**See also**

`setWhichEigenpairs`, `PEPGetWhichEigenpairs`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:602` `<slepc4py/SLEPc/PEP.pyx#L602>`

**reset()**

Reset the PEP object.

Collective.

**See also**

`PEPReset`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:338` `<slepc4py/SLEPc/PEP.pyx#L338>`

**Return type**

*None*

**setBV(bv)**

Set a basis vectors object associated to the eigensolver.

Collective.

**Parameters**

**bv** (*BV*) – The basis vectors context.

**Return type**

*None*

**See also**

`getBV`, `PEPSetBV`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1270` `<slepc4py/SLEPc/PEP.pyx#L1270>`

### **setBasis**(*basis*)

Set the type of polynomial basis used.

Logically collective.

Set the type of polynomial basis used to describe the polynomial eigenvalue problem.

#### **Parameters**

**basis** ([Basis](#)) – The basis to be set.

#### **Return type**

[None](#)

### **Notes**

By default, the coefficient matrices passed via [setOperators\(\)](#) are expressed in the monomial basis, i.e.  $P(\lambda) = A_0 + \lambda A_1 + \lambda^2 A_2 + \cdots + \lambda^d A_d$ . Other polynomial bases may have better numerical behavior, but the user must then pass the coefficient matrices accordingly.

#### **See also**

[getBasis](#), [setOperators](#), [PEPSetBasis](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:524](#) <[slepc4py/SLEPc/PEP.pyx#L524](#)>

### **setCISSExtraction**(*extraction*)

Set the extraction technique used in the CISS solver.

Logically collective.

#### **Parameters**

**extraction** ([CISSExtraction](#)) – The extraction technique.

#### **Return type**

[None](#)

#### **See also**

[getCISSExtraction](#), [PEPCISSSetExtraction](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2711](#) <[slepc4py/SLEPc/PEP.pyx#L2711](#)>

### **setCISSRefinement**(*inner=None*, *blsize=None*)

Set the values of various refinement parameters in the CISS solver.

Logically collective.

#### **Parameters**

- **inner** ([int](#) / [None](#)) – Number of iterative refinement iterations (inner loop).
- **blsize** ([int](#) / [None](#)) – Number of iterative refinement iterations (blocksize loop).

#### **Return type**

[None](#)

#### See also

[getCISSRefinement](#), [PEPCISSSetRefinement](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2881 <slepc4py/SLEPc/PEP.pyx#L2881>`

**setCISSSizes**(*ip=None, bs=None, ms=None, npart=None, bsmax=None, realmats=False*)

Set the values of various size parameters in the CISS solver.

Logically collective.

#### Parameters

- **ip** (*int* | *None*) – Number of integration points.
- **bs** (*int* | *None*) – Block size.
- **ms** (*int* | *None*) – Moment size.
- **npart** (*int* | *None*) – Number of partitions when splitting the communicator.
- **bsmax** (*int* | *None*) – Maximum block size.
- **realmats** (*bool*) – True if A and B are real.

#### Return type

*None*

#### Notes

The default number of partitions is 1. This means the internal `petsc4py.PETSc.KSP` object is shared among all processes of the *PEP* communicator. Otherwise, the communicator is split into `npart` communicators, so that `npart` `petsc4py.PETSc.KSP` solves proceed simultaneously.

#### See also

[getCISSSizes](#), [setCISSThreshold](#), [setCISSRefinement](#), [PEPCISSSetSizes](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2748 <slepc4py/SLEPc/PEP.pyx#L2748>`

**setCISSThreshold**(*delta=None, spur=None*)

Set the values of various threshold parameters in the CISS solver.

Logically collective.

#### Parameters

- **delta** (*float* | *None*) – Threshold for numerical rank.
- **spur** (*float* | *None*) – Spurious threshold (to discard spurious eigenpairs).

#### Return type

*None*

#### See also

[getCISSThreshold](#), [PEPCISSSetThreshold](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2836 <slepc4py/SLEPc/PEP.pyx#L2836>`

### **setConvergenceTest**(*conv*)

Set how to compute the error estimate used in the convergence test.

Logically collective.

#### **Parameters**

**conv** (*Conv*) – The method used to compute the error estimate used in the convergence test.

#### **Return type**

*None*

#### **See also**

*getConvergenceTest*, *PEPSetConvergenceTest*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:840 <slepc4py/SLEPc/PEP.pyx#L840>`

### **setDS**(*ds*)

Set a direct solver object associated to the eigensolver.

Collective.

#### **Parameters**

**ds** (*DS*) – The direct solver context.

#### **Return type**

*None*

#### **See also**

*getDS*, *PEPSetDS*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1344 <slepc4py/SLEPc/PEP.pyx#L1344>`

### **setDimensions**(*nev=None*, *ncv=None*, *mpd=None*)

Set the number of eigenvalues to compute and the dimension of the subspace.

Logically collective.

#### **Parameters**

- **nev** (*int* / *None*) – Number of eigenvalues to compute.
- **ncv** (*int* / *None*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int* / *None*) – Maximum dimension allowed for the projected problem.

#### **Return type**

*None*

### **Notes**

Use *DETERMINE* for *ncv* and *mpd* to assign a reasonably good value, which is dependent on the solution method.

The parameters *ncv* and *mpd* are intimately related, so that the user is advised to set one of them at most. Normal usage is the following:



- In cases where `nev` is small, the user sets `ncv` (a reasonable default is  $2 * nev$ ).
- In cases where `nev` is large, the user sets `mpd`.

The value of `ncv` should always be between `nev` and  $(nev + mpd)$ , typically `ncv = nev + mpd`. If `nev` is not too large, `mpd = nev` is a reasonable choice, otherwise a smaller value should be used.

#### See also

`getDimensions`, `PEPSetDimensions`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1063 <slepc4py/SLEPc/PEP.pyx#L1063>`

**setEigenvalueComparison**(*comparison*, *args=None*, *kargs=None*)

Set an eigenvalue comparison function.

Logically collective.

#### Notes

This eigenvalue comparison function is used when `setWhichEigenpairs()` is set to `PEP.Which.USER`.

#### See also

`getEigenvalueComparison`, `PEPSetEigenvalueComparison`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1500 <slepc4py/SLEPc/PEP.pyx#L1500>`

#### Parameters

- **comparison** (`PEPEigenvalueComparison` / `None`)
- **args** (`tuple[Any, ...]` / `None`)
- **kargs** (`dict[str, Any]` / `None`)

#### Return type

`None`

**setExtract**(*extract*)

Set the extraction strategy to be used.

Logically collective.

#### Parameters

**extract** (`Extract`) – The extraction strategy.

#### Return type

`None`

#### Notes

This is relevant for solvers based on linearization. Once the solver has converged, the polynomial eigenvectors can be extracted from the eigenvectors of the linearized problem in different ways.

#### See also

`getExtract`, `PEPSetExtract`

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:954 <slepc4py/SLEPc/PEP.pyx#L954>``

### **setFromOptions()**

Set PEP options from the options database.

Collective.

### **Notes**

To see all options, run your program with the `-help` option.

This routine must be called before `setUp()` if the user is to be allowed to set the solver type.

#### **See also**

`setOptionsPrefix`, `PEPSetFromOptions`

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:486 <slepc4py/SLEPc/PEP.pyx#L486>``

### **Return type**

`None`

### **setInitialSpace(space)**

Set the initial space from which the eigensolver starts to iterate.

Collective.

### **Parameters**

**space** (`Vec` / `list[Vec]`) – The initial space.

### **Return type**

`None`

### **Notes**

Some solvers start to iterate on a single vector (initial vector). In that case, only the first vector is taken into account and the other vectors are ignored.

These vectors do not persist from one `solve()` call to the other, so the initial space should be set every time.

The vectors do not need to be mutually orthonormal, since they are explicitly orthonormalized internally.

Common usage of this function is when the user can provide a rough approximation of the wanted eigenspace. Then, convergence may be faster.

#### **See also**

`setUp`, `PEPSetInitialSpace`

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:1420 <slepc4py/SLEPc/PEP.pyx#L1420>``

### **setInterval(inta, intb)**

Set the computational interval for spectrum slicing.

Logically collective.

### **Parameters**

- **inta** (`float`) – The left end of the interval.

- **intb** (*float*) – The right end of the interval.

**Return type**

None

**Notes**

Spectrum slicing is a technique employed for computing all eigenvalues of symmetric quadratic eigenproblems in a given interval. This function provides the interval to be considered. It must be used in combination with *PEP.Which.ALL*, see *setWhichEigenpairs()*. Note that in polynomial eigenproblems spectrum slicing is implemented in *STOAR* only.

**See also**

*getInterval*, *PEPSetInterval*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:790 <slepc4py/SLEPc/PEP.pyx#L790>`

**setJDFix**(*fix*)

Set the threshold for changing the target in the correction equation.

Logically collective.

**Parameters**

**fix** (*float*) – Threshold for changing the target.

**Return type**

None

**Notes**

The target in the correction equation is fixed at the first iterations. When the norm of the residual vector is lower than the fix value, the target is set to the corresponding eigenvalue.

**See also**

*getJDFix*, *PEPJDSetsFix*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2542 <slepc4py/SLEPc/PEP.pyx#L2542>`

**setJDMINimalityIndex**(*flag*)

Set the maximum allowed value for the minimality index.

Logically collective.

**Parameters**

**flag** (*int*) – The maximum minimality index.

**Return type**

None

**Notes**

The default value is equal to the degree of the polynomial. A smaller value can be used if the wanted eigenvectors are known to be linearly independent.

#### See also

[`getJDMinimalityIndex`](#), [`PEPJDSetsMinimalityIndex`](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2629 <slepc4py/SLEPc/PEP.pyx#L2629>`

#### **setJDProjection**(*proj*)

Set the type of projection to be used in the Jacobi-Davidson solver.

Logically collective.

##### Parameters

**proj** ([`JDProjection`](#)) – The type of projection.

##### Return type

[`None`](#)

#### See also

[`getJDProjection`](#), [`PEPJDSetsProjection`](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2672 <slepc4py/SLEPc/PEP.pyx#L2672>`

#### **setJDRestart**(*keep*)

Set the restart parameter for the Jacobi-Davidson method.

Logically collective.

Set the restart parameter for the Jacobi-Davidson method, in particular the proportion of basis vectors that must be kept after restart.

##### Parameters

**keep** ([`float`](#)) – The number of vectors to be kept at restart.

##### Return type

[`None`](#)

#### Notes

Allowed values are in the range [0.1,0.9]. The default is 0.5.

#### See also

[`getJDRestart`](#), [`PEPJDSetsRestart`](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2497 <slepc4py/SLEPc/PEP.pyx#L2497>`

#### **setJDReusePreconditioner**(*flag*)

Set a flag indicating whether the preconditioner must be reused or not.

Logically collective.

##### Parameters

**flag** ([`bool`](#)) – The reuse flag.

##### Return type

[`None`](#)

## Notes

The default value is `False`. If set to `True`, the preconditioner is built only at the beginning, using the target value. Otherwise, it may be rebuilt (depending on the `fix` parameter) at each iteration from the Ritz value.

### See also

[`setJDFix`](#), [`getJDReusePreconditioner`](#), [`PEPJDSetsReusePreconditioner`](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2585 <slepc4py/SLEPc/PEP.pyx#L2585>](#)

### **setLinearEPS**(*eps*)

Set an eigensolver object associated to the polynomial eigenvalue solver.

Collective.

#### Parameters

**eps** (*EPS*) – The linear eigensolver.

#### Return type

`None`

### See also

[`getLinearEPS`](#), [`PEPLinearSetEPS`](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1895 <slepc4py/SLEPc/PEP.pyx#L1895>](#)

### **setLinearExplicitMatrix**(*flag*)

Set flag to explicitly build the matrices for the linearization.

Logically collective.

#### Parameters

**flag** (*bool*) – Boolean flag indicating if the matrices are built explicitly.

#### Return type

`None`

### See also

[`getLinearExplicitMatrix`](#), [`PEPLinearSetExplicitMatrix`](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1975 <slepc4py/SLEPc/PEP.pyx#L1975>](#)

### **setLinearLinearization**(*alpha=1.0, beta=0.0*)

Set the coefficients that define the linearization of a quadratic eigenproblem.

Logically collective.

#### Parameters

- **alpha** (*float*) – First parameter of the linearization.
- **beta** (*float*) – Second parameter of the linearization.

#### Return type

`None`

#### See also

[getLinearLinearization](#), [PEPLinearSetLinearization](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1932 <slepc4py/SLEPc/PEP.pyx#L1932>`

**setMonitor**(*monitor*, *args=None*, *kargs=None*)

Append a monitor function to the list of monitors.

Logically collective.

#### See also

[getMonitor](#), [cancelMonitor](#), [PEPMonitorSet](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1547 <slepc4py/SLEPc/PEP.pyx#L1547>`

#### Parameters

- **monitor** ([PEPMonitorFunction](#) / *None*)
- **args** ([tuple](#)[*Any*, ...] / *None*)
- **kargs** ([dict](#)[*str*, *Any*] / *None*)

#### Return type

*None*

**setOperators**(*operators*)

Set the matrices associated with the eigenvalue problem.

Collective.

#### Parameters

**operators** ([list](#)[[Mat](#)]) – The matrices associated with the eigensystem.

#### Return type

*None*

#### Notes

The polynomial eigenproblem is defined as  $P(\lambda)x = 0$ , where  $\lambda$  is the eigenvalue,  $x$  is the eigenvector, and  $P$  is defined as  $P(\lambda) = A_0 + \lambda A_1 + \dots + \lambda^d A_d$ , with  $d = \text{nmat}-1$  (the degree of  $P$ ). For non-monomial bases, this expression is different.

#### See also

[getOperators](#), [PEPSetOperators](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1387 <slepc4py/SLEPc/PEP.pyx#L1387>`

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all PEP options in the database.

Logically collective.

#### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all PEP option requests.

**Return type**

None

**Notes**

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

For example, to distinguish between the runtime options for two different PEP contexts, one could call:

```
P1.setOptionsPrefix("pep1_")
P2.setOptionsPrefix("pep2_")
```

**See also**

[\*appendOptionsPrefix\*](#), [\*getOptionsPrefix\*](#), [\*PEPGetOptionsPrefix\*](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:436 <slepc4py/SLEPc/PEP.pyx#L436>](#)

**setProblemType**(*problem\_type*)

Set the type of the polynomial eigenvalue problem.

Logically collective.

**Parameters**

**problem\_type** ([\*ProblemType\*](#)) – The problem type to be set.

**Return type**

None

**Notes**

This function is used to instruct SLEPc to exploit certain structure in the polynomial eigenproblem. By default, no particular structure is assumed.

If the problem matrices are Hermitian (symmetric in the real case) or Hermitian/skew-Hermitian then the solver can exploit this fact to perform less operations or provide better stability. Hyperbolic problems are a particular case of Hermitian problems, some solvers may treat them simply as Hermitian.

**See also**

[\*setOperators\*](#), [\*setType\*](#), [\*getProblemType\*](#), [\*PEPSetProblemType\*](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:572 <slepc4py/SLEPc/PEP.pyx#L572>](#)

**setQArnoldiLocking**(*lock*)

Toggle between locking and non-locking variants of the Q-Arnoldi method.

Logically collective.

**Parameters**

**lock** ([\*bool\*](#)) – True if the locking variant must be selected.

**Return type**

None

## Notes

The default is to lock converged eigenpairs when the method restarts. This behavior can be changed so that all directions are kept in the working subspace even if already converged to working accuracy (the non-locking variant).

### See also

[`getQArnoldiLocking`](#), [`PEPQArnoldiSetLocking`](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2059](#) <[slepc4py/SLEPc/PEP.pyx#L2059](#)>

## **setQArnoldiRestart**(*keep*)

Set the restart parameter for the Q-Arnoldi method.

Logically collective.

Set the restart parameter for the Q-Arnoldi method, in particular the proportion of basis vectors that must be kept after restart.

### Parameters

**keep** (*float*) – The number of vectors to be kept at restart.

### Return type

*None*

## Notes

Allowed values are in the range [0.1,0.9]. The default is 0.5.

### See also

[`getQArnoldiRestart`](#), [`PEPQArnoldiSetRestart`](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2014](#) <[slepc4py/SLEPc/PEP.pyx#L2014](#)>

## **setRG**(*rg*)

Set a region object associated to the eigensolver.

Collective.

### Parameters

**rg** (*RG*) – The region context.

### Return type

*None*

### See also

[`getRG`](#), [`PEPSetRG`](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1307](#) <[slepc4py/SLEPc/PEP.pyx#L1307](#)>

## **setRefine**(*ref*, *npart=None*, *tol=None*, *its=None*, *scheme=None*)

Set the refinement strategy used by the PEP object.

Logically collective.



Set the refinement strategy used by the PEP object, and the associated parameters.

#### Parameters

- **ref** ([Refine](#)) – The refinement type.
- **npart** ([int](#) / [None](#)) – The number of partitions of the communicator.
- **tol** ([float](#) / [None](#)) – The convergence tolerance.
- **its** ([int](#) / [None](#)) – The maximum number of refinement iterations.
- **scheme** ([RefineScheme](#) / [None](#)) – Scheme for solving linear systems.

#### Return type

[None](#)

#### See also

[getRefine](#), [PEPSetRefine](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:890 <slepc4py/SLEPc/PEP.pyx#L890>](#)

### **setST**(*st*)

Set a spectral transformation object associated to the eigensolver.

Collective.

#### Parameters

**st** ([ST](#)) – The spectral transformation.

#### Return type

[None](#)

#### See also

[getST](#), [PEPSetST](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1134 <slepc4py/SLEPc/PEP.pyx#L1134>](#)

### **setSTOARCheckEigenvalueType**(*flag*)

Set flag to check if all eigenvalues have the same definite type.

Logically collective.

Set a flag to check that all the eigenvalues obtained throughout the spectrum slicing computation have the same definite type.

#### Parameters

**flag** ([bool](#)) – Whether the eigenvalue type is checked or not.

#### Return type

[None](#)

### Notes

This option is relevant only for spectrum slicing computations, but is ignored in [slepc4py.SLEPc.PEP.ProblemType.HYPERBOLIC](#) problems.

This flag is turned on by default, to guarantee that the computed eigenvalues have the same type (otherwise the computed solution might be wrong). But since the check is computationally quite expensive, the check may be turned off if the user knows for sure that all eigenvalues in the requested interval have the same type.

#### See also

[`getSTOARCheckEigenvalueType`](#), [`PEPSTOARSetCheckEigenvalueType`](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2444 <slepc4py/SLEPc/PEP.pyx#L2444>`

**setSTOARDetectZeros**(*detect*)

Set flag to enforce detection of zeros during the factorizations.

Logically collective.

Set a flag to enforce detection of zeros during the factorizations throughout the spectrum slicing computation.

#### Parameters

**detect** (*bool*) – True if zeros must be checked for.

#### Return type

`None`

#### Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with [`setInterval\(\)`](#) and `SINVERT` is set.

A zero in the factorization indicates that a shift coincides with an eigenvalue.

This flag is turned off by default, and may be necessary in some cases. This feature currently requires an external package for factorizations with support for zero detection, e.g. MUMPS.

#### See also

[`setInterval`](#), [`getSTOARDetectZeros`](#), [`PEPSTOARSetDetectZeros`](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2283 <slepc4py/SLEPc/PEP.pyx#L2283>`

**setSTOARDimensions**(*nev=None*, *ncv=None*, *mpd=None*)

Set the dimensions used for each subsolve step.

Logically collective.

#### Parameters

- **nev** (*int* / *None*) – Number of eigenvalues to compute.
- **ncv** (*int* / *None*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int* / *None*) – Maximum dimension allowed for the projected problem.

#### Return type

`None`

## Notes

This call makes sense only for spectrum slicing runs, that is, when an interval has been given with `setInterval()` and `SINVERT` is set.

The meaning of the parameters is the same as in `setDimensions()`, but the ones here apply to every subsolve done by the child `PEP` object.

### See also

`setInterval`, `setDimensions`, `getSTOARDimensions`, `PEPSTOARSetDimensions`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2335 <slepc4py/SLEPc/PEP.pyx#L2335>`

**setSTOARLinearization**(*alpha*=1.0, *beta*=0.0)

Set the coefficients that define the linearization of a quadratic eigenproblem.

Logically collective.

### Parameters

- **alpha** (*float*) – First parameter of the linearization.
- **beta** (*float*) – Second parameter of the linearization.

### Return type

None

### See also

`getSTOARLinearization`, `PEPSTOARSetLinearization`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2196 <slepc4py/SLEPc/PEP.pyx#L2196>`

**setSTOARLocking**(*lock*)

Toggle between locking and non-locking variants of the STOAR method.

Logically collective.

### Parameters

- **lock** (*bool*) – True if the locking variant must be selected.

### Return type

None

## Notes

The default is to lock converged eigenpairs when the method restarts. This behavior can be changed so that all directions are kept in the working subspace even if already converged to working accuracy (the non-locking variant).

### See also

`getSTOARLocking`, `PEPSTOARSetLocking`

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2239 <slepc4py/SLEPc/PEP.pyx#L2239>`

**setScale**(*scale*, *alpha*=None, *Dl*=None, *Dr*=None, *its*=None, *lbda*=None)

Set the scaling strategy to be used.

Collective.

#### Parameters

- **scale** (*Scale*) – The scaling strategy.
- **alpha** (*float* / *None*) – The scaling factor.
- **Dl** (*petsc4py.PETSc.Vec* / *None*) – The left diagonal matrix.
- **Dr** (*petsc4py.PETSc.Vec* / *None*) – The right diagonal matrix.
- **its** (*int* / *None*) – The number of iterations of diagonal scaling.
- **lbda** (*float* / *None*) – Approximation of the wanted eigenvalues (modulus).

#### Return type

None

#### See also

[getScale](#), [PEPSetScale](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1204 <slepc4py/SLEPc/PEP.pyx#L1204>](#)

**setStoppingTest**(*stopping*, *args*=None, *kargs*=None)

Set a function to decide when to stop the outer iteration of the eigensolver.

Logically collective.

#### See also

[getStoppingTest](#), [PEPSetStoppingTestFunction](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:1459 <slepc4py/SLEPc/PEP.pyx#L1459>](#)

#### Parameters

- **stopping** (*PEPStoppingFunction* / *None*)
- **args** (*tuple*[*Any*, ...] / *None*)
- **kargs** (*dict*[*str*, *Any*] / *None*)

#### Return type

None

**setTOARLocking**(*lock*)

Toggle between locking and non-locking variants of the TOAR method.

Logically collective.

#### Parameters

- **lock** (*bool*) – True if the locking variant must be selected.

#### Return type

None

## Notes

The default is to lock converged eigenpairs when the method restarts. This behavior can be changed so that all directions are kept in the working subspace even if already converged to working accuracy (the non-locking variant).

### See also

[getTOARLocking](#), [PEPTOARSetLocking](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2150 <slepc4py/SLEPc/PEP.pyx#L2150>](#)

## **setTOARRestart**(*keep*)

Set the restart parameter for the TOAR method.

Logically collective.

Set the restart parameter for the TOAR method, in particular the proportion of basis vectors that must be kept after restart.

### Parameters

**keep** (*float*) – The number of vectors to be kept at restart.

### Return type

*None*

## Notes

Allowed values are in the range [0.1,0.9]. The default is 0.5.

### See also

[getTOARRestart](#), [PEPTOARSetRestart](#)

**:sources:** [Source code at slepc4py/SLEPc/PEP.pyx:2105 <slepc4py/SLEPc/PEP.pyx#L2105>](#)

## **setTarget**(*target*)

Set the value of the target.

Logically collective.

### Parameters

**target** (*Scalar*) – The value of the target.

### Return type

*None*

## Notes

The target is a scalar value used to determine the portion of the spectrum of interest. It is used in combination with [setWhichEigenpairs\(\)](#).

When PETSc is built with real scalars, it is not possible to specify a complex target.

### See also

[getTarget](#), [setWhichEigenpairs](#), [PEPSetTarget](#)

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:681 <slepc4py/SLEPc/PEP.pyx#L681>`

**setTolerances**(*tol=None, max\_it=None*)

Set the tolerance and maximum iteration count.

Logically collective.

Set the tolerance and maximum iteration count used by the default PEP convergence tests.

**Parameters**

- **tol** (*float* / *None*) – The convergence tolerance.
- **max\_it** (*int* / *None*) – The maximum number of iterations

**Return type**

*None*

**Notes**

Use *DETERMINE* for *max\_it* to assign a reasonably good value, which is dependent on the solution method.

**See also**

*getTolerances*, *PEPSetTolerances*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:733 <slepc4py/SLEPc/PEP.pyx#L733>`

**setTrackAll**(*trackall*)

Set flag to compute the residual of all approximate eigenpairs.

Logically collective.

Set if the solver must compute the residual of all approximate eigenpairs or not.

**Parameters**

- **trackall** (*bool*) – Whether to compute all residuals or not.

**Return type**

*None*

**See also**

*getTrackAll*, *PEPSetTrackAll*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1017 <slepc4py/SLEPc/PEP.pyx#L1017>`

**setType**(*pep\_type*)

Set the particular solver to be used in the PEP object.

Logically collective.

**Parameters**

- **pep\_type** (*Type* / *str*) – The solver to be used.

**Return type**

*None*

## Notes

The default is *TOAR*. Normally, it is best to use *setFromOptions()* and then set the PEP type from the options database rather than by using this routine. Using the options database provides the user with maximum flexibility in evaluating the different available methods.

### See also

*getType*, *PEPSetType*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:371 <slepc4py/SLEPc/PEP.pyx#L371>`

## setUp()

Set up all the internal data structures.

Collective.

## Notes

Sets up all the internal data structures necessary for the execution of the eigensolver. This includes the setup of the internal *ST* object.

This function need not be called explicitly in most cases, since *solve()* calls it. It can be useful when one wants to measure the set-up time separately from the solve time.

### See also

*solve*, *PEPSetUp*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1604 <slepc4py/SLEPc/PEP.pyx#L1604>`

### Return type

None

## setWhichEigenpairs(*which*)

Set which portion of the spectrum is to be sought.

Logically collective.

### Parameters

**which** (*Which*) – The portion of the spectrum to be sought by the solver.

### Return type

None

## Notes

Not all eigensolvers implemented in PEP account for all the possible values. Also, some values make sense only for certain types of problems. If SLEPc is compiled for real numbers *PEP.Which.LARGEST\_IMAGINARY* and *PEP.Which.SMALLEST\_IMAGINARY* use the absolute value of the imaginary part for eigenvalue selection.

