### User Documentation for KINSOL v2.3.0

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# Chapter 1

## Introduction

KINSOL is part of a software family called SUNDIALS: SUite of Nonlinear and DIfferential/ALgebraic equation Solvers. This suite consists of CVODE, KINSOL, and IDA, and variants of these. KINSOL is a general-purpose nonlinear system solver based on Newton-Krylov solver technology.

### 1.1 Historical Background

The first nonlinear solver packages based on Newton-Krylov methods were written in FORTRAN. In particular, the NKSOL package, written at LLNL, was the first Newton-Krylov solver package written for solution of systems arising in solution of partial differential equations [3]. This FORTRAN code made use of Newton's method to solve the discrete nonlinear systems and applied a preconditioned Krylov linear solver for solution of the Jacobian system at each nonlinear iteration. The key to the Newton-Krylov method was that the matrix-vector multiplies required by the Krylov method could effectively be approximated by a finite difference of the nonlinear system-defining function, preventing a requirement for the formation of the actual Jacobian matrix. Significantly less memory was required for the solver as a result.

In the late 1990's, there was a push at LLNL to rewrite the nonlinear solver into C and port it to distributed memory parallel machines. Both Newton and Krylov methods are easily implemented in parallel, and this effort gave rise to the KINSOL package. KINSOL is similar to NKSOL in functionality, except that it provides for more options in the choice of linear system tolerances and has a more modular design to provide flexibility for future enhancements.

There are several motivations for choosing the C language for KINSOL. First, a general movement away from FORTRAN and toward C in scientific computing is apparent. Second, the pointer, structure, and dynamic memory allocation features in C are extremely useful in software of this complexity, with the great variety of method options offered. Finally, we prefer C over C++ for KINSOL because of the wider availability of C compilers, the potentially greater efficiency of C, and the greater ease of interfacing the solver to applications written in FORTRAN.

In the process of translating NKSOL into C, the overall KINSOL organization has been changed considerably. One key feature of the KINSOL organization is that a separate module devoted to vector operations has been created. This module facilitated extension to multiprosessor environments with minimal impact on the rest of the solver. The new vector module design is shared across the SUNDIALS suite. This NVECTOR module is written in terms of abstract vector operations with the actual routines attached by a particular implementation (such as serial or parallel) of NVECTOR. This allows writing the SUNDIALS solvers in a manner independent of the actual NVECTOR implementation (which can be user-supplied), as well as allowing more than one NVECTOR module linked into an executable file.

2 Introduction

### 1.2 Changes from previous versions

### Changes in v2.3.0

The user interface has been further refined. Several functions used for setting optional inputs were combined into a single one. Additionally, to resolve potential variable scope issues, all SUNDIALS solvers release user data right after its use. The build systems has been further improved to make it more robust.

### Changes in v2.2.1

The changes in this minor SUNDIALS release affect only the build system.

### Changes in v2.2.0

The major changes from the previous version involve a redesign of the user interface across the entire SUNDIALS suite. We have eliminated the mechanism of providing optional inputs and extracting optional statistics from the solver through the iopt and ropt arrays. Instead, KINSOL now provides a set of routines (with prefix KINSet) to change the default values for various quantities controlling the solver and a set of extraction routines (with prefix KINGet) to extract statistics after return from the main solver routine. Similarly, each linear solver module provides its own set of set- and get-type routines. For more details see Chapter 5.

Additionally, the interfaces to several user-supplied routines (such as those providing Jacobian-vector products and preconditioner information) were simplified by reducing the number of arguments. The same information that was previously accessible through such arguments can now be obtained through set-type functions.

### 1.3 Reading this User Guide

The structure of this document is as follows:

- The next chapter discusses how to install the KINSOLpackage.
- In Chapter 3, we provide short descriptions of the numerical methods implemented by KINSOL for the solution of nonlinear systems.
- The following chapter describes the structure of the SUNDIALS suite of solvers (§4.1) and the software organization of the KINSOL solver (§4.2).
- In Chapter 5, we give an overview of the usage of KINSOL, as well as a complete description of the user interface and of the user-defined routines for solution of nonlinear systems.
- Chapter 6 gives a brief overview of the generic NVECTOR module shared among the various components of SUNDIALS, as well as details of the two NVECTOR implementations provided with SUNDIALS: a serial implementation (§6.1) and a parallel implementation, based on MPI (§6.2).
- Chapter 7 describes the interfaces to the linear solver modules, so that a user can provide his/her own such module.
- Chapter 8 describes the generic linear solvers shared by all SUNDIALS solvers.
- Finally, Chapter 9 lists the constants used for input to and output from KINSOL.

Finally, the reader should be aware of the following notational conventions in this user guide: program listings and identifiers (such as KINMalloc) within textual explanations appear in typewriter type style; fields in C structures (such as *content*) appear in italics; and packages or modules are written in all capitals.

**Acknowledgments.** We wish to acknowledge the contributions to previous versions of the KINSOL code and user guide of Allan G. Taylor.

# Chapter 2

## KINSOL Installation Procedure

The installation of KINSOL is accomplished by installing the SUNDIALS suite as a whole, according to the instructions that follow. The same procedure applies whether or not the downloaded file contains solvers other than KINSOL.

Generally speaking, the installation procedure outlined in §2.1 below will work on commodity LINUX/UNIX systems without modification. Users are still encouraged, however, to carefully read the entire chapter before attempting to install the SUNDIALS suite, in case non-default choices are desired for compilers, compilation options, or the like. In lieu of reading the option list below, the user may invoke the configuration script with the help flag to view a complete listing of available options, which may be done by issuing

### % ./configure --help

from within the sundials directory.

In the descriptions below, build\_tree refers to the directory under which the user wants to build and/or install the SUNDIALS package. By default, the SUNDIALS libraries and header files are installed under the subdirectories build\_tree/lib and build\_tree/include, respectively. Also, source\_tree refers to the directory where the SUNDIALS source code is located. The chosen build\_tree may be different from the source\_tree, thus allowing for multiple installations of the SUNDIALS suite with different configuration options.

Concerning the installation procedure outlined below, after invoking the tar command with the appropriate options, the contents of the SUNDIALS archive (or the source\_tree) will be extracted to a directory named sundials. Since the name of the extracted directory is not version-specific it is recommended that the user refrain from extracting the archive to a directory containing a previous version/release of the SUNDIALS suite. If the user is only upgrading and the previous installation of SUNDIALS is not needed, then the user may remove the previous installation by issuing

#### % rm -rf sundials

from a shell command prompt.

Even though the installation procedure given below presupposes that the user will use the default vector modules supplied with the distribution, using the SUNDIALS suite with a user-supplied vector module normally will not require any changes to the build procedure.

### 2.1 Installation steps

To install the SUNDIALS suite, given a downloaded file named <code>sundials\_file.tar.gz</code>, issue the following commands from a shell command prompt, while within the directory where <code>source\_tree</code> is to be located. The names of installed libraries and header files are listed in Table 2.1 for reference. (For brevity, the corresponding <code>.c</code> files are not listed.) Regarding the file extension <code>.lib</code> appearing in Table 2.1, shared libraries generally have an extension of <code>.so</code> and static libraries have an extension of <code>.a.</code> (See <code>Options for library support</code> for additional details.)

```
1. gunzip sundials\_file.tar.gz
```

2. tar -xf sundials\_file.tar [creates sundials directory]

3. cd build\_tree

4. path\_to\_source\_tree/configure options [options can be absent]

5. make

6. make install

7. make examples

8. If system storage space conservation is a priority, then issue

% make clean

and/or

% make examples\_clean

from a shell command prompt to remove unneeded object files.

### 2.2 Configuration options

The installation procedure given above will generally work without modification; however, if the system includes multiple MPI implementations, then certain configure script-related options may be used to indicate which MPI implementation should be used. Also, if the user wants to use non-default language compilers, then, again, the necessary shell environment variables must be appropriately redefined. The remainder of this section provides explanations of available configure script options.

### General options

### --prefix=PREFIX

Location for architecture-independent files.

Default: PREFIX=build\_tree

### --includedir=DIR

Alternate location for installation of header files.

Default: DIR=PREFIX/include

### --libdir=DIR

Alternate location for installation of libraries.

Default: DIR=PREFIX/lib

### --disable-examples

All available example programs are automatically built unless this option is given. The example executables are stored under the following subdirectories of the associated solver:

build\_tree/solver/examples\_ser : serial C examples

build\_tree/solver/examples\_par: parallel C examples (MPI-enabled)

build\_tree/solver/fcmix/examples\_ser : serial FORTRAN examples

build\_tree/solver/fcmix/examples\_par: parallel Fortran examples (MPI-enabled)

*Note*: Some of these subdirectories may not exist depending upon the solver and/or the configuration options given.

Table 2.1: SUNDIALS libraries and header files

Module	Libraries	Header files
SHARED	libsundials_shared.lib	sundialstypes.h
		sundialsmath.h
		sundials_config.h
		dense.h
		smalldense.h
		band.h
		spgmr.h
		iterative.h
		nvector.h
NVECTOR_SERIAL	${\tt libsundials\_nvecserial.}\ lib$	nvector_serial.h
	libsundials_fnvecserial.a	
NVECTOR_PARALLEL	${ t libsundials\_nvecparallel.}\ lib$	nvector_parallel.h
	libsundials_fnvecparallel.a	
CVODE	${ t libsundials\_cvode.}\ lib$	cvode.h
	libsundials_fcvode.a	cvdense.h
		cvband.h
		cvdiag.h
		cvspgmr.h
		cvbandpre.h
		cvbbdpre.h
CVODES	${ t libsundials\_cvodes.}\ lib$	cvodes.h
		cvodea.h
		cvdense.h
		cvband.h
		cvdiag.h
		cvspgmr.h
		cvbandpre.h
		cvbbdpre.h
IDA	libsundials_ida.lib	ida.h
		idadense.h
		idaband.h
		idaspgmr.h
		idabbdpre.h
KINSOL	libsundials_kinsol. $\it lib$	kinsol.h
	libsundials_fkinsol.a	kinspgmr.h
		kinbbdpre.h

### --disable-solver

Although each existing solver module is built by default, support for a given solver can be explicitly disabled using this option. The valid values for *solver* are: cvode, cvodes, ida, and kinsol.

### --with-cppflags=ARG

Specify additional C preprocessor flags (e.g., ARG=-I<include\_dir> if necessary header files are located in nonstandard locations).

### --with-cflags=ARG

Specify additional C compilation flags.

### --with-ldflags=ARG

Specify additional linker flags (e.g., ARG=-L<lib\_dir> if required libraries are located in non-standard locations).

### --with-libs=ARG

Specify additional libraries to be used (e.g., ARG=-1<foo> to link with the library named libfoo.so).

### --with-precision=ARG

By default, SUNDIALS will define a real number (internally referred to as realtype) to be a double-precision floating-point numeric data type (double C-type); however, this option may be used to build SUNDIALS with realtype alternatively defined as a single-precision floating-point numeric data type (float C-type) if ARG=single, or as a long double C-type if ARG=extended.

Default: ARG=double

### Options for Fortran support

#### --disable-f77

Using this option will disable all FORTRAN support. The FCVODE, FKINSOL and FNVECTOR modules will not be built regardless of availability.

### --with-fflags=ARG

Specify additional FORTRAN compilation flags.

The configuration script will attempt to automatically determine the function name mangling scheme required by the specified FORTRAN compiler, but the following two options may be used to override the default behavior.

### --with-f77underscore=ARG

This option pertains to the FKINSOL, FCVODE and FNVECTOR FORTRAN-C interface modules and is used to specify the number of underscores to append to function names so FORTRAN routines can properly link with the associated SUNDIALS libraries. Valid values for ARG are: none, one and two.

Default: ARG=one

### --with-f77case=ARG

Use this option to specify whether the external names of the FKINSOL, FCVODE and FNVECTOR FORTRAN-C interface functions should be lowercase or uppercase so FORTRAN routines can properly link with the associated SUNDIALS libraries. Valid values for ARG are: lower and upper.

Default: ARG=lower

### Options for MPI support

The following configuration options are only applicable to the parallel SUNDIALS packages:

### --disable-mpi

Using this option will completely disable MPI support.

--with-mpicc=ARG

### --with-mpif77=ARG

By default, the configuration utility script will use the MPI compiler scripts named mpic and mpif77 to compile the parallelized SUNDIALS subroutines; however, for reasons of compatibility, different executable names may be specified via the above options. Also, ARG=no can be used to disable the use of MPI compiler scripts, thus causing the serial C and FORTRAN compilers to be used to compile the parallelized SUNDIALS functions and examples.

### --with-mpi-root=MPIDIR

This option may be used to specify which MPI implementation should be used. The SUNDIALS configuration script will automatically check under the subdirectories MPIDIR/include and MPIDIR/lib for the necessary header files and libraries. The subdirectory MPIDIR/bin will also be searched for the C and FORTRAN MPI compiler scripts, unless the user uses --with-mpicc=no or --with-mpif77=no.

```
--with-mpi-incdir=INCDIR
```

--with-mpi-libdir=LIBDIR

### --with-mpi-libs=LIBS

These options may be used if the user would prefer not to use a preexisting MPI compiler script, but instead would rather use a serial complier and provide the flags necessary to compile the MPI-aware subroutines in SUNDIALS.

Often an MPI implementation will have unique library names and so it may be necessary to specify the appropriate libraries to use (e.g., LIBS=-lmpich).