The target is a scalar value provided with *setTarget()*.

The criterion *PEP.Which.TARGET\_IMAGINARY* is available only in case PETSc and SLEPc have been built with complex scalars.

*PEP.Which.ALL* is intended for use in combination with an interval (see *setInterval()*), when all eigenvalues within the interval are requested, or in the context of the *PEP.Type.CISS* solver for computing all eigenvalues in a region.

**See also**

*getWhichEigenpairs*, *setTarget*, *setInterval*, *PEPSetWhichEigenpairs*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:621 <slepc4py/SLEPc/PEP.pyx#L621>`

**solve()**

Solve the polynomial eigenproblem.

Collective.

**Notes**

The problem matrices are specified with *setOperators()*.

*solve()* will return without generating an error regardless of whether all requested solutions were computed or not. Call *getConverged()* to get the actual number of computed solutions, and *getConvergedReason()* to determine if the solver converged or failed and why.

**See also**

*setUp*, *setOperators*, *getConverged*, *getConvergedReason*, *PEPSolve*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1626 <slepc4py/SLEPc/PEP.pyx#L1626>`

**Return type**

None

**valuesView(viewer=None)**

Display the computed eigenvalues in a viewer.

Collective.

**Parameters**

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

None

**See also**

*solve*, *vectorsView*, *errorView*, *PEPValuesView*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1855 <slepc4py/SLEPc/PEP.pyx#L1855>`

**vectorsView(viewer=None)**

Output computed eigenvectors to a viewer.

Collective.



#### Parameters

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

#### Return type

*None*

#### See also

*solve*, *valuesView*, *errorView*, *PEPVectorsView*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:1874 <slepc4py/SLEPc/PEP.pyx#L1874>`

**view**(*viewer=None*)

Print the PEP data structure.

Collective.

#### Parameters

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

#### Return type

*None*

#### See also

*PEPView*

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:305 <slepc4py/SLEPc/PEP.pyx#L305>`

### Attributes Documentation

**bv**

The basis vectors (*BV*) object associated.

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:3011 <slepc4py/SLEPc/PEP.pyx#L3011>`

**ds**

The direct solver (*DS*) object associated.

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:3025 <slepc4py/SLEPc/PEP.pyx#L3025>`

**extract**

The type of extraction technique to be employed.

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2976 <slepc4py/SLEPc/PEP.pyx#L2976>`

**max\_it**

The maximum iteration count.

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2990 <slepc4py/SLEPc/PEP.pyx#L2990>`

**problem\_type**

The type of the eigenvalue problem.

**:sources:** `Source code at slepc4py/SLEPc/PEP.pyx:2955 <slepc4py/SLEPc/PEP.pyx#L2955>`

**rg**

The region (*RG*) object associated.

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:3018 <slepc4py/SLEPc/PEP.pyx#L3018>``

**st**

The spectral transformation (*ST*) object associated.

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:3004 <slepc4py/SLEPc/PEP.pyx#L3004>``

**target**

The value of the target.

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:2969 <slepc4py/SLEPc/PEP.pyx#L2969>``

**tol**

The tolerance.

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:2983 <slepc4py/SLEPc/PEP.pyx#L2983>``

**track\_all**

Compute the residual norm of all approximate eigenpairs.

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:2997 <slepc4py/SLEPc/PEP.pyx#L2997>``

**which**

The portion of the spectrum to be sought.

**:sources:** ``Source code at slepc4py/SLEPc/PEP.pyx:2962 <slepc4py/SLEPc/PEP.pyx#L2962>``

## slepc4py.SLEPc.RG

**class** slepc4py.SLEPc.RG

Bases: `Object`

Region.

The *RG* package provides a way to define a region of the complex plane. This is used in various eigensolvers to specify where the wanted eigenvalues are located.

## Enumerations

<i>QuadRule</i>	RG quadrature rule for contour integral methods.
<i>Type</i>	RG type.

## slepc4py.SLEPc.RG.QuadRule

**class** slepc4py.SLEPc.RG.QuadRule

Bases: `object`

RG quadrature rule for contour integral methods.

- *TRAPEZOIDAL*: Trapezoidal rule.
- *CHEBYSHEV*: Chebyshev points.

See also
<a href="#">RGQuadRule</a>

## Attributes Summary

<a href="#"><i>CHEBYSHEV</i></a>	Constant CHEBYSHEV of type <a href="#">int</a>
<a href="#"><i>TRAPEZOIDAL</i></a>	Constant TRAPEZOIDAL of type <a href="#">int</a>

## Attributes Documentation

**CHEBYSHEV:** [int](#) = CHEBYSHEV

Constant CHEBYSHEV of type [int](#)

**TRAPEZOIDAL:** [int](#) = TRAPEZOIDAL

Constant TRAPEZOIDAL of type [int](#)

## slepc4py.SLEPc.RG.Type

**class** slepc4py.SLEPc.RG.Type

Bases: [object](#)

RG type.

- [\*INTERVAL\*](#): A (generalized) interval.
- [\*POLYGON\*](#): A polygonal region defined by its vertices.
- [\*ELLIPSE\*](#): An ellipse defined by its center, radius and vertical scale.
- [\*RING\*](#): A ring region.

See also
<a href="#">RGType</a>

## Attributes Summary

<a href="#"><i>ELLIPSE</i></a>	Object ELLIPSE of type <a href="#">str</a>
<a href="#"><i>INTERVAL</i></a>	Object INTERVAL of type <a href="#">str</a>
<a href="#"><i>POLYGON</i></a>	Object POLYGON of type <a href="#">str</a>
<a href="#"><i>RING</i></a>	Object RING of type <a href="#">str</a>

## Attributes Documentation

**ELLIPSE:** [str](#) = ELLIPSE

Object ELLIPSE of type [str](#)

**INTERVAL:** [str](#) = INTERVAL

Object INTERVAL of type [str](#)

**POLYGON:** `str` = POLYGON

Object POLYGON of type `str`

**RING:** `str` = RING

Object RING of type `str`

## Methods Summary

<code>appendOptionsPrefix([prefix])</code>	Append to the prefix used for searching for all RG options in the database.
<code>canUseConjugates([realmats])</code>	Half of integration points can be avoided (use their conjugates).
<code>checkInside(a)</code>	Determine if a set of given points are inside the region or not.
<code>computeBoundingBox()</code>	Compute box containing the region.
<code>computeContour(n)</code>	Compute points on the contour of the region.
<code>computeQuadrature(quad, n)</code>	Compute the values of the parameters used in a quadrature rule.
<code>create([comm])</code>	Create the RG object.
<code>destroy()</code>	Destroy the RG object.
<code>getComplement()</code>	Get the flag indicating whether the region is complemented or not.
<code>getEllipseParameters()</code>	Get the parameters that define the ellipse region.
<code>getIntervalEndpoints()</code>	Get the parameters that define the interval region.
<code>getOptionsPrefix()</code>	Get the prefix used for searching for all RG options in the database.
<code>getPolygonVertices()</code>	Get the parameters that define the interval region.
<code>getRingParameters()</code>	Get the parameters that define the ring region.
<code>getScale()</code>	Get the scaling factor.
<code>getType()</code>	Get the RG type of this object.
<code>isAxisymmetric([vertical])</code>	Determine if the region is axisymmetric.
<code>isTrivial()</code>	Tell whether it is the trivial region (whole complex plane).
<code>setComplement([comp])</code>	Set a flag to indicate that the region is the complement of the specified one.
<code>setEllipseParameters(center, radius[, vscale])</code>	Set the parameters defining the ellipse region.
<code>setFromOptions()</code>	Set RG options from the options database.
<code>setIntervalEndpoints(a, b, c, d)</code>	Set the parameters defining the interval region.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all RG options in the database.
<code>setPolygonVertices(v)</code>	Set the vertices that define the polygon region.
<code>setRingParameters(center, radius, vscale, ...)</code>	Set the parameters defining the ring region.
<code>setScale([sfactor])</code>	Set the scaling factor to be used.
<code>setType(rg_type)</code>	Set the type for the RG object.
<code>view([viewer])</code>	Print the RG data structure.

## Attributes Summary

<code>complement</code>	If the region is the complement of the specified one.
<code>scale</code>	The scaling factor to be used.

## Methods Documentation

### **appendOptionsPrefix**(*prefix=None*)

Append to the prefix used for searching for all RG options in the database.

Logically collective.

#### **Parameters**

**prefix** (*str* / *None*) – The prefix string to prepend to all RG option requests.

#### **Return type**

*None*

#### **See also**

*setOptionsPrefix*, *getOptionsPrefix*, *RGAppendOptionsPrefix*

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:190 <slepc4py/SLEPc/RG.pyx#L190>](#)

### **canUseConjugates**(*realmats=True*)

Half of integration points can be avoided (use their conjugates).

Not collective.

Used in contour integral methods to determine whether half of integration points can be avoided (use their conjugates).

#### **Parameters**

**realmats** (*bool*) – True if the problem matrices are real.

#### **Returns**

Whether it is possible to use conjugates.

#### **Return type**

*bool*

## **Notes**

If some integration points are the conjugates of other points, then the associated computational cost can be saved. This depends on the problem matrices being real and also the region being symmetric with respect to the horizontal axis. The result is *false* if using real arithmetic or in the case of a flat region (height equal to zero).

#### **See also**

*isAxisymmetric*, *RGCanUseConjugates*

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:458 <slepc4py/SLEPc/RG.pyx#L458>](#)

### **checkInside**(*a*)

Determine if a set of given points are inside the region or not.

Not collective.

#### **Parameters**

**a** (*Sequence[complex]*) – The coordinates of the points.

**Returns**

Computed result for each point (1=inside, 0=on the contour, -1=outside).

**Return type**

*ArrayInt*

**Notes**

If a scaling factor was set, the points are scaled before checking.

**See also**

*setScale, setComplement, RGCheckInside*

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:355 <slepc4py/SLEPc/RG.pyx#L355>`

**computeBoundingBox()**

Compute box containing the region.

Not collective.

Determine the endpoints of a rectangle in the complex plane that contains the region.

**Returns**

- **a** (*float*) – The left endpoint of the bounding box in the real axis.
- **b** (*float*) – The right endpoint of the bounding box in the real axis.
- **c** (*float*) – The bottom endpoint of the bounding box in the imaginary axis.
- **d** (*float*) – The top endpoint of the bounding box in the imaginary axis.

**Return type**

*tuple*[*float*, *float*, *float*, *float*]

**See also**

*computeContour, setScale, RGComputeBoundingBox*

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:430 <slepc4py/SLEPc/RG.pyx#L430>`

**computeContour(*n*)**

Compute points on the contour of the region.

Not collective.

Compute the coordinates of several points lying on the contour of the region.

**Parameters**

**n** (*int*) – The number of points to compute.

**Returns**

Computed points.

**Return type**

*list* of *complex*

#### See also

[computeBoundingBox](#), [setScale](#), [RGComputeContour](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:395 <slepc4py/SLEPc/RG.pyx#L395>](#)

**computeQuadrature**(*quad*, *n*)

Compute the values of the parameters used in a quadrature rule.

Not collective.

Compute the values of the parameters used in a quadrature rule for a contour integral around the boundary of the region.

#### Parameters

- **quad** ([QuadRule](#)) – The type of quadrature.
- **n** ([int](#)) – The number of quadrature points to compute.

#### Returns

- **z** ([ArrayScalar](#)) – Quadrature points.
- **zn** ([ArrayScalar](#)) – Normalized quadrature points.
- **w** ([ArrayScalar](#)) – Quadrature weights.

#### Return type

[tuple](#)[[ArrayScalar](#), [ArrayScalar](#), [ArrayScalar](#)]

#### Notes

In complex scalars, the values returned in **z** are often the same as those computed by [computeContour\(\)](#), but this is not the case in real scalars where all output arguments are real.

The computed values change for different quadrature rules.

#### See also

[computeContour](#), [RGComputeQuadrature](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:494 <slepc4py/SLEPc/RG.pyx#L494>](#)

**create**(*comm=None*)

Create the RG object.

Collective.

#### Parameters

**comm** ([Comm](#) | [None](#)) – MPI communicator; if not provided, it defaults to all processes.

#### Return type

[Self](#)

#### See also

[RGCreate](#)

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:87 <slepc4py/SLEPc/RG.pyx#L87>`

### **destroy()**

Destroy the RG object.

Collective.

#### **See also**

`RGDestroy`

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:73 <slepc4py/SLEPc/RG.pyx#L73>`

### **Return type**

*Self*

### **getComplement()**

Get the flag indicating whether the region is complemented or not.

Not collective.

### **Returns**

Whether the region is complemented or not.

### **Return type**

`bool`

#### **See also**

`setComplement`, `RGGetComplement`

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:277 <slepc4py/SLEPc/RG.pyx#L277>`

### **getEllipseParameters()**

Get the parameters that define the ellipse region.

Not collective.

### **Returns**

- **center** (*Scalar*) – The center.
- **radius** (`float`) – The radius.
- **vscale** (`float`) – The vertical scale.

### **Return type**

`tuple[Scalar, float, float]`

#### **See also**

`setEllipseParameters`, `RGEllipseGetParameters`

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:572 <slepc4py/SLEPc/RG.pyx#L572>`



### **getIntervalEndpoints()**

Get the parameters that define the interval region.

Not collective.

#### **Returns**

- **a** (`float`) – The left endpoint in the real axis.
- **b** (`float`) – The right endpoint in the real axis.
- **c** (`float`) – The bottom endpoint in the imaginary axis.
- **d** (`float`) – The top endpoint in the imaginary axis.

#### **Return type**

`tuple[float, float, float, float]`

#### **See also**

`setIntervalEndpoints`, `RGIntervalGetEndpoints`

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:634 <slepc4py/SLEPc/RG.pyx#L634>`

### **getOptionsPrefix()**

Get the prefix used for searching for all RG options in the database.

Not collective.

#### **Returns**

The prefix string set for this RG object.

#### **Return type**

`str`

#### **See also**

`setOptionsPrefix`, `appendOptionsPrefix`, `RGGetOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:171 <slepc4py/SLEPc/RG.pyx#L171>`

### **getPolygonVertices()**

Get the parameters that define the interval region.

Not collective.

#### **Returns**

The vertices.

#### **Return type**

`ArrayComplex`

#### **See also**

`setPolygonVertices`, `RGPolygonGetVertices`

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:690 <slepc4py/SLEPc/RG.pyx#L690>`

### **getRingParameters()**

Get the parameters that define the ring region.

Not collective.

#### **Returns**

- **center** (*Scalar*) – The center.
- **radius** (*float*) – The radius.
- **vscale** (*float*) – The vertical scale.
- **start\_ang** (*float*) – The right-hand side angle.
- **end\_ang** (*float*) – The left-hand side angle.
- **width** (*float*) – The width of the ring.

#### **Return type**

`tuple[Scalar, float, float, float, float, float]`

#### **See also**

`setRingParameters`, `RGRingGetParameters`

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:775 <slepc4py/SLEPc/RG.pyx#L775>`

### **getScale()**

Get the scaling factor.

Not collective.

#### **Returns**

The scaling factor.

#### **Return type**

`float`

#### **See also**

`setScale`, `RGGetScale`

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:336 <slepc4py/SLEPc/RG.pyx#L336>`

### **getType()**

Get the RG type of this object.

Not collective.

#### **Returns**

The region type currently being used.

#### **Return type**

`str`

#### **See also**

`setType`, `RGGetType`

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:127](#) <slepc4py/SLEPc/RG.pyx#L127>

**isAxisymmetric**(*vertical=False*)

Determine if the region is axisymmetric.

Not collective.

Determine if the region is symmetric with respect to the real or imaginary axis.

**Parameters**

**vertical** (*bool*) – True if symmetry must be checked against the vertical axis.

**Returns**

True if the region is axisymmetric.

**Return type**

*bool*

**See also**

[canUseConjugates](#), [RGIsAxisymmetric](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:249](#) <slepc4py/SLEPc/RG.pyx#L249>

**isTrivial**()

Tell whether it is the trivial region (whole complex plane).

Not collective.

**Returns**

True if the region is equal to the whole complex plane, e.g., an interval region with all four endpoints unbounded or an ellipse with infinite radius.

**Return type**

*bool*

**See also**

[checkInside](#), [RGIsTrivial](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:228](#) <slepc4py/SLEPc/RG.pyx#L228>

**setComplement**(*comp=True*)

Set a flag to indicate that the region is the complement of the specified one.

Logically collective.

**Parameters**

**comp** (*bool*) – Activate/deactivate the complementation of the region.

**Return type**

*None*

**See also**

[getComplement](#), [RGSetComplement](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:296](#) <slepc4py/SLEPc/RG.pyx#L296>

**setEllipseParameters**(*center, radius, vscale=None*)

Set the parameters defining the ellipse region.

Logically collective.

**Parameters**

- **center** (*Scalar*) – The center.
- **radius** (*float*) – The radius.
- **vscale** (*float* / *None*) – The vertical scale.

**Return type**

*None*

**Notes**

When PETSc is built with real scalars, the center is restricted to a real value.

**See also**

*getEllipseParameters*, *RGEllipseSetParameters*

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:542 <slepc4py/SLEPc/RG.pyx#L542>`

**setFromOptions**()

Set RG options from the options database.

Collective.

**Notes**

To see all options, run your program with the `-help` option.

**See also**

*setOptionsPrefix*, *RGSetFromOptions*

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:209 <slepc4py/SLEPc/RG.pyx#L209>`

**Return type**

*None*

**setIntervalEndpoints**(*a, b, c, d*)

Set the parameters defining the interval region.

Logically collective.

**Parameters**

- **a** (*float*) – The left endpoint in the real axis.
- **b** (*float*) – The right endpoint in the real axis.
- **c** (*float*) – The bottom endpoint in the imaginary axis.
- **d** (*float*) – The top endpoint in the imaginary axis.

**Return type**

*None*

## Notes

The region is defined as  $[a, b]x[c, d]$ . Particular cases are an interval on the real axis ( $c = d = 0$ ), similarly for the imaginary axis ( $a = b = 0$ ), the whole complex plane ( $a = -\infty, b = \infty, c = -\infty, d = \infty$ ), and so on.

When PETSc is built with real scalars, the region must be symmetric with respect to the real axis.

### See also

[`getIntervalEndpoints`](#), [`RGIntervalSetEndpoints`](#)

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:597 <slepc4py/SLEPc/RG.pyx#L597>`

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all RG options in the database.

Logically collective.

#### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all RG option requests.

#### Return type

*None*

## Notes

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

### See also

[`appendOptionsPrefix`](#), [`getOptionsPrefix`](#), [`RGGetOptionsPrefix`](#)

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:146 <slepc4py/SLEPc/RG.pyx#L146>`

**setPolygonVertices**(*v*)

Set the vertices that define the polygon region.

Logically collective.

#### Parameters

**v** (*Sequence[float]* / *Sequence[Scalar]*) – The vertices.

#### Return type

*None*

### See also

[`getPolygonVertices`](#), [`RGPolygonSetVertices`](#)

**:sources:** `Source code at slepc4py/SLEPc/RG.pyx:662 <slepc4py/SLEPc/RG.pyx#L662>`

**setRingParameters**(*center, radius, vscale, start\_ang, end\_ang, width*)

Set the parameters defining the ring region.

Logically collective.

### Parameters

- **center** (*Scalar*) – The center.
- **radius** (*float*) – The radius.
- **vscale** (*float*) – The vertical scale.
- **start\_ang** (*float*) – The right-hand side angle.
- **end\_ang** (*float*) – The left-hand side angle.
- **width** (*float*) – The width of the ring.

### Return type

None

### Notes

The values of **center**, **radius** and **vscale** have the same meaning as in the ellipse region. The **start\_ang** and **end\_ang** define the span of the ring (by default it is the whole ring), while the **width** is the separation between the two concentric ellipses (above and below the radius by  $\text{width}/2$ ).

The start and end angles are expressed as a fraction of the circumference. The allowed range is  $[0, \dots, 1]$ , with 0 corresponding to 0 radians, 0.25 to  $\pi/2$  radians, and so on. It is allowed to have **start\_ang** > **end\_ang**, in which case the ring region crosses over the zero angle.

When PETSc is built with real scalars, the center is restricted to a real value, and the start and end angles must be such that the region is symmetric with respect to the real axis.

#### See also

[\*getRingParameters\*](#), [\*RGRingSetParameters\*](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:716](#) <[slepc4py/SLEPc/RG.pyx#L716](#)>

### **setScale**(*sfactor=None*)

Set the scaling factor to be used.

Logically collective.

Set the scaling factor to be used when checking that a point is inside the region and when computing the contour.

### Parameters

- **sfactor** (*float*) – The scaling factor (default=1).

### Return type

None

#### See also

[\*getScale\*](#), [\*checkInside\*](#), [\*computeContour\*](#), [\*RGSetScale\*](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:314](#) <[slepc4py/SLEPc/RG.pyx#L314](#)>

### **setType**(*rg\_type*)

Set the type for the RG object.

Logically collective.

#### Parameters

**rg\_type** (*Type* / *str*) – The region type to be used.

#### Return type

None

#### See also

[getType](#), [RGSetType](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:108 <slepc4py/SLEPc/RG.pyx#L108>](#)

**view**(*viewer=None*)

Print the RG data structure.

Collective.

#### Parameters

**viewer** (*Viewer* / *None*) – Visualization context; if not provided, the standard output is used.

#### Return type

None

#### See also

[RGView](#)

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:54 <slepc4py/SLEPc/RG.pyx#L54>](#)

### Attributes Documentation

#### **complement**

If the region is the complement of the specified one.

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:811 <slepc4py/SLEPc/RG.pyx#L811>](#)

#### **scale**

The scaling factor to be used.

**:sources:** [Source code at slepc4py/SLEPc/RG.pyx:818 <slepc4py/SLEPc/RG.pyx#L818>](#)

### **slepc4py.SLEPc.ST**

**class** `slepc4py.SLEPc.ST`

Bases: [Object](#)

Spectral Transformation.

The Spectral Transformation ([ST](#)) class encapsulates the functionality required for acceleration techniques based on the transformation of the spectrum. The eigensolvers implemented in [EPS](#) work by applying an operator to a set of vectors and this operator can adopt different forms. The [ST](#) object handles all the different possibilities in a uniform way, so that the solver can proceed without knowing which transformation has been selected. Polynomial eigensolvers in [PEP](#) also support spectral transformation.

## Enumerations

<i>FilterDamping</i>	ST filter damping.
<i>FilterType</i>	ST filter type.
<i>MatMode</i>	ST matrix mode.
<i>Type</i>	ST type.

### slepc4py.SLEPc.ST.FilterDamping

**class** slepc4py.SLEPc.ST.**FilterDamping**

Bases: `object`

ST filter damping.

- *NONE*: No damping
- *JACKSON*: Jackson damping
- *LANCZOS*: Lanczos damping
- *FEJER*: Fejer damping

#### See also

STFilterDamping

### Attributes Summary

<i>FEJER</i>	Constant FEJER of type <code>int</code>
<i>JACKSON</i>	Constant JACKSON of type <code>int</code>
<i>LANCZOS</i>	Constant LANCZOS of type <code>int</code>
<i>NONE</i>	Constant NONE of type <code>int</code>

### Attributes Documentation

**FEJER:** `int` = **FEJER**

Constant FEJER of type `int`

**JACKSON:** `int` = **JACKSON**

Constant JACKSON of type `int`

**LANCZOS:** `int` = **LANCZOS**

Constant LANCZOS of type `int`

**NONE:** `int` = **NONE**

Constant NONE of type `int`

### slepc4py.SLEPc.ST.FilterType

**class** slepc4py.SLEPc.ST.**FilterType**

Bases: `object`

ST filter type.



- **FILTLAN**: An adapted implementation of the Filtered Lanczos Package.
- **CHEBYSHEV**: A polynomial filter based on a truncated Chebyshev series.

#### See also

[STFilterType](#)

### Attributes Summary

<i>CHEBYSHEV</i>	Constant CHEBYSHEV of type <code>int</code>
<i>FILTLAN</i>	Constant FILTLAN of type <code>int</code>

### Attributes Documentation

**CHEBYSHEV**: `int` = **CHEBYSHEV**

Constant CHEBYSHEV of type `int`

**FILTLAN**: `int` = **FILTLAN**

Constant FILTLAN of type `int`

### `slepc4py.SLEPc.ST.MatMode`

**class** `slepc4py.SLEPc.ST.MatMode`

Bases: `object`

ST matrix mode.

- *COPY*: A working copy of the matrix is created.
- *INPLACE*: The operation is computed in-place.
- *SHELL*: The matrix  $A - \sigma B$  is handled as an implicit matrix.

#### See also

[STMatMode](#)

### Attributes Summary

<i>COPY</i>	Constant COPY of type <code>int</code>
<i>INPLACE</i>	Constant INPLACE of type <code>int</code>
<i>SHELL</i>	Constant SHELL of type <code>int</code>

### Attributes Documentation

**COPY**: `int` = **COPY**

Constant COPY of type `int`

**INPLACE**: `int` = **INPLACE**

Constant INPLACE of type `int`

**SHELL:** `int` = **SHELL**  
Constant SHELL of type `int`

### `slepc4py.SLEPc.ST.Type`

**class** `slepc4py.SLEPc.ST.Type`

Bases: `object`

ST type.

- *SHIFT*: Shift from origin.
- *SINVERT*: Shift-and-invert.
- *CAYLEY*: Cayley transform.
- *PRECOND*: Preconditioner.
- *FILTER*: Polynomial filter.
- *SHELL*: User-defined.

#### See also

`STType`

### Attributes Summary

<i>CAYLEY</i>	Object CAYLEY of type <code>str</code>
<i>FILTER</i>	Object FILTER of type <code>str</code>
<i>PRECOND</i>	Object PRECOND of type <code>str</code>
<i>SHELL</i>	Object SHELL of type <code>str</code>
<i>SHIFT</i>	Object SHIFT of type <code>str</code>
<i>SINVERT</i>	Object SINVERT of type <code>str</code>

### Attributes Documentation

**CAYLEY:** `str` = **CAYLEY**

Object CAYLEY of type `str`

**FILTER:** `str` = **FILTER**

Object FILTER of type `str`

**PRECOND:** `str` = **PRECOND**

Object PRECOND of type `str`

**SHELL:** `str` = **SHELL**

Object SHELL of type `str`

**SHIFT:** `str` = **SHIFT**

Object SHIFT of type `str`

**SINVERT:** `str` = **SINVERT**

Object SINVERT of type `str`

## Methods Summary

<code>appendOptionsPrefix([prefix])</code>	Append to the prefix used for searching for all ST options in the database.
<code>apply(x, y)</code>	Apply the spectral transformation operator to a vector.
<code>applyHermitianTranspose(x, y)</code>	Apply the Hermitian-transpose of the operator to a vector.
<code>applyMat(X, Y)</code>	Apply the spectral transformation operator to a matrix.
<code>applyTranspose(x, y)</code>	Apply the transpose of the operator to a vector.
<code>create([comm])</code>	Create the ST object.
<code>destroy()</code>	Destroy the ST object.
<code>getCayleyAntishift()</code>	Get the value of the anti-shift for the Cayley spectral transformation.
<code>getFilterDamping()</code>	Get the type of damping used in the polynomial filter.
<code>getFilterDegree()</code>	Get the degree of the filter polynomial.
<code>getFilterInterval()</code>	Get the interval containing the desired eigenvalues.
<code>getFilterRange()</code>	Get the interval containing all eigenvalues.
<code>getFilterType()</code>	Get the method to be used to build the polynomial filter.
<code>getKSP()</code>	Get the KSP object associated with the spectral transformation.
<code>getMatMode()</code>	Get a flag that indicates how the matrix is being shifted.
<code>getMatStructure()</code>	Get the internal matrix structure attribute.
<code>getMatrices()</code>	Get the matrices associated with the eigenvalue problem.
<code>getOperator()</code>	Get a shell matrix that represents the operator of the spectral transformation.
<code>getOptionsPrefix()</code>	Get the prefix used for searching for all ST options in the database.
<code>getPreconditionerMat()</code>	Get the matrix previously set by <code>setPreconditionerMat()</code> .
<code>getShift()</code>	Get the shift associated with the spectral transformation.
<code>getSplitPreconditioner()</code>	Get the matrices to be used to build the preconditioner.
<code>getTransform()</code>	Get the flag indicating whether the transformed matrices are computed or not.
<code>getType()</code>	Get the ST type of this object.
<code>reset()</code>	Reset the ST object.
<code>restoreOperator(op)</code>	Restore the previously seized operator matrix.
<code>setCayleyAntishift(mu)</code>	Set the value of the anti-shift for the Cayley spectral transformation.
<code>setFilterDamping(damping)</code>	Set the type of damping to be used in the polynomial filter.
<code>setFilterDegree(deg)</code>	Set the degree of the filter polynomial.
<code>setFilterInterval(inta, intb)</code>	Set the interval containing the desired eigenvalues.
<code>setFilterRange(left, right)</code>	Set the numerical range (or field of values) of the matrix.
<code>setFilterType(filter_type)</code>	Set the method to be used to build the polynomial filter.

continues on next page

Table 95 – continued from previous page

<code>setFromOptions()</code>	Set ST options from the options database.
<code>setKSP(ksp)</code>	Set the KSP object associated with the spectral transformation.
<code>setMatMode(mode)</code>	Set a flag related to management of transformed matrices.
<code>setMatStructure(structure)</code>	Set the matrix structure attribute.
<code>setMatrices(operators)</code>	Set the matrices associated with the eigenvalue problem.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all ST options in the database.
<code>setPreconditionerMat([P])</code>	Set the matrix to be used to build the preconditioner.
<code>setShift(shift)</code>	Set the shift associated with the spectral transformation.
<code>setSplitPreconditioner(operators[, structure])</code>	Set the matrices to be used to build the preconditioner.
<code>setTransform([flag])</code>	Set a flag to indicate whether the transformed matrices are computed or not.
<code>setType(st_type)</code>	Set the particular spectral transformation to be used.
<code>setUp()</code>	Prepare for the use of a spectral transformation.
<code>view([viewer])</code>	Print the ST data structure.

## Attributes Summary

<code>ksp</code>	KSP object associated with the spectral transformation.
<code>mat_mode</code>	How the transformed matrices are being stored in the ST.
<code>mat_structure</code>	Relation of the sparsity pattern of all ST matrices.
<code>shift</code>	Value of the shift.
<code>transform</code>	If the transformed matrices are computed.

## Methods Documentation

### `appendOptionsPrefix(prefix=None)`

Append to the prefix used for searching for all ST options in the database.

Logically collective.

#### Parameters

**prefix** (`str` / `None`) – The prefix string to prepend to all ST option requests.

#### Return type

`None`

#### See also

`setOptionsPrefix`, `getOptionsPrefix`, `STAppendOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:256 <slepc4py/SLEPc/ST.pyx#L256>`

### `apply(x, y)`

Apply the spectral transformation operator to a vector.

Collective.

Apply the spectral transformation operator to a vector, for instance  $y = (A - \sigma B)^{-1} Bx$  in the case of the shift-and-invert transformation and generalized eigenproblem.

#### Parameters

- **x** (*Vec*) – The input vector.
- **y** (*Vec*) – The result vector.

#### Return type

None

#### See also

[applyTranspose](#), [applyHermitianTranspose](#), [applyMat](#), [STApply](#)

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:758 <slepc4py/SLEPc/ST.pyx#L758>](#)

### **applyHermitianTranspose**(*x*, *y*)

Apply the Hermitian-transpose of the operator to a vector.

Collective.

Apply the Hermitian-transpose of the operator to a vector, for instance  $y = B^*(A - \sigma B)^{-*}x$  in the case of the shift-and-invert transformation and generalized eigenproblem.

#### Parameters

- **x** (*Vec*) – The input vector.
- **y** (*Vec*) – The result vector.

#### Return type

None

#### See also

[apply](#), [applyTranspose](#), [STApplyHermitianTranspose](#)

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:804 <slepc4py/SLEPc/ST.pyx#L804>](#)

### **applyMat**(*X*, *Y*)

Apply the spectral transformation operator to a matrix.

Collective.

Apply the spectral transformation operator to a matrix, for instance  $Y = (A - \sigma B)^{-1} BX$  in the case of the shift-and-invert transformation and generalized eigenproblem.

#### Parameters

- **X** (*Mat*) – The input matrix.
- **Y** (*Mat*) – The result matrix.

#### Return type

None

**See also**[apply](#), [STApplyMat](#)

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:827 <slepc4py/SLEPc/ST.pyx#L827>](#)

**applyTranspose**(*x*, *y*)

Apply the transpose of the operator to a vector.

Collective.

Apply the transpose of the operator to a vector, for instance  $y = B^T(A - \sigma B)^{-T}x$  in the case of the shift-and-invert transformation and generalized eigenproblem.

**Parameters**

- **x** (*Vec*) – The input vector.
- **y** (*Vec*) – The result vector.

**Return type**

*None*

**See also**[apply](#), [applyHermitianTranspose](#), [STApplyTranspose](#)

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:781 <slepc4py/SLEPc/ST.pyx#L781>](#)

**create**(*comm=None*)

Create the ST object.

Collective.

**Parameters**

**comm** (*Comm* / *None*) – MPI communicator; if not provided, it defaults to all processes.

**Return type**

*Self*

**See also**[STCreate](#)

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:145 <slepc4py/SLEPc/ST.pyx#L145>](#)

**destroy**()

Destroy the ST object.

Collective.

**See also**[STDestroy](#)

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:119 <slepc4py/SLEPc/ST.pyx#L119>](#)

**Return type***Self***getCayleyAntishift()**

Get the value of the anti-shift for the Cayley spectral transformation.

Not collective.

**Returns**

The anti-shift.

**Return type***Scalar***See also***setCayleyAntishift, STCayleyGetAntishift*

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:942](#) <slepc4py/SLEPc/ST.pyx#L942>

**getFilterDamping()**

Get the type of damping used in the polynomial filter.

Not collective.

**Returns**

The type of damping.

**Return type***FilterDamping***See also***setFilterDamping, STFilterGetDamping*

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1164](#) <slepc4py/SLEPc/ST.pyx#L1164>

**getFilterDegree()**

Get the degree of the filter polynomial.

Not collective.

**Returns**

The polynomial degree.

**Return type***int***See also***setFilterDegree, STFilterGetDegree*

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1123](#) <slepc4py/SLEPc/ST.pyx#L1123>

### **getFilterInterval()**

Get the interval containing the desired eigenvalues.

Not collective.

#### **Returns**

- **inta** (*float*) – The left end of the interval.
- **intb** (*float*) – The right end of the interval.

#### **Return type**

*tuple*[*float*, *float*]

#### **See also**

*setFilterInterval*, *STFilterGetInterval*

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1031 <slepc4py/SLEPc/ST.pyx#L1031>](#)

### **getFilterRange()**

Get the interval containing all eigenvalues.

Not collective.

#### **Returns**

- **left** (*float*) – The left end of the spectral range.
- **right** (*float*) – The right end of the spectral range.

#### **Return type**

*tuple*[*float*, *float*]

#### **See also**

*getFilterInterval*, *STFilterGetRange*

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1083 <slepc4py/SLEPc/ST.pyx#L1083>](#)

### **getFilterType()**

Get the method to be used to build the polynomial filter.

Not collective.

#### **Returns**

The type of filter.

#### **Return type**

*FilterType*

#### **See also**

*setFilterType*, *STFilterGetType*

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:979 <slepc4py/SLEPc/ST.pyx#L979>](#)



### **getKSP()**

Get the KSP object associated with the spectral transformation.

Collective.

#### **Returns**

The linear solver object.

#### **Return type**

`petsc4py.PETSc.KSP`

#### **See also**

`setKSP`, `STGetKSP`

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:585 <slepc4py/SLEPc/ST.pyx#L585>](#)

### **getMatMode()**

Get a flag that indicates how the matrix is being shifted.

Not collective.

Get a flag that indicates how the matrix is being shifted in the shift-and-invert and Cayley spectral transformations.

#### **Returns**

The mode flag.

#### **Return type**

`MatMode`

#### **See also**

`setMatMode`, `STGetMatMode`

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:430 <slepc4py/SLEPc/ST.pyx#L430>](#)

### **getMatStructure()**

Get the internal matrix structure attribute.

Not collective.

Get the internal `petsc4py.PETSc.Mat.Structure` attribute to indicate which is the relation of the sparsity pattern of the matrices.

#### **Returns**

The structure flag.

#### **Return type**

`petsc4py.PETSc.Mat.Structure`

#### **See also**

`setMatStructure`, `STGetMatStructure`

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:545 <slepc4py/SLEPc/ST.pyx#L545>](#)

## getMatrices()

Get the matrices associated with the eigenvalue problem.

Collective.

### Returns

The matrices associated with the eigensystem.

### Return type

list of `petsc4py.PETSc.Mat`

### See also

`setMatrices`, `STGetNumMatrices`, `STGetMatrix`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:486 <slepc4py/SLEPc/ST.pyx#L486>`

## getOperator()

Get a shell matrix that represents the operator of the spectral transformation.

Collective.

### Returns

Operator matrix.

### Return type

`petsc4py.PETSc.Mat`

## Notes

The operator is defined in linear eigenproblems only, not in polynomial ones, so the call will fail if more than 2 matrices were passed in `setMatrices()`.

The returned shell matrix is essentially a wrapper to the `apply()` and `applyTranspose()` operations. The operator can often be expressed as

$$Op = DK^{-1}MD^{-1}$$

where  $D$  is the balancing matrix, and  $M$  and  $K$  are two matrices corresponding to the numerator and denominator for spectral transformations that represent a rational matrix function.

The preconditioner matrix  $K$  typically depends on the value of the shift, and its inverse is handled via an internal KSP object. Normal usage does not require explicitly calling `getOperator()`, but it can be used to force the creation of  $K$  and  $M$ , and then  $K$  is passed to the KSP. This is useful for setting options associated with the PCFactor (to set MUMPS options, for instance).

The returned matrix must NOT be destroyed by the user. Instead, when no longer needed it must be returned with `restoreOperator()`. In particular, this is required before modifying the `ST` matrices or the shift.

### See also

`apply`, `setMatrices`, `setShift`, `restoreOperator`, `STGetOperator`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:850 <slepc4py/SLEPc/ST.pyx#L850>`

### **getOptionsPrefix()**

Get the prefix used for searching for all ST options in the database.

Not collective.

#### **Returns**

The prefix string set for this ST object.

#### **Return type**

`str`

#### **See also**

`setOptionsPrefix`, `appendOptionsPrefix`, `STGetOptionsPrefix`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:237 <slepc4py/SLEPc/ST.pyx#L237>`

### **getPreconditionerMat()**

Get the matrix previously set by `setPreconditionerMat()`.

Not collective.

#### **Returns**

The matrix that will be used in constructing the preconditioner.

#### **Return type**

`petsc4py.PETSc.Mat`

#### **See also**

`setPreconditionerMat`, `STGetPreconditionerMat`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:649 <slepc4py/SLEPc/ST.pyx#L649>`

### **getShift()**

Get the shift associated with the spectral transformation.

Not collective.

#### **Returns**

The value of the shift.

#### **Return type**

`Scalar`

#### **See also**

`setShift`, `STGetShift`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:323 <slepc4py/SLEPc/ST.pyx#L323>`

### **getSplitPreconditioner()**

Get the matrices to be used to build the preconditioner.

Not collective.

#### **Returns**

- `list` of `petsc4py.PETSc.Mat` – The list of matrices associated with the preconditioner.
- `petsc4py.PETSc.Mat.Structure` – The structure flag.

**Return type**

`tuple[list[petsc4py.PETSc.Mat], petsc4py.PETSc.Mat.Structure]`

**See also**

`STGetSplitPreconditionerInfo`, `STGetSplitPreconditionerTerm`

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:716 <slepc4py/SLEPc/ST.pyx#L716>](#)

**getTransform()**

Get the flag indicating whether the transformed matrices are computed or not.

Not collective.

**Returns**

This flag is intended for the case of polynomial eigenproblems solved via linearization. If this flag is `False` (default) the spectral transformation is applied to the linearization (handled by the eigensolver), otherwise it is applied to the original problem.

**Return type**

`bool`

**See also**

`setTransform`, `STGetTransform`

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:364 <slepc4py/SLEPc/ST.pyx#L364>](#)

**getType()**

Get the ST type of this object.

Not collective.

**Returns**

The spectral transformation currently being used.

**Return type**

`str`

**See also**

`setType`, `STGetType`

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:193 <slepc4py/SLEPc/ST.pyx#L193>](#)

**reset()**

Reset the ST object.

Collective.

<b>See also</b>
<a href="#">STReset</a>

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:133 <slepc4py/SLEPc/ST.pyx#L133>](#)

**Return type**

None

**restoreOperator**(*op*)

Restore the previously seized operator matrix.

Logically collective.

**Parameters**

**op** (*Mat*) – Operator matrix previously obtained with [getOperator\(\)](#).

**Return type**

None

<b>See also</b>
<a href="#">getOperator</a> , <a href="#">STRestoreOperator</a>

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:898 <slepc4py/SLEPc/ST.pyx#L898>](#)

**setCayleyAntishift**(*mu*)

Set the value of the anti-shift for the Cayley spectral transformation.

Logically collective.

**Parameters**

**mu** (*Scalar*) – The anti-shift.

**Return type**

None

**Notes**

In the generalized Cayley transform, the operator can be expressed as  $(A - \sigma B)^{-1}(A + \mu B)$ . This function sets the value of *mu*. Use [setShift\(\)](#) for setting  $\sigma$ .

<b>See also</b>
<a href="#">setShift</a> , <a href="#">getCayleyAntishift</a> , <a href="#">STCayleySetAntishift</a>

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:918 <slepc4py/SLEPc/ST.pyx#L918>](#)

**setFilterDamping**(*damping*)

Set the type of damping to be used in the polynomial filter.

Logically collective.

## Parameter

### damping

The type of damping.

## Notes

Only used in *FilterType.CHEBYSHEV* filters.

### See also

*getFilterDamping*, *STFilterSetDamping*

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:1142 <slepc4py/SLEPc/ST.pyx#L1142>`

### Parameters

**damping** (*FilterDamping*)

### Return type

*None*

### setFilterDegree(*deg*)

Set the degree of the filter polynomial.

Logically collective.

### Parameters

**deg** (*int*) – The polynomial degree.

### Return type

*None*

### See also

*getFilterDegree*, *STFilterSetDegree*

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:1105 <slepc4py/SLEPc/ST.pyx#L1105>`

### setFilterInterval(*inta*, *intb*)

Set the interval containing the desired eigenvalues.

Logically collective.

### Parameters

- **inta** (*float*) – The left end of the interval.
- **intb** (*float*) – The right end of the interval.

### Return type

*None*

## Notes

The filter will be configured to emphasize eigenvalues contained in the given interval, and damp out eigenvalues outside it. If the interval is open, then the filter is low- or high-pass, otherwise it is mid-pass.

Common usage is to set the interval in *EPS* with *EPS.setInterval()*.

The interval must be contained within the numerical range of the matrix, see `setFilterRange()`.

**See also**

`getFilterInterval`, `setFilterRange`, `STFilterSetInterval`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:998 <slepc4py/SLEPc/ST.pyx#L998>`

**setFilterRange**(*left*, *right*)

Set the numerical range (or field of values) of the matrix.

Logically collective.

Set the numerical range (or field of values) of the matrix, that is, the interval containing all eigenvalues.

**Parameters**

- **left** (*float*) – The left end of the spectral range.
- **right** (*float*) – The right end of the spectral range.

**Return type**

None

**Notes**

The filter will be most effective if the numerical range is tight, that is, `left` and `right` are good approximations to the leftmost and rightmost eigenvalues, respectively.

**See also**

`setFilterInterval`, `getFilterRange`, `STFilterSetRange`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:1053 <slepc4py/SLEPc/ST.pyx#L1053>`

**setFilterType**(*filter\_type*)

Set the method to be used to build the polynomial filter.

Logically collective.

**Parameter**

**filter\_type**

The type of filter.

**See also**

`getFilterType`, `STFilterSetType`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:961 <slepc4py/SLEPc/ST.pyx#L961>`

**Parameters**

**filter\_type** (*FilterType*)

**Return type**

None

## **setFromOptions()**

Set ST options from the options database.

Collective.

### **Notes**

To see all options, run your program with the `-help` option.

This routine must be called before `setUp()` if the user is to be allowed to set the solver type.

#### **See also**

`setOptionsPrefix`, `STSetFromOptions`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:275 <slepc4py/SLEPc/ST.pyx#L275>`

#### **Return type**

None

## **setKSP(*ksp*)**

Set the KSP object associated with the spectral transformation.

Collective.

#### **Parameters**

**ksp** (*KSP*) – The linear solver object.

#### **Return type**

None

#### **See also**

`getKSP`, `STSetKSP`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:568 <slepc4py/SLEPc/ST.pyx#L568>`

## **setMatMode(*mode*)**

Set a flag related to management of transformed matrices.

Logically collective.

The flag indicates how the transformed matrices are being stored in the spectral transformation.

#### **Parameters**

**mode** (*MatMode*) – The mode flag.

#### **Return type**

None

### **Notes**

By default (`ST.MatMode.COPY`), a copy of matrix  $A$  is made and then this copy is modified explicitly, e.g.,  $A \leftarrow (A - \sigma B)$ .

With `ST.MatMode.INPLACE`, the original matrix  $A$  is modified at `setUp()` and reverted at the end of the computations. With respect to the previous one, this mode avoids a copy of matrix  $A$ . However, a drawback is that the recovered matrix might be slightly different from the original one (due to roundoff).



With `ST.MatMode.SHELL`, the solver works with an implicit shell matrix that represents the shifted matrix. This mode is the most efficient in creating the transformed matrix but it places serious limitations to the linear solves performed in each iteration of the eigensolver (typically, only iterative solvers with Jacobi preconditioning can be used).

In the two first modes the efficiency of this computation can be controlled with `setMatStructure()`.

#### See also

`setMatrices`, `setMatStructure`, `getMatMode`, `STSetMatMode`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:387 <slepc4py/SLEPc/ST.pyx#L387>`

**setMatStructure**(*structure*)

Set the matrix structure attribute.

Logically collective.

Set an internal `petsc4py.PETSc.Mat.Structure` attribute to indicate which is the relation of the sparsity pattern of all the `ST` matrices.

#### Parameters

**structure** (`petsc4py.PETSc.Mat.Structure`) – The matrix structure specification.

#### Return type

`None`

#### Notes

By default, the sparsity patterns are assumed to be different. If the patterns are equal or a subset then it is recommended to set this attribute for efficiency reasons (in particular, for internal `Mat.axy()` operations).

This function has no effect in the case of standard eigenproblems.

In case of polynomial eigenproblems, the flag applies to all matrices relative to the first one.

#### See also

`getMatStructure`, `setMatrices`, `STSetMatStructure`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:512 <slepc4py/SLEPc/ST.pyx#L512>`

**setMatrices**(*operators*)

Set the matrices associated with the eigenvalue problem.

Collective.

#### Parameters

**operators** (`list[Mat]`) – The matrices associated with the eigensystem.

#### Return type

`None`

#### Notes

It must be called before `setUp()`. If it is called again after `setUp()` then the `ST` object is reset.

In standard eigenproblems only one matrix is passed, while in generalized problems two matrices are provided. The number of matrices is larger in polynomial eigenproblems.

In normal usage, matrices are provided via the corresponding [EPS](#) of [PEP](#) interface function.

#### See also

[getMatrices](#), [setUp](#), [reset](#), [STSetMatrices](#)

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:452 <slepc4py/SLEPc/ST.pyx#L452>`

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all ST options in the database.

Logically collective.

#### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all ST option requests.

#### Return type

*None*

#### Notes

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

#### See also

[appendOptionsPrefix](#), [getOptionsPrefix](#), [STGetOptionsPrefix](#)

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:212 <slepc4py/SLEPc/ST.pyx#L212>`

**setPreconditionerMat**(*P=None*)

Set the matrix to be used to build the preconditioner.

Collective.

#### Parameters

**P** (*Mat* / *None*) – The matrix that will be used in constructing the preconditioner.

#### Return type

*None*

#### Notes

This matrix will be passed to the internal KSP object (via the last argument of `KSP.setOperators()`) as the matrix to be used when constructing the preconditioner. If no matrix is set then  $A - \sigma B$  will be used to build the preconditioner, being  $\sigma$  the value set by [setShift\(\)](#).

More precisely, this is relevant for spectral transformations that represent a rational matrix function, and use a KSP object for the denominator. It includes also the [PRECOND](#) case. If the user has a good approximation to matrix that can be used to build a cheap preconditioner, it can be passed with this function. Note that it affects only the `Pmat` argument of `KSP.setOperators()`, not the `Amat` argument.

If a preconditioner matrix is set, the default is to use an iterative KSP rather than a direct method.

An alternative to pass an approximation of  $A - \sigma B$  with this function is to provide approximations of  $A$  and  $B$  via [setSplitPreconditioner\(\)](#). The difference is that when  $\sigma$  changes the preconditioner is recomputed.

A call with no matrix argument will remove a previously set matrix.

#### See also

[`getPreconditionerMat`](#), [`STSetPreconditionerMat`](#)

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:605 <slepc4py/SLEPc/ST.pyx#L605>`

#### **setShift**(*shift*)

Set the shift associated with the spectral transformation.

Collective.

##### Parameters

**shift** ([`Scalar`](#)) – The value of the shift.

##### Return type

[`None`](#)

#### Notes

In some spectral transformations, changing the shift may have associated a lot of work, for example recomputing a factorization.

This function is normally not directly called by users, since the shift is indirectly set by [`EPS.setTarget\(\)`](#).

#### See also

[`getShift`](#), [`STSetShift`](#)

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:296 <slepc4py/SLEPc/ST.pyx#L296>`

#### **setSplitPreconditioner**(*operators*, *structure=None*)

Set the matrices to be used to build the preconditioner.

Collective.

##### Parameters

- **operators** ([`list\[petsc4py.PETSc.Mat\]`](#)) – The matrices associated with the preconditioner.
- **structure** ([`petsc4py.PETSc.Mat.Structure`](#) / [`None`](#)) – The matrix structure specification.

##### Return type

[`None`](#)

#### Notes

The number of matrices passed here must be the same as in [`setMatrices\(\)`](#).

For linear eigenproblems, the preconditioner matrix is computed as  $P(\sigma) = A_0 - \sigma B_0$ , where  $A_0, B_0$  are approximations of  $A, B$  (the eigenproblem matrices) provided via the **operators** argument in this function. Compared to [`setPreconditionerMat\(\)`](#), this function allows setting a preconditioner in a way that is independent of the shift  $\sigma$ . Whenever the value of  $\sigma$  changes the preconditioner is recomputed.

Similarly, for polynomial eigenproblems the matrix for the preconditioner is expressed as  $P(\sigma) = \sum_i P_i \phi_i(\sigma)$ , for  $i = 1, \dots, n$ , where  $P_i$  are given in operators and the  $\phi_i$ 's are the polynomial basis functions.

The structure flag provides information about the relative nonzero pattern of the operators matrices, in the same way as in `setMatStructure()`.

#### See also

`getSplitPreconditioner`, `setPreconditionerMat`, `STSetSplitPreconditioner`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:669 <slepc4py/SLEPc/ST.pyx#L669>`

**setTransform**(*flag=True*)

Set a flag to indicate whether the transformed matrices are computed or not.

Logically collective.

#### Parameters

**flag** (*bool*) – This flag is intended for the case of polynomial eigenproblems solved via linearization. If this flag is `False` (default) the spectral transformation is applied to the linearization (handled by the eigensolver), otherwise it is applied to the original problem.

#### Return type

`None`

#### See also

`getTransform`, `STSetTransform`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:342 <slepc4py/SLEPc/ST.pyx#L342>`

**setType**(*st\_type*)

Set the particular spectral transformation to be used.

Logically collective.

#### Parameters

**st\_type** (*Type* / *str*) – The spectral transformation to be used.

#### Return type

`None`

### Notes

The default is `SHIFT` with a zero shift. Normally, it is best to use `setFromOptions()` and then set the ST type from the options database rather than by using this routine. Using the options database provides the user with maximum flexibility in evaluating the different available methods.

#### See also

`getType`, `STSetType`

**:sources:** `Source code at slepc4py/SLEPc/ST.pyx:166 <slepc4py/SLEPc/ST.pyx#L166>`

## setUp()

Prepare for the use of a spectral transformation.

Collective.

### See also

[apply](#), [STSetUp](#)

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:746 <slepc4py/SLEPc/ST.pyx#L746>](#)

### Return type

None

## view(viewer=None)

Print the ST data structure.

Collective.