Default: INCDIR=MPIDIR/include, LIBDIR=MPIDIR/lib and LIBS=-lmpi

### --with-mpi-flags=ARG

Specify additional MPI-specific flags.

### Options for library support

By default, only static libraries are built, but the following option may be used to build shared libraries on supported platforms.

### --enable-shared

Using this particular option will result in both static and shared versions of the available SUN-DIALS libraries being built if the system supports shared libraries. To build only shared libraries also specify --disable-static.

Note: The FCVODE and FKINSOL libraries can only be built as static libraries because they contain references to externally defined symbols, namely user-supplied FORTRAN subroutines. Although the FORTRAN interfaces to the serial and parallel implementations of the supplied NVECTOR module do not contain any unresolvable external symbols, the libraries are still built as static libraries for the purpose of consistency.

### Options for cross-compilation

If the SUNDIALS suite will be cross-compiled (meaning the build procedure will not be completed on the actual destination system, but rather on an alternate system with a different architecture) then the following two options should be used:

### --build=BUILD

This particular option is used to specify the canonical system/platform name for the build system.

#### --host=HOST

If cross-compiling, then the user must use this option to specify the canonical system/platform name for the destination system.

### Environment variables

The following environment variables can be locally (re)defined for use during the configuration of SUNDIALS. See the next section for illustrations of these.

CC

F77

Since the configuration script uses the first C and FORTRAN compilers found in the current executable search path, then each relevant shell variable (CC and F77) must be locally (re)defined in order to use a different compiler. For example, to use xcc (executable name of chosen compiler) as the C language compiler, use CC=xcc in the configure step.

**CFLAGS** 

**FFLAGS** 

Use these environment variables to override the default C and FORTRAN compilation flags.

### 2.3 Configuration examples

The following examples are meant to help demonstrate proper usage of the configure options:

The above example builds SUNDIALS using gcc as the serial C compiler, g77 as the serial FORTRAN compiler, mpicc as the parallel C compiler, mpif77 as the parallel FORTRAN compiler, and appends the -g3 compilaton flag to the list of default flags.

This example again builds SUNDIALS using gcc as the serial C compiler, but the --with-mpicc=no option explicitly disables the use of the corresponding MPI compiler script. In addition, since the --with-mpi-root option is given, the compilation flags -I/usr/apps/mpich/1.2.4/include and -L/usr/apps/mpich/1.2.4/lib are passed to gcc when compiling the MPI-enabled functions. The --disable-examples option disables the examples (which means a FORTRAN compiler is not required). The --with-mpi-libs option is still needed so that the configure script can check if gcc can link with the appropriate MPI library as -lmpi is the internal default.

# Chapter 3

# Mathematical Considerations

KINSOL solves nonlinear algebraic systems in real space, which we write as

$$F(u) = 0, \quad F: \mathbf{R}^N \to \mathbf{R}^N,$$
 (3.1)

given an initial guess  $u_0$ .

KINSOL employs the Inexact Newton method developed in [1, 3, 5] and further described in [6, 8], resulting in the following iteration:

### **Inexact Newton iteration**

- 1. Set  $u_0 = \text{an initial guess}$
- 2. For  $n = 0, 1, 2, \dots$  until convergence do:
  - (a) Approximately solve  $J(u_n)\delta_n = -F(u_n)$
  - (b) Set  $u_{n+1} = u_n + \lambda \delta_n$ ,  $0 < \lambda < 1$
  - (c) Test for convergence

Here,  $u_n$  is the *n*th iterate to u, and J(u) = F'(u) is the system Jacobian. As this code module is anticipated for use on large systems, only iterative methods are provided to solve the system in step 2(a). These solutions are only approximate. At each stage in the iteration process, a scalar multiple of the approximate solution,  $\delta_n$ , is added to  $u_n$  to produce a new iterate,  $u_{n+1}$ . A test for convergence is made before the iteration continues.

The linear iterative method currently implemented is one of the class of Krylov methods, GMRES [2, 9], provided through the SPGMR module common to all SUNDIALS codes. Use of SPGMR provides a linear solver which, by default, is applied in a matrix-free manner, with matrix-vector products Jv obtained by either finite difference quotients or a user-supplied routine. In the case where finite differences are used, the matrix-vector product J(u)v is approximated by a quotient of the form given by

$$J(u)v \approx [F(u+\sigma v) - F(u)]/\sigma \tag{3.2}$$

where u is the current approximation to a root of (3.1), and  $\sigma$  is a scalar. The choice of  $\sigma$  is taken from [3] and is given by

$$\sigma = \frac{\max\{|u^T v|, \operatorname{typ} u^T | v|\}}{\|v\|_2} \operatorname{sign}(u^T v) \sqrt{U}, \qquad (3.3)$$

where  $\operatorname{typ} u$  is a vector of typical values for the absolute values of the solution (and can be taken to be inverses of the scale factors given for u as described below), and U is unit roundoff. Convergence of the Newton method is maintained as long as the value of  $\sigma$  remains appropriately small as shown in [1].

To the above methods are added scaling and preconditioning. Scaling is allowed for both the solution vector and the system function vector. For scaling to be used, the user should supply values

 $D_u$ , which are diagonal elements of the scaling matrix such that  $D_u u_n$  has all components roughly the same magnitude when  $u_n$  is close to a solution, and  $D_F$ , which are diagonal scaling matrix elements such that  $D_F F$  has all components roughly the same magnitude when  $u_n$  is not too close to a solution. In the text below, we use the following scaled norms:

$$||z||_{D_u} = ||D_u z||_2, \quad ||z||_{D_F} = ||D_F z||_2, \quad ||z||_{D_u,\infty} = ||D_u z||_\infty, \quad \text{and} \quad ||z||_{D_F,\infty} = ||D_F z||_\infty$$
 (3.4)

where  $\|\cdot\|_{\infty}$  is the max norm. When scaling values are provided for the solution vector, these values are automatically incorporated into the calculation of  $\sigma$  in (3.3). Additionally, right preconditioning is provided if the preconditioning setup and solve routines are supplied by the user. In this case, GMRES is applied to the linear systems  $(JP^{-1})(P\delta) = -F$ , where P denotes the right preconditioning matrix.

Two methods of applying a computed step  $\delta_n$  to the previously computed solution vector are implemented. The first and simplest is the Inexact Newton strategy which applies step 2(b) as above with  $\lambda$  always set to 1. The other method is a global strategy, which attempts to use the direction implied by  $\delta_n$  in the most efficient way for furthering convergence of the nonlinear problem. This technique is implemented in the second strategy, called Linesearch. This option employs both the  $\alpha$  and  $\beta$  conditions of the Goldstein-Armijo linesearch given in [6] for step 2(b), where  $\lambda$  is chosen to guarantee a sufficient decrease in F relative to the step length as well as a minimum step length relative to the initial rate of decrease of F. One property of the algorithm is that the full Newton step tends to be taken close to the solution. For more details, the reader is referred to [6].

Stopping criteria for the Newton method are applied to both of the nonlinear residual and the step length. For the former, the Newton iteration must pass a stopping test

$$||F(u_n)||_{D_F,\infty} < \text{FTOL},$$

where FTOL is an input scalar tolerance with a default value of  $U^{1/3}$ . For the latter, the Newton method will terminate when the maximum scaled step is below a given tolerance

$$\|\lambda \delta_n\|_{D_n,\infty} < \text{STEPTOL},$$

where STEPTOL is an input scalar tolerance with a default value of  $U^{2/3}$ . Only the first condition (small residual) is considered a successful completion of KINSOL. The second condition (small step) may indicate that the iteration is stalled near a point for which the residual is still unacceptable.

Three options for stopping criteria for the linear system solve are implemented, including the two algorithms of Eisenstat and Walker [7]. The Krylov iteration must pass a stopping test

$$||J\delta_n + F||_{D_F} < (\eta_n + U)||F||_{D_F}$$

where  $\eta_n$  is one of:

• Eisenstat and Walker Choice 1

$$\eta_n = \frac{| \|F(u_n)\|_{D_F} - \|F(u_{n-1}) + J(u_{n-1})\delta_n\|_{D_F} |}{\|F(u_{n-1})\|_{D_F}},$$

• Eisenstat and Walker Choice 2

$$\eta_n = \gamma \left( \frac{\|F(u_n)\|_{D_F}}{\|F(u_{n-1})\|_{D_F}} \right)^{\alpha},$$

where default values of  $\gamma$  and  $\alpha$  are 0.9 and 2, respectively.

•  $\eta_n = \text{constant with } 0.1 \text{ as the default.}$ 

The default is Eisenstat and Walker Choice 1. For both options 1 and 2, appropriate safeguards are incorporated to ensure that  $\eta$  does not decrease too quickly [7].

As a user option, KINSOL permits the application of inequality constraints,  $u^i > 0$  and  $u^i < 0$ , as well as  $u^i \geq 0$  and  $u^i \leq 0$ , where  $u^i$  is the *i*th component of u. Any such constraint, or no constraint, may be imposed on each component. KINSOL will reduce step lengths in order to ensure that no constraint is violated. Specifically, if a new Newton iterate will violate a constraint, the maximum (over all i) step length along the Newton direction that will satisfy all constraints is found and  $\delta_n$  in Step 2(b) is scaled to take a step of that length.

# Chapter 4

# Code Organization

### 4.1 SUNDIALS organization

The family of solvers referred to as SUNDIALS consists of the solvers CVODE (for ODE systems), KINSOL (for nonlinear algebraic systems), and IDA (for differential-algebraic systems). In addition, variants of these which also do sensitivity analysis calculations are available or in development. CVODES, an extension of CVODE that provides both forward and adjoint sensitivity capabilities is available, while IDAS is currently in development.

The various solvers of this family share many subordinate modules. For this reason, it is organized as a family, with a directory structure that exploits that sharing (see Fig. 4.1). The following is a list of the solver packages presently available:

- CVODE, a solver for stiff and nonstiff ODEs dy/dt = f(t, y);
- CVODES, a solver for stiff and nonstiff ODEs dy/dt = f(t, y, p) with sensitivity analysis capabilities;
- KINSOL, a solver for nonlinear algebraic systems F(u) = 0;
- IDA, a solver for differential-algebraic systems F(t, y, y') = 0.

### 4.2 KINSOL organization

The KINSOL package is written in the ANSI C language. This section summarizes the basic structure of the package, although knowledge of this structure is not necessary for its use.

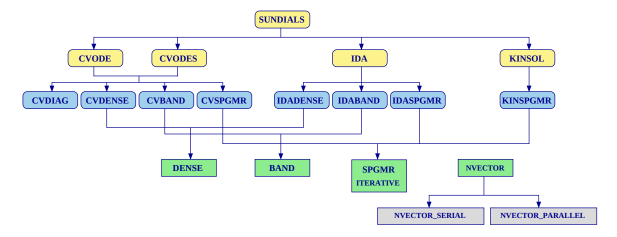
The overall organization of the KINSOL package is shown in Figure 4.2. The central solver module, implemented in the files kinsol.h and kinsol.c, deals with the solution of a nonlinear algebraic system using either an Inexact Newton method or a line search method for the global strategy. Although this module contains logic for the Newton iteration, it has no knowledge of the method used to solve the linear systems that arise. For any given user problem, the user must specify which linear solver module to use.

At present, the package includes the following KINSOL linear system module:

• KINSPGMR: scaled preconditioned GMRES method.

This set of linear solver modules is intended to be expanded in the future as new algorithms are developed.

The KINSPGMR package includes an algorithm for the approximation by difference quotients of the product between the Jacobian matrix and a vector of appropriate length. The user has the option of providing a routine for this operation. With KINSPGMR, the preconditioning must be supplied by the user, in two phases: setup (preprocessing of preconditioner data) and solve.



(a) High-level diagram

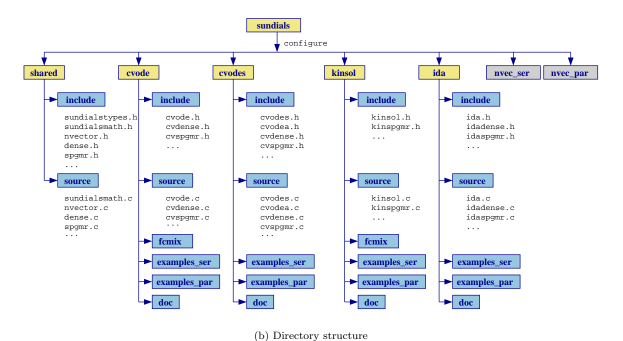


Figure 4.1: Organization of the SUNDIALS suite

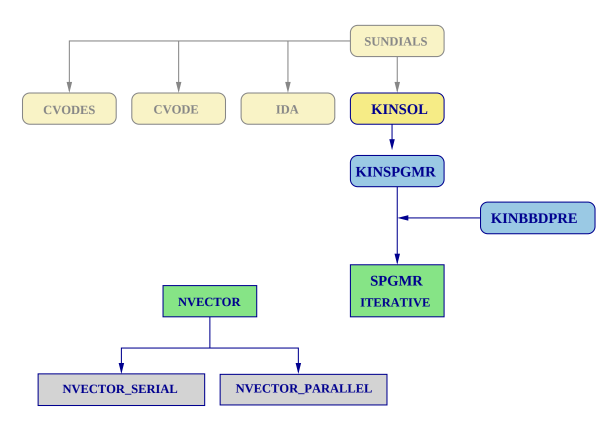


Figure 4.2: Overall structure diagram of the KINSOL package. Modules specific to KINSOL are distinguished by rounded boxes, while generic solver and auxiliary modules are in rectangular boxes. Grayed boxes refer to the encompassing SUNDIALS structure.