### Parameters

**viewer** ([Viewer](#) / [None](#)) – Visualization context; if not provided, the standard output is used.

### Return type

None

### See also

[STView](#)

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:100 <slepc4py/SLEPc/ST.pyx#L100>](#)

## Attributes Documentation

### ksp

KSP object associated with the spectral transformation.

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1213 <slepc4py/SLEPc/ST.pyx#L1213>](#)

### mat\_mode

How the transformed matrices are being stored in the ST.

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1199 <slepc4py/SLEPc/ST.pyx#L1199>](#)

### mat\_structure

Relation of the sparsity pattern of all ST matrices.

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1206 <slepc4py/SLEPc/ST.pyx#L1206>](#)

### shift

Value of the shift.

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1185 <slepc4py/SLEPc/ST.pyx#L1185>](#)

### transform

If the transformed matrices are computed.

**:sources:** [Source code at slepc4py/SLEPc/ST.pyx:1192 <slepc4py/SLEPc/ST.pyx#L1192>](#)

## slepc4py.SLEPc.SVD

**class** slepc4py.SLEPc.SVD

Bases: [Object](#)

Singular Value Decomposition Solver.

The Singular Value Decomposition Solver ([SVD](#)) is very similar to the [EPS](#) object, but intended for the computation of the partial SVD of a rectangular matrix. With this type of object, the user can specify an SVD problem and solve it with any of the different solvers encapsulated by the package. Some of these solvers are actually implemented through calls to [EPS](#) eigensolvers.

### Enumerations

<a href="#">Conv</a>	SVD convergence test.
<a href="#">ConvergedReason</a>	SVD convergence reasons.
<a href="#">ErrorType</a>	SVD error type to assess accuracy of computed solutions.
<a href="#">ProblemType</a>	SVD problem type.
<a href="#">Stop</a>	SVD stopping test.
<a href="#">TRLanczosGBidiag</a>	SVD TRLanczos bidiagonalization choices for the GSVD case.
<a href="#">Type</a>	SVD type.
<a href="#">Which</a>	SVD desired part of spectrum.

## slepc4py.SLEPc.SVD.Conv

**class** slepc4py.SLEPc.SVD.Conv

Bases: [object](#)

SVD convergence test.

- [ABS](#): Absolute convergence test.
- [REL](#): Convergence test relative to the singular value.
- [NORM](#): Convergence test relative to the matrix norms.
- [MAXIT](#): No convergence until maximum number of iterations has been reached.
- [USER](#): User-defined convergence test.

### See also

[SVDConv](#)

### Attributes Summary

<a href="#">ABS</a>	Constant ABS of type <a href="#">int</a>
<a href="#">MAXIT</a>	Constant MAXIT of type <a href="#">int</a>
<a href="#">NORM</a>	Constant NORM of type <a href="#">int</a>
<a href="#">REL</a>	Constant REL of type <a href="#">int</a>
<a href="#">USER</a>	Constant USER of type <a href="#">int</a>

## Attributes Documentation

**ABS:** `int` = `ABS`

Constant ABS of type `int`

**MAXIT:** `int` = `MAXIT`

Constant MAXIT of type `int`

**NORM:** `int` = `NORM`

Constant NORM of type `int`

**REL:** `int` = `REL`

Constant REL of type `int`

**USER:** `int` = `USER`

Constant USER of type `int`

## `slepc4py.SLEPc.SVD.ConvergedReason`

**class** `slepc4py.SLEPc.SVD.ConvergedReason`

Bases: `object`

SVD convergence reasons.

- `CONVERGED_TOL`: All eigenpairs converged to requested tolerance.
- `CONVERGED_USER`: User-defined convergence criterion satisfied.
- `CONVERGED_MAXIT`: Maximum iterations completed in case MAXIT convergence criterion.
- `DIVERGED_ITS`: Maximum number of iterations exceeded.
- `DIVERGED_BREAKDOWN`: Solver failed due to breakdown.
- `DIVERGED_SYMMETRY_LOST`: Underlying indefinite eigensolver was not able to keep symmetry.
- `CONVERGED_ITERATING`: Iteration not finished yet.

### See also

`SVDConvergedReason`

## Attributes Summary

<code>CONVERGED_ITERATING</code>	Constant <code>CONVERGED_ITERATING</code> of type <code>int</code>
<code>CONVERGED_MAXIT</code>	Constant <code>CONVERGED_MAXIT</code> of type <code>int</code>
<code>CONVERGED_TOL</code>	Constant <code>CONVERGED_TOL</code> of type <code>int</code>
<code>CONVERGED_USER</code>	Constant <code>CONVERGED_USER</code> of type <code>int</code>
<code>DIVERGED_BREAKDOWN</code>	Constant <code>DIVERGED_BREAKDOWN</code> of type <code>int</code>
<code>DIVERGED_ITS</code>	Constant <code>DIVERGED_ITS</code> of type <code>int</code>
<code>DIVERGED_SYMMETRY_LOST</code>	Constant <code>DIVERGED_SYMMETRY_LOST</code> of type <code>int</code>
<code>ITERATING</code>	Constant <code>ITERATING</code> of type <code>int</code>

Attributes Documentation

**CONVERGED\_ITERATING:** `int` = `CONVERGED_ITERATING`  
Constant `CONVERGED_ITERATING` of type `int`

**CONVERGED\_MAXIT:** `int` = `CONVERGED_MAXIT`  
Constant `CONVERGED_MAXIT` of type `int`

**CONVERGED\_TOL:** `int` = `CONVERGED_TOL`  
Constant `CONVERGED_TOL` of type `int`

**CONVERGED\_USER:** `int` = `CONVERGED_USER`  
Constant `CONVERGED_USER` of type `int`

**DIVERGED\_BREAKDOWN:** `int` = `DIVERGED_BREAKDOWN`  
Constant `DIVERGED_BREAKDOWN` of type `int`

**DIVERGED\_ITS:** `int` = `DIVERGED_ITS`  
Constant `DIVERGED_ITS` of type `int`

**DIVERGED\_SYMMETRY\_LOST:** `int` = `DIVERGED_SYMMETRY_LOST`  
Constant `DIVERGED_SYMMETRY_LOST` of type `int`

**ITERATING:** `int` = `ITERATING`  
Constant `ITERATING` of type `int`

`slepc4py.SLEPc.SVD.ErrorType`

`class slepc4py.SLEPc.SVD.ErrorType`  
Bases: `object`

SVD error type to assess accuracy of computed solutions.

- *ABSOLUTE*: Absolute error.
- *RELATIVE*: Relative error.
- *NORM*: Error relative to the matrix norm.

See also
<code>SVDErrorType</code>

Attributes Summary

<i>ABSOLUTE</i>	Constant <code>ABSOLUTE</code> of type <code>int</code>
<i>NORM</i>	Constant <code>NORM</code> of type <code>int</code>
<i>RELATIVE</i>	Constant <code>RELATIVE</code> of type <code>int</code>

Attributes Documentation

**ABSOLUTE:** `int` = `ABSOLUTE`  
Constant `ABSOLUTE` of type `int`

**NORM:** `int` = `NORM`  
Constant `NORM` of type `int`



**RELATIVE:** `int` = **RELATIVE**  
Constant RELATIVE of type `int`

### `slepc4py.SLEPc.SVD.ProblemType`

**class** `slepc4py.SLEPc.SVD.ProblemType`

Bases: `object`

SVD problem type.

- *STANDARD*: Standard SVD.
- *GENERALIZED*: Generalized singular value decomposition (GSVD).
- *HYPERBOLIC* : Hyperbolic singular value decomposition (HSVD).

#### See also

`SVDProblemType`

### Attributes Summary

<i>GENERALIZED</i>	Constant GENERALIZED of type <code>int</code>
<i>HYPERBOLIC</i>	Constant HYPERBOLIC of type <code>int</code>
<i>STANDARD</i>	Constant STANDARD of type <code>int</code>

### Attributes Documentation

**GENERALIZED:** `int` = **GENERALIZED**  
Constant GENERALIZED of type `int`

**HYPERBOLIC:** `int` = **HYPERBOLIC**  
Constant HYPERBOLIC of type `int`

**STANDARD:** `int` = **STANDARD**  
Constant STANDARD of type `int`

### `slepc4py.SLEPc.SVD.Stop`

**class** `slepc4py.SLEPc.SVD.Stop`

Bases: `object`

SVD stopping test.

- *BASIC*: Default stopping test.
- *USER*: User-defined stopping test.
- *THRESHOLD*: Threshold stopping test.

#### See also

`SVDSStop`

## Attributes Summary

<i>BASIC</i>	Constant BASIC of type <code>int</code>
<i>THRESHOLD</i>	Constant THRESHOLD of type <code>int</code>
<i>USER</i>	Constant USER of type <code>int</code>

## Attributes Documentation

**BASIC:** `int` = BASIC

Constant BASIC of type `int`

**THRESHOLD:** `int` = THRESHOLD

Constant THRESHOLD of type `int`

**USER:** `int` = USER

Constant USER of type `int`

## slepc4py.SLEPc.SVD.TRLanczosGBidiag

**class** slepc4py.SLEPc.SVD.TRLanczosGBidiag

Bases: `object`

SVD TRLanczos bidiagonalization choices for the GSVD case.

- *SINGLE*: Single bidiagonalization ( $Q_A$ ).
- *UPPER*: Joint bidiagonalization, both  $Q_A$  and  $Q_B$  in upper bidiagonal form.
- *LOWER*: Joint bidiagonalization,  $Q_A$  lower bidiagonal,  $Q_B$  upper bidiagonal.

### See also

SVDTRLanczosGBidiag

## Attributes Summary

<i>LOWER</i>	Constant LOWER of type <code>int</code>
<i>SINGLE</i>	Constant SINGLE of type <code>int</code>
<i>UPPER</i>	Constant UPPER of type <code>int</code>

## Attributes Documentation

**LOWER:** `int` = LOWER

Constant LOWER of type `int`

**SINGLE:** `int` = SINGLE

Constant SINGLE of type `int`

**UPPER:** `int` = UPPER

Constant UPPER of type `int`

## slepc4py.SLEPc.SVD.Type

**class** slepc4py.SLEPc.SVD.Type

Bases: `object`

SVD type.

Native singular value solvers.

- *CROSS*: Eigenproblem with the cross-product matrix.
- *CYCLIC*: Eigenproblem with the cyclic matrix.
- *LANCZOS*: Explicitly restarted Lanczos.
- *TRLANCZOS*: Thick-restart Lanczos.
- *RANDOMIZED*: Iterative RSVD for low-rank matrices.

Wrappers to external SVD solvers (should be enabled during installation of SLEPc).

- *LAPACK*: Sequential dense SVD solver.
- *SCALAPACK*: Parallel dense SVD solver.
- *KSVD*: Parallel dense SVD solver.
- *ELEMENTAL*: Parallel dense SVD solver.
- *PRIMME*: Iterative SVD solvers of Davidson type.

### See also

SVDType

## Attributes Summary

<i>CROSS</i>	Object CROSS of type <code>str</code>
<i>CYCLIC</i>	Object CYCLIC of type <code>str</code>
<i>ELEMENTAL</i>	Object ELEMENTAL of type <code>str</code>
<i>KSVD</i>	Object KSVD of type <code>str</code>
<i>LANCZOS</i>	Object LANCZOS of type <code>str</code>
<i>LAPACK</i>	Object LAPACK of type <code>str</code>
<i>PRIMME</i>	Object PRIMME of type <code>str</code>
<i>RANDOMIZED</i>	Object RANDOMIZED of type <code>str</code>
<i>SCALAPACK</i>	Object SCALAPACK of type <code>str</code>
<i>TRLANCZOS</i>	Object TRLANCZOS of type <code>str</code>

## Attributes Documentation

**CROSS:** `str` = **CROSS**

Object CROSS of type `str`

**CYCLIC:** `str` = **CYCLIC**

Object CYCLIC of type `str`

**ELEMENTAL:** `str` = **ELEMENTAL**

Object ELEMENTAL of type `str`

**KSVD:** `str = KSVD`  
Object KSVD of type `str`

**LANCZOS:** `str = LANCZOS`  
Object LANCZOS of type `str`

**LAPACK:** `str = LAPACK`  
Object LAPACK of type `str`

**PRIMME:** `str = PRIMME`  
Object PRIMME of type `str`

**RANDOMIZED:** `str = RANDOMIZED`  
Object RANDOMIZED of type `str`

**SCALAPACK:** `str = SCALAPACK`  
Object SCALAPACK of type `str`

**TRLANCZOS:** `str = TRLANCZOS`  
Object TRLANCZOS of type `str`

**slepc4py.SLEPc.SVD.Which**

**class** `slepc4py.SLEPc.SVD.Which`  
Bases: `object`

SVD desired part of spectrum.

- *LARGEST*: Largest singular values.
- *SMALLEST*: Smallest singular values.

See also	
SVDWhich	

**Attributes Summary**

<i>LARGEST</i>	Constant LARGEST of type <code>int</code>
<i>SMALLEST</i>	Constant SMALLEST of type <code>int</code>

**Attributes Documentation**

**LARGEST:** `int = LARGEST`  
Constant LARGEST of type `int`

**SMALLEST:** `int = SMALLEST`  
Constant SMALLEST of type `int`

**Methods Summary**

<i>appendOptionsPrefix</i> ([prefix])	Append to the prefix used for searching for all SVD options in the database.
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<code>cancelMonitor()</code>	Clear all monitors for an <i>SVD</i> object.
<code>computeError(i[, etype])</code>	Compute the error associated with the <i>i</i> -th singular triplet.
<code>create([comm])</code>	Create the SVD object.
<code>destroy()</code>	Destroy the SVD object.
<code>errorView([etype, viewer])</code>	Display the errors associated with the computed solution.
<code>getBV()</code>	Get the basis vectors objects associated to the SVD object.
<code>getConverged()</code>	Get the number of converged singular triplets.
<code>getConvergedReason()</code>	Get the reason why the <code>solve()</code> iteration was stopped.
<code>getConvergenceTest()</code>	Get the method used to compute the error estimate used in the convergence test.
<code>getCrossEPS()</code>	Get the eigensolver object associated to the singular value solver.
<code>getCrossExplicitMatrix()</code>	Get the flag indicating if $A^*A$ is built explicitly.
<code>getCyclicEPS()</code>	Get the eigensolver object associated to the singular value solver.
<code>getCyclicExplicitMatrix()</code>	Get the flag indicating if $H(A)$ is built explicitly.
<code>getDS()</code>	Get the direct solver associated to the singular value solver.
<code>getDimensions()</code>	Get the number of singular values to compute and the dimension of the subspace.
<code>getImplicitTranspose()</code>	Get the mode used to handle the transpose of the associated matrix.
<code>getIterationNumber()</code>	Get the current iteration number.
<code>getLanczosOneSide()</code>	Get if the variant of the Lanczos method to be used is one-sided or two-sided.
<code>getMonitor()</code>	Get the list of monitor functions.
<code>getOperators()</code>	Get the matrices associated with the singular value problem.
<code>getOptionsPrefix()</code>	Get the prefix used for searching for all SVD options in the database.
<code>getProblemType()</code>	Get the problem type from the SVD object.
<code>getSignature([omega])</code>	Get the signature matrix defining a hyperbolic singular value problem.
<code>getSingularTriplet(i[, U, V])</code>	Get the <i>i</i> -th triplet of the singular value decomposition.
<code>getStoppingTest()</code>	Get the stopping test function.
<code>getTRLanczosExplicitMatrix()</code>	Get the flag indicating if $Z = [A^*, B^*]^*$ is built explicitly.
<code>getTRLanczosGBidiag()</code>	Get bidiagonalization choice used in the GSVD TR-Lanczos solver.
<code>getTRLanczosKSP()</code>	Get the linear solver object associated with the SVD solver.
<code>getTRLanczosLocking()</code>	Get the locking flag used in the thick-restart Lanczos method.
<code>getTRLanczosOneSide()</code>	Get if the variant of the method to be used is one-sided or two-sided.
<code>getTRLanczosRestart()</code>	Get the restart parameter used in the thick-restart Lanczos method.

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Table 106 – continued from previous page

<code>getThreshold()</code>	Get the threshold used in the threshold stopping test.
<code>getTolerances()</code>	Get the tolerance and maximum iteration count.
<code>getTrackAll()</code>	Get the flag indicating if all residual norms must be computed or not.
<code>getType()</code>	Get the SVD type of this object.
<code>getValue(i)</code>	Get the i-th singular value as computed by <code>solve()</code> .
<code>getVectors(i, U, V)</code>	Get the i-th left and right singular vectors as computed by <code>solve()</code> .
<code>getWhichSingularTriplets()</code>	Get which singular triplets are to be sought.
<code>isGeneralized()</code>	Tell if the SVD corresponds to a generalized singular value problem.
<code>isHyperbolic()</code>	Tell whether the SVD object corresponds to a hyperbolic singular value problem.
<code>reset()</code>	Reset the SVD object.
<code>setBV(V[, U])</code>	Set basis vectors objects associated to the SVD solver.
<code>setConvergenceTest(conv)</code>	Set how to compute the error estimate used in the convergence test.
<code>setCrossEPS(eps)</code>	Set an eigensolver object associated to the singular value solver.
<code>setCrossExplicitMatrix([flag])</code>	Set if the eigensolver operator $A^*A$ must be computed.
<code>setCyclicEPS(eps)</code>	Set an eigensolver object associated to the singular value solver.
<code>setCyclicExplicitMatrix([flag])</code>	Set if the eigensolver operator $H(A)$ must be computed explicitly.
<code>setDS(ds)</code>	Set a direct solver object associated to the singular value solver.
<code>setDimensions([nsv, ncv, mpd])</code>	Set the number of singular values to compute and the dimension of the subspace.
<code>setFromOptions()</code>	Set SVD options from the options database.
<code>setImplicitTranspose(mode)</code>	Set how to handle the transpose of the associated matrix.
<code>setInitialSpace([spaceright, spaceleft])</code>	Set the initial spaces from which the SVD solver starts to iterate.
<code>setLanczosOneSide([flag])</code>	Set if the variant of the Lanczos method to be used is one-sided or two-sided.
<code>setMonitor(monitor[, args, kargs])</code>	Append a monitor function to the list of monitors.
<code>setOperators(A[, B])</code>	Set the matrices associated with the singular value problem.
<code>setOptionsPrefix([prefix])</code>	Set the prefix used for searching for all SVD options in the database.
<code>setProblemType(problem_type)</code>	Set the type of the singular value problem.
<code>setSignature([omega])</code>	Set the signature matrix defining a hyperbolic singular value problem.
<code>setStoppingTest(stopping[, args, kargs])</code>	Set a function to decide when to stop the outer iteration of the eigensolver.
<code>setTRLanczosExplicitMatrix([flag])</code>	Set if the matrix $Z = [A^*, B^*]^*$ must be built explicitly.
<code>setTRLanczosGBidiag(bidiag)</code>	Set the bidiagonalization choice to use in the GSVD TRLanczos solver.
<code>setTRLanczosKSP(ksp)</code>	Set a linear solver object associated to the SVD solver.

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<code>setTRLanczosLocking(lock)</code>	Toggle between locking and non-locking variants of TRLanczos.
<code>setTRLanczosOneSide([flag])</code>	Set if the variant of the method to be used is one-sided or two-sided.
<code>setTRLanczosRestart(keep)</code>	Set the restart parameter for the thick-restart Lanczos method.
<code>setThreshold(thres[, rel])</code>	Set the threshold used in the threshold stopping test.
<code>setTolerances([tol, max_it])</code>	Set the tolerance and maximum iteration count used.
<code>setTrackAll(trackall)</code>	Set flag to compute the residual of all singular triplets.
<code>setType(svd_type)</code>	Set the particular solver to be used in the SVD object.
<code>setUp()</code>	Set up all the internal data structures.
<code>setWhichSingularTriplets(which)</code>	Set which singular triplets are to be sought.
<code>solve()</code>	Solve the singular value problem.
<code>valuesView([viewer])</code>	Display the computed singular values in a viewer.
<code>vectorsView([viewer])</code>	Output computed singular vectors to a viewer.
<code>view([viewer])</code>	Print the SVD data structure.

### Attributes Summary

<code>ds</code>	The direct solver ( <i>DS</i> ) object associated.
<code>max_it</code>	The maximum iteration count.
<code>problem_type</code>	The type of the eigenvalue problem.
<code>tol</code>	The tolerance.
<code>track_all</code>	Compute the residual norm of all approximate eigenpairs.
<code>transpose_mode</code>	How to handle the transpose of the matrix.
<code>which</code>	The portion of the spectrum to be sought.

### Methods Documentation

#### `appendOptionsPrefix(prefix=None)`

Append to the prefix used for searching for all SVD options in the database.

Logically collective.

##### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all SVD option requests.

##### Return type

*None*

#### See also

`setOptionsPrefix`, `getOptionsPrefix`, `SVDAppendOptionsPrefix`

**:sources:** ``Source code at slepc4py/SLEPc/SVD.pyx:357 <slepc4py/SLEPc/SVD.pyx#L357>``

#### `cancelMonitor()`

Clear all monitors for an *SVD* object.

Logically collective.

<b>See also</b>
<a href="#">SVDMonitorCancel</a>

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1149 <slepc4py/SLEPc/SVD.pyx#L1149>`

**Return type**

`None`

**computeError**(*i*, *etype*=`None`)

Compute the error associated with the *i*-th singular triplet.

Collective.

Compute the error (based on the residual norm) associated with the *i*-th singular triplet.

**Parameters**

- **i** (`int`) – Index of the solution to be considered.
- **etype** (`ErrorType` / `None`) – The error type to compute.

**Returns**

The error bound, computed in various ways from the residual norm  $\sqrt{\eta_1^2 + \eta_2^2}$  where  $\eta_1 = \|Av - \sigma u\|_2$ ,  $\eta_2 = \|A^*u - \sigma v\|_2$ ,  $\sigma$  is the approximate singular value,  $u$  and  $v$  are the left and right singular vectors.

**Return type**

`float`

**Notes**

The index *i* should be a value between 0 and `nconv-1` (see [getConverged\(\)](#)).

In the case of the GSVD, the two components of the residual norm are  $\eta_1 = \|s^2 A^* u - c B^* B x\|_2$  and  $\eta_2 = \|c^2 B^* v - s A^* A x\|_2$ , where  $(\sigma, u, v, x)$  is the approximate generalized singular quadruple, with  $\sigma = c/s$ .

<b>See also</b>
<a href="#">solve</a> , <a href="#">SVDComputeError</a>

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1377 <slepc4py/SLEPc/SVD.pyx#L1377>`

**create**(*comm*=`None`)

Create the SVD object.

Collective.

**Parameters**

**comm** (`Comm` / `None`) – MPI communicator; if not provided, it defaults to all processes.

**Return type**

`Self`

<b>See also</b>
<a href="#">SVDCreate</a>



**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:240 <slepc4py/SLEPc/SVD.pyx#L240>](#)

### **destroy()**

Destroy the SVD object.

Collective.

#### **See also**

[SVDDestroy](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:214 <slepc4py/SLEPc/SVD.pyx#L214>](#)

### **Return type**

*Self*

### **errorView(etype=None, viewer=None)**

Display the errors associated with the computed solution.

Collective.

Display the errors and the singular values.

#### **Parameters**

- **etype** ([ErrorType](#) / *None*) – The error type to compute.
- **viewer** ([petsc4py.PETSc.Viewer](#) / *None*) – Visualization context; if not provided, the standard output is used.

### **Return type**

*None*

### **Notes**

By default, this function checks the error of all singular triplets and prints the singular values if all of them are below the requested tolerance. If the viewer has format ASCII\_INFO\_DETAIL then a table with singular values and corresponding errors is printed.

#### **See also**

[solve](#), [valuesView](#), [vectorsView](#), [SVDErrorView](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1424 <slepc4py/SLEPc/SVD.pyx#L1424>](#)

### **getBV()**

Get the basis vectors objects associated to the SVD object.

Not collective.

#### **Returns**

- **V** (*BV*) – The basis vectors context for right singular vectors.
- **U** (*BV*) – The basis vectors context for left singular vectors.

### **Return type**

*tuple[BV, BV]*

**See also**

[setBV](#), [SVDGetBV](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:829](#) <[slepc4py/SLEPc/SVD.pyx#L829](#)>

**getConverged()**

Get the number of converged singular triplets.

Not collective.

**Returns**

**nconv** – Number of converged singular triplets.

**Return type**

`int`

**Notes**

This function should be called after [solve\(\)](#) has finished.

The value `nconv` may be different from the number of requested solutions `nsv`, but not larger than `ncv`, see [setDimensions\(\)](#).

**See also**

[setDimensions](#), [solve](#), [getValue](#), [SVDGetConverged](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1248](#) <[slepc4py/SLEPc/SVD.pyx#L1248](#)>

**getConvergedReason()**

Get the reason why the [solve\(\)](#) iteration was stopped.

Not collective.

**Returns**

Negative value indicates diverged, positive value converged.

**Return type**

[ConvergedReason](#)

**See also**

[setTolerances](#), [solve](#), [SVDGetConvergedReason](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1229](#) <[slepc4py/SLEPc/SVD.pyx#L1229](#)>

**getConvergenceTest()**

Get the method used to compute the error estimate used in the convergence test.

Not collective.

**Returns**

The method used to compute the error estimate used in the convergence test.

**Return type**

[Conv](#)

**See also**

[`setConvergenceTest`](#), [`SVDGetConvergenceTest`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:674 <slepc4py/SLEPc/SVD.pyx#L674>`

**getCrossEPS()**

Get the eigensolver object associated to the singular value solver.

Collective.

**Returns**

The eigensolver object.

**Return type**

[`EPS`](#)

**See also**

[`setCrossEPS`](#), [`SVDCrossGetEPS`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1513 <slepc4py/SLEPc/SVD.pyx#L1513>`

**getCrossExplicitMatrix()**

Get the flag indicating if  $A^*A$  is built explicitly.

Not collective.

**Returns**

True if  $A^*A$  is built explicitly.

**Return type**

[`bool`](#)

**See also**

[`setCrossExplicitMatrix`](#), [`SVDCrossGetExplicitMatrix`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1560 <slepc4py/SLEPc/SVD.pyx#L1560>`

**getCyclicEPS()**

Get the eigensolver object associated to the singular value solver.

Collective.

**Returns**

The eigensolver object.

**Return type**

[`EPS`](#)

**See also**

[`setCyclicEPS`](#), [`SVDCyclicGetEPS`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1596 <slepc4py/SLEPc/SVD.pyx#L1596>`

### **getCyclicExplicitMatrix()**

Get the flag indicating if  $H(A)$  is built explicitly.

Not collective.

Get the flag indicating if  $H(A) = \begin{bmatrix} 0 & A; A^T & 0 \end{bmatrix}$  is built explicitly.

#### **Returns**

True if  $H(A)$  is built explicitly.

#### **Return type**

`bool`

#### **See also**

`setCyclicExplicitMatrix`, `SVDCyclicGetExplicitMatrix`

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1645](#) <[slepc4py/SLEPc/SVD.pyx#L1645](#)>

### **getDS()**

Get the direct solver associated to the singular value solver.

Not collective.

#### **Returns**

The direct solver context.

#### **Return type**

`DS`

#### **See also**

`setDS`, `SVDGetDS`

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:874](#) <[slepc4py/SLEPc/SVD.pyx#L874](#)>

### **getDimensions()**

Get the number of singular values to compute and the dimension of the subspace.

Not collective.

#### **Returns**

- `nsv (int)` – Number of singular values to compute.
- `ncv (int)` – Maximum dimension of the subspace to be used by the solver.
- `mpd (int)` – Maximum dimension allowed for the projected problem.

#### **Return type**

`tuple[int, int, int]`

#### **See also**

`setDimensions`, `SVDGetDimensions`

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:753](#) <[slepc4py/SLEPc/SVD.pyx#L753](#)>

### **getImplicitTranspose()**

Get the mode used to handle the transpose of the associated matrix.

Not collective.

#### **Returns**

How to handle the transpose (implicitly or not).

#### **Return type**

`bool`

#### **See also**

`setImplicitTranspose`, `SVDGetImplicitTranspose`

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:478 <slepc4py/SLEPc/SVD.pyx#L478>](#)

### **getIterationNumber()**

Get the current iteration number.

Not collective.

If the call to `solve()` is complete, then it returns the number of iterations carried out by the solution method.

#### **Returns**

Iteration number.

#### **Return type**

`int`

#### **See also**

`getConvergedReason`, `setTolerances`, `SVDGetIterationNumber`

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1207 <slepc4py/SLEPc/SVD.pyx#L1207>](#)

### **getLanczosOneSide()**

Get if the variant of the Lanczos method to be used is one-sided or two-sided.

Not collective.

#### **Returns**

True if the method is one-sided.

#### **Return type**

`bool`

#### **See also**

`setLanczosOneSide`, `SVDLanczosGetOneSide`

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1693 <slepc4py/SLEPc/SVD.pyx#L1693>](#)

### **getMonitor()**

Get the list of monitor functions.

Not collective.

#### **Returns**

The list of monitor functions.

#### **Return type**

*SVDMonitorFunction*

#### **See also**

*setMonitor*

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1132 <slepc4py/SLEPc/SVD.pyx#L1132>](#)

### **getOperators()**

Get the matrices associated with the singular value problem.

Collective.

#### **Returns**

- **A** (`petsc4py.PETSc.Mat`) – The matrix associated with the singular value problem.
- **B** (`petsc4py.PETSc.Mat`) – The second matrix in the case of GSVD.

#### **Return type**

`tuple[Mat, Mat] | tuple[Mat, None]`

#### **See also**

*setOperators*, *SVDGetOperators*

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:911 <slepc4py/SLEPc/SVD.pyx#L911>](#)

### **getOptionsPrefix()**

Get the prefix used for searching for all SVD options in the database.

Not collective.

#### **Returns**

The prefix string set for this SVD object.

#### **Return type**

`str`

#### **See also**

*setOptionsPrefix*, *appendOptionsPrefix*, *SVDGetOptionsPrefix*

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:307 <slepc4py/SLEPc/SVD.pyx#L307>](#)

### **getProblemType()**

Get the problem type from the SVD object.

Not collective.

**Returns**

The problem type that was previously set.

**Return type**

*ProblemType*

**See also**

*setProblemType*, *SVDGetProblemType*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:395 <slepc4py/SLEPc/SVD.pyx#L395>`

**getSignature**(*omega=None*)

Get the signature matrix defining a hyperbolic singular value problem.

Collective.

**Parameters**

**omega** (*Vec* / *None*) – Optional vector to store the diagonal elements of the signature matrix.

**Returns**

A vector containing the diagonal elements of the signature matrix.

**Return type**

*petsc4py.PETSc.Vec*

**See also**

*setSignature*, *SVDGetSignature*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:958 <slepc4py/SLEPc/SVD.pyx#L958>`

**getSingularTriplet**(*i*, *U=None*, *V=None*)

Get the *i*-th triplet of the singular value decomposition.

Collective.

Get the *i*-th triplet of the singular value decomposition as computed by *solve()*. The solution consists of the singular value and its left and right singular vectors.

**Parameters**

- **i** (*int*) – Index of the solution to be obtained.
- **U** (*Vec* / *None*) – Placeholder for the returned left singular vector.
- **V** (*Vec* / *None*) – Placeholder for the returned right singular vector.

**Returns**

The computed singular value.

**Return type**

*float*

**Notes**

The index *i* should be a value between 0 and *nconv*−1 (see *getConverged()*. Singular triplets are indexed according to the ordering criterion established with *setWhichSingularTriplets()*.

**See also**

*getConverged, setWhichSingularTriplets, SVDGetSingularTriplet*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1334 <slepc4py/SLEPc/SVD.pyx#L1334>`

**getStoppingTest()**

Get the stopping test function.

Not collective.

**Returns**

The stopping test function.

**Return type**

*SVDStoppingFunction*

**See also**

*setStoppingTest*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1088 <slepc4py/SLEPc/SVD.pyx#L1088>`

**getTRLanczosExplicitMatrix()**

Get the flag indicating if  $Z = [A^*, B^*]^*$  is built explicitly.

Not collective.

**Returns**

True if  $Z = [A^*, B^*]^*$  is built explicitly.

**Return type**

*bool*

**See also**

*setTRLanczosExplicitMatrix, SVDTRLanczosGetExplicitMatrix*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1948 <slepc4py/SLEPc/SVD.pyx#L1948>`

**getTRLanczosGBidiag()**

Get bidiagonalization choice used in the GSVD TRLanczos solver.

Not collective.

**Returns**

The bidiagonalization choice.

**Return type**

*TRLanczosGBidiag*

**See also**

*setTRLanczosGBidiag, SVDTRLanczosGetGBidiag*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1780 <slepc4py/SLEPc/SVD.pyx#L1780>`



### **getTRLanczosKSP()**

Get the linear solver object associated with the SVD solver.

Collective.

#### **Returns**

The linear solver object.

#### **Return type**

`petsc4py.PETSc.KSP`

#### **See also**

`setTRLanczosKSP`, `SVDTRLanczosGetKSP`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1905 <slepc4py/SLEPc/SVD.pyx#L1905>`

### **getTRLanczosLocking()**

Get the locking flag used in the thick-restart Lanczos method.

Not collective.

#### **Returns**

The locking flag.

#### **Return type**

`bool`

#### **See also**

`setTRLanczosLocking`, `SVDTRLanczosGetLocking`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1869 <slepc4py/SLEPc/SVD.pyx#L1869>`

### **getTRLanczosOneSide()**

Get if the variant of the method to be used is one-sided or two-sided.

Not collective.

Get if the variant of the thick-restart Lanczos method to be used is one-sided or two-sided.

#### **Returns**

True if the method is one-sided.

#### **Return type**

`bool`

#### **See also**

`setTRLanczosOneSide`, `SVDLanczosGetOneSide`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1740 <slepc4py/SLEPc/SVD.pyx#L1740>`

### **getTRLanczosRestart()**

Get the restart parameter used in the thick-restart Lanczos method.

Not collective.

**Returns**

The number of vectors to be kept at restart.

**Return type**

`float`

**See also**

[`setTRLanczosRestart`](#), [`SVDTRLanczosGetRestart`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1825 <slepc4py/SLEPc/SVD.pyx#L1825>`

**getThreshold()**

Get the threshold used in the threshold stopping test.

Not collective.

**Returns**

- **thres** (`float`) – The threshold.
- **rel** (`bool`) – Whether the threshold is relative or not.

**Return type**

`tuple[float, bool]`

**See also**

[`setThreshold`](#), [`SVDGetThreshold`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:561 <slepc4py/SLEPc/SVD.pyx#L561>`

**getTolerances()**

Get the tolerance and maximum iteration count.

Not collective.

Get the tolerance and maximum iteration count used by the default SVD convergence tests.

**Returns**

- **tol** (`float`) – The convergence tolerance.
- **max\_it** (`int`) – The maximum number of iterations.

**Return type**

`tuple[float, int]`

**See also**

[`setTolerances`](#), [`SVDGetTolerances`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:618 <slepc4py/SLEPc/SVD.pyx#L618>`

**getTrackAll()**

Get the flag indicating if all residual norms must be computed or not.

Not collective.

**Returns**

Whether the solver computes all residuals or not.

**Return type**

`bool`

**See also**

`setTrackAll`, `SVDGetTrackAll`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:713 <slepc4py/SLEPc/SVD.pyx#L713>`

**getType()**

Get the SVD type of this object.

Not collective.

**Returns**

The solver currently being used.

**Return type**

`str`

**See also**

`setType`, `SVDGetType`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:288 <slepc4py/SLEPc/SVD.pyx#L288>`

**getValue(i)**

Get the i-th singular value as computed by `solve()`.

Collective.

**Parameters**

**i** (`int`) – Index of the solution to be obtained.

**Returns**

The computed singular value.

**Return type**

`float`

**Notes**

The index `i` should be a value between 0 and `nconv-1` (see `getConverged()`). Singular triplets are indexed according to the ordering criterion established with `setWhichSingularTriplets()`.

**See also**

`getConverged`, `setWhichSingularTriplets`, `SVDGetSingularTriplet`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1274 <slepc4py/SLEPc/SVD.pyx#L1274>`

### **getVectors(*i*, *U*, *V*)**

Get the *i*-th left and right singular vectors as computed by [solve\(\)](#).

Collective.

#### **Parameters**

- **i** (*int*) – Index of the solution to be obtained.
- **U** (*Vec*) – Placeholder for the returned left singular vector.
- **V** (*Vec*) – Placeholder for the returned right singular vector.

#### **Return type**

*None*

### **Notes**

The index *i* should be a value between 0 and `nconv-1` (see [getConverged\(\)](#)). Singular triplets are indexed according to the ordering criterion established with [setWhichSingularTriplets\(\)](#).

#### **See also**

[getConverged](#), [setWhichSingularTriplets](#), [SVDGetSingularTriplet](#)

**:sources:** ``Source code at slepc4py/SLEPc/SVD.pyx:1305 <slepc4py/SLEPc/SVD.pyx#L1305>``

### **getWhichSingularTriplets()**

Get which singular triplets are to be sought.

Not collective.

#### **Returns**

The singular values to be sought (either largest or smallest).

#### **Return type**

*Which*

#### **See also**

[setWhichSingularTriplets](#), [SVDGetWhichSingularTriplets](#)

**:sources:** ``Source code at slepc4py/SLEPc/SVD.pyx:524 <slepc4py/SLEPc/SVD.pyx#L524>``

### **isGeneralized()**

Tell if the SVD corresponds to a generalized singular value problem.

Not collective.

#### **Returns**

True if two matrices were set with [setOperators\(\)](#).

#### **Return type**

*bool*

#### **See also**

[setProblemType](#), [isHyperbolic](#), [SVDIsGeneralized](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:438](#) <slepc4py/SLEPc/SVD.pyx#L438>

### **isHyperbolic()**

Tell whether the SVD object corresponds to a hyperbolic singular value problem.

Not collective.

#### **Returns**

True if the problem was specified as hyperbolic.

#### **Return type**

bool

#### **See also**

[setProblemType](#), [isGeneralized](#), [SVDIsHyperbolic](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:457](#) <slepc4py/SLEPc/SVD.pyx#L457>

### **reset()**

Reset the SVD object.

Collective.

#### **See also**

[SVDReset](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:228](#) <slepc4py/SLEPc/SVD.pyx#L228>

#### **Return type**

None

### **setBV(*V*, *U*=None)**

Set basis vectors objects associated to the SVD solver.

Collective.

#### **Parameters**

- **V** ([BV](#)) – The basis vectors context for right singular vectors.
- **U** ([BV](#) / *None*) – The basis vectors context for left singular vectors.

#### **Return type**

None

#### **See also**

[getBV](#), [SVDSetBV](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:853](#) <slepc4py/SLEPc/SVD.pyx#L853>

### **setConvergenceTest(*conv*)**

Set how to compute the error estimate used in the convergence test.

Logically collective.

#### Parameters

**conv** ([Conv](#)) – The method used to compute the error estimate used in the convergence test.

#### Return type

None

#### See also

[getConvergenceTest](#), [SVDSetConvergenceTest](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:694](#) <[slepc4py/SLEPc/SVD.pyx#L694](#)>

#### **setCrossEPS**(*eps*)

Set an eigensolver object associated to the singular value solver.

Collective.

#### Parameters

**eps** ([EPS](#)) – The eigensolver object.

#### Return type

None

#### See also

[getCrossEPS](#), [SVDCrossSetEPS](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1496](#) <[slepc4py/SLEPc/SVD.pyx#L1496](#)>

#### **setCrossExplicitMatrix**(*flag=True*)

Set if the eigensolver operator  $A^*A$  must be computed.

Logically collective.

#### Parameters

**flag** ([bool](#)) – True to build  $A^*A$  explicitly.

#### Return type

None

#### Notes

In GSVD there are two cross product matrices,  $A^*A$  and  $B^*B$ . In HSVD the expression for the cross product matrix is different,  $A^*\Omega A$ .

By default the matrices are not built explicitly, but handled as shell matrices

#### See also

[getCrossExplicitMatrix](#), [SVDCrossSetExplicitMatrix](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1533](#) <[slepc4py/SLEPc/SVD.pyx#L1533](#)>

#### **setCyclicEPS**(*eps*)

Set an eigensolver object associated to the singular value solver.

Collective.

**Parameters**

**eps** (*EPS*) – The eigensolver object.

**Return type**

None

**See also**

*getCyclicEPS*, *SVDCyclicSetEPS*

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1579 <slepc4py/SLEPc/SVD.pyx#L1579>](#)

**setCyclicExplicitMatrix**(*flag=True*)

Set if the eigensolver operator  $H(A)$  must be computed explicitly.

Logically collective.

Set if the eigensolver operator  $H(A) = [0 \ A; A^T \ 0]$  must be computed explicitly.

**Parameters**

**flag** (*bool*) – True if  $H(A)$  must be built explicitly.

**Return type**

None

**Notes**

In GSVD and HSVD the equivalent eigenvalue problem has generalized form, and hence two matrices are built.

By default the matrices are not built explicitly, but handled as shell matrices.

**See also**

*getCyclicExplicitMatrix*, *SVDCyclicSetExplicitMatrix*

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1616 <slepc4py/SLEPc/SVD.pyx#L1616>](#)

**setDS**(*ds*)

Set a direct solver object associated to the singular value solver.

Collective.

**Parameters**

**ds** (*DS*) – The direct solver context.

**Return type**

None

**See also**

*getDS*, *SVDSetDS*

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:894 <slepc4py/SLEPc/SVD.pyx#L894>](#)

**setDimensions**(*nsv=None, ncv=None, mpd=None*)

Set the number of singular values to compute and the dimension of the subspace.

Logically collective.

#### Parameters

- **nsv** (*int* / *None*) – Number of singular values to compute.
- **ncv** (*int* / *None*) – Maximum dimension of the subspace to be used by the solver.
- **mpd** (*int* / *None*) – Maximum dimension allowed for the projected problem.

#### Return type

*None*

#### Notes

Use *DETERMINE* for *ncv* and *mpd* to assign a reasonably good value, which is dependent on the solution method.

The parameters *ncv* and *mpd* are intimately related, so that the user is advised to set one of them at most. Normal usage is the following:

- In cases where *nsv* is small, the user sets *ncv* (a reasonable default is  $2 * nsv$ ).
- In cases where *nsv* is large, the user sets *mpd*.

The value of *ncv* should always be between *nsv* and (*nsv* + *mpd*), typically *ncv* = *nsv* + *mpd*. If *nsv* is not too large, *mpd* = *nsv* is a reasonable choice, otherwise a smaller value should be used.

#### See also

*getDimensions*, *SVDSetDimensions*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:778 <slepc4py/SLEPc/SVD.pyx#L778>`

**setFromOptions()**

Set SVD options from the options database.

Collective.

#### Notes

To see all options, run your program with the `-help` option.

This routine must be called before *setUp()* if the user is to be allowed to set the solver type.

#### See also

*setOptionsPrefix*, *SVDSetFromOptions*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:376 <slepc4py/SLEPc/SVD.pyx#L376>`

#### Return type

*None*



### **setImplicitTranspose(*mode*)**

Set how to handle the transpose of the associated matrix.

Logically collective.

#### **Parameters**

- **impl** – How to handle the transpose (implicitly or not).
- **mode** (*bool*)

#### **Return type**

*None*

#### **Notes**

By default, the transpose of the matrix is explicitly built (if the matrix has defined the `Mat.transpose()` operation).

If this flag is set to `True`, the solver does not build the transpose, but handles it implicitly via `Mat.multTranspose()` (or `Mat.multHermitianTranspose()` in the complex case).

#### **See also**

`getImplicitTranspose`, `SVDSetImplicitTranspose`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:497 <slepc4py/SLEPc/SVD.pyx#L497>`

### **setInitialSpace(*spaceright=None, spaceleft=None*)**

Set the initial spaces from which the SVD solver starts to iterate.

Collective.

#### **Parameters**

- **spaceright** (*list[Vec] | None*) – The right initial space.
- **spaceleft** (*list[Vec] | None*) – The left initial space.

#### **Return type**

*None*

#### **Notes**

The initial right and left spaces are rough approximations to the right and/or left singular subspaces from which the solver starts to iterate. It is not necessary to provide both sets of vectors.

Some solvers start to iterate on a single vector (initial vector). In that case, the other vectors are ignored.

These vectors do not persist from one `solve()` call to the other, so the initial spaces should be set every time.

The vectors do not need to be mutually orthonormal, since they are explicitly orthonormalized internally.

Common usage of this function is when the user can provide a rough approximation of the wanted singular spaces. Then, convergence may be faster.

#### **See also**

`SVDSetInitialSpaces`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1007 <slepc4py/SLEPc/SVD.pyx#L1007>`

**setLanczosOneSide**(*flag=True*)

Set if the variant of the Lanczos method to be used is one-sided or two-sided.

Logically collective.

**Parameters**

**flag** (*bool*) – True if the method is one-sided.

**Return type**

*None*

**Notes**

By default, a two-sided variant is selected, which is sometimes slightly more robust. However, the one-sided variant is faster because it avoids the orthogonalization associated to left singular vectors. It also saves the memory required for storing such vectors.

**See also**

*getLanczosOneSide*, *SVDLanczosSetOneSide*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1667 <slepc4py/SLEPc/SVD.pyx#L1667>`

**setMonitor**(*monitor*, *args=None*, *kargs=None*)

Append a monitor function to the list of monitors.

Logically collective.

**See also**

*getMonitor*, *cancelMonitor*, *SVDMonitorSet*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1107 <slepc4py/SLEPc/SVD.pyx#L1107>`

**Parameters**

- **monitor** (*SVDMonitorFunction* | *None*)
- **args** (*tuple*[*Any*, ...] | *None*)
- **kargs** (*dict*[*str*, *Any*] | *None*)

**Return type**

*None*

**setOperators**(*A*, *B=None*)

Set the matrices associated with the singular value problem.

Collective.

**Parameters**

- **A** (*Mat*) – The matrix associated with the singular value problem.
- **B** (*Mat* | *None*) – The second matrix in the case of GSVD.

**Return type**

*None*

#### See also

[getOperators](#), [SVDSetOperators](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:938 <slepc4py/SLEPc/SVD.pyx#L938>](#)

**setOptionsPrefix**(*prefix=None*)

Set the prefix used for searching for all SVD options in the database.

Logically collective.

#### Parameters

**prefix** (*str* / *None*) – The prefix string to prepend to all SVD option requests.

#### Return type

*None*

#### Notes

A hyphen (-) must NOT be given at the beginning of the prefix name. The first character of all runtime options is AUTOMATICALLY the hyphen.

For example, to distinguish between the runtime options for two different SVD contexts, one could call:

```
S1.setOptionsPrefix("svd1_")
S2.setOptionsPrefix("svd2_")
```

#### See also

[appendOptionsPrefix](#), [getOptionsPrefix](#), [SVDGetOptionsPrefix](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:326 <slepc4py/SLEPc/SVD.pyx#L326>](#)

**setProblemType**(*problem\_type*)

Set the type of the singular value problem.

Logically collective.

#### Parameters

**problem\_type** ([ProblemType](#)) – The problem type to be set.

#### Return type

*None*

#### Notes

The GSVD requires that two matrices have been passed via [setOperators\(\)](#). The HSVD requires that a signature matrix has been passed via [setSignature\(\)](#).

#### See also

[setOperators](#), [setSignature](#), [getProblemType](#), [SVDSetProblemType](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:414 <slepc4py/SLEPc/SVD.pyx#L414>](#)

**setSignature**(*omega=None*)

Set the signature matrix defining a hyperbolic singular value problem.

Collective.

**Parameters**

**omega** (*Vec* / *None*) – A vector containing the diagonal elements of the signature matrix.

**Return type**

*None*

**See also**

*getSignature*, *SVDSetSignature*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:987 <slepc4py/SLEPc/SVD.pyx#L987>`

**setStoppingTest**(*stopping, args=None, kargs=None*)

Set a function to decide when to stop the outer iteration of the eigensolver.

Logically collective.

**See also**

*getStoppingTest*, *SVDSetStoppingTestFunction*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1064 <slepc4py/SLEPc/SVD.pyx#L1064>`

**Parameters**

- **stopping** (*SVDStoppingFunction* / *None*)
- **args** (*tuple*[*Any*, ...] / *None*)
- **kargs** (*dict*[*str*, *Any*] / *None*)

**Return type**

*None*

**setTRLanczosExplicitMatrix**(*flag=True*)

Set if the matrix  $Z = [A^*, B^*]^*$  must be built explicitly.

Logically collective.

**Parameters**

**flag** (*bool*) – True if  $Z = [A^*, B^*]^*$  is built explicitly.

**Return type**

*None*

**Notes**

This option is relevant for the GSVD case only.  $Z$  is the coefficient matrix of the least-squares solver used internally.

**See also**

*getTRLanczosExplicitMatrix*, *SVDTRLanczosSetExplicitMatrix*

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1925 <slepc4py/SLEPc/SVD.pyx#L1925>`

**setTRLanczosGBidiag**(*bidiag*)

Set the bidiagonalization choice to use in the GSVD TRLanczos solver.

Logically collective.

**Parameters**

**bidiag** (*TRLanczosGBidiag*) – The bidiagonalization choice.

**Return type**

None

**See also**

[getTRLanczosGBidiag](#), [SVDTRLanczosSetGBidiag](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1762 <slepc4py/SLEPc/SVD.pyx#L1762>`

**setTRLanczosKSP**(*ksp*)

Set a linear solver object associated to the SVD solver.

Collective.

**Parameters**

**ksp** (*KSP*) – The linear solver object.

**Return type**

None

**See also**

[getTRLanczosKSP](#), [SVDTRLanczosSetKSP](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1888 <slepc4py/SLEPc/SVD.pyx#L1888>`

**setTRLanczosLocking**(*lock*)

Toggle between locking and non-locking variants of TRLanczos.

Logically collective.

**Parameters**

**lock** (*bool*) – True if the locking variant must be selected.

**Return type**

None

**Notes**

The default is to lock converged singular triplets when the method restarts. This behavior can be changed so that all directions are kept in the working subspace even if already converged to working accuracy (the non-locking variant).

**See also**

[getTRLanczosLocking](#), [SVDTRLanczosSetLocking](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1844 <slepc4py/SLEPc/SVD.pyx#L1844>`

**setTRLanczosOneSide**(*flag=True*)

Set if the variant of the method to be used is one-sided or two-sided.

Logically collective.

Set if the variant of the thick-restart Lanczos method to be used is one-sided or two-sided.

**Parameters**

**flag** (*bool*) – True if the method is one-sided.

**Return type**

None

**Notes**

By default, a two-sided variant is selected, which is sometimes slightly more robust. However, the one-sided variant is faster because it avoids the orthogonalization associated to left singular vectors.

**See also**

[`getTRLanczosOneSide`](#), [`SVDLanczosSetOneSide`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1712 <slepc4py/SLEPc/SVD.pyx#L1712>`

**setTRLanczosRestart**(*keep*)

Set the restart parameter for the thick-restart Lanczos method.

Logically collective.

Set the restart parameter for the thick-restart Lanczos method, in particular the proportion of basis vectors that must be kept after restart.

**Parameters**

**keep** (*float*) – The number of vectors to be kept at restart.

**Return type**

None

**Notes**

Allowed values are in the range [0.1,0.9]. The default is 0.5.

**See also**

[`getTRLanczosRestart`](#), [`SVDTRLanczosSetRestart`](#)

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1799 <slepc4py/SLEPc/SVD.pyx#L1799>`

**setThreshold**(*thres, rel=False*)

Set the threshold used in the threshold stopping test.

Logically collective.

**Parameters**

- **thres** (*float*) – The threshold.
- **rel** (*bool*) – Whether the threshold is relative or not.

**Return type**

None

**Notes**

This function internally sets a special stopping test based on the threshold, where singular values are computed in sequence until one of the computed singular values is below/above the threshold (depending on whether largest or smallest singular values are computed).

In the case of largest singular values, the threshold can be made relative with respect to the largest singular value (i.e., the matrix norm).

The details are given in `SVDSetThreshold`.

**See also**

`setStoppingTest`, `getThreshold`, `SVDSetThreshold`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:583 <slepc4py/SLEPc/SVD.pyx#L583>`

**setTolerances**(*tol=None, max\_it=None*)

Set the tolerance and maximum iteration count used.

Logically collective.

Set the tolerance and maximum iteration count used by the default SVD convergence tests.

**Parameters**

- **tol** (*float* | *None*) – The convergence tolerance.
- **max\_it** (*int* | *None*) – The maximum number of iterations

**Return type**

None

**Notes**

Use `DETERMINE` for `max_it` to assign a reasonably good value, which is dependent on the solution method.