A KINSOL linear solver module consists of four routines, devoted to (1) memory allocation and initialization, (2) setup of the matrix data involved, (3) solution of the system, and (4) freeing of memory. The setup and solution phases are separate because the evaluation of Jacobians and preconditioners is done only periodically during the integration, as required to achieve convergence. The call list within the central KINSOL module to each of the associated functions is fixed, thus allowing the central module to be completely independent of the linear system method.

Linear solver modules are also decomposed in another way. The module KINSPGMR is a set of interface routines built on top of a generic solver module SPGMR. The interface deals with the use of these methods in the KINSOL context, whereas the generic solver is independent of the context. While the generic solvers here were generated with SUNDIALS in mind, our intention is that they be usable in other applications as general-purpose solvers. This separation also allows for any generic solver to be replaced by an improved version, with no necessity to revise the KINSOL package elsewhere.

KINSOL also provides a preconditioner module called KINBBDPRE which works in conjunction with NVECTOR\_PARALLEL to generate a preconditioner that is a block-diagonal matrix with each block being a band matrix, as further described in §5.6.

All state information used by KINSOL to solve a given problem is saved in a structure, and a pointer to that structure is returned to the user. There is no global data in the KINSOL package, and so in this respect it is reentrant. State information specific to the linear solver is saved in a separate structure, a pointer to which resides in the KINSOL memory structure. The reentrancy of KINSOL was motivated by the anticipated multicomputer extension, but is also essential in a uniprocessor setting where two or more different problems are solved by intermixed calls to the package from one user program.

# Chapter 5

# Using KINSOL

This chapter is concerned with the use of KINSOL for the solution of nonlinear systems. The following subsections treat the header files, the layout of the user's main program, description of the KINSOL user-callable routines, and user-supplied functions. The listings of the sample programs in the companion document [4] may also be helpful. Those codes may be used as templates (with the removal of some lines involved in testing), and are included in the KINSOL package.

KINSOL uses various constants for both input and output. These are defined as needed in this chapter, but for convenience are also listed separately in Chapter 9.

### 5.1 Data types

The sundialstypes.h file contains the definition of the type realtype, which is used by the SUNDIALS solvers for all floating-point data. The type realtype can be float, double, or long double, with the default being double. The user can change the precision of the SUNDIALS solvers arithmetic at the configuration stage (see §2.2).

Additionally, based on the current precision, sundialstypes.h defines BIG\_REAL to be the largest value representable as a realtype, SMALL\_REAL to be the smallest value representable as a realtype, and UNIT\_ROUNDOFF to be the difference between 1.0 and the minimum realtype greater than 1.0.

Within SUNDIALS, real constants are set by way of a macro called RCONST. It is this macro that needs the ability to branch on the definition realtype. In ANSI C, a floating-point constant with no suffix is stored as a double. Placing the suffix "F" at the end of a floating point constant makes it a float, whereas using the suffix "L" makes it a long double. For example,

```
#define A 1.0
#define B 1.0F
#define C 1.0L
```

defines A to be a double constant equal to 1.0, B to be a float constant equal to 1.0, and C to be a long double constant equal to 1.0. The macro call RCONST(1.0) automatically expands to 1.0 if realtype is double, to 1.0F if realtype is float, or to 1.0L if realtype is long double. SUNDIALS uses the RCONST macro internally to declare all of its floating-point constants.

A user program which uses the type realtype and the RCONST macro to handle floating-point constants is precision-independent except for any calls to precision-specific standard math library functions. (Our example programs use both realtype and RCONST.) Users can, however, use the type double, float, or long double in their code (assuming the typedef for realtype matches this choice). Thus, a previously existing piece of ANSI C code can use SUNDIALS without modifying the code to use realtype, so long as the SUNDIALS libraries use the correct precision (for details see §2.2).

### 5.2 Header Files

The calling program must include several header files so that various macros and data types can be used. The header file that is always required is:

• kinsol.h, the header file for KINSOL, which defines several types and various constants, and includes function prototypes.

kinsol.h also includes sundialstypes.h, which defines the types realtype and booleantype and constants FALSE and TRUE.

The calling program must also include an NVECTOR implementation header file (see Chapter 6 for details). For the two NVECTOR implementations that are included in the KINSOL package, the corresponding header files are:

- nvector\_serial.h, which defines the serial implementation, NVECTOR\_SERIAL;
- nvector\_parallel.h, which defines the parallel MPI implementation, NVECTOR\_PARALLEL.

Note that both of these files include in turn the header file nvector.h, which defines the abstract N\_Vector type.

Finally, a linear solver module header file is required. At the present time, KINSOL offers only a Krylov linear solver, KINSPGMR, whose corresponding header file is kinspgmr.h, This in turn includes a header file (iterative.h) which enumerates the kind of preconditioning and the choices for the Gram-Schmidt process.

Other headers may be needed, according to the choice of preconditioner, etc. For example, in the kinwebs example [4], preconditioning is done with a block-diagonal matrix. For this, the header smalldense.h is included.

### 5.3 A Skeleton of the User's Main Program

A high-level view of the combined user program and KINSOL package is shown in Figure 5.1. The following is a skeleton of the user's main program (or calling program) for the solution of a nonlinear problem. Most steps are independent of the NVECTOR implementation used; where this is not the case, usage specifications are given for the two implementations provided with KINSOL: steps marked with [P] correspond to NVECTOR\_PARALLEL, while steps marked with [S] correspond to NVECTOR\_SERIAL.

### 1. Initialize MPI

[P] MPI\_Init(&argc, &argv); to initialize MPI if used by the user's program, aside from the internal use in NVECTOR\_PARALLEL. Here argc and argv are the command line argument counter and array received by main.

### 2. Set problem dimensions

[S] Set N, the problem size N.

[P] Set Nlocal, the local vector length (the sub-vector length for this process); N, the global vector length (the problem size N, and the sum of all the values of Nlocal); and the active set of processes.

### 3. Set vector with initial guess

To set the vector  $\mathbf{u}$  of initial values, use functions defined by a particular NVECTOR implementation. If a realtype array udata already exists, containing the initial guess of  $u_0$ , make the call:

```
[S] u = NV_Make_Serial(N, udata);
[P] u = NV_Make_Parallel(comm, Nlocal, N, udata);
```

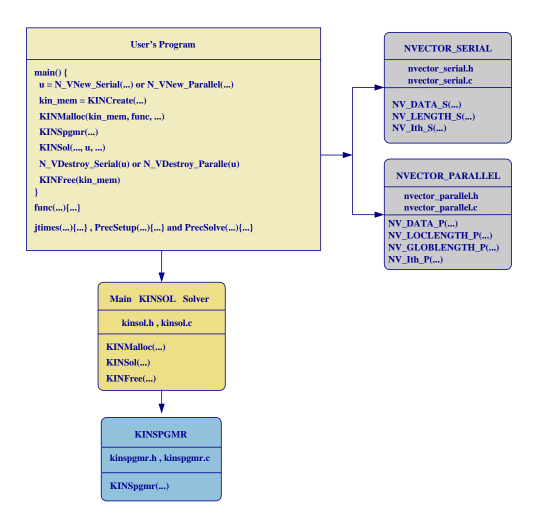


Figure 5.1: Diagram of the user program and KINSOL package for the solution of nonlinear systems

Otherwise, make the call:

```
[S] u = NV_New_Serial(N);
```

[P] u = NV\_New\_Parallel(comm, Nlocal, N);

and load initial values into the structure defined by:

[S] NV\_DATA\_S(u)

[P] NV\_DATA\_P(u)

Here comm is the MPI communicator, set in one of two ways: If a proper subset of active processes is to be used, comm must be set by suitable MPI calls. Otherwise, to specify that all processes are to be used, comm must be MPI\_COMM\_WORLD.

### 4. Create KINSOL object

Call kin\_mem = KINCreate(); to create the KINSOL memory block. KINCreate returns a pointer to the KINSOL memory structure.

### 5. Set optional inputs

Call KINSet\* routines to change any optional inputs that control the behavior of KINSOL from their default values.

### 6. Allocate internal memory

Call KINMalloc(...); to specify the problem defining function F, allocate internal memory for KINSOL, and initialize KINSOL. KINMalloc returns an error flag to indicate success or an illegal argument value (for details see  $\S5.4.1$ ).

### 7. Attach linear solver module

Initialize the linear solver module by calling KINSpgmr(...); to specify the maximum dimension of the Krylov subspace.

### 8. Set linear solver optional inputs

Call KINSpgmrSet\* routines to change optional inputs for the KINSPGMR linear solver.

### 9. Solve problem

Call KINSol(...); to solve the nonlinear problem for a given initial guess (see §5.4.3 for details).

### 10. Get optional outputs

Call KINGet\* functions to obtain optional output from KINSOL, and call KINSpgmrGet\* functions for optional outputs from KINSPGMR. See §5.4.5.

### 11. Deallocate memory for solution vector

Upon completion of the solution, deallocate memory for the vector **u** by calling the destructor function defined by the NVECTOR implementation:

```
[S] NV_Destroy_Serial(u);
```

[P] NV\_Destroy\_Parallel(u);

#### 12. Free solver memory

Call KINFree(kin\_mem); to free the memory allocated for KINSOL.

### 13. [P] Finalize MPI

Call MPI\_Finalize(); to terminate MPI.

### 5.4 User-callable functions

This section describes the KINSOL functions that are called by the user to set up and solve a nonlinear problem. Some of these are required. However, starting with §5.4.4, the functions listed involve optional inputs/outputs or restarting, and those paragraphs can be skipped for a casual use of KINSOL. In any case, refer to §5.3 for the correct order of these calls.

### 5.4.1 KINSOL initialization and deallocation functions

The following three functions must be called in the order listed. The last one is to be called only after the problem solution is complete, as it frees the KINSOL memory block created and allocated by the first two calls.

### KINCreate

Call kin\_mem = KINCreate();

Description The function KINCreate instantiates a KINSOL solver object.

Arguments This function has no arguments.

 $Return\ value\ If\ successful,\ \texttt{KINCreate}\ returns\ a\ pointer\ to\ the\ newly\ created\ \texttt{KINSOL}\ memory\ block$ 

(of type void \*). If an error occurred, KINCreate prints an error message to stderr

and returns NULL.

### KINMalloc

Call flag = KINMalloc(kin\_mem, func, tmpl);

Description The function KINMalloc specifies the problem-defining function, allocates internal

memory, and initializes KINSOL.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block returned by KINCreate.

func (KINSysFn) is the C function which computes F in the nonlinear problem. This function has the form func(u, fval, f\_data) (for full details

see  $\S 5.5.1$ ).

tmpl (N\_Vector) is an N\_Vector which is used as a template to create (by cloning)

necessary vectors in kin\_mem.

Return value The return flag flag (of type int) will be one of the following:

KIN\_SUCCESS The call to KINMalloc was successful.

KIN\_MEM\_NULL The kinsol memory block was not initialized through a previous call

to KINCreate.

KIN\_MEM\_FAIL A memory allocation request has failed.

KIN\_ILL\_INPUT An input argument to KINMalloc has an illegal value.

Notes If an error occurred, KINMalloc also prints an error message to the file specified by

the optional input errfp.

### KINFree

Call KINFree(kin\_mem);

Description The function KINFree frees the pointer allocated by a previous call to KINMalloc.

Arguments The argument is the pointer to the KINSOL memory block (of type void \*).

Return value The function KINFree has no return value.

### 5.4.2 Linear solver specification function

As previously explained, Newton iteration requires the solution of linear systems of the form (2). At the present time there is only one solver available for this task, KINSPGMR. This is an iterative solver that uses a scaled preconditioned GMRES method.

To attach the KINSPGMR linear solver, after the call to KINCreate but before any call to KINSol, the user's program must call KINSpgmr, as documented below. The first argument passed to this function is the KINSOL memory pointer returned by KINCreate. The call to this function links the linear solver to the main KINSOL memory block and allows the user to specify parameters for KINSPGMR

The KINSPGMR linear solver is actually built on top of a generic linear system solver, which may be of interest in itself. This generic solver, SPGMR, is described separately in Chapter 8.

KINSpgmr

Call flag = KINSpgmr(kin\_mem, maxl);

Description The function KINSpgmr selects the KINSPGMR linear solver.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

maxl (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value KINSPGMR\_MAXL= 5.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The KINSPGMR initialization was successful.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_ILL\_INPUT The NVECTOR module used does not implement a required operation.

KINSPGMR\_MEM\_FAIL A memory allocation request failed.

Notes The KINSPGMR solver uses a scaled preconditioned GMRES iterative method to solve the linear system (2).

Within KINSOL, only right preconditioning is available. For specification of the preconditioner, see §5.4.4 and §5.5.

If preconditioning is done, user-supplied functions define the right preconditioner matrices P, which approximate the Newton matrix from (2).

### 5.4.3 KINSOL solver function

KINSol

Call flag = KINSol(kin\_mem, u, strategy, u\_scale, f\_scale);

Description The function KINSol computes an approximate solution of the nonlinear system.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

u (N\_Vector) vector set to initial guess by user before calling KINSo1, but which upon return contains an approximate solution of the nonlinear system F(u) = 0 the computed solution vector.

strategy globalization strategy applied to the Newton method. It must be one of KIN\_INEXACT\_NEWTON or KIN\_LINESEARCH.

u\_scale vector containing diagonal elements of scaling matrix  $D_u$  for vector u chosen so that the components of  $D_u \cdot \mathbf{u}$  (as a matrix multiplication) all have about the same magnitude when u is close to a root of F(u).