**See also**

`getTolerances`, `SVDSetTolerances`

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:643 <slepc4py/SLEPc/SVD.pyx#L643>`

**setTrackAll**(*trackall*)

Set flag to compute the residual of all singular triplets.

Logically collective.

Set if the solver must compute the residual of all approximate singular triplets or not.

**Parameters**

- **trackall** (*bool*) – Whether to compute all residuals or not.

**Return type**

None

**See also**

[getTrackAll](#), [SVDSetTrackAll](#)

**:sources:** ``Source code at slepc4py/SLEPc/SVD.pyx:732 <slepc4py/SLEPc/SVD.pyx#L732>``

**setType**(*svd\_type*)

Set the particular solver to be used in the SVD object.

Logically collective.

**Parameters**

**svd\_type** ([Type](#) / *str*) – The solver to be used.

**Return type**

[None](#)

**Notes**

The default is [CROSS](#). Normally, it is best to use [setFromOptions\(\)](#) and then set the SVD type from the options database rather than by using this routine. Using the options database provides the user with maximum flexibility in evaluating the different available methods.

**See also**

[getType](#), [SVDSetType](#)

**:sources:** ``Source code at slepc4py/SLEPc/SVD.pyx:261 <slepc4py/SLEPc/SVD.pyx#L261>``

**setUp()**

Set up all the internal data structures.

Collective.

**Notes**

Sets up all the internal data structures necessary for the execution of the singular value solver.

This function need not be called explicitly in most cases, since [solve\(\)](#) calls it. It can be useful when one wants to measure the set-up time separately from the solve time.

**See also**

[solve](#), [SVDSetUp](#)

**:sources:** ``Source code at slepc4py/SLEPc/SVD.pyx:1164 <slepc4py/SLEPc/SVD.pyx#L1164>``

**Return type**

[None](#)

**setWhichSingularTriplets**(*which*)

Set which singular triplets are to be sought.

Logically collective.

**Parameters**

**which** ([Which](#)) – The singular values to be sought (either largest or smallest).



**Return type**

None

**See also**

[\*getWhichSingularTriplets\*](#), [\*SVDSetWhichSingularTriplets\*](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:543](#) <[slepc4py/SLEPc/SVD.pyx#L543](#)>

**solve()**

Solve the singular value problem.

Collective.

**Notes**

The problem matrices are specified with [\*setOperators\(\)\*](#).

[\*solve\(\)\*](#) will return without generating an error regardless of whether all requested solutions were computed or not. Call [\*getConverged\(\)\*](#) to get the actual number of computed solutions, and [\*getConvergedReason\(\)\*](#) to determine if the solver converged or failed and why.

**See also**

[\*setUp\*](#), [\*setOperators\*](#), [\*getConverged\*](#), [\*getConvergedReason\*](#), [\*SVDSolve\*](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1185](#) <[slepc4py/SLEPc/SVD.pyx#L1185](#)>

**Return type**

None

**valuesView(viewer=None)**

Display the computed singular values in a viewer.

Collective.

**Parameters**

**viewer** ([\*Viewer\*](#) / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

None

**See also**

[\*solve\*](#), [\*vectorsView\*](#), [\*errorView\*](#), [\*SVDValuesView\*](#)

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1456](#) <[slepc4py/SLEPc/SVD.pyx#L1456](#)>

**vectorsView(viewer=None)**

Output computed singular vectors to a viewer.

Collective.

**Parameters**

**viewer** ([\*Viewer\*](#) / *None*) – Visualization context; if not provided, the standard output is used.

**Return type**

None

**See also**[solve](#), [valuesView](#), [errorView](#), [SVDVectorsView](#)**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1475 <slepc4py/SLEPc/SVD.pyx#L1475>](#)**view**(viewer=None)

Print the SVD data structure.

Collective.

**Parameters****viewer** ([Viewer](#) / [None](#)) – Visualization context; if not provided, the standard output is used.**Return type**

None

**See also**[SVDView](#)**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:195 <slepc4py/SLEPc/SVD.pyx#L195>](#)**Attributes Documentation****ds**The direct solver (*DS*) object associated.**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:2013 <slepc4py/SLEPc/SVD.pyx#L2013>](#)**max\_it**

The maximum iteration count.

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1999 <slepc4py/SLEPc/SVD.pyx#L1999>](#)**problem\_type**

The type of the eigenvalue problem.

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1971 <slepc4py/SLEPc/SVD.pyx#L1971>](#)**tol**

The tolerance.

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1992 <slepc4py/SLEPc/SVD.pyx#L1992>](#)**track\_all**

Compute the residual norm of all approximate eigenpairs.

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:2006 <slepc4py/SLEPc/SVD.pyx#L2006>](#)**transpose\_mode**

How to handle the transpose of the matrix.

**:sources:** [Source code at slepc4py/SLEPc/SVD.pyx:1978 <slepc4py/SLEPc/SVD.pyx#L1978>](#)

which

The portion of the spectrum to be sought.

**:sources:** `Source code at slepc4py/SLEPc/SVD.pyx:1985 <slepc4py/SLEPc/SVD.pyx#L1985>`

## slepc4py.SLEPc.Sys

**class** `slepc4py.SLEPc.Sys`

Bases: `object`

System utilities.

### Methods Summary

<code>getVersion([devel, date, author])</code>	Return SLEPc version information.
<code>getVersionInfo()</code>	Return SLEPc version information.
<code>hasExternalPackage(package)</code>	Return whether SLEPc has support for external package.
<code>isFinalized()</code>	Return whether SLEPc has been finalized.
<code>isInitialized()</code>	Return whether SLEPc has been initialized.

### Methods Documentation

**classmethod** `getVersion(devel=False, date=False, author=False)`

Return SLEPc version information.

Not collective.

#### Parameters

- **devel** (`bool`) – Additionally, return whether using an in-development version.
- **date** (`bool`) – Additionally, return date information.
- **author** (`bool`) – Additionally, return author information.

#### Returns

- **major** (`int`) – Major version number.
- **minor** (`int`) – Minor version number.
- **micro** (`int`) – Micro (or patch) version number.

#### Return type

`tuple[int, int, int]`

#### See also

`SlepcGetVersion`, `SlepcGetVersionNumber`

**:sources:** `Source code at slepc4py/SLEPc/Sys.pyx:8 <slepc4py/SLEPc/Sys.pyx#L8>`

**classmethod** `getVersionInfo()`

Return SLEPc version information.

Not collective.

**Returns**

**info** – Dictionary with version information.

**Return type**

dict

**See also**

[SlepcGetVersion](#), [SlepcGetVersionNumber](#)

**:sources:** [Source code at slepc4py/SLEPc/Sys.pyx:64 <slepc4py/SLEPc/Sys.pyx#L64>](#)

**classmethod** **hasExternalPackage**(*package*)

Return whether SLEPc has support for external package.

Not collective.

**Parameters**

**package** (*str*) – The external package name.

**Return type**

bool

**See also**

[SlepcHasExternalPackage](#)

**:sources:** [Source code at slepc4py/SLEPc/Sys.pyx:118 <slepc4py/SLEPc/Sys.pyx#L118>](#)

**classmethod** **isFinalized**()

Return whether SLEPc has been finalized.

Not collective.

**See also**

[isInitialized](#)

**:sources:** [Source code at slepc4py/SLEPc/Sys.pyx:103 <slepc4py/SLEPc/Sys.pyx#L103>](#)

**Return type**

bool

**classmethod** **isInitialized**()

Return whether SLEPc has been initialized.

Not collective.

**See also**

[isFinalized](#)

**:sources:** [Source code at slepc4py/SLEPc/Sys.pyx:90 <slepc4py/SLEPc/Sys.pyx#L90>](#)

**Return type**  
bool

## slepc4py.SLEPc.Util

**class** slepc4py.SLEPc.Util

Bases: `object`

Other utilities such as the creation of structured matrices.

## Methods Summary

<code>createMatBSE(R, C)</code>	Create a matrix that can be used to define a BSE type problem.
<code>createMatHamiltonian(A, B, C)</code>	Create matrix to be used for a structured Hamiltonian eigenproblem.

## Methods Documentation

**classmethod** `createMatBSE(R, C)`

Create a matrix that can be used to define a BSE type problem.

Collective.

Create a matrix that can be used to define a structured eigenvalue problem of type BSE (Bethe-Salpeter Equation).

### Parameters

- **R** (`petsc4py.PETSc.Mat`) – The matrix for the diagonal block (resonant).
- **C** (`petsc4py.PETSc.Mat`) – The matrix for the off-diagonal block (coupling).

### Returns

The matrix with the block form  $H = \begin{bmatrix} R & C \\ -C^* & -R^T \end{bmatrix}$ .

### Return type

`petsc4py.PETSc.Mat`

### See also

`MatCreateBSE`

**:sources:** `Source code at slepc4py/SLEPc/Util.pyx:8 <slepc4py/SLEPc/Util.pyx#L8>`

**classmethod** `createMatHamiltonian(A, B, C)`

Create matrix to be used for a structured Hamiltonian eigenproblem.

Collective.

### Parameters

- **A** (`petsc4py.PETSc.Mat`) – The matrix for (0,0) block.
- **B** (`petsc4py.PETSc.Mat`) – The matrix for (0,1) block, must be real symmetric or Hermitian.
- **C** (`petsc4py.PETSc.Mat`) – The matrix for (1,0) block, must be real symmetric or Hermitian.

### Returns

The matrix with the block form  $H = \begin{bmatrix} A & B \\ C & -A^* \end{bmatrix}$ .

### Return type

`petsc4py.PETSc.Mat`

### See also

`MatCreateHamiltonian`

**:sources:** `Source code at slepc4py/SLEPc/Util.pyx:38 <slepc4py/SLEPc/Util.pyx#L38>`

### Attributes

<i>DECIDE</i>	Constant DECIDE of type <code>int</code>
<i>DEFAULT</i>	Constant DEFAULT of type <code>int</code>
<i>DETERMINE</i>	Constant DETERMINE of type <code>int</code>
<i>CURRENT</i>	Constant CURRENT of type <code>int</code>

### `slepc4py.SLEPc.DECIDE`

`slepc4py.SLEPc.DECIDE: int = DECIDE`

Constant DECIDE of type `int`

### `slepc4py.SLEPc.DEFAULT`

`slepc4py.SLEPc.DEFAULT: int = DEFAULT`

Constant DEFAULT of type `int`

### `slepc4py.SLEPc.DETERMINE`

`slepc4py.SLEPc.DETERMINE: int = DETERMINE`

Constant DETERMINE of type `int`

### `slepc4py.SLEPc.CURRENT`

`slepc4py.SLEPc.CURRENT: int = CURRENT`

Constant CURRENT of type `int`

## 3.7 slepc4py demos

### ex1.py: Standard symmetric eigenproblem for the 1-D Laplacian

This example computes eigenvalues and eigenvectors of the discrete Laplacian on a one-dimensional domain with finite differences.

The full source code for this demo can be [downloaded here](#).

The first thing to do is initialize the libraries. This is normally not required, as it is done automatically at import time. However, if you want to gain access to the facilities for setting command-line options, the following lines must be executed by the main script prior to any `petsc4py` or `slepc4py` calls:

```
import sys, slepc4py
slepc4py.init(sys.argv)
```

Next, we have to import the relevant modules. Normally, both PETSc and SLEPc modules have to be imported in all slepc4py programs. It may be useful to import NumPy as well:

```
from petsc4py import PETSc
from slepc4py import SLEPc
import numpy
```

At this point, we can use any petsc4py and slepc4py operations. For instance, the following lines allow the user to specify an integer command-line argument `n` with a default value of 30 (see below for example usage of command-line options):

```
opts = PETSc.Options()
n = opts.getInt('n', 30)
```

It is necessary to build a matrix to define an eigenproblem (or two in the case of generalized eigenproblems). The following fragment of code creates the matrix object and then fills the non-zero elements one by one. The matrix of this particular example is tridiagonal, with value 2 in the diagonal, and -1 in off-diagonal positions. See petsc4py documentation for details about matrix objects:

```
A = PETSc.Mat(); A.create()
A.setSizes([n, n])
A.setFromOptions()

rstart, rend = A.getOwnershipRange()

# first row
if rstart == 0:
    A[0, :2] = [2, -1]
    rstart += 1
# last row
if rend == n:
    A[n-1, -2:] = [-1, 2]
    rend -= 1
# other rows
for i in range(rstart, rend):
    A[i, i-1:i+2] = [-1, 2, -1]

A.assemble()
```

The solver object is created in a similar way as other objects in petsc4py:

```
E = SLEPc.EPS(); E.create()
```

Once the object is created, the eigenvalue problem must be specified. At least one matrix must be provided. The problem type must be indicated as well, in this case it is HEP (Hermitian eigenvalue problem). Apart from these, other settings could be provided here (for instance, the tolerance for the computation). After all options have been set, the user should call the `setFromOptions()` operation, so that any options specified at run time in the command line are passed to the solver object:

```
E.setOperators(A)
E.setProblemType(SLEPc.EPS.ProblemType.HEP)
```

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```
history = []
def monitor(eps, its, nconv, eig, err):
    if nconv < len(err): history.append(err[nconv])
E.setMonitor(monitor)

E.setFromOptions()
```

After that, the `solve()` method will run the selected eigensolver, keeping the solution stored internally:

```
E.solve()
```

Once the computation has finished, we are ready to print the results. First, some informative data can be retrieved from the solver object:

```
Print = PETSc.Sys.Print

Print()
Print("*****")
Print("*** SLEPc Solution Results ***")
Print("*****")
Print()

its = E.getIterationNumber()
Print( "Number of iterations of the method: %d" % its )

eps_type = E.getType()
Print( "Solution method: %s" % eps_type )

nev, ncv, mpd = E.getDimensions()
Print( "Number of requested eigenvalues: %d" % nev )

tol, maxit = E.getTolerances()
Print( "Stopping condition: tol=%.4g, maxit=%d" % (tol, maxit) )
```

For retrieving the solution, it is necessary to find out how many eigenpairs have converged to the requested precision:

```
nconv = E.getConverged()
Print( "Number of converged eigenpairs %d" % nconv )
```

For each of the `nconv` eigenpairs, we can retrieve the eigenvalue `k`, and the eigenvector, which is represented by means of two `petsc4py` vectors `vr` and `vi` (the real and imaginary part of the eigenvector, since for real matrices the eigenvalue and eigenvector may be complex). In this example we know that both the eigenvalue and eigenvector are real, so only one vector `v` is needed. We also compute the corresponding relative errors in order to make sure that the computed solution is indeed correct:

```
if nconv > 0:
    # Create the results vectors
    v, _ = A.createVecs()
    #
    Print()
    Print("          k          ||Ax-kx||/||kx|| ")
    Print("-----")
```

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```
for i in range(nconv):
    k = E.getEigenpair(i, v)
    error = E.computeError(i)
    Print( " %12f          %12g" % (k, error) )
Print()
```

### Example of command-line usage

Now we illustrate how to specify command-line options in order to extract the full potential of slepc4py.

A simple execution of the demo/ex1.py script will result in the following output:

```
$ python demo/ex1.py

*****
*** SLEPc Solution Results ***
*****

Number of iterations of the method: 4
Solution method: krylovschur
Number of requested eigenvalues: 1
Stopping condition: tol=1e-07, maxit=100
Number of converged eigenpairs 4

  k          ||Ax-kx||/||kx||
-----
  3.989739      5.76012e-09
  3.959060      1.41957e-08
  3.908279      6.74118e-08
  3.837916      8.34269e-08
```

For specifying different setting for the solver parameters, we can use SLEPc command-line options with the `-eps` prefix. For instance, to change the number of requested eigenvalues and the tolerance:

```
$ python demo/ex1.py -eps_nev 10 -eps_tol 1e-11
```

The method used by the solver object can also be set at run time:

```
$ python demo/ex1.py -eps_type subspace
```

All the above settings can also be changed within the source code by making use of the appropriate slepc4py method. Since options can be set from within the code and the command-line, it is often useful to view the particular settings that are currently being used:

```
$ python demo/ex1.py -eps_view

EPS Object: 1 MPI process
  type: krylovschur
    50% of basis vectors kept after restart
    using the locking variant
  problem type: symmetric eigenvalue problem
  selected portion of the spectrum: largest eigenvalues in magnitude
  number of eigenvalues (nev): 1
```

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```

number of column vectors (ncv): 16
maximum dimension of projected problem (mpd): 16
maximum number of iterations: 100
tolerance: 1e-08
convergence test: relative to the eigenvalue
BV Object: 1 MPI process
  type: mat
  17 columns of global length 30
  orthogonalization method: classical Gram-Schmidt
  orthogonalization refinement: if needed (eta: 0.7071)
  block orthogonalization method: GS
  doing matmult as a single matrix-matrix product
DS Object: 1 MPI process
  type: hep
  solving the problem with: Implicit QR method (_steqr)
ST Object: 1 MPI process
  type: shift
  shift: 0
  number of matrices: 1

```

Note that for computing eigenvalues of smallest magnitude we can use the option `-eps_smallest_magnitude`, but for interior eigenvalues things are not so straightforward. One possibility is to try with harmonic extraction, for instance to get the eigenvalues closest to 0.6:

```
$ python demo/ex1.py -eps_harmonic -eps_target 0.6
```

Depending on the problem, harmonic extraction may fail to converge. In those cases, it is necessary to specify a spectral transformation other than the default. In the command-line, this is indicated with the `-st_` prefix. For example, shift-and-invert with a value of the shift equal to 0.6 would be:

```
$ python demo/ex1.py -st_type sinvert -eps_target 0.6
```

## ex2.py: Standard symmetric eigenproblem for the 2-D Laplacian

This example computes eigenvalues and eigenvectors of the discrete Laplacian on a two-dimensional domain with finite differences.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```

try: range = xrange
except: pass

import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc

Print = PETSc.Sys.Print

```

In this example we have organized the code in several functions. This one builds the finite-difference Laplacian matrix by computing the indices of each entry. An alternative would be to use the functionality offered by [DMDA](#).

```

def construct_operator(m, n):
    # Create matrix for 2D Laplacian operator
    A = PETSc.Mat().create()
    A.setSizes([m*n, m*n])
    A.setFromOptions()
    # Fill matrix
    hx = 1.0/(m-1) # x grid spacing
    hy = 1.0/(n-1) # y grid spacing
    diagv = 2.0*hy/hx + 2.0*hx/hy
    offdx = -1.0*hy/hx
    offdy = -1.0*hx/hy
    Istart, Iend = A.getOwnershipRange()
    for I in range(Istart, Iend) :
        A[I,I] = diagv
        i = I//n # map row number to
        j = I - i*n # grid coordinates
        if i> 0 : J = I-n; A[I,J] = offdx
        if i< m-1: J = I+n; A[I,J] = offdx
        if j> 0 : J = I-1; A[I,J] = offdy
        if j< n-1: J = I+1; A[I,J] = offdy
    A.assemble()
    return A

```

This function receives the matrix and the problem type, then solves the eigenvalue problem and prints information about the computed solution. Although we know that eigenvalues and eigenvectors are real in this example, the function is prepared to solve it as a non-symmetric problem, by passing `SLEPc.EPS.ProblemType.NHEP`, that is why the code handles possibly complex eigenvalues and eigenvectors.

```

def solve_eigensystem(A, problem_type=SLEPc.EPS.ProblemType.HEP):
    # Create the result vectors
    xr, xi = A.createVecs()

    # Setup the eigensolver
    E = SLEPc.EPS().create()
    E.setOperators(A, None)
    E.setDimensions(3, PETSc.DECIDE)
    E.setProblemType(problem_type)
    E.setFromOptions()

    # Solve the eigensystem
    E.solve()

    Print("")
    its = E.getIterationNumber()
    Print("Number of iterations of the method: %i" % its)
    sol_type = E.getType()
    Print("Solution method: %s" % sol_type)
    nev, ncv, mpd = E.getDimensions()
    Print("Number of requested eigenvalues: %i" % nev)
    tol, maxit = E.getTolerances()
    Print("Stopping condition: tol=%.4g, maxit=%d" % (tol, maxit))
    nconv = E.getConverged()
    Print("Number of converged eigenpairs: %d" % nconv)

```

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```
if nconv > 0:
    Print("")
    Print("          k          ||Ax-kx||/||kx|| ")
    Print("-----")
    for i in range(nconv):
        k = E.getEigenpair(i, xr, xi)
        error = E.computeError(i)
        if k.imag != 0.0:
            Print(" %9f%+9f j %12g" % (k.real, k.imag, error))
        else:
            Print(" %12f %12g" % (k.real, error))
    Print("")
```

The main program simply processes three user-defined command-line options and calls the other two functions.

```
def main():
    opts = PETSc.Options()
    N = opts.getInt('N', 32)
    m = opts.getInt('m', N)
    n = opts.getInt('n', m)
    Print("Symmetric Eigenproblem (sparse matrix), "
          "N=%d (%dx%d grid)" % (m*n, m, n))
    A = construct_operator(m,n)
    solve_eigensystem(A)

if __name__ == '__main__':
    main()
```

### ex3.py: Matrix-free eigenproblem for the 2-D Laplacian

This example solves the eigenproblem for the 2-D discrete Laplacian without building the matrix explicitly.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```
import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc
import numpy as np
```

In this case, the program cannot be run in parallel, so we check that the number of MPI processes is 1. In order to enable parallelism, we should implement a parallel matrix-vector operation ourselves, which is not done in this example.

```
assert PETSc.COMM_WORLD.getSize() == 1

Print = PETSc.Sys.Print
```

This function computes the matrix-vector product  $f = L \cdot x$  where the Laplacian  $L$  is not built explicitly, and the vectors  $x, f$  are viewed as two-dimensional arrays associated to grid points.

```
def laplace2d(U, x, f):
    U[:, :] = 0
    U[1:-1, 1:-1] = x
    # Grid spacing
    m, n = x.shape
    hx = 1.0/(m-1) # x grid spacing
    hy = 1.0/(n-1) # y grid spacing
    # Setup 5-points stencil
    u = U[1:-1, 1:-1] # center
    uN = U[1:-1, :-2] # north
    uS = U[1:-1, 2:] # south
    uW = U[: -2, 1:-1] # west
    uE = U[2:, 1:-1] # east
    # Apply Laplacian
    f[:, :] = \
        (2*u - uE - uW) * (hy/hx) \
        + (2*u - uN - uS) * (hx/hy) \
```

For a matrix-free solution in slepc4py we have to create a class that wraps the matrix-vector operation and optionally other operations of the matrix. In this case, we provide the constructor and the mult operation, that simply calls the laplace2d function above.

```
class Laplacian2D(object):

    def __init__(self, m, n):
        self.m, self.n = m, n
        scalar = PETSc.ScalarType
        self.U = np.zeros([m+2, n+2], dtype=scalar)

    def mult(self, A, x, y):
        m, n = self.m, self.n
        xx = x.getArray(readonly=1).reshape(m,n)
        yy = y.getArray(readonly=0).reshape(m,n)
        laplace2d(self.U, xx, yy)
```

In this example, building the matrix amounts to creating an object of the class defined above, and passing it to a special petsc4py matrix with `createPython()`.

```
def construct_operator(m, n):
    # Create shell matrix
    context = Laplacian2D(m,n)
    A = PETSc.Mat().createPython([m*n,m*n], context)
    return A
```

This function receives the matrix and the problem type, then solves the eigenvalue problem and prints information about the computed solution. Although we know that eigenvalues and eigenvectors are real in this example, the function is prepared to solve it as a non-symmetric problem, by passing `SLEPc.EPS.ProblemType.NHEP`, that is why the code handles possibly complex eigenvalues and eigenvectors.

```
def solve_eigensystem(A, problem_type=SLEPc.EPS.ProblemType.HEP):
    # Create the result vectors
    xr, xi = A.createVecs()
```

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```

# Setup the eigensolver
E = SLEPc.EPS().create()
E.setOperators(A, None)
E.setDimensions(3, PETSc.DECIDE)
E.setProblemType(problem_type)
E.setFromOptions()

# Solve the eigensystem
E.solve()
Print("")
its = E.getIterationNumber()
Print("Number of iterations of the method: %i" % its)
sol_type = E.getType()
Print("Solution method: %s" % sol_type)
nev, ncv, mpd = E.getDimensions()
Print("Number of requested eigenvalues: %i" % nev)
tol, maxit = E.getTolerances()
Print("Stopping condition: tol=%.4g, maxit=%d" % (tol, maxit))
nconv = E.getConverged()
Print("Number of converged eigenpairs: %d" % nconv)
if nconv > 0:
    Print("")
    Print("          k          ||Ax-kx||/||kx|| ")
    Print("-----")
    for i in range(nconv):
        k = E.getEigenpair(i, xr, xi)
        error = E.computeError(i)
        if k.imag != 0.0:
            Print(" %9f%+9f j %12g" % (k.real, k.imag, error))
        else:
            Print(" %12f %12g" % (k.real, error))
    Print("")

```

The main program simply processes three user-defined command-line options and calls the other two functions.