**f\_scale** vector containing diagonal elements of scaling matrix  $D_F$  for F(u) chosen so that the components of  $D_F \cdot F(\mathbf{u})$  (as a matrix multiplication) all have roughly the same magnitude when  $\mathbf{u}$  is not too near a root of F(u).

Return value On return, KINSol returns the approximate solution in the vector **u**. The return value flag (of type int) will be one of the following:

### KIN\_SUCCESS

KINSol succeeded; the scaled norm of F(u) is less than fnormtol.

### KIN\_INITIAL\_GUESS\_OK

The guess  $\mathbf{u} = u_0$  satisfied the system F(u) = 0 within the tolerances specified.

### KIN\_STEP\_LT\_STPTOL

KINSOL stopped based on scaled step length. This means that the current iterate may be an approximate solution of the given nonlinear system, but it is also quite possible that the algorithm is "stalled" (making insufficient progress) near an invalid solution, or that the scalar scsteptol is too large (see KINSetScaledStepTol in §5.4.4 to change scsteptol from its default value).

#### KIN\_MEM\_NULL

The KINSOL memory block pointer was NULL.

### KIN\_ILL\_INPUT

An input parameter was invalid.

#### KIN\_NO\_MALLOC

The KINSOL memory was not allocated by a call to KINMalloc.

#### KIN\_LINESEARCH\_NONCONV

The line search algorithm was unable to find an iterate sufficiently distinct from the current iterate, or could not find an iterate satisfying the sufficient decrease condition.

Failure to satisfy the sufficient decrease condition could mean the current iterate is "close" to an approximate solution of the given nonlinear system, the finite difference approximation of the matrix-vector product J(u)v is inaccurate, or the real scalar scsteptol is too large.

#### KIN\_MAXITER\_REACHED

The maximum number of nonlinear iterations has been reached.

#### KIN\_MXNEWT\_5X\_EXCEEDED

Five consecutive steps have been taken that satisfy the inequality  $||D_u p||_{L2} > 0.99$  mxnewtstep, where p denotes the current step and mxnewtstep is a scalar upper bound on the scaled step length.

Such a failure may mean that  $||D_F F(u)||_{L^2}$  asymptotes from above to a finite value, or the real scalar mxnewtstep is too small.

### KIN\_LINESEARCH\_BCFAIL

The line search algorithm was unable to satisfy the "beta-condition" for MXNBCF + 1 nonlinear iterations (not necessarily consecutive), which may indicate the algorithm is making poor progress.

### KIN\_LINSOLV\_NO\_RECOVERY

The user-supplied routine psolve encountered a recoverable error, but the preconditioner is already current.

### KIN\_LINIT\_FAIL

The linear solver initialization routine (linit) encountered an error.

### KIN\_LSETUP\_FAIL

The user-supplied routine pset (used to set up the preconditioner data) encountered an unrecoverable error.

### KIN\_LSOLVE\_FAIL

Either the user-supplied routine psolve (used to to solve the preconditioned linear system) encountered an unrecoverable error, or the linear solver routine (lsolve) encountered an error condition.

Optional input	Function name	Default		
KINSOL main solver				
Pointer to an error file	KINSetErrFile	stderr		
Pointer to an info file	KINSetInfoFile	stdout		
Data for problem-defining function	KINSetFdata	NULL		
Verbosity level of output	KINSetPrintLevel	0		
Max. number of nonlinear iterations	KINSetNumMaxIters	200		
No initial preconditioner setup	KINSetNoPrecInit	FALSE		
Max. iterations without prec. setup	KINSetMaxPrecCalls	10		
Form of $\eta$ coefficient	KINSetEtaForm	KIN_ETACHOICE1		
Constant value of $\eta$	KINSetEtaConstValue	0.1		
Values of $\gamma$ and $\alpha$	KINSetEtaParams	0.9  and  2.0		
Lower bound on $\epsilon$	KINSetNoMinEps	FALSE		
Max. scaled length of Newton step	KINSetMaxNewtonStep	$1000  D_u u_0  _2$		
Rel. error for F.D. $Jv$	KINSetRelErrFunc	$\sqrt{\text{uround}}$		
Function-norm stopping tolerance	KINSetFuncNormTol	<sup>3</sup> √uround		
Scaled-step stopping tolerance	KINSetScaledSteptol	$uround^{2/3}$		
Inequality constraints on solution	KINSetConstraints	NULL		
Nonlinear system function	KINSetSysFunc	none		
KINSPGMR linear solver				
Max. number of restarts	KINSpgmrSetMaxRestarts	0		
Preconditioner functions and data	KINSpgmrSetPreconditioner	NULL, NULL, NULL		
Jacobian-vector product function and data	KINSpgmrSetJacTimesVecFn	internal DQ, NULL		

Table 5.1: Optional inputs for KINSOL and KINSPGMR

Notes

The components of vectors u\_scale and f\_scale should be strictly positive.

KIN\_SUCCESS = 0, KIN\_INITIAL\_GUESS\_OK = 1, and KIN\_STEP\_LT\_STPTOL = 2. All remaining return values are negative and therefore a test flag < 0 will trap all KINSol failures.

### 5.4.4 Optional input functions

KINSOL provides an extensive list of functions that can be used to change from their default values various optional input parameters that control the behavior of the KINSOL solver. Table 5.1 lists all optional input functions in KINSOL, which are then described in detail in the remainder of this section. For the most casual use of KINSOL, the reader can skip to §5.5.

We note that, on error return, all of these functions also print an error message to **stderr** (or to the file pointed to by **errfp** if already specified). We also note that all error return values are negative, so a test flag < 0 will catch any error.

### Main solver optional input functions

The calls listed here can be executed in any order. However, if KINSetErrFile is to be called, that call should be first, in order to take effect for any later error message.

KINSetErrFile	l
---------------	---

Call flag = KINSetErrFile(kin\_mem, errfp);

Description The function KINSetErrFile specifies the pointer to the file where all KINSOL error

messages should be directed.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

errfp (FILE \*) pointer to output file.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

Notes The default value for errfp is stderr.

Passing a value NULL disables all future error message output (except for the case in which the KINSOL memory pointer is NULL).

If KINSetErrFile is to be called, it should be called before any other optional input functions, in order to take effect for any later error message.

### KINSetInfoFile

Call flag = KINSetInfoFile(kin\_mem, infofp);

Description The function KINSetInfoFile specifies the pointer to the file where all informative

messages should be directed.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

infofp (FILE \*) pointer to output file.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

Notes The default value for infofp is stderr.

### KINSetPrintLevel

Call flag = KINSetPrintLevel(kin\_mem, printfl);

Description The function KINSetPrintLevel specifies the level of verbosity of the output.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

printfl (int) flag indicating the level of verbosity. Must be one of:

0 no information displayed.

- 1 for each nonlinear iteration display the following information: the scaled Euclidean  $\ell_2$  norm of the system function evaluated at the current iterate, the scaled norm of the Newton step (only if using KIN\_INEXACT\_NEWTON), and the number of function evaluations performed so far.
- 2 display level 1 output and the following values for each iteration:  $\|F(u)\|_{D_F}$  (only for KIN\_INEXACT\_NEWTON).  $\|F(u)\|_{D_F,\infty}$  (for KIN\_INEXACT\_NEWTON and KIN\_LINESEARCH).
- 3 display level 2 output plus additional values used by the global strategy (only if using KIN\_LINESEARCH), and statistical information for the linear solver.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

KIN\_ILL\_INPUT The argument printfl had an illegal value.

Notes The default value for printfl is 0.

### KINSetFdata

Call flag = KINSetFdata(kin\_mem, f\_data);

Description The function KINSetFdata specifies the pointer to user-defined memory that is to be

passed to the user-supplied function implementing the nonlinear system residual.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

f\_data (void \*) pointer to the user-defined memory.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

Notes The default value for f\_data is NULL.

### KINSetNumMaxIters

Call flag = KINSetNumMaxIters(kin\_mem, mxiter);

Description The function KINSetNumMaxIters specifies the maximum number of nonlinear itera-

tions allowed.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

mxiter (long int) maximum number of nonlinear iterations.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

KIN\_ILL\_INPUT The maximum number of iterations was non-positive.

Notes The default value for mxiter is MXITER\_DEFAULT = 200.

### KINSetNoPrecInit

Call flag = KINSetNoPrecInit(kin\_mem, noPrecInit);

Description The function KINSetNoPrecInit specifies whether an initial call to the preconditioner

setup function should be made or not.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

noPrecInit (booleantype) flag controlling whether or not an initial call to the pre-

conditioner setup function is made.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

Notes The default value for noPrecInit is FALSE, meaning that an initial call to the precon-

ditioner setup function will be made.

### KINSetMaxPrecCalls

Call flag = KINSetMaxPrecCalls(kin\_mem, msbpre);

Description The function KINSetMaxPrecCalls specifies the maximum number of nonlinear iter-

ations that can be performed between calls to the preconditioner setup function.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

msbpre (long int) maximum number of nonlinear iterations without a call to the

preconditioner setup function.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

KIN\_ILL\_INPUT The argument msbpre was negative.

Notes The default value for msbpre is MSBPRE = 10.

### KINSetEtaForm

Call flag = KINSetEtaForm(kin\_mem, etachoice);

Description – The function KINSetEtaForm specifies the method for computing the value of the  $\eta$ 

coefficient used in the calculation of the linear solver convergence tolerance.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

etachoice (int) flag indicating the method for computing  $\eta$ . etachoice must be one of KIN\_ETACHOICE1, KIN\_ETACHOICE2, or KIN\_ETACONSTANT (see Chapter

3 for details).

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

KIN\_ILL\_INPUT The argument etachoice had an illegal value.

Notes The default value for etachoice is KIN\_ETACHOICE1.

### KINSetEtaConstValue

Call flag = KINSetEtaConstValue(kin\_mem, eta);

Description The function KINSetEtaConstValue specifies the constant value for  $\eta$  in the case

etachoice = KIN\_ETACONSTANT.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

eta (realtype) constant value for  $\eta$ .

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

KIN\_ILL\_INPUT The argument eta had an illegal value

Notes The default value for eta is 0.1. The valid values are  $0.0 < \text{eta} \le 1.0$ .

### KINSetEtaParams

Call flag = KINSetEtaParams(kin\_mem, egamma, ealpha);

Description The function KINSetEtaParams specifies the parameters  $\gamma$  and  $\alpha$  in the formula for  $\eta$ ,

in the case etachoice = KIN\_ETACHOICE2.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

egamma (realtype) value of the  $\gamma$  parameter. ealpha (realtype) value of the  $\alpha$  parameter.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional values have been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

KIN\_ILL\_INPUT One of the arguments egamma or ealpha had an illegal value.

Notes The default values for egamma and ealpha are 0.9 and 2.0, respectively.

The valid values for ealpha are  $1.0 < \text{ealpha} \le 2.0$ . If ealpha = 0.0, then its value is set to 2.0.

The valid values for egamma are  $0.0 < \text{egamma} \le 1.0$ . If egamma = 0.0, then its value is set to 0.9.

### KINSetNoMinEps

Call flag = KINSetNoMinEps(kin\_mem, noMinEps);

Description The function KINSetNoMinEps specifies a flag that controls whether or not the value

of  $\epsilon$ , the scaled linear residual tolerance, is bounded from below.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

noMinEps (booleantype) flag controlling the bound on  $\epsilon$ .

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

Notes The default value for noMinEps is FALSE.

### KINSetMaxNewtonStep

Call flag = KINSetMaxNewtonStep(kin\_mem, mxnewtstep);

Description The function KINSetMaxNewtonStep specifies the maximum allowable scaled length of

the Newton step.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

mxnewtstep (realtype) maximum scaled step length.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

KIN\_ILL\_INPUT The maximum step was non-positive.

Notes The default value of mxnewtstep is  $1000 \|u_0\|_{D_u}$ , where  $u_0$  is the initial guess.

### KINSetRelErrFunc

Call flag = KINSetRelErrFunc(kin\_mem, relfunc);

Description The function KINSetRelErrFunc specifies the relative error in computing F(u), which

is used in the difference quotient approximation of the Jacobian-vector product.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

relfunc (realtype) relative error in F(u).

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

 ${\tt KIN\_MEM\_NULL} \quad {\tt The \; kin\_mem \; pointer \; is \; NULL}.$ 

KIN\_ILL\_INPUT The relative error was non-positive.

Notes The default value for relfunc is  $\sqrt{\text{unit roundoff}}$ .

#### KINSetFuncNormTol

Call flag = KINSetFuncNormTol(kin\_mem, fnormtol);

Description The function KINSetFuncNormTol specifies the scalar used as a stopping tolerance on

the scaled maximum norm of the system function F(u).

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

fnormtol (realtype) tolerance for stopping based on scaled function norm.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL. KIN\_ILL\_INPUT The tolerance was non-positive.

Notes The default value for fnormtol is  $\sqrt[3]{\text{unit roundoff}}$ .

#### ${\tt KINSetScaledStepTol}$

Call flag = KINSetScaledStepTol(kin\_mem, scsteptol);

Description The function KINSetScaledStepTol specifies the scalar used as a stopping tolerance

on the minimum scaled step length.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

scsteptol (realtype) tolerance for stopping based on scaled step length..

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.
KIN\_ILL\_INPUT The tolerance was non-positive.

Notes The default value for scsteptol is (unit roundoff) $^{2/3}$ .

#### KINSetConstraints

Call flag = KINSetConstraints(kin\_mem, constraints);

Description The function KINSetConstraints specifies a vector that defines inequality constraints

for each component of the solution vector u.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

constraints (N\_Vector) vector of constraint flags. If constraints[i] is

0.0 then no constraint is imposed on  $u_i$ .

1.0 then  $u_i$  will be constrained to be  $u_i > 0.0$ .

-1.0 then  $u_i$  will be constrained to be  $u_i < 0.0$ .

2.0 then  $u_i$  will be constrained to be  $u_i \geq 0.0$ .

-2.0 then  $u_i$  will be constrained to be  $u_i \leq 0.0$ .

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

KIN\_ILL\_INPUT The constraint vector contains illegal values.

Notes The presence of a non-NULL constraints vector that is not 0.0 in all components will

cause constraint checking to be performed.

The function creates a private copy of the constraints vector. Consequently, the user-supplied vector can be freed after the function call, and the constraints can only be

changed by calling this function.

#### KINSetSysFunc

Call flag = KINSetSysFunc(kin\_mem, func);

Description The function KINSetSysFunc specifies the user-provided function that evaluates the

nonlinear system function F(u).

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

func (KINSysFn) user-supplied function that evaluates F(u).

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.
KIN\_ILL\_INPUT The argument func was NULL.

Notes The nonlinear system function is initially specified through KINMalloc. The option

of changing the system function is provided for a user who wishes to solve several

problems of the same size but with different functions.

#### Linear solver optional input functions

The KINSPGMR linear solver module allows for various optional inputs, which are described here. The call to KINSpgmr is used to communicate the maximum dimension of the Krylov subspace to be used (max1).

If preconditioning is to be done within the SPGMR method, then the user must supply a preconditioner solve function psolve and specify it through a call to KINSpgmrSetPrecSolveFn. The evaluation and preprocessing of any Jacobian-related data needed by the user's preconditioner solve function is done in the optional user-supplied function psetup. Both of these functions are fully specified in §5.5. If used, the psetup function should be specified through a call to KINSpgmrSetPrecSetupFn. Optionally, the KINSPGMR solver passes the pointer it receives through KINSpgmrSetPrecData to the preconditioner setup and solve functions. This allows the user to create an arbitrary structure with relevant problem data and access it during the execution of the user-supplied preconditioner functions without using global data in the program. The pointer prec\_data may be identical to f\_data, if the latter was specified through KINSetFdata.

The KINSPGMR solver requires a function to compute an approximation to the product between the Jacobian matrix J(u) and a vector v. The user can supply his/her own Jacobian-vector product approximation function, or use the difference quotient function KINSpgmrDQJtimes that comes with the KINSPGMR solver. A user-defined Jacobian-vector function must be of type KINSpgmrJtimesFn and can be specified through a call to KINSpgmrSetJacTimesVecFn (see §5.5 for specification details). As with the preconditioner user data structure prec\_data, the user can specify, through a call to KINSpgmrSetJacData, a pointer to a user-defined data structure, jac\_data, which the KINSPGMR solver passes to the Jacobian-vector product function jtimes each time it is called. The pointer jac\_data may be identical to prec\_data and/or f\_data.

#### KINSpgmrSetMaxRestarts

Call flag = KINSpgmrSetMaxRestarts(kin\_mem, maxrs);

Description The function KINSpgmrSetMaxRestarts specifies the maximum number of times the

SPGMR linear solver can be restarted.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

maxrs (int) maximum number of restarts.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional value has been successfully set.

KINSPGMR\_ILL\_INPUT The maximum number of restarts specified is negative.

 ${\tt KINSPGMR\_MEM\_NULL} \quad {\tt The \; kin\_mem \; pointer \; is \; NULL}.$ 

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

#### ${\tt KINSpgmrSetPreconditioner}$

Call flag = KINSpgmrSetPreconditioner(kin.mem, psetup, psolve, prec\_data);

Description The function KINSpgmrSetPreconditioner specifies the preconditioner setup and

solve function and the pointer to user data.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

psetup (KINSpgmrPrecSetupFn) user-defined preconditioner setup function. psolve (KINSpgmrPrecSolveFn) user-defined preconditioner solve function.

prec\_data (void \*) pointer to the user-defined data structure.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional value has been successfully set.

 ${\tt KINSPGMR\_MEM\_NULL} \quad {\tt The \; kin\_mem \; pointer \; is \; NULL}.$ 

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

Notes The function type KINSpgmrPrecSetupFn is described in §5.5.4. The function type

KINSpgmrPrecSolveFn is described in §5.5.3.

#### KINSpgmrSetJacTimesVecFn

Call flag = KINSpgmrSetJacTimesVecFn(kin\_mem, jtimes, jac\_data);

Description The function KINSpgmrSetJacTimesVecFn specifies the Jacobian-vector product func-

tion to be used and the pointer to user data.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

jtimes (KINSpgmrJacTimesVecFn) user-defined Jacobian-vector product function.

jac\_data (void \*) pointer to the user-defined data structure.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional value has been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

Notes By default, KINSPGMR uses the difference quotient function KINSpgmrDQJtimes. If

NULL is passed to jtimes, this default function is used.

The function type KINSpgmrJacTimesVecFn is described in §5.5.2.

#### 5.4.5 Optional output functions

KINSOL provides an extensive list of functions that can be used to obtain solver performance information. Table 5.2 lists all optional output functions in KINSOL, which are then described in detail in the remainder of this section.

#### Main solver optional output functions

KINSOL provides several user-callable functions that can be used to obtain different quantities that may be of interest to the user, such as solver workspace requirements and solver performance statistics. These optional output functions are described next.

#### KINGetWorkSpace

Call flag = KINGetWorkSpace(kin\_mem, &lenrw, &leniw);

Description The function KINGetWorkSpace returns the KINSOL integer and real workspace sizes.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

lenrw (long int) the number of realtype values in the KINSOL workspace.

Table 5.2: Optional outputs from KINSOL and KINSPGMR

Optional output	Function name		
KINSOL main solve	r		
Size of Kinsol real and integer workspaces	KINGetWorkSpace		
Number of function evaluations	KINGetNumFuncEvals		
Number of nonlinear iterations	KINGetNumNolinSolvIters		
Number of $\beta$ -condition failures	KINGetNumBetaCondFails		
Number of backtrack operations	KINGetNumBacktrackOps		
Scaled norm of $F$	KINGetFuncNorm		
Scaled norm of the step	KINGetStepLength		
KINSPGMR linear solver			
Size of Kinspgmr real and integer workspaces	KINSpgmrGetWorkSpace		
No. of linear iterations	KINSpgmrGetNumLinIters		
No. of linear convergence failures	KINSpgmrGetNumConvFails		
No. of preconditioner evaluations	KINSpgmrGetNumPrecEvals		
No. of preconditioner solves	KINSpgmrGetNumPrecSolves		
No. of Jacobian-vector product evaluations	KINSpgmrGetNumJtimesEvals		
No. of fct. calls for finite diff. Jacobian-vector evals.	KINSpgmrGetNumFuncEvals		
Last return from a KINSPGMR function	KINSpgmrGetLastFlag		

leniw (long int) the number of integer values in the KINSOL workspace.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output values have been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

Notes

In terms of the problem size N, the actual size of the real workspace is 17 + 5N realtype words. The real workspace is increased by an additional N words if constraint checking is enabled (see KINSetConstraints).

The actual size of the integer workspace (without distinction between int and long int) is 22 + 5N (increased by N if constraint checking is enabled).

#### KINGetNumFuncEvals

Call flag = KINGetNumFuncEvals(kin\_mem, &nfevals);

Description The function KINGetNumFuncEvals returns the number of evaluations of the system

function.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

**nfevals** (long int) number of calls to the user-supplied function that evaluates F(u).

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

#### KINGetNumNonlinSolvIters

Call flag = KINGetNumNonlinSolvIters(kin\_mem, &nniters);

Description The function KINGetNumNonlinSolvIters returns the number of nonlinear iterations.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

nniters (long int) number of nonlinear iterations.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

#### KINGetNumBetaCondFails

Call flag = KINGetNumBetaCondFails(kin\_mem, &nbcfails);

Description The function KINGetNumBetaCondFails returns the number of  $\beta$ -condition failures.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

nbcfails (long int) number of  $\beta$ -condition failures.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

#### KINGetNumBacktrackOps

Call flag = KINGetNumBacktrackOps(kin\_mem, &nbacktr);

Description The function KINGetNumBacktrackOps returns the number of backtrack operations

(step length adjustments) performed by the line search algorithm.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

nbacktr (long int) number of backtrack operations.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

#### KINGetFuncNorm

Call flag = KINGetFuncNorm(kin\_mem, &fnorm);

Description The function KINGetFuncNorm returns the scaled Euclidean  $\ell_2$  norm of the nonlinear

system function F(u) evaluated at the current iterate.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

fnorm (realtype) current scaled norm of F(u).

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

#### KINGetStepLength

Call flag = KINGetStepLength(kin\_mem, &steplength);

Description The function KINGetStepLength returns the scaled Euclidean  $\ell_2$  norm of the step

used during the previous iteration.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

steplength (realtype) scaled norm of the Newton step.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output value has been successfully set.

KIN\_MEM\_NULL The kin\_mem pointer is NULL.

#### Linear solver optional output functions

The functions available to access various optional outputs that describe the performance of the KINSPGMR module are described below.

#### KINSpgmrGetWorkSpace

Call flag = KINSpgmrGetWorkSpace(kin\_mem, &lenrwSG, &leniwSG);

Description The function KINSpgmrGetWorkSpace returns the real and integer workspace sizes

used by KINSPGMR.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

lenrwSG (long int) the number of realtype values in the KINSPGMR workspace.

leniwSG (long int) the number of integer values in the KINSPGMR workspace.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional output values have been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

Notes In terms of the problem size N and maximum subspace size max1, the actual size of

the real workspace is  $(\max 1+5) * N + \max 1 * (\max 1+4) + 1$  realtype words. (In a

parallel setting, this value is global - summed over all processes.)

#### KINSpgmrGetNumLinIters

Call flag = KINSpgmrGetNumLinIters(kin\_mem, &nliters);

Description The function KINSpgmrGetNumLinIters returns the cumulative number of linear iter-

ations.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

nliters (long int) the current number of linear iterations.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional output value has been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

#### KINSpgmrGetNumConvFails

Call flag = KINSpgmrGetNumConvFails(kin\_mem, &nlcfails);

Description The function KINSpgmrGetNumConvFails returns the cumulative number of linear

convergence failures.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

nlcfails (long int) the current number of linear convergence failures.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional output value has been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

#### ${\tt KINSpgmrGetNumPrecEvals}$

Call flag = KINSpgmrGetNumPrecEvals(kin\_mem, &npevals);

Description The function KINSpgmrGetNumPrecEvals returns the number of preconditioner eval-

uations, i.e., the number of calls made to psetup.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

npevals (long int) the current number of calls to psetup.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional output value has been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

#### KINSpgmrGetNumPrecSolves

Call flag = KINSpgmrGetNumPrecSolves(kin\_mem, &npsolves);

Description The function KINSpgmrGetNumPrecSolves returns the cumulative number of calls

made to the preconditioner solve function, psolve.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

npsolves (long int) the current number of calls to psolve.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional output value has been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

#### KINSpgmrGetNumJtimesEvals

Call flag = KINSpgmrGetNumJtimesEvals(kin\_mem, &njvevals);

Description The function KINSpgmrGetNumJtimesEvals returns the cumulative number made to

the Jacobian-vector product function, jtimes.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

njvevals (long int) the current number of calls to jtimes.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional output value has been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

#### KINSpgmrGetNumRhsEvals

Call flag = KINSpgmrGetNumRhsEvals(kin\_mem, &nfevalsSG);

Description The function KINSpgmrGetNumRhsEvals returns the number of calls to the user right-

hand side function for finite difference Jacobian-vector product approximations.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

**nfevalsSG** (long int) the number of calls to the user right-hand side function.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional output value has been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

Notes The value nfevalsSG is incremented only if the default KINSpgmrDQJtimes difference

quotient function is used.

#### KINSpgmrGetLastFlag

Call flag = KINSpgmrGetLastFlag(kin\_mem, &flag);

Description The function KINSpgmrGetLastFlag returns the last return value from a KINSPGMR

routine.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

flag (int) the value of the last return flag from a KINSPGMR function.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The optional output value has been successfully set.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_LMEM\_NULL The KINSPGMR linear solver has not been initialized.

Notes If the KINSPGMR setup function failed (KINSOL returned KINLSETUP\_FAIL), flag con-

tains the return value of the preconditioner setup function psetup.

If the KINSPGMR solve function failed (KINSOL returned KIN\_LSOLVE\_FAIL), flag contains the error return flag from SpgmrSolve and will be one of: SPGMR\_CONV\_FAIL, indicating a failure to converge; SPGMR\_QRFACT\_FAIL, indicating a singular matrix found during the QR factorization; SPGMR\_PSOLVE\_FAIL\_REC, indicating that the preconditioner solve function psolve failed recoverably; SPGMR\_MEM\_NULL, indicating that the SPGMR memory is NULL; SPGMR\_ATIMES\_FAIL, indicating a failure in the Jacobian-vector product function; SPGMR\_PSOLVE\_FAIL\_UNREC, indicating that the preconditioner solve function psolve failed unrecoverably; SPGMR\_GS\_FAIL, indicating a failure in the Gram-Schmidt procedure; or SPGMR\_QRSOL\_FAIL, indicating that the matrix R was found to be singular during the QR solve phase.