```

def main():
    opts = PETSc.Options()
    N = opts.getInt('N', 32)
    m = opts.getInt('m', N)
    n = opts.getInt('n', m)
    Print("Symmetric Eigenproblem (matrix-free), "
          "N=%d (%dx%d grid)" % (m*n, m, n))
    A = construct_operator(m,n)
    solve_eigensystem(A)

if __name__ == '__main__':
    main()

```

#### ex4.py: Singular value decomposition of the Lauchli matrix

This example illustrates the use of the SVD solver in slepc4py. It computes singular values and vectors of the Lauchli matrix, whose condition number depends on a parameter  $\mu$ .

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```
try: range = xrange
except: pass

import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc
```

This example takes two command-line arguments, the matrix size  $n$  and the  $\mu$  parameter.

```
opts = PETSc.Options()
n = opts.getInt('n', 30)
mu = opts.getReal('mu', 1e-6)

PETSc.Sys.Print( "Lauchli singular value decomposition, (%d x %d) mu=%g\n" % (n+1, n, mu) )
```

Create the matrix and fill its nonzero entries. Every MPI process will insert its locally owned part only.

```
A = PETSc.Mat(); A.create()
A.setSizes([n+1, n])
A.setFromOptions()

rstart, rend = A.getOwnershipRange()

for i in range(rstart, rend):
    if i==0:
        for j in range(n):
            A[0,j] = 1.0
    else:
        A[i,i-1] = mu

A.assemble()
```

The singular value solver is similar to the eigensolver used in previous examples. In this case, we select the thick-restart Lanczos bidiagonalization method.

```
S = SLEPc.SVD(); S.create()

S.setOperator(A)
S.setType(S.Type.TRLANCZOS)
S.setFromOptions()

S.solve()
```

After solve, we print some informative data and extract the computed solution, showing the list of singular values and the corresponding residual errors.

```

Print = PETSc.Sys.Print

Print( "*****" )
Print( "*** SLEPc Solution Results ***" )
Print( "*****\n" )

svd_type = S.getType()
Print( "Solution method: %s" % svd_type )

its = S.getIterationNumber()
Print( "Number of iterations of the method: %d" % its )

nsv, ncv, mpd = S.getDimensions()
Print( "Number of requested singular values: %d" % nsv )

tol, maxit = S.getTolerances()
Print( "Stopping condition: tol=%.4g, maxit=%d" % (tol, maxit) )

nconv = S.getConverged()
Print( "Number of converged approximate singular triplets %d" % nconv )

if nconv > 0:
    v, u = A.createVecs()
    Print()
    Print("      sigma      residual norm ")
    Print("-----")
    for i in range(nconv):
        sigma = S.getSingularTriplet(i, u, v)
        error = S.computeError(i)
        Print( "      %6f      %12g" % (sigma, error) )
    Print()

```

### ex5.py: Simple quadratic eigenvalue problem

This example solves a polynomial eigenvalue problem of degree 2, which is the most commonly found in applications. Larger degree polynomials are handled similarly. The coefficient matrices in this example do not come from an application, they are just simple matrices that are easy to build, just to illustrate how it works. The eigenvalues in this example are purely imaginary and come in conjugate pairs.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```

import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc

Print = PETSc.Sys.Print

```

A function to build the matrices. The lowest degree coefficient is the 2-D Laplacian, the highest degree one is the identity matrix, and the other matrix is set to zero (which means that this problem could have been solved as a linear eigenproblem).



```

def construct_operators(m,n):
    Print("Quadratic Eigenproblem, N=%d (%dx%d grid)"% (m*n, m, n))
    # K is the 2-D Laplacian
    K = PETSc.Mat().create()
    K.setSizes([n*m, n*m])
    K.setFromOptions()
    Istart, Iend = K.getOwnershipRange()
    for I in range(Istart,Iend):
        v = -1.0; i = I//n; j = I-i*n;
        if i>0:
            J=I-n; K[I,J] = v
        if i<m-1:
            J=I+n; K[I,J] = v
        if j>0:
            J=I-1; K[I,J] = v
        if j<n-1:
            J=I+1; K[I,J] = v
        v=4.0; K[I,I] = v
    K.assemble()
    # C is the zero matrix
    C = PETSc.Mat().create()
    C.setSizes([n*m, n*m])
    C.setFromOptions()
    C.assemble()
    # M is the identity matrix
    M = PETSc.Mat().createConstantDiagonal([n*m, n*m], 1.0)

    return M, C, K

```

The polynomial eigenvalue solver is similar to the linear eigensolver used in previous examples. The main difference is that we must provide a list of matrices, from lowest to highest degree.

```

def solve_eigensystem(M, C, K):
    # Setup the eigensolver
    Q = SLEPc.PEP().create()
    Q.setOperators([K, C, M])
    Q.setDimensions(6)
    Q.setProblemType(SLEPc.PEP.ProblemType.GENERAL)
    Q.setFromOptions()
    # Solve the eigensystem
    Q.solve()
    # Create the result vectors
    xr, xi = K.createVecs()

    its = Q.getIterationNumber()
    Print("Number of iterations of the method: %i" % its)
    sol_type = Q.getType()
    Print("Solution method: %s" % sol_type)
    nev, ncv, mpd = Q.getDimensions()
    Print("")
    Print("Number of requested eigenvalues: %i" % nev)
    tol, maxit = Q.getTolerances()
    Print("Stopping condition: tol=%.4g, maxit=%d" % (tol, maxit))

```

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```
nconv = Q.getConverged()
Print("Number of converged approximate eigenpairs: %d" % nconv)
if nconv > 0:
    Print("")
    Print("          k          ||(k^2M+Ck+K)x||/||kx|| ")
    Print("-----")
    for i in range(nconv):
        k = Q.getEigenpair(i, xr, xi)
        error = Q.computeError(i)
        if k.imag != 0.0:
            Print("%9f%+9f j    %12g" % (k.real, k.imag, error))
        else:
            Print("%12f          %12g" % (k.real, error))
    Print("")
```

The main program simply processes two user-defined command-line options (the dimensions of the mesh) and calls the other two functions.

```
if __name__ == '__main__':
    opts = PETSc.Options()
    m = opts.getInt('m', 32)
    n = opts.getInt('n', m)
    M, C, K = construct_operators(m,n)
    solve_eigensystem(M, C, K)
    M = C = K = None
```

### ex6.py: Compute $\exp(tA)v$ for a matrix from a Markov model

This example illustrates the functionality in slepc4py for computing matrix functions, or more precisely, the application of a matrix function on a given vector. The example works with the exponential function, which is most commonly found in applications.

The main focus of slepc4py is eigenvalue and singular value problems, but it has some codes to deal with matrix functions, which sometimes are needed in the context of eigenproblems, but have interest on their own.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```
import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc

Print = PETSc.Sys.Print
```

This function builds a matrix that implements a Markov model of a random walk on a triangular grid. The entries of the matrix represent probabilities of moving to neighboring cells in the grid.

```
def build_matrix(m):
    N = m*(m+1)/2
    Print("Markov y=exp(t*A)*e_1, N=%d (m=%d)" % (N, m))
    A = PETSc.Mat().create()
```

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```
A.setSizes([N, N])
A.setFromOptions()
Istart, Iend = A.getOwnershipRange()
ix = 0
cst = 0.5/(m-1)
for i in range(1,m+1):
    jmax = m-i+1
    for j in range(1,jmax+1):
        ix = ix + 1
        if ix-1<Istart or ix>Iend:
            continue # compute only owned rows
        if j!=jmax:
            pd = cst*(i+j-1)
            # north
            if i==1:
                A[ix-1,ix] = 2*pd
            else:
                A[ix-1,ix] = pd
            # east
            if j==1:
                A[ix-1,ix+jmax-1] = 2*pd
            else:
                A[ix-1,ix+jmax-1] = pd
            # south
            pu = 0.5 - cst*(i+j-3)
            if j>1:
                A[ix-1,ix-2] = pu
            # west
            if i>1:
                A[ix-1,ix-jmax-2] = pu

A.assemble()
return A
```

The following function solves the problem. This case is quite different from eigenproblems, and is more similar to solving a linear system of equations with `KSP`. To configure the problem we must provide the matrix and the function (the exponential in this case). Note how the internal `FN` object is extracted from the `MFN` solver. Also, it is often necessary to specify a scale factor, which in this case represents the time for which we want to obtain the evolved state. Once the solver is set up, we call `solve()` passing the right-hand side vector `b` and the solution vector `x`.

```
def solve_exp(t, A, b, x):
    # Setup the solver
    M = SLEPc.MFN().create()
    M.setOperator(A)
    f = M.getFN()
    f.setType(SLEPc.FN.Type.EXP)
    f.setScale(t)
    M.setTolerances(1e-7)
    M.setFromOptions()
    # Solve the problem
    M.solve(b,x)
```

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```
its = M.getIterationNumber()
Print("Number of iterations of the method: %i" % its)
sol_type = M.getType()
Print("Solution method: %s" % sol_type)
ncv = M.getDimensions()
Print("")
Print("Subspace dimension: %i" % ncv)
tol, maxit = M.getTolerances()
Print("Stopping condition: tol=%.4g, maxit=%d" % (tol, maxit))
Print("Computed vector at time t=%.4g has norm %g" % (t.real, x.norm()))
Print("")
```

The main program processes the command-line option `m` (size of the grid), builds the matrix and calls the solver. Note how the vectors are created from the matrix. In this case, the right-hand side vector is the first element of the canonical basis.

```
if __name__ == '__main__':
    opts = PETSc.Options()
    m = opts.getInt('m', 15)
    A = build_matrix(m) # transition probability matrix
    x, b = A.createVecs()
    x.set(0)
    b.set(0)
    b[0] = 1
    b.assemble()
    t = 2
    solve_exp(t, A, b, x) # compute x=exp(t*A)*b
    A = None
    b = x = None
```

### ex7.py: Nonlinear eigenproblem with callback functions

This example solves a nonlinear eigenvalue problem arising from the discretization of a PDE on a one-dimensional domain with finite differences. The nonlinearity comes from the boundary conditions. The PDE is

$$-u'' = \lambda u$$

defined on the interval  $[0,1]$  and subject to the boundary conditions

$$u(0) = 0, u'(1) = u(1)\lambda \frac{\kappa}{\kappa - \lambda},$$

where  $\lambda$  is the eigenvalue and  $\kappa$  is a parameter.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```
import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc
from numpy import sqrt, sin

Print = PETSc.Sys.Print
```

When implementing a nonlinear eigenproblem with callback functions we must provide code that builds the function matrix  $T(\lambda)$  for a given  $\lambda$  and optionally the Jacobian matrix  $T'(\lambda)$ , i.e., the derivative with respect to the eigenvalue.

In `slepc4py` the callbacks are integrated in a class. In this example, apart from the constructor, we have three methods:

- `formFunction` to fill the function matrix `F`. Note that `F` is received as an argument and we just need to fill its entries using the value of the parameter `mu`. Matrix `B` is used to build the preconditioner, and is usually equal to `F`.
- `formJacobian` to fill the Jacobian matrix `J`. Some eigensolvers do not need this, but it is recommended to implement it.
- `checkSolution` is just a convenience method to check that a given solution satisfies the PDE.

```
class MyPDE(object):

    def __init__(self, kappa, h):
        self.kappa = kappa
        self.h      = h

    def formFunction(self, nep, mu, F, B):
        n, m = F.getSize()
        Istart, Iend = F.getOwnershipRange()
        i1 = Istart
        if Istart==0: i1 = i1 + 1
        i2 = Iend
        if Iend==n: i2 = i2 - 1
        h = self.h
        c = self.kappa/(mu-self.kappa)
        d = n

        # Interior grid points
        for i in range(i1,i2):
            val = -d-mu*h/6.0
            F[i,i-1] = val
            F[i,i]   = 2.0*(d-mu*h/3.0)
            F[i,i+1] = val

        # Boundary points
        if Istart==0:
            F[0,0] = 2.0*(d-mu*h/3.0)
            F[0,1] = -d-mu*h/6.0
        if Iend==n:
            F[n-1,n-2] = -d-mu*h/6.0
            F[n-1,n-1] = d-mu*h/3.0+mu*c

        F.assemble()
        if B != F: B.assemble()
        return PETSc.Mat.Structure.SAME_NONZERO_PATTERN

    def formJacobian(self, nep, mu, J):
        n, m = J.getSize()
        Istart, Iend = J.getOwnershipRange()
        i1 = Istart
        if Istart==0: i1 = i1 + 1
```

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```
i2 = Iend
if Iend==n: i2 = i2 - 1
h = self.h
c = self.kappa/(mu-self.kappa)

# Interior grid points
for i in range(i1,i2):
    J[i,i-1] = -h/6.0
    J[i,i]   = -2.0*h/3.0
    J[i,i+1] = -h/6.0

# Boundary points
if Istart==0:
    J[0,0] = -2.0*h/3.0
    J[0,1] = -h/6.0
if Iend==n:
    J[n-1,n-2] = -h/6.0
    J[n-1,n-1] = -h/3.0-c*c

J.assemble()
return PETSc.Mat.Structure.SAME_NONZERO_PATTERN

def checkSolution(self, mu, y):
    nu = sqrt(mu)
    u = y.duplicate()
    n = u.getSize()
    Istart, Iend = J.getOwnershipRange()
    h = self.h
    for i in range(Istart,Iend):
        x = (i+1)*h
        u[i] = sin(nu*x);
    u.assemble()
    u.normalize()
    u.axpy(-1.0,y)
    return u.norm()
```

We use an auxiliary function `FixSign` to force the computed eigenfunction to be real and positive, since some eigensolvers may return the eigenvector multiplied by a complex number of modulus one.

```
def FixSign(x):
    comm = x.getComm()
    rank = comm.getRank()
    n = 1 if rank == 0 else 0
    aux = PETSc.Vec().createMPI((n, PETSc.DECIDE), comm=comm)
    if rank == 0: aux[0] = x[0]
    aux.assemble()
    x0 = aux.sum()
    sign = x0/abs(x0)
    x.scale(1.0/sign)
```

The main program processes two command-line options, `n` (size of the grid) and `kappa` (the parameter of the PDE), then creates an object of the class we have defined previously.

```

opts = PETSc.Options()
n = opts.getInt('n', 128)
kappa = opts.getReal('kappa', 1.0)
pde = MyPDE(kappa, 1.0/n)

```

In order to set up the solver we have to pass the two callback functions (methods of the class) together with the matrix objects that will be used every time these methods are called. In this simple example we can do a preallocation of the matrices, although this is not necessary.

```

nep = SLEPc.NEP().create()
F = PETSc.Mat().create()
F.setSizes([n, n])
F.setType('aij')
F.setPreallocationNNZ(3)
nep.setFunction(pde.formFunction, F)

J = PETSc.Mat().create()
J.setSizes([n, n])
J.setType('aij')
J.setPreallocationNNZ(3)
nep.setJacobian(pde.formJacobian, J)

```

After setting some options, we can solve the problem. Here we also illustrate how to pass an initial guess to the solver.

```

nep.setTolerances(tol=1e-9)
nep.setDimensions(1)
nep.setFromOptions()

x = F.createVecs('right')
x.set(1.0)
nep.setInitialSpace(x)
nep.solve()

```

Once the solver has finished, we print some information together with the computed solution. For each computed eigenpair, we print the residual norm and also the error estimated with the class method `checkSolution`.

```

its = nep.getIterationNumber()
Print("Number of iterations of the method: %i" % its)
sol_type = nep.getType()
Print("Solution method: %s" % sol_type)
nev, ncv, mpd = nep.getDimensions()
Print("")
Print("Subspace dimension: %i" % ncv)
tol, maxit = nep.getTolerances()
Print("Stopping condition: tol=%.4g" % tol)
Print("")

nconv = nep.getConverged()
Print("Number of converged eigenpairs %d" % nconv)

if nconv > 0:
    Print()
    Print("          k          ||T(k)x||          error ")

```

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```
Print("-----")
for i in range(nconv):
    k = nep.getEigenpair(i, x)
    FixSign(x)
    res = nep.computeError(i)
    error = pde.checkSolution(k.real, x)
    if k.imag != 0.0:
        Print( " %9f%+9f j %12g      %12g" % (k.real, k.imag, res, error) )
    else:
        Print( " %12f      %12g      %12g" % (k.real, res, error) )
Print()
```

### ex8.py: Nonlinear eigenproblem with split form

This example solves a nonlinear eigenvalue problem where the nonlinear function is expressed in split form.

We want to solve the following parabolic partial differential equation with time delay  $\tau$

$$\begin{aligned}u_t &= u_{xx} + au(t) + bu(t - \tau) \\ u(0, t) &= u(\pi, t) = 0\end{aligned}$$

with  $a = 20$  and  $b(x) = -4.1 + x(1 - e^{x-\pi})$ .

Discretization leads to a DDE of dimension  $n$

$$-u' = Au(t) + Bu(t - \tau)$$

which results in the nonlinear eigenproblem

$$(-\lambda I + A + e^{-\tau\lambda}B)u = 0.$$

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples. In this case we also need to import some math symbols.

```
import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc
from numpy import exp
from math import pi

Print = PETSc.Sys.Print
```

This script has two command-line options: the discretization size  $n$  and the time delay  $\tau$ .

```
opts = PETSc.Options()
n = opts.getInt('n', 128)
tau = opts.getReal('tau', 0.001)
a = 20
h = pi/(n+1)
```

Next we have to set up the solver. In this case, we are going to represent the nonlinear problem in split form, i.e., as a sum of terms made of a constant matrix multiplied by a scalar nonlinear function.



```
nep = SLEPc.NEP().create()
```

The first term involves the identity matrix.

```
Id = PETSc.Mat().createConstantDiagonal([n, n], 1.0)
```

The second term has a tridiagonal matrix obtained from the discretization,  $A = \frac{1}{h^2} \text{tridiag}(1, -2, 1) + aI$ .

```
A = PETSc.Mat().create()
A.setSizes([n, n])
A.setFromOptions()
rstart, rend = A.getOwnershipRange()
vd = -2.0/(h*h)+a
vo = 1.0/(h*h)
if rstart == 0:
    A[0, :2] = [vd, vo]
    rstart += 1
if rend == n:
    A[n-1, -2:] = [vo, vd]
    rend -= 1
for i in range(rstart, rend):
    A[i, i-1:i+2] = [vo, vd, vo]
A.assemble()
```

The third term includes a diagonal matrix  $B = \text{diag}(b(x_i))$ .

```
B = PETSc.Mat().create()
B.setSizes([n, n])
B.setFromOptions()
rstart, rend = B.getOwnershipRange()
for i in range(rstart, rend):
    xi = (i+1)*h
    B[i, i] = -4.1+xi*(1.0-exp(xi-pi));
B.assemble()
B.setOption(PETSc.Mat.Option.HERMITIAN, True)
```

Apart from the matrices, we have to create the functions, represented with `FN` objects:  $f_1 = -\lambda, f_2 = 1, f_3 = \exp(-\tau\lambda)$ .

```
f1 = SLEPc.FN().create()
f1.setType(SLEPc.FN.Type.RATIONAL)
f1.setRationalNumerator([-1, 0])
f2 = SLEPc.FN().create()
f2.setType(SLEPc.FN.Type.RATIONAL)
f2.setRationalNumerator([1])
f3 = SLEPc.FN().create()
f3.setType(SLEPc.FN.Type.EXP)
f3.setScale(-tau)
```

Put all the information together to define the split operator. Note that `A` is passed first so that `SUBSET nonzero_pattern` can be used.

```
nep.setSplitOperator([A, Id, B], [f2, f1, f3], PETSc.Mat.Structure.SUBSET)
```

Now we can set some options and call the solver.

```

nep.setTolerances(tol=1e-9)
nep.setDimensions(1)
nep.setFromOptions()

nep.solve()

```

Once the solver has finished, we print some information together with the computed solution. For each computed eigenpair, we print the eigenvalue and the residual norm.

```

its = nep.getIterationNumber()
Print("Number of iterations of the method: %i" % its)
sol_type = nep.getType()
Print("Solution method: %s" % sol_type)
nev, ncv, mpd = nep.getDimensions()
Print("")
Print("Subspace dimension: %i" % ncv)
tol, maxit = nep.getTolerances()
Print("Stopping condition: tol=%.4g" % tol)
Print("")

nconv = nep.getConverged()
Print( "Number of converged eigenpairs %d" % nconv )

if nconv > 0:
    x = Id.createVecs('right')
    x.set(1.0)
    Print()
    Print("          k          ||T(k)x||")
    Print("-----")
    for i in range(nconv):
        k = nep.getEigenpair(i, x)
        res = nep.computeError(i)
        if k.imag != 0.0:
            Print( " %9f%+9f j %12g" % (k.real, k.imag, res) )
        else:
            Print( " %12f          %12g" % (k.real, res) )
    Print()

```

### ex9.py: Generalized symmetric-definite eigenproblem

This example computes eigenvalues and eigenvectors of a generalized symmetric-definite eigenvalue problem, where the first matrix is the discrete Laplacian in two dimensions and the second matrix is quasi diagonal.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```

try: range = xrange
except: pass

import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc

```

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```
from slepc4py import SLEPc

Print = PETSc.Sys.Print
```

This function builds the discretized Laplacian operator in 2 dimensions.

```
def Laplacian2D(m, n):
    # Create matrix for 2D Laplacian operator
    A = PETSc.Mat().create()
    A.setSizes([m*n, m*n])
    A.setFromOptions()
    # Fill matrix
    hx = 1.0/(m-1) # x grid spacing
    hy = 1.0/(n-1) # y grid spacing
    diagv = 2.0*hy/hx + 2.0*hx/hy
    offdx = -1.0*hy/hx
    offdy = -1.0*hx/hy
    Istart, Iend = A.getOwnershipRange()
    for I in range(Istart, Iend):
        A[I,I] = diagv
        i = I//n # map row number to
        j = I - i*n # grid coordinates
        if i> 0 : J = I-n; A[I,J] = offdx
        if i< m-1: J = I+n; A[I,J] = offdx
        if j> 0 : J = I-1; A[I,J] = offdy
        if j< n-1: J = I+1; A[I,J] = offdy
    A.assemble()
    return A
```

This function builds a quasi-diagonal matrix. It is two times the identity matrix except for the 2x2 leading submatrix  $\begin{bmatrix} 6 & -1 \\ -1 & 1 \end{bmatrix}$ .

```
def QuasiDiagonal(N):
    # Create matrix
    B = PETSc.Mat().create()
    B.setSizes([N, N])
    B.setFromOptions()
    # Fill matrix
    Istart, Iend = B.getOwnershipRange()
    for I in range(Istart, Iend):
        B[I,I] = 2.0
    if Istart==0:
        B[0,0] = 6.0
        B[0,1] = -1.0
        B[1,0] = -1.0
        B[1,1] = 1.0
    B.assemble()
    return B
```

The following function receives the two matrices and solves the eigenproblem. In this example we illustrate how to pass objects that have been created beforehand, instead of extracting the internal objects. We are using a spectral transformation of type `ST.Type.PRECOND` and a Block Jacobi preconditioner. We want to compute the leftmost eigenvalues. The selected eigensolver is LOBPCG, which is appropriate for this use case. After the solve, we print the computed

solution.

```
def solve_eigensystem(A, B, problem_type=SLEPc.EPS.ProblemType.GHEP):
    # Create the results vectors
    xr, xi = A.createVecs()

    pc = PETSc.PC().create()
    # pc.setType(pc.Type.HYPRE)
    pc.setType(pc.Type.BJACOBI)

    ksp = PETSc.KSP().create()
    ksp.setType(ksp.Type.PREONLY)
    ksp.setPC( pc )

    F = SLEPc.ST().create()
    F.setType(F.Type.PRECOND)
    F.setKSP( ksp )
    F.setShift(0)

    # Setup the eigensolver
    E = SLEPc.EPS().create()
    E.setST(F)
    E.setOperators(A,B)
    E.setType(E.Type.LOBPCG)
    E.setDimensions(10,PETSc.DECIDE)
    E.setWhichEigenpairs(E.Which.SMALLEST_REAL)
    E.setProblemType( problem_type )
    E.setFromOptions()

    # Solve the eigensystem
    E.solve()

    Print("")
    its = E.getIterationNumber()
    Print("Number of iterations of the method: %i" % its)
    sol_type = E.getType()
    Print("Solution method: %s" % sol_type)
    nev, ncv, mpd = E.getDimensions()
    Print("Number of requested eigenvalues: %i" % nev)
    tol, maxit = E.getTolerances()
    Print("Stopping condition: tol=%.4g, maxit=%d" % (tol, maxit))
    nconv = E.getConverged()
    Print("Number of converged eigenpairs: %d" % nconv)
    if nconv > 0:
        Print("")
        Print("          k          ||Ax-kx||/||kx|| ")
        Print("-----")
        for i in range(nconv):
            k = E.getEigenpair(i, xr, xi)
            error = E.computeError(i)
            if k.imag != 0.0:
                Print(" %9f%+9f j %12g" % (k.real, k.imag, error))
            else:
```

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```
Print(" %12f      %12g" % (k.real, error))
Print("")
```

The main program simply processes three user-defined command-line options and calls the other functions.

```
def main():
    opts = PETSc.Options()
    N = opts.getInt('N', 10)
    m = opts.getInt('m', N)
    n = opts.getInt('n', m)
    Print("Symmetric-definite Eigenproblem, N=%d (%dx%d grid)" % (m*n, m, n))
    A = Laplacian2D(m,n)
    B = QuasiDiagonal(m*n)
    solve_eigensystem(A,B)

if __name__ == '__main__':
    main()
```

### ex10.py: Laplace problem using the Proper Orthogonal Decomposition

This example program solves the Laplace problem using the Proper Orthogonal Decomposition (POD) reduced-order modeling technique. For a full description of the technique the reader is referred to<sup>1,2</sup>.

The method is split into an offline (computationally intensive) and an online (computationally cheap) phase. This has many applications including real-time simulation, uncertainty quantification and inverse problems, where similar models must be evaluated quickly and many times.

Offline phase:

1. A set of solution snapshots of the 1D Laplace problem in the full problem space are constructed and assembled into the columns of a dense matrix  $S$ .
2. A standard eigenvalue decomposition is performed on the matrix  $S^T S$ .
3. The eigenvectors and eigenvalues are projected back to the original eigenvalue problem  $S$ .
4. The leading eigenvectors then form the POD basis.

Online phase:

1. The operator corresponding to the discrete Laplacian is projected onto the POD basis.
2. The operator corresponding to the right-hand side is projected onto the POD basis.
3. The reduced (dense) problem expressed in the POD basis is solved.
4. The reduced solution is projected back to the full problem space.

Contributed by: Elisa Schenone, Jack S. Hale.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples, but importing some additional modules.

<sup>1</sup> K. Kunisch and S. Volkwein. Galerkin proper orthogonal decomposition methods for a general equation in fluid dynamics. SIAM Journal on Numerical Analysis, 40(2):492-515, 2003.

<sup>2</sup> S. Volkwein, Optimal control of a phase-field model using the proper orthogonal decomposition, Z. Angew. Math. Mech., 81:83-97, 2001.

```

try: range = xrange
except: pass

import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc
import numpy

import random
import math

```

This function builds the discrete Laplacian operator in 1 dimension with homogeneous Dirichlet boundary conditions.

```

def construct_operator(m):
    # Create matrix for 1D Laplacian operator
    A = PETSc.Mat().create(PETSc.COMM_SELF)
    A.setSizes([m, m])
    A.setFromOptions()
    # Fill matrix
    hx = 1.0/(m-1) # x grid spacing
    diagv = 2.0/hx
    offdx = -1.0/hx
    Istart, Iend = A.getOwnershipRange()
    for i in range(Istart, Iend):
        if i != 0 and i != (m - 1):
            A[i, i] = diagv
            if i > 1: A[i, i - 1] = offdx
            if i < m - 2: A[i, i + 1] = offdx
        else:
            A[i, i] = 1.

    A.assemble()

    return A

```

Set bell-shape function as the solution of the Laplacian problem in 1 dimension with homogeneous Dirichlet boundary conditions and compute the associated discrete RHS.