### 5.5 User-supplied functions

The user-supplied functions consist of one function defining the nonlinear system, (optionally) a function that provides Jacobian-related information for the linear solver, and (optionally) one or two functions that define the preconditioner for use in the SPGMR algorithm.

#### 5.5.1 Problem-defining function

The user must provide a function of type KINSysFn defined as follows:

KINSysFn

Definition typedef void (\*KINSysFn)(N\_Vector u, N\_Vector fval, void \*f\_data );

Purpose This function computes F(u) for a given value of the vector u.

Arguments u is the current value of the variable vector, u.

fval is the output vector F(u).

f\_data is a pointer to user data, same as the pointer f\_data passed to KINSetFdata.

Return value A KINSysFn function type does not have a return value.

Notes Allocation of memory for fval is handled within KINSOL.

#### 5.5.2 Jacobian information (SPGMR matrix-vector product)

The user may provide a function of type KINSpgmrJacTimesVecFn to evaluate Jacobian-vector products for the KINSPGMR linear solver module. This function has the following form:

#### ${\tt KINSpgmrJacTimesVecFn}$

Definition typedef int (\*KINSpgmrJacTimesVecFn)(N\_Vector v, N\_Vector Jv,

N\_Vector u, booleantype \*new\_u,

void \*jac\_data);

Purpose This function computes the product  $Jv = (\partial F/\partial u)v$  (or an approximation to it).

Arguments v is the vector by which the Jacobian must be multiplied to the right.

Jv is the output vector computed.

u is the current (unscaled) value of the iterate.

new\_u is a flag (reset by user) indicating if the iterate u has been updated in the

interim. The Jacobian-vector product needs to be updated/reevaluated, if

appropriate, unless new\_u = FALSE.

jac\_data is a pointer to user data, the same as the jac\_data parameter passed to

KINSpgmrSetJacData.

Return value The value to be returned by the Jacobian-vector product function should be 0 if

successful. Any other return value will result in an unrecoverable error of the SPGMR

generic solver, in which case the solution process is halted.

Notes If a user-defined routine is not given, then an internal kinspgmr function, using

difference quotient approximations, is used.

If the user-provided KINSpgmrJacTimesVec function needs the unit roundoff, this can be accessed as UNIT\_ROUNDOFF defined in sundialstypes.h.



The user is responsible for resetting the value of new\_u to FALSE.

### 5.5.3 Preconditioning (SPGMR linear system solution)

If preconditioning is used, then the user must provide a C function to solve the linear system Pz = r where P is the preconditioner matrix. This function must be of type KINSpgmrPrecSolveFn, defined as follows:

#### KINSpgmrPrecSolveFn

Definition typedef int (\*KINSpgmrPrecSolveFn)(N\_Vector u, N\_Vector uscale,

N\_Vector fval, N\_Vector fscale,
N\_Vector v, void \*prec\_data,

N\_Vector tmp);

Purpose This function solves the preconditioning system Pz = r.

Arguments u is the current (unscaled) value of the iterate.

uscale is a vector containing diagonal elements of the scaling matrix for u.

fval is the vector F(u) evaluated at u.

fscale is a vector containing diagonal elements of the scaling matrix for fval.

v on input, v is set to the right-hand side vector of the linear system,  ${\tt r}.$  On

output, v must contain the solution z of the linear system Pz = r.

prec\_data is a pointer to user data - the same as the prec\_data parameter passed to

the function KINSpgmrSetPrecData.

 $\verb|tmp| is a pointer to memory allocated for a variable of type $\tt N\_Vector$ which can$ 

be used for work space.

Return value The value to be returned by the preconditioner solve function is a flag indicating whether it was successful. This value should be 0 if successful, positive for a recoverable

error, and negative for an unrecoverable error.

#### 5.5.4 Preconditioning (SPGMR Jacobian data)

If the user's preconditioner requires that any Jacobian-related data be evaluated or preprocessed, then this needs to be done in a user-supplied C function of type KINSpgmrPrecSetupFn, defined as follows:

#### KINSpgmrPrecSetupFn

Definition typedef int (\*KINSpgmrPrecSetupFn) (N\_Vector u, N\_Vector uscale,

N\_Vector fval, N\_Vector fscale,
void \*prec\_data, N\_Vector tmp1,
N\_Vector tmp2);

N\_Vector tmp2)

Purpose This function evaluates and/or preprocesses Jacobian-related data needed by the pre-

conditioner.

Arguments The arguments of a KINSpgmrPrecSetupFn are as follows:

u is the current (unscaled) value of the iterate.

uscale is a vector containing diagonal elements of the scaling matrix for u.

fval is the vector F(u) evaluated at **u**.

fscale is a vector containing diagonal elements of the scaling matrix for fval.

prec\_data is a pointer to user data - the same as the prec\_data parameter passed to

the function KINSpgmrSetPrecData.

tmp1

tmp2 are pointers to memory allocated for variables of type N\_Vector which can

be used by KINSpgmrPrecSetupFn as temporary storage or work space.

Return value The value to be returned by the preconditioner setup function is a flag indicating whether it was successful. This value should be 0 if successful, positive for a recoverable

error, and negative for an unrecoverable error.

Notes The user-supplied preconditioner setup subroutine should compute the right preconditioner matrix P (stored in the memory block referenced by the prec\_data pointer)

used to form the scaled preconditioned linear system

$$(D_F J(u) P^{-1} D_u^{-1}) \cdot (D_u P x) = -D_F F(u) ,$$

where  $D_u$  and  $D_F$  denote the diagonal scaling matrices whose diagonal elements are stored in the vectors uscale and fscale, respectively.

The preconditioner setup routine will not be called prior to every call made to the preconditioner solve function, but will instead be called only as often as necessary to achieve convergence of the Newton iteration.

If the preconditioner solve routine requires no preparation, then a preconditioner setup function need not be given.

## 5.6 A parallel band-block-diagonal preconditioner module

The efficiency of Krylov iterative methods for the solution of linear systems can be greatly enhanced through preconditioning. For problems in which the user cannot define a more effective, problem-specific preconditioner, KINSOL provides a band-block-diagonal preconditioner module KINBBDPRE, to be used with the parallel N\_Vector module described in §6.2.

This module provides a preconditioner matrix for KINSOL that is block-diagonal with banded blocks. The blocking corresponds to the distribution of the dependent variable vector u amongst the processes. Each preconditioner block is generated from the Jacobian of the local part (associated with the current process) of a given function G(u) approximating F(u) (G = F is allowed). The

blocks are generated by each process via a difference quotient scheme, utilizing a specified banded structure. This structure is given by upper and lower half-bandwidths, mu and ml, defined as the number of non-zero diagonals above and below the main diagonal, respectively.

This pair of parameters need not be the true half-bandwidths of the Jacobian of the local block of G, if smaller values provide a more efficient preconditioner. Such an efficiency gain may occur if the couplings in the system outside a certain bandwidth are considerably weaker than those within the band. Reducing mu and ml lumps the outer Jacobian elements into the computed elements within the narrower band. This loss of accuracy in the Jacobian may (or may not) be offset by the lower cost of the narrower band matrices, so users should experiment with the values of mu and ml.

The KINBBDPRE module calls two user-provided functions to construct P: a required function Gloc (of type KINLocalFn) which approximates the nonlinear system function function  $G(u) \approx F(u)$  and which is computed locally, and an optional function Gcomm (of type KINCommFn) which performs all interprocess communication necessary to evaluate the approximate function G. These are in addition to the user-supplied nonlinear system function that evaluates F(u). Both functions take as input the same pointer f\_data as that passed by the user to KINSetFdata and passed to the user's function func, and neither function has a return value. The user is responsible for providing space (presumably within f\_data) for components of u that are communicated by Gcomm from the other processes, and that are then used by Gloc, which is not expected to do any communication.

#### KINLocalFn

Definition typedef void (\*KINLocalFn)(long int Nlocal, N\_Vector u, N\_Vector gval, void \*f\_data);

Purpose This function computes  $G(\mathbf{u})$ , and outputs the resulting vector as gval.

Arguments Nlocal is the local vector length.

u is the current value of the iterate.

gval is the output vector.

f\_data is a pointer to user data - the same as the f\_data parameter passed to KINSetFdata.

NINSetruata.

Return value A KINLocalFn function type does not have a return value.

Notes This function assumes that all interprocess communication of data needed to calculate

gval has already been done, and this data is accessible within f\_data.

Memory for u and gval is handled within the preconditioner module.

The case where G is mathematically identical to F is allowed.

#### ${\tt KINCommFn}$

Definition typedef void (\*KINCommFn)(long int Nlocal, N\_Vector u, void \*f\_data);

Purpose This function performs all interprocess communications necessary for the execution of

the gloc function above, using the input vector u.

Arguments Nlocal is the local vector length.

u is the current value of the iterate.

f\_data is a pointer to user data - the same as the f\_data parameter passed to KINSetFdata.

Return value A KINCommFn function type does not have a return value.

Notes The Gcomm function is expected to save communicated data in space defined within the structure f\_data.

Each call to the Gcomm function is preceded by a call to the system function func with the same u argument. Thus Gcomm can omit any communications done by func if relevant to the evaluation of Gloc. If all necessary communication was done in func, then Gcomm = NULL can be passed in the call to KINBBDPrecAlloc (see below).

Besides the header files required for the solution of a nonlinear problem (see §5.2), to use the KINBBDPRE module, the main program must include the header file kinbbdpre.h which declares the needed function prototypes.

The following is a summary of the usage of this module and describes the sequence of calls in the user main program. Steps that are unchanged from the user main program presented in §5.3 are graved out.

- 1. Initialize MPI
- 2. Set problem dimensions
- 3. Set vector with initial guess
- 4. Create KINSOL object
- 5. Set optional inputs
- 6. Allocate internal memory

#### 7. Initialize the KINBBDPRE preconditioner module

Specify the upper and lower half-bandwidths mu, ml and call

to allocate memory for and initialize a data structure bbd\_data to be passed to the KINSPGMR linear solver. The last two arguments of KINBBDPrecAlloc are the two user-supplied functions described above.

#### 8. Attach the KINSPGMR linear solver

```
flag = KINBBDSpgmr(kin_mem, maxl, bbd_data);
```

The function KINBBDSpgmr is a wrapper around the KINSPGMR specification function KINSpgmr and performs the following actions:

- •Attaches the KINSPGMR linear solver to the main CVODE solver memory;
- •Sets the preconditioner data structure for KINBBDPRE;
- •Sets the preconditioner setup function for KINBBDPRE;
- •Sets the preconditioner solve function for KINBBDPRE;

The argument maxl is described below. The last argument of KINBBDSpgmr is the pointer to the KINBBDPRE data returned by KINBBDPrecAlloc.

#### 9. Set linear solver optional inputs

Note that the user should not overwrite the preconditioner data, setup function, or solve function through calls to KINSPGMR optional input functions.

- 10. Solve problem
- 11. Get optional output
- 12. Deallocate memory for solution vector

#### 13. Free the KINBBDPRE data structure

KINBBDPrecFree(bbd\_data);

14. Free solver memory

#### 15. Finalize MPI

The three user-callable functions that initialize, attach, and deallocate the KINBBDPRE preconditioner module (steps 7, 8, and 13 above) are described next.

#### KINBBDPrecAlloc

Call bbd\_data = KINBBDPrecAlloc(kin\_mem, Nlocal, mu, ml, dq\_rel\_u, Gloc, Gcomm);

Description The function KINBBDPrecAlloc initializes and allocates memory for the KINBBDPRE preconditioner.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

Nlocal (long int) local vector length.

mu (long int) upper half-bandwidth to be used in the difference quotient Jacobian approximation.

ml (long int) lower half-bandwidth to be used in the difference quotient Jacobian approximation.

dq\_rel\_u (realtype) the relative increment in components of u used in the difference quotient approximations. The default is dq\_rel\_u= √unit roundoff, which can be specified by passing dq\_rel\_u= 0.0.

Gloc (KINLocalFn) the C function which computes the approximation  $G(u) \approx F(u)$ .

Gcomm (KINCommFn) the optional C function which performs all interprocess communication required for the computation of G(u).

Return value If successful, KINBBDPrecAlloc returns a pointer to the newly created KINBBDPRE memory block (of type void \*). If an error occurred, KINBBDPrecAlloc returns NULL.

Notes The half-bandwidths mu and ml need not be the true half-bandwidths of the Jacobian of the local block of G, when smaller values may provide a greater efficiency.

Moreover, the half-bandwidth values need not be the same for every process.

#### KINBBDSpgmr

Call flag = KINBBDSpgmr(kin\_mem, maxl, bbd\_data);

Description The function KINBBDSpgmr links the KINBBDPRE data to the KINSPGMR linear solver and attaches the latter to the KINSOL memory block.

Arguments kin\_mem (void \*) pointer to the KINSOL memory block.

maxl (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value KINSPGMR\_MAXL= 5.

bbd\_data (void \*) pointer to the KINBBDPRE data structure.

Return value The return value flag (of type int) is one of:

KINSPGMR\_SUCCESS The KINSPGMR initialization was successful.

KINSPGMR\_MEM\_NULL The kin\_mem pointer is NULL.

KINSPGMR\_ILL\_INPUT The NVECTOR module used does not implement a required operation.

KINSPGMR\_MEM\_FAIL A memory allocation request failed.

KIN\_PDATA\_NULL The KINBBDPRE preconditioner has not been initialized.

#### KINBBDPrecFree

Call KINBBDPrecFree(bbd\_data);

Description The function KINBBDPrecFree frees the pointer allocated by KINBBDPrecAlloc.

Arguments The only argument of KINBBDPrecFree is the pointer to the KINBBDPRE data structure

(of type void \*).

Return value The function KINBBDPrecFree has no return value.

The following two optional output functions are available for use with the KINBBDPRE module:

#### KINBBDPrecGetWorkSpace

Call flag = KINBBDPrecGetWorkSpace(bbd\_data, &lenrwBBDP, &leniwBBDP);

Description The function KINBBDPrecGetWorkSpace returns the local KINBBDPRE real and integer

workspace sizes.

Arguments bbd\_data (void \*) pointer to the KINBBDPRE data structure.

 $\verb|lenrwBBDP| (long int) local number of \verb|realtype| values in the \verb|KINBBDPRE| workspace.$ 

leniwBBDP (long int) local number of integer values in the KINBBDPRE workspace.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output values have been successfully set.

KIN\_PDATA\_NULL The KINBBDPRE preconditioner has not been initialized.

Notes In terms of the local vector dimension  $N_l$ , the actual size of the real workspace is  $N_l$  (2

ml + mu + smu + 2) realtype words, where  $smu = min(N_l - 1, mu + ml)$ .

The actual size of the integer workspace is  $N_l$  integer words.

#### KINBBDPrecGetNumGfnEvals

Call flag = KINBBDPrecGetNumGfnEvals(bbd\_data, &ngevalsBBDP);

Description The function KINBBDPrecGetNumGfnEvals returns the number of calls to the user

Gloc function due to the finite difference approximation of the Jacobian blocks used

within KINBBDPRE's preconditioner setup function.

Arguments bbd\_data (void \*) pointer to the KINBBDPRE data structure.

ngevalsBBDP (long int) the number of calls to the user Gloc function.

Return value The return value flag (of type int) is one of:

KIN\_SUCCESS The optional output value has been successfully set.

KIN\_PDATA\_NULL The KINBBDPRE preconditioner has not been initialized.

## 5.7 FKINSOL, a FORTRAN-C interface module

The FKINSOL interface module is a package of C functions which support the use of the KINSOL solver, for the solution nonlinear systems F(u) = 0, in a mixed FORTRAN/C setting. While KINSOL is written in C, it is assumed here that the user's calling program and user-supplied problem-defining routines are written in FORTRAN. This package provides the necessary interface to KINSOL for both the serial and the parallel NVECTOR implementations, and may also be used with user-supplied mixed FORTRAN/C NVECTOR implementations.

#### 5.7.1 FKINSOL routines

The user-callable functions, with the corresponding KINSOL functions, are as follows:

- Interface to the NVECTOR modules
  - FNVINITS (defined by NVECTOR\_SERIAL) interfaces to NV\_New\_Serial.
  - FNVINITP (defined by NVECTOR\_PARALLEL) interfaces to NV\_New\_Parallel.
  - FNVFREES (defined by NVECTOR\_SERIAL) interface to NV\_Destroy\_Serial.
  - FNVFREEP (defined by NVECTOR\_PARALLEL) interfaces to NV\_Destroy\_Parallel.
- Interface to the main KINSOL module
  - FKINMALLOC interfaces to KINCreate, KINSet\* functions, and KINMalloc.
  - FKINSOL interfaces to KINSol, KINGet\* functions, and to the optional output functions for the KINSPGMR linear solver module.
  - FKINFREE interfaces to KINFree.
- Interface to the KINSPGMR solver module
  - FKINSPGMR interfaces to KINSpgmr and SPGMR optional input functions.
  - FKINSPGMRSETJAC interfaces to KINSpgmrSetJacTimesVecFn.
  - FKINSPGMRSETPREC interfaces to KINSpgmrSetPrecSetupFn and KINSpgmrSetPrecSolveFn.

The user-supplied functions, each listed with the corresponding interface function which calls it (and its type within KINSOL), are as follows:

FKINSOL routine (FORTRAN)	KINSOL function (C)	KINSOL function type	
FKFUN	FKINfunc	KINSysFn	
FKPSOL	FKINPSol	KINSpgmrPrecSolveFn	
FKPSET	FKINPSet	KINSpgmrPrecSetupFn	
FKJTIMES	FKINJtimes	${\tt KINSpgmrJacTimesVecFn}$	

In contrast to the case of direct use of KINSOL, the names of all user-supplied routines here are fixed, in order to maximize portability for the resulting mixed-language program.

#### Important note on portability

In this package, the names of the interface functions, and the names of the FORTRAN user routines called by them, appear as dummy names which are mapped to actual values by a series of definitions in the header files fkinsol.h and fkinbbd.h. By default, those mapping definitions depend in turn on the C macro F77\_FUNC defined in the header file config.h by configure. However, the set of flags - SUNDIALS\_CASE\_UPPER, SUNDIALS\_CASE\_LOWER, SUNDIALS\_UNDERSCORE\_NONE, SUNDIALS\_UNDERSCORE\_ONE, and SUNDIALS\_UNDERSCORE\_TWO can be explicitly defined in config.h when configuring SUNDIALS via the --with-f77underscore and --with-f77case options to override the default behavior if necessary (see Chapter 2). Either way, the names into which the dummy names are mapped are in upper or lower case and have up to two underscores appended.

The user must also ensure that variables in the user FORTRAN code are declared in a manner consistent with their counterparts in KINSOL. All real variables must be declared as REAL, DOUBLE PRECISION, or perhaps as REAL\*n, where n denotes the number of bytes, depending on whether KINSOL was built in single, double or extended precision (see Chapter 2). Moreover, some of the FORTRAN integer variables must be declared as INTEGER\*4 or INTEGER\*8 according to the C type long int. These integer variables include: the array of integer optional inputs and outputs (IOPT), problem dimensions (NEQ, NLOCAL, NGLOBAL), and Jacobian half-bandwidths (MU and ML). This is particularly important when using KINSOL and the FKINSOL package on 64-bit architectures.

Table 5.3: Description of the FKINSOL optional input-output arrays IOPT and ROPT

Integer input-output array IOPT

Index	Optional input	Optional output	KINSOL function		
	KINSOL main solver				
1	PRINTFL		KINSetPrintLevel		
2	MXITER		KINSetNumMaxIters		
3	PRECOND_NO_INIT		KINSetNoPrecInit		
4		NNI	KINGetNumNonlinSolvIters		
5		NFE	KINGetNumFuncEvals		
6		NBCF	${\tt KINGetNumBetaCondFails}$		
7		NBKTRK	KINGetNumBacktrackOps		
8	ETACHOICE		KINSetEtaForm		
9	NO_MIN_EPS		KINSetNoMinEps		
		KINSPGMR linear solv	ver		
11		NLI	KINSpgmrGetNumLinIters		
12		NPE	KINSpgmrGetNumPrecEvals		
13		NPS	KINSpgmrGetNumPrecSolves		
14		NCFL	KINSpgmrGetNumConvFails		
15		LS_FLAG	KINSpgmrGetLastFlag		

Real input-output array ROPT

Index	Optional input	Optional output	KINSOL function	
1	MXNEWTSTEP		KINSetMaxNewtonStep	
2	RELFUNC	KINSetRelErrFunc		
3		FNORM KINGetFuncNorm		
4		STEPL KINGetStepLength		
5	ETACONST	KINSetEtaConstValue		
6	ETAGAMMA	KINSetEtaParams		
7	ETAALPHA		KINSetEtaParams	

#### 5.7.2 FKINSOL optional input and output

In order to keep the number of user-callable FKINSOL interface routines to a minimum, optional inputs and outputs to the KINSOL solver and to related modules are not accessed through individual functions, but rather through a pair of arrays, IOPT of integer type and ROPT of real type. Table 5.3 lists the entries in these two arrays and specifies the FKINSOL user-callable routine which sets/accesses the corresponding optional variable, as well as the KINSOL optional function which is actually called. For more details on the optional inputs and outputs, see §5.4.4 and §5.4.5.

#### 5.7.3 Usage of the FKINSOL interface module

The usage of FKINSOL requires calls to several interface functions, depending on the method options selected, and one or more user-supplied routines which define the problem to be solved. These function calls and user routines are summarized separately below. Some details are omitted, and the user is referred to the description of the corresponding KINSOL functions for information on the arguments of any given user-callable interface routine, or of a given user-supplied function called by an interface function. The usage of FKINSOL with the band-block-diagonal preconditioner module KINBBDPRE is described in the next subsection.

Steps marked with [S] in the instructions below apply to the serial NVECTOR implementation (NVECTOR\_SERIAL) only, while those marked with [P] apply to NVECTOR\_PARALLEL.

#### 1. Nonlinear system function specification

The user must in all cases supply the following FORTRAN routine

SUBROUTINE FKFUN(U, FVAL) DIMENSION U(\*), FVAL(\*)

It must set the FVAL array to F(u), the system function, as a function of the array U. Here U and FVAL are arrays representing vectors, which are distributed vectors in the parallel case.

#### 2. NVECTOR module initialization

[S] To initialize the serial NVECTOR module, the user must make the following call:

CALL FNVINITS (NEQ, IER)

where NEQ is the size of vectors and IER is a return completion flag which is set to 0 on success and -1 if a failure occurred.

[P] To initialize the parallel vector module, the user must make the following call:

CALL FNVINITP(NLOCAL, NGLOBAL, IER)

in which the arguments are: NLOCAL the local size of vectors for this process, NGLOBAL the system size (and the global size of vectors, that is the sum of all values of NLOCAL). The return completion flag IER is set to 0 upon successful return and to -1 otherwise. Note that if MPI was initialized by the user, the communicator must be set to MPI\_COMM\_WORLD. If not, this routine initializes MPI and sets the communicator equal to MPI\_COMM\_WORLD.

#### 3. Problem specification

To set various problem and solution parameters and allocate internal memory, make the following call:

DIZT	TARKE	$T \cap A$
HKI	${\sf NMAL}$	1 1111

Call	CALL FKINM	ALLOC(MSBPRE, FNORMTOL, SCSTEPTOL, CONSTRAINTS,
	&	OPTIN, IOPT, ROPT, IER)
Description		provides required problem and solution specifications, specifies opallocates internal memory, and initializes ${\tt KINSOL}.$
Arguments	MSBPRE	is the maximum number of preconditioning solve calls without calling the preconditioning setup routine. A value of $0$ indicates the default.
	FNORMTOL	is the tolerance on the scaled maximum norm of $F(u)$ to accept
		convergence.
	SCSTEPTOL	is the tolerance on minimum scaled step size.
	CONSTRAINTS	is an array of constraint values on the components of the solution
		u.
	OPTIN	is an integer flag indicating whether input values in IOPT and/or
		ROPT are to be used for input. A value of 0 means $no$ and a value of 1 indicates $yes$ .
	IOPT	is an array of integer optional inputs and outputs (must be declared
		as INTEGER*4 or INTEGER*8 according to the C type long int).
	ROPT	is an array of real optional inputs and outputs.
Return value	IER is the re	eturn completion flag. Its possible values are $0$ indicating success or

-1 indicating failure.

Notes

The optional inputs and outputs associated with the main KINSOL integrator are listed in Table 5.3. If any of the optional inputs are used, the others must be set to zero to indicate default values.

Since KINSOL maintains a private copy of the constraints vector, the CONSTRAINTS array passed as an argument to FKINMALLOC can be reused after the function call.

If OPTIN is set to 1, then FKINSOL will set the applicable optional outputs before returning, so the IOPT and ROPT arrays should not be reused.

#### 4. Linear solver specification

The solution method in KINSOL involves the solution of linear systems related to the Jacobian of the nonlinear system.

For the Scaled Preconditioned GMRES solution of the linear systems, the user must make the call:

```
CALL FKINSPGMR (MAXL, MAXLRST, IER)
```

The arguments are as follows. MAXL is the maximum Krylov subspace dimension (0 indicates default). MAXLRST is the maximum number of linear system restarts (0 indicates default). IER is the return completion flag (possible values are 0: success and -1: failure).

As an option when using the SPGMR linear solver, the user may supply a routine that computes the product of the system Jacobian  $J = \partial F/\partial u$  and a given vector v. If supplied, it must have the following form:

```
SUBROUTINE FKJTIMES(V, Z, NEWU, U, IER)
DIMENSION V(*), Z(*), U(*)
```

This must set the array Z to the product Jv, where J is the Jacobian matrix  $J=\partial F/\partial u$ , and V is a given array. Here U is an array containing the current value of the unknown vector u. NEWU is an input integer indicating whether U has changed since FKJTIMES was last called (1=yes,0=no). If FKJTIMES computes and saves Jacobian data, then no such computation is necessary when NEWU = 0. The arguments V, Z, and U are arrays of length NEQ, the problem size, or the local length of all distributed vectors in the parallel case. FKJTIMES should return IER = 0 if successful, or a nonzero IER otherwise.

If the user program includes the FKJTIMES routine for the evaluation of the Jacobian vector product, the following call must be made:

```
CALL FKINSPGMRSETJAC(FLAG, IER)
```

with  $FLAG \neq 0$  to specify use of the user-supplied Jacobian times vector approximation. The argument IER is an error return flag which can be 0 for success or nonzero if an error occurred.