```

def set_problem_rhs(m):
    # Create 1D mass matrix operator
    M = PETSc.Mat().create(PETSc.COMM_SELF)
    M.setSizes([m, m])
    M.setFromOptions()
    # Fill matrix
    hx = 1.0/(m-1) # x grid spacing
    diagv = hx/3
    offdx = hx/6
    Istart, Iend = M.getOwnershipRange()
    for i in range(Istart, Iend):
        if i != 0 and i != (m - 1):
            M[i, i] = 2*diagv

```

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```

else:
    M[i, i] = diagv
    if i > 1: M[i, i - 1] = offdx
    if i < m - 2: M[i, i + 1] = offdx

M.assemble()

x_0 = 0.3
x_f = 0.7
mu = x_0 + (x_f - x_0)*random.random()
sigma = 0.1**2
uex, f = M.createVecs()
for j in range(Istart, Iend):
    value = 2/sigma * math.exp(-(hx*j - mu)**2/sigma) * (1 - 2/sigma * (hx*j -
↪mu)**2 )
    f.setValue(j, value)
    value = math.exp(-(hx*j - mu)**2/sigma)
    uex.setValue(j, value)
f.assemble()
uex.assemble()

RHS = f.duplicate()
M.mult(f, RHS)
RHS.setValue(0, 0.)
RHS.setValue(m-1, 0.)
RHS.assemble()

return RHS, uex

```

Solve 1D Laplace problem with FEM.

```

def solve_laplace_problem(A, RHS):
    u = A.createVecs('right')
    r, c = A.getOrdering("natural")
    A.factorILU(r, c)
    A.solve(RHS, u)
    A.setUnfactored()
    return u

```

Solve 1D Laplace problem with POD (dense matrix).

```

def solve_laplace_problem_pod(A, RHS, u):
    ksp = PETSc.KSP().create(PETSc.COMM_SELF)
    ksp.setOperators(A)
    ksp.setType('preonly')
    pc = ksp.getPC()
    pc.setType('none')
    ksp.setFromOptions()

    ksp.solve(RHS, u)

    return u

```

Set  $N$  solution of the 1D Laplace problem as columns of a matrix (snapshot matrix).

Note: For simplicity we do not perform a linear solve, but use some analytical solution:  $z(x) = \exp(-(x - \mu)^2/\sigma)$ .

```
def construct_snapshot_matrix(A, N, m):
    snapshots = PETSc.Mat().create(PETSc.COMM_SELF)
    snapshots.setSizes([m, N])
    snapshots.setType('seqdense')

    Istart, Iend = snapshots.getOwnershipRange()
    hx = 1.0/(m - 1)
    x_0 = 0.3
    x_f = 0.7
    sigma = 0.1**2
    for i in range(N):
        mu = x_0 + (x_f - x_0)*random.random()
        for j in range(Istart, Iend):
            value = math.exp(-(hx*j - mu)**2/sigma)
            snapshots.setValue(j, i, value)
    snapshots.assemble()

    return snapshots
```

Solve the eigenvalue problem: the eigenvectors of this problem form the POD basis.

```
def solve_eigenproblem(snapshots, N):
    print('Solving POD basis eigenproblem using eigensolver...')

    Es = SLEPc.EPS()
    Es.create(PETSc.COMM_SELF)
    Es.setDimensions(N)
    Es.setProblemType(SLEPc.EPS.ProblemType.NHEP)
    Es.setTolerances(1.0e-8, 500);
    Es.setKrylovSchurRestart(0.6)
    Es.setWhichEigenpairs(SLEPc.EPS.Which.LARGEST_REAL)
    Es.setOperators(snapshots)
    Es.setFromOptions()

    Es.solve()
    print('Solved POD basis eigenproblem.')
    return Es
```

Function to project  $S^T S$  eigenvectors to  $S$  eigenvectors.

```
def project_STS_eigenvectors_to_S_eigenvectors(bvEs, S):
    sizes = S.getSizes()[0]
    N = bvEs.getActiveColumns()[1]
    bv = SLEPc.BV().create(PETSc.COMM_SELF)
    bv.setSizes(sizes, N)
    bv.setActiveColumns(0, N)
    bv.setFromOptions()

    tmpvec2 = S.createVecs('left')
    for i in range(N):
```

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```
    tmpvec = bvEs.getColumn(i)
    S.mult(tmpvec, tmpvec2)
    bv.insertVec(i, tmpvec2)
    bvEs.restoreColumn(i, tmpvec)

    return bv
```

Function to project the reduced space to the full space.

```
def project_reduced_to_full_space(alpha, bv):
    uu = bv.getColumn(0)
    uPOD = uu.duplicate()
    bv.restoreColumn(0,uu)

    scatter, Wr = PETSc.Scatter.toAll(alpha)
    scatter.begin(alpha, Wr, PETSc.InsertMode.INSERT, PETSc.ScatterMode.FORWARD)
    scatter.end(alpha, Wr, PETSc.InsertMode.INSERT, PETSc.ScatterMode.FORWARD)
    PODcoeff = Wr.getArray(readonly=1)

    bv.multVec(1., 0., uPOD, PODcoeff)

    return uPOD
```

The main function realizes the full procedure.

```
def main():
    problem_dim = 200
    num_snapshots = 30
    num_pod_basis_functions = 8

    assert(num_pod_basis_functions <= num_snapshots)

    A = construct_operator(problem_dim)
    S = construct_snapshot_matrix(A, num_snapshots, problem_dim)

    # Instead of solving the SVD of S, we solve the standard
    # eigenvalue problem on S.T*S
    STS = S.transposeMatMult(S)

    Es = solve_eigenproblem(STS, num_pod_basis_functions)
    nconv = Es.getConverged()
    print('Number of converged eigenvalues: %i' % nconv)
    Es.view()

    # get the EPS solution in a BV object
    bvEs = Es.getBV()
    bvEs.setActiveColumns(0, num_pod_basis_functions)

    # set the bv POD basis
    bv = project_STS_eigenvectors_to_S_eigenvectors(bvEs, S)
    # rescale the eigenvectors
    for i in range(num_pod_basis_functions):
```

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```
ll = Es.getEigenvalue(i)
print('Eigenvalue '+str(i)+' : '+str(ll.real))
bv.scaleColumn(i,1.0/math.sqrt(ll.real))

print('-----')
# Verify that the active columns of bv form an orthonormal subspace, i.e. that  $X^H X_L$ 
↪= Id
print('Check that bv.dot(bv) is close to the identity matrix')
XtX = bv.dot(bv)
XtX.view()
XtX_array = XtX.getDenseArray()
n,m = XtX_array.shape
assert numpy.allclose(XtX_array, numpy.eye(n, m))
print('-----')
print('Solve the problem with POD')

# Project the linear operator A
Ared = bv.matProject(A,bv)

# Set the RHS and the exact solution
RHS, uex = set_problem_rhs(problem_dim)

# Project the RHS on the POD basis
RHSred = PETSc.Vec().createWithArray(bv.dotVec(RHS))

# Solve the problem with POD
alpha = Ared.createVecs('right')
alpha = solve_laplace_problem_pod(Ared,RHSred,alpha)

# Project the POD solution back to the FE space
uPOD = project_reduced_to_full_space(alpha, bv)

# Compute the L2 and Linf norm of the error
error = uex.copy()
error.axpy(-1,uPOD)
errorL2 = math.sqrt(error.dot(error).real)
print('The L2-norm of the error is: '+str(errorL2))

print("NORMAL END")

if __name__ == '__main__':
    main()
```

### ex11.py: 2-D Laplacian eigenproblem solved with contour integral

This example is similar to ex2.py, but employs a contour integral solver. It illustrates how to define a region of the complex plane using an *RG* object.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```
try: range = xrange
```

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```
except: pass

import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc

Print = PETSc.Sys.Print
```

Build the finite-difference 2-D Laplacian matrix.

```
def construct_operator(m, n):
    # Create matrix for 2D Laplacian operator
    A = PETSc.Mat().create()
    A.setSizes([m*n, m*n])
    A.setFromOptions()
    # Fill matrix
    hx = 1.0/(m-1) # x grid spacing
    hy = 1.0/(n-1) # y grid spacing
    diagv = 2.0*hy/hx + 2.0*hx/hy
    offdx = -1.0*hy/hx
    offdy = -1.0*hx/hy
    Istart, Iend = A.getOwnershipRange()
    for I in range(Istart, Iend) :
        A[I,I] = diagv
        i = I//n # map row number to
        j = I - i*n # grid coordinates
        if i> 0 : J = I-n; A[I,J] = offdx
        if i< m-1: J = I+n; A[I,J] = offdx
        if j> 0 : J = I-1; A[I,J] = offdy
        if j< n-1: J = I+1; A[I,J] = offdy
    A.assemble()
    return A
```

In the main function, first two command-line options are processed to set the grid dimensions. Then the matrix is built and passed to the solver object. In this case, the solver is configured to use the contour integral method. Next, the region of interest is defined, in this case an ellipse centered at the origin, with radius 0.2 and vertical scaling of 0.1. Finally, the solver is run. In this example, we illustrate how to print the solution using the solver method `errorView()`.

```
def main():
    opts = PETSc.Options()
    n = opts.getInt('n', 32)
    m = opts.getInt('m', 32)
    Print("2-D Laplacian Eigenproblem solved with contour integral, "
          "N=%d (%dx%d grid)\n" % (m*n, m, n))
    A = construct_operator(m,n)

    E = SLEPc.EPS().create()
    E.setOperators(A)
    E.setProblemType(SLEPc.EPS.ProblemType.HEP)
    E.setType(SLEPc.EPS.Type.CISS)
```

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```

R = E.getRG()
R.setType(SLEPc.RG.Type.ELLIPSE)
R.setEllipseParameters(0.0,0.2,0.1)
E.setFromOptions()

E.solve()

vw = PETSc.Viewer.STDOUT()
vw.pushFormat(PETSc.Viewer.Format.ASCII_INFO_DETAIL)
E.errorView(viewer=vw)
vw.popFormat()

if __name__ == '__main__':
    main()

```

### ex12.py: Illustrate the use of arbitrary selection

This example solves a simple tridiagonal eigenproblem. It illustrates how to set up the arbitrary selection of eigenvalues, where the decision of which is the preferred eigenvalue is made based not only on the value of the approximate eigenvalue but also on the approximate eigenvector.

In this example, the selection criterion is based on the projection of the approximate eigenvector onto a precomputed eigenvector. That is why we solve the problem twice.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```

import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc
import numpy

```

The matrix size  $n$  can be specified at the command line.

```

opts = PETSc.Options()
n = opts.getInt('n', 30)

```

Create the matrix `tridiag([-1 0 -1])`.

```

A = PETSc.Mat(); A.create()
A.setSizes([n, n])
A.setFromOptions()
rstart, rend = A.getOwnershipRange()
for i in range(rstart, rend):
    if i>0: A[i, i-1] = -1
    if i<n-1: A[i, i+1] = -1
A.assemble()

```

Configure the linear eigensolver initially to compute leftmost eigenvalues.

```

E = SLEPc.EPS(); E.create()
E.setOperators(A)
E.setProblemType(SLEPc.EPS.ProblemType.HEP)
E.setWhichEigenpairs(SLEPc.EPS.Which.SMALLEST_REAL)
E.setFromOptions()

```

Solve the eigenproblem and store the first computed eigenvector in `sx` to be used later. For the second solve, we configure the solver to select largest magnitude values with an arbitrary selection callback function `myArbitrarySel()`. It means that instead of sorting eigenvalues, the solver will sort the approximations according to the largest values of the result of `myArbitrarySel()` evaluated on the approximate eigenvectors. In this way, the same eigenvalue should be computed again.

```

E.solve()
nconv = E.getConverged()
Print = PETSc.Sys.Print
vw = PETSc.Viewer.STDOUT()
if nconv>0:
    sx, _ = A.createVecs()
    E.getEigenpair(0, sx)
    vw.pushFormat(PETSc.Viewer.Format.ASCII_INFO_DETAIL)
    E.errorView(viewer=vw)
    def myArbitrarySel(evalue, xr, xi, sx):
        return abs(xr.dot(sx))
    E.setArbitrarySelection(myArbitrarySel,sx)
    E.setWhichEigenpairs(SLEPc.EPS.Which.LARGEST_MAGNITUDE)
    E.solve()
    E.errorView(viewer=vw)
    vw.popFormat()
else:
    Print( "No eigenpairs converged" )

```

### ex13.py: Nonlinear eigenproblem with contour integral

This example solves a nonlinear eigenvalue problem with the *NEP* module configured to apply a contour integral method.

The problem arises from the PDE

$$u_{xx}(x) + n_c^2 \lambda^2 u(x) + g(\lambda) D_0 \lambda^2 u(x) = 0$$

where

$$\begin{aligned}
 g(\lambda) &= g_t / (\lambda - k_a + i g_t), \\
 k_a &= 8.0, \\
 g_t &= 0.5, \\
 D_0 &= 0.5, \\
 n_c &= 1.2,
 \end{aligned}$$

and the boundary conditions are

$$\begin{aligned}
 u(0) &= 0 \\
 u_x(1) &= i \lambda u(1).
 \end{aligned}$$

For the discretization,  $n$  grid points are used,  $x_1 = 0, \dots, x_n = 1$ , with step size  $h = 1/(n - 1)$ . Hence,

$$\begin{aligned} u_{xx}(x_i) &= \frac{1}{h^2}(u_{i-1} - 2u_i + u_{i+1}) \\ &= \frac{1}{h^2}[1, -2, 1][u_{i-1}, u_i, u_{i+1}]^T. \end{aligned}$$

The boundary condition at  $x = 0$  is  $u_1 = 0$ , and at  $x = 1$ :

$$\begin{aligned} u'(1) &= 1/2(u'(1 + h/2) + u'(1 - h/2)) \\ &= 1/2((u_{n+1} - u_n)/h + (u_n - u_{n-1})/h) \\ &= 1/(2h)(u_{n+1} - u_{n-1}) = i\lambda u_n, \end{aligned}$$

and therefore

$$u_{n+1} = 2ih\lambda u_n + u_{n-1}.$$

The Laplace term for  $u_n$  is

$$\begin{aligned} &= \frac{1}{h^2}(u_{n-1} - 2u_n + u_{n+1}) \\ &= \frac{1}{h^2}(u_{n-1} - 2u_n + 2ih\lambda u_n + u_{n-1}) \\ &= \frac{1}{h^2}(2u_{n-1} + (2ih\lambda - 2)u_n) \\ &= \frac{1}{h^2}[2, 2ih\lambda - 2][u_{n-1}, u_n]^T. \end{aligned}$$

The above discretization allows us to write the nonlinear PDE in the following split-operator form

$$\{A + \lambda^2 n_c^2 I_d + g(\lambda)\lambda^2 D_0 I_d + 2i\lambda/h D\}u = 0$$

so  $f_1 = 1$ ,  $f_2 = n_c^2 \lambda^2$ ,  $f_3 = g(\lambda)\lambda^2 D_0$ ,  $f_4 = 2i\lambda/h$ , with coefficient matrices

$$A = \begin{bmatrix} 1 & 0 & 0 & \dots & 0 \\ 0 & * & * & \dots & 0 \\ 0 & * & * & \dots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \dots & * \end{bmatrix}, I_d = \begin{bmatrix} 0 & 0 & 0 & \dots & 0 \\ 0 & 1 & 0 & \dots & 0 \\ 0 & 0 & 1 & \dots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \dots & 0 \end{bmatrix}, D = \begin{bmatrix} 0 & 0 & 0 & \dots & 0 \\ 0 & 0 & 0 & \dots & 0 \\ 0 & 0 & 0 & \dots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \dots & 1 \end{bmatrix}.$$

Contributed by: Thomas Hisch.

The full source code for this demo can be [downloaded here](#).

Initialization by importing slepc4py, petsc4py, numpy and scipy.

```
import sys

import slepc4py

slepc4py.init(sys.argv) # isort:skip

import numpy as np

try:
    import scipy
```

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```
import scipy.optimize
except ImportError:
    scipy = None

from petsc4py import PETSc
from slepc4py import SLEPc

Print = PETSc.Sys.Print
```

Check that the selected PETSc/SLEPc are built with complex scalars.

```
if not np.issubdtype(PETSc.ScalarType, np.complexfloating):
    Print("Demo should only be executed with complex PETSc scalars")
    exit(0)
```

Long function that defines the nonlinear eigenproblem and solves it.

```
def solve(n):
    L = 1.0
    h = L / (n - 1)
    nc = 1.2
    ka = 10.0
    gt = 4.0
    D0 = 0.5

    A = PETSc.Mat().create()
    A.setSizes([n, n])
    A.setFromOptions()
    A.setOption(PETSc.Mat.Option.HERMITIAN, False)

    rstart, rend = A.getOwnershipRange()
    d0, d1, d2 = (
        1 / h**2,
        -2 / h**2,
        1 / h**2,
    )
    Print(f"dterms={d0, d1, d2}")

    if rstart == 0:
        # dirichlet boundary condition at the left lead
        A[0, 0] = 1.0
        A[0, 1] = 0.0
        A[1, 0] = 0.0

        A[1, 1] = d1
        A[1, 2] = d2
        rstart += 2
    if rend == n:
        # at x=1.0 neumann boundary condition (not handled here but in a
        # different matrix (D))
        A[n - 1, n - 2] = 2.0 / h**2
        A[n - 1, n - 1] = (-2) / h**2 # + 2j*k*h / h**2 (neumann)
```

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```

    rend -= 1

    for i in range(rstart, rend):
        A[i, i - 1 : i + 2] = [d0, d1, d2]

    A.assemble()

    Id = PETSc.Mat().create()
    Id.setSizes([n, n])
    Id.setFromOptions()
    Id.setOption(PETSc.Mat.Option.HERMITIAN, True)
    rstart, rend = Id.getOwnershipRange()
    if rstart == 0:
        # due to dirichlet BC
        rstart += 1
    for i in range(rstart, rend):
        Id[i, i] = 1.0
    Id.assemble()

    D = PETSc.Mat().create()
    D.setSizes([n, n])
    D.setFromOptions()
    D.setOption(PETSc.Mat.Option.HERMITIAN, True)
    _, rend = D.getOwnershipRange()
    if rend == n:
        D[n - 1, n - 1] = 1
    D.assemble()

    Print(f'DOF: {A.getInfo()['nz_used']}, MEM: {A.getInfo()['memory']}')

    f1 = SLEPc.FN().create()
    f1.setType(SLEPc.FN.Type.RATIONAL)
    f1.setRationalNumerator([1.0])
    f2 = SLEPc.FN().create()
    f2.setType(SLEPc.FN.Type.RATIONAL)
    f2.setRationalNumerator([nc**2, 0.0, 0.0])
    f3 = SLEPc.FN().create()
    f3.setType(SLEPc.FN.Type.RATIONAL)
    f3.setRationalNumerator([D0 * gt, 0.0, 0.0])
    f3.setRationalDenominator([1.0, -ka + 1j * gt])
    f4 = SLEPc.FN().create()
    f4.setType(SLEPc.FN.Type.RATIONAL)
    f4.setRationalNumerator([2j / h, 0])

    # Setup the solver
    nep = SLEPc.NEP().create()

    nep.setSplitOperator(
        [A, Id, Id, D],
        [f1, f2, f3, f4],
        PETSc.Mat.Structure.SUBSET,
    )

```

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```

# Customize options
nep.setTolerances(tol=1e-7)

nep.setDimensions(nev=24)
nep.setType(SLEPc.NEP.Type.CISS)

# the rg params are chosen s.t. the singularity at  $k = ka - 1j*gt$  is
# outside of the contour.
radius = 3 * gt
vscale = 0.5 * gt / radius
rg_params = (ka, 3 * gt, vscale)
R = nep.getRG()
R.setType(SLEPc.RG.Type.ELLIPSE)
Print(f"RG params: {rg_params}")
R.setEllipseParameters(*rg_params)

nep.setFromOptions()

# Solve the problem
nep.solve()

its = nep.getIterationNumber()
Print("Number of iterations of the method: %i" % its)
sol_type = nep.getType()
Print("Solution method: %s" % sol_type)
nev, ncv, mpd = nep.getDimensions()
Print("")
Print("Subspace dimension: %i" % ncv)
tol, maxit = nep.getTolerances()
Print("Stopping condition: tol=%.4g" % tol)
Print("")

nconv = nep.getConverged()
Print("Number of converged eigenpairs %d" % nconv)

x = A.createVecs("right")

evals = []
modes = []
if nconv > 0:
    Print()
    Print("          lam          ||T(lam)x||    |lam-lam_exact|/|lam_exact| ")
    Print("-----")
    for i in range(nconv):
        lam = nep.getEigenpair(i, x)
        error = nep.computeError(i)

        def eigenvalue_error_term(k):
            gkmu = gt / (k - ka + 1j * gt)
            nceff = np.sqrt(nc**2 + gkmu * D0)
            return -1j / np.tan(nceff * k * L) - 1 / nceff

```

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```

    # compute the expected_eigenvalue

    # we assume that the numerically calculated eigenvalue is close to
    # the exact one, which we can determine using a Newton-Raphson
    # method.
    if scipy:
        expected_lam = scipy.optimize.newton(
            eigenvalue_error_term, np.complex128(lam), rtol=1e-11
        )
        rel_err = abs(lam - expected_lam) / abs(expected_lam)
        rel_err = "%6g" % rel_err
    else:
        rel_err = "scipy not installed"

    Print(" %9f%+9f j %12g  %s" % (lam.real, lam.imag, error, rel_err))

    evals.append(lam)
    modes.append(x.getArray().copy())
    Print()

    return np.asarray(evals), rg_params, ka, gt

```

The main function reads the problem size  $n$  from the command line, solves the problem with the above function, and then plots the computed eigenvalues.

```

def main():
    opts = PETSc.Options()
    n = opts.getInt("n", 256)
    Print(f"n={n}")

    evals, rg_params, ka, gt = solve(n)

    if not opts.getBool("ploteigs", True) or PETSc.COMM_WORLD.getRank():
        return

    try:
        import matplotlib.pyplot as plt
        from matplotlib.patches import Ellipse
    except ImportError:
        print("plot is not shown, because matplotlib is not installed")
    else:
        fig, ax = plt.subplots()
        ax.plot(evals.real, evals.imag, "x")

        height = 2 * rg_params[1] * rg_params[2]
        ellipse = Ellipse(
            xy=(rg_params[0], 0.0),
            width=rg_params[1] * 2,
            height=height,
            edgecolor="r",
            fc="None",

```

```

        lw=2,
    )
    ax.add_patch(ellipse)

    ax.grid()
    ax.legend()
    plt.show()

if __name__ == "__main__":
    main()

```

### ex14.py: Lyapunov equation with the shifted 2-D Laplacian

This example illustrates the use of slepc4py linear matrix equations solver object. In particular, a Lyapunov equation is solved, where the coefficient matrix is the shifted 2-D Laplacian.

The full source code for this demo can be [downloaded here](#).

Initialization is similar to previous examples.

```

import sys, slepc4py
slepc4py.init(sys.argv)

from petsc4py import PETSc
from slepc4py import SLEPc

Print = PETSc.Sys.Print

```

Once again, a function to build the discretized Laplacian operator in two dimensions. In this case, we build the shifted negative Laplacian because the Lyapunov equation requires the coefficient matrix being stable.

```

def Laplacian2D(m, n, sigma):
    # Create matrix for 2D Laplacian operator
    A = PETSc.Mat().create()
    A.setSizes([m*n, m*n])
    A.setFromOptions()
    # Fill matrix
    Istart, Iend = A.getOwnershipRange()
    for I in range(Istart, Iend):
        A[I,I] = -4.0-sigma
        i = I//n    # map row number to
        j = I - i*n # grid coordinates
        if i> 0 : J = I-n; A[I,J] = 1.0
        if i< m-1: J = I+n; A[I,J] = 1.0
        if j> 0 : J = I-1; A[I,J] = 1.0
        if j< n-1: J = I+1; A[I,J] = 1.0
    A.assemble()
    return A

```

The following function solves the continuous-time Lyapunov equation

$$AX + XA^* = -C$$

where  $C$  is supposed to have low rank. The solution  $X$  also has low rank, which will be restricted to `rk` if it was provided as an argument of the function. After solving the equation, this function computes and prints a residual norm.

```
def solve_lyap(A, C, rk=0):
    # Setup the solver
    L = SLEPc.LME().create()
    L.setProblemType(SLEPc.LME.ProblemType.LYAPUNOV)
    L.setCoefficients(A)
    L.setRHS(C)

    N = C.size[0]
    if rk>0:
        X1 = PETSc.Mat().createDense((N,rk))
        X1.assemble()
        X = PETSc.Mat().createLRC(None, X1, None, None)
        L.setSolution(X)

    L.setTolerances(1e-7)
    L.setFromOptions()

    # Solve the problem
    L.solve()

    if rk==0:
        X = L.getSolution()

    its = L.getIterationNumber()
    Print(f'Number of iterations of the method: {its}')
    sol_type = L.getType()
    Print(f'Solution method: {sol_type}')
    tol, maxit = L.getTolerances()
    Print(f'Stopping condition: tol={tol}, maxit={maxit}')
    Print(f'Error estimate reported by the solver: {L.getErrorEstimate()}')
    if N<500:
        Print(f'Computed residual norm: {L.computeError()}')
    Print('')

    return X
```

The main function processes several command-line arguments such as the grid dimension (`n`, `m`), the shift `sigma` and the rank of the solution, `rank`.

The coefficient matrix  $A$  is built with the function above, while the right-hand side matrix  $C$  must be built as a low-rank matrix using `Mat.createLRC()`.

```
if __name__ == '__main__':
    opts = PETSc.Options()
    n = opts.getInt('n', 15)
    m = opts.getInt('m', n)
    N = m*n
    sigma = opts.getScalar('sigma', 0.0)
    rk = opts.getInt('rank', 0)

    # Coefficient matrix A
```

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```

A = Laplacian2D(m, n, sigma)

# Create a low-rank Mat to store the right-hand side C = C1*C1'
C1 = PETSc.Mat().createDense((N,2))
rstart, rend = C1.getOwnershipRange()
v = C1.getDenseArray()
for i in range(rstart,rend):
    if i<N//2: v[i-rstart,0] = 1.0
    if i==0: v[i-rstart,1] = -2.0
    if i==1 or i==2: v[i-rstart,1] = -1.0
C1.assemble()
C = PETSc.Mat().createLRC(None, C1, None, None)

X = solve_lyap(A, C, rk)

```

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