If preconditioning is to be done, then, following the call to FKINSPGMR, the user must call

```
CALL FKINSPGMRSETPREC(FLAG, IER)
```

with  $FLAG \neq 0$ , and the user program must include the following routine for solution of the preconditioner linear system:

```
SUBROUTINE FKPSOL (U, USCALE, FVAL, FSCALE, VTEM, FTEM, IER)
DIMENSION U(*), USCALE(*), FVAL(*), FSCALE(*), VTEM(*), FTEM(*)
```

Typically this routine will use only U, FVAL, VTEM and FTEM. It must solve the preconditioned linear system Pz=r, where r= VTEM is input, and store the solution z in VTEM as well. Here P is the right preconditioner. If scaling is being used, the routine supplied must also account for scaling on either coordinate or function value, as given in the arrays USCALE and FSCALE, respectively.

If the user's preconditioner requires that any Jacobian-related data be evaluated or preprocessed, then the following routine can be used for the evaluation and preprocessing of the preconditioner:

```
SUBROUTINE FKPSET (U, USCALE, FVAL, FSCALE, VTEMP1, VTEMP2, IER)
DIMENSION U(*), USCALE(*), FVAL(*), FSCALE(*), VTEMP1(*), VTEMP2(*)
```

It must perform any evaluation of Jacobian-related data and preprocessing needed for the solution of the preconditioned linear systems by FKPSOL. The variables U through FSCALE are for use in the preconditioning setup process. Typically, the system function FKFUN is called before any calls to FKPSET, so that FVAL will have been updated. U is the current solution iterate. The arrays VTEMP1 and VTEMP2 are available for work space. If scaling is being used, USCALE and FSCALE are available for those operations requiring scaling. NEQ is the problem size.

On return, set IER = 0 if FKPSET was successful or set IER = 1 if an error occurred.

If the user calls FKINSPGMRSETPREC, the routine FKINPSET must be provided, even if it's empty.

5. **Problem solution** Solving the nonlinear system is accomplished by making the following call:

```
CALL FKINSOL(U, GLOBALSTRAT, USCALE, FSCALE, IER)
```

The arguments are as follows. U is an array containing the initial guess on input, and the solution on return. GLOBALSTRAT is an integer (type INTEGER) defining the global strategy choice (1 specifies Inexact Newton, while 2 indicates line search). USCALE is an array of scaling factors for the U vector. FSCALE is an array of scaling factors for the FVAL vector. IER is an integer completion flag and will have one of the following values: 0 to indicate success, 1 to indicate that the initial guess satisfies F(u) = 0 within tolerances, 2 to indicate apparent stalling (small step), or a negative value to indicate an error or failure. The possible negative return values and the corresponding KINSol return values (see §5.4.3) are: -1: KIN\_MEM\_NULL, -2: KIN\_ILL\_INPUT, -3: KIN\_NO\_MALLOC, -4: KIN\_MEM\_FAIL, -5: KIN\_LINESEARCH\_NONCONV, -6: KIN\_MAXITER\_REACHED, -7: KIN\_MXNEWT\_5X\_EXCEEDED, -8: KIN\_LINESEARCH\_BCFAIL, -9: KIN\_LINSOLV\_NO\_RECOVERY, -10: KIN\_LINIT\_FAIL, -11: KIN\_LSETUP\_FAIL, -12: KIN\_LSOLVE\_FAIL, and -13: KIN\_PDATA\_NULL.

The current values of the optional outputs are available in IOPT and ROPT (see Table 5.3).

6. **Memory deallocation** To free the internal memory created by the call to FKINMALLOC, make the call

```
CALL FKINFREE
```

and then, depending on the NVECTOR version (serial or parallel), either

CALL FNVFREES

or

CALL FNVFREEP

respectively.

#### 5.7.4 Usage of the FKINBBD interface to KINBBDPRE

The FKINBBD interface sub-module is a package of C functions which, as part of the FKINSOL interface module, support the use of the KINSOL solver with the parallel NVECTOR\_PARALLEL module and the KINBBDPRE preconditioner module (see §5.6), for the solution of nonlinear problems in a mixed FORTRAN/C setting.

The user-callable functions in this package, with the corresponding KINSOL and KINBBDPRE functions, are as follows:

- FKINBBDINIT interfaces to KINBBDPrecAlloc.
- FKINBBDSPGMR interfaces to KINBBDSpgmr and SPGMR optional input functions.
- FKINBBDOPT interfaces to KINBBDPRE optional output functions.
- FKINBBDFREE interfaces to KINBBDPrecFree.

In addition to the FORTRAN right-hand side function FKFUN, the user-supplied functions used by this package, are listed below, each with the corresponding interface function which calls it (and its type within KINBBDPRE or KINSOL):

FKINBBD routine (FORTRAN)	KINSOL function (C)	KINSOL function type
FKLOCFN	FKINgloc	KINLocalFn
FKCOMMF	FKINgcomm	KINCommFn
FKJTIMES	FKINJtimes	${\tt KINSpgmrJacTimesVecFn}$

As with the rest of the FKINSOL routines, the names of the user-supplied routines are mapped to actual values through a series of definitions in the header file fkinbbd.h (see §5.7).

The following is a summary of the usage of this module. Steps that are unchanged from the main program described in §5.7.3 are grayed out.

- 1. Nonlinear system function specification
- 2. NVECTOR module initialization
- 3. Problem specification

#### 4. Linear solver specification

To initialize the KINBBDPRE preconditioner, make the following call:

The arguments are as follows. NLOCAL is the local size of vectors for this process. MU and ML are the upper and lower half-bandwidths to be used in the computation of the local Jacobian blocks by difference quotients. These may be smaller than the true half-bandwidths of the Jacobian of the local block of G, when smaller values may provide greater efficiency. IER is a return completion flag. A value of 0 indicates success, while a value of -1 indicates that a memory failure occurred or that an input had an illegal value.

To specify the SPGMR linear system solver and use the KINBBDPRE preconditioner, make the following call:

```
CALL FKINBBDSPGMR (MAXL, MAXLRST, IER)
```

Its arguments are the same as those of FKINSPGMR (see step 4 in  $\S5.7.3$ ).

Optionally, to specify that SPGMR should use the supplied FKJTIMES, make the call

CALL FKINSPGMRSETJAC(FLAG, IER)

with FLAG  $\neq 0$ .

#### 5. Problem solution

#### 6. KINBBDPRE Optional outputs

To obtain the optional outputs associated with the KINBBDPRE module, make the following call:

CALL FKINBBDOPT(LENRPW, LENIPW, NGE)

The arguments returned are as follows. LENRPW is the length of real preconditioner work space, in realtype words. This size is local to the current process. LENIPW is the length of integer preconditioner work space, in integer words. This size is local to the current process. NGE is the cumulative number of G(u) evaluations (calls to FKLOCFN).

#### 7. Memory deallocation

To free the internal memory created by the call to FKINBBDINIT, before calling FKINFREE and FNVFREEP, the user must call

CALL FKINBBDFREE

# Chapter 6

# Description of the NVECTOR module

The SUNDIALS solvers are written in a data-independent manner. They all operate on generic vectors (of type N\_Vector) through a set of operations defined by the particular NVECTOR implementation. Users can provide their own specific implementation of the NVECTOR module or use one of two provided within SUNDIALS, a serial and an MPI parallel implementations.

The generic N\_Vector type is a pointer to a structure that has an implementation-dependent content field containing the description and actual data of the vector, and an ops field pointing to a structure with generic vector operations. The type N\_Vector is defined as

```
typedef struct _generic_N_Vector *N_Vector;
struct _generic_N_Vector {
    void *content;
    struct _generic_N_Vector_Ops *ops;
};
```

The \_generic\_N\_Vector\_Ops structure is essentially a list of pointers to the various actual vector operations, and is defined as

```
struct _generic_N_Vector_Ops {
 N_Vector
              (*nvclone)(N_Vector);
 N_Vector
              (*nvcloneempty)(N_Vector);
 void
              (*nvdestroy)(N_Vector);
              (*nvspace)(N_Vector, long int *, long int *);
  void
              (*nvgetarraypointer)(N_Vector);
 realtype*
              (*nvsetarraypointer)(realtype *, N_Vector);
  void
              (*nvlinearsum)(realtype, N_Vector, realtype, N_Vector, N_Vector);
  void
              (*nvconst)(realtype, N_Vector);
  void
              (*nvprod)(N_Vector, N_Vector, N_Vector);
  void
              (*nvdiv)(N_Vector, N_Vector, N_Vector);
  void
              (*nvscale)(realtype, N_Vector, N_Vector);
  void
  void
              (*nvabs)(N_Vector, N_Vector);
  void
              (*nvinv)(N_Vector, N_Vector);
              (*nvaddconst)(N_Vector, realtype, N_Vector);
 void
              (*nvdotprod)(N_Vector, N_Vector);
 realtype
 realtype
              (*nvmaxnorm)(N_Vector);
              (*nvwrmsnorm)(N_Vector, N_Vector);
 realtype
 realtype
              (*nvwrmsnormmask)(N_Vector, N_Vector, N_Vector);
 realtype
              (*nvmin)(N_Vector);
```

```
realtype (*nvwl2norm)(N_Vector, N_Vector);
realtype (*nvl1norm)(N_Vector);
void (*nvcompare)(realtype, N_Vector, N_Vector);
booleantype (*nvinvtest)(N_Vector, N_Vector);
booleantype (*nvconstrmask)(N_Vector, N_Vector, N_Vector);
realtype (*nvminquotient)(N_Vector, N_Vector);
};
```

The generic NVECTOR module also defines and implements the vector operations acting on N\_Vector. These routines are nothing but wrappers for the vector operations defined by a particular NVECTOR implementation, which are accessed through the *ops* field of the N\_Vector structure. To illustrate this point we show below the implementation of a typical vector operation from the generic NVECTOR module, namely N\_VScale, which performs the scaling of a vector x by a scalar c:

```
void N_VScale(realtype c, N_Vector x, N_Vector z)
{
   z->ops->nvscale(c, x, z);
}
```

Table 6.1 contains a complete list of all vector operations defined by the generic NVECTOR module. Finally, note that the generic NVECTOR module defines the functions N\_VCloneVectorArray and N\_VCloneEmptyVectorArray. Both functions create (by cloning) an array of count variables of type N\_Vector, each of the same type as an existing N\_Vector. Their prototypes are

```
N_Vector *N_VCloneVectorArray(int count, N_Vector w);
N_Vector *N_VCloneEmptyVectorArray(int count, N_Vector w);
```

and their definitions are based on the implementation-specific N\_VClone and N\_VCloneEmpty operations, respectively. An array of variables of type N\_Vector can be destroyed by calling N\_VDestroyVectorArray, whose prototype is

```
void N_VDestroyVectorArray(N_Vector *vs, int count);
```

and whose definition is based on the implementation-specific N\_VDestroy operation.

A particular implementation of the NVECTOR module must:

- Specify the *content* field of N\_Vector.
- Define and implement the vector operations. Note that the names of these routines should be unique to that implementation in order to permit using more than one NVECTOR module (each with different N\_Vector internal data representations) in the same code.
- Define and implement user-callable constructor and destructor routines to create and free an N\_Vector with the new *content* field and with *ops* pointing to the new vector operations.
- Optionally, define and implement additional user-callable routines acting on the newly defined N\_Vector (e.g., a routine to print the content for debugging purposes).
- Optionally, provide accessor macros as needed for that particular implementation to be used to access different parts in the *content* field of the newly defined N\_Vector.

Table 6.1: Description of the NVECTOR operations

Name	Usage and Description
N_VClone	<pre>v = N_VClone(w); Creates a new N_Vector of the same type as an existing vector w and sets the ops field. It does not copy the vector, but rather allocates storage for the new vector.</pre>
N_VCloneEmpty	<pre>v = N_VCloneEmpty(w); Creates a new N_Vector of the same type as an existing vector w and sets the ops field. It does not allocate storage for the data array.</pre>
N_VDestroy	N_VDestroy(v); Destroys the N_Vector v and frees memory allocated for its internal data.
N_VSpace	N_VSpace(nvSpec, &lrw, &liw); Returns storage requirements for one N_Vector. lrw contains the number of realtype words and liw contains the number of integer words.
N_VGetArrayPointer	<pre>vdata = N_VGetArrayPointer(v); Returns a pointer to a realtype array from the N_Vector v. Note that this assumes that the internal data in N_Vector is a contiguous array of realtype. This routine is only used in the solver-specific interfaces to the dense and banded linear solvers, as well as the interfaces to the banded preconditioners provided with SUNDIALS.</pre>
N_VSetArrayPointer	N_VSetArrayPointer(vdata, v); Overwrites the data in an N_Vector with a given array of realtype. Note that this assumes that the internal data in N_Vector is a contiguous array of realtype. This routine is only used in the interfaces to the dense linear solver.
N_VLinearSum	N_VLinearSum(a, x, b, y, z); Performs the operation $z = ax + by$ , where a and b are scalars and x and y are of type N_Vector: $z_i = ax_i + by_i$ , $i = 0, \ldots, n-1$ .
N_VConst	N_VConst(c, z); Sets all components of the N_Vector z to c: $z_i=c,\ i=0,\ldots,n-1.$
N_VProd	N_VProd(x, y, z); Sets the N_Vector z to be the component-wise product of the N_Vector inputs x and y: $z_i = x_i y_i$ , $i = 0, \ldots, n-1$ .
N_VDiv	N_VDiv(x, y, z); Sets the N_Vector z to be the component-wise ratio of the N_Vector inputs x and y: $z_i = x_i/y_i$ , $i = 0,, n-1$ . The $y_i$ may not be tested for 0 values. It should only be called with an x that is guaranteed to have all nonzero components. $continued on next page$

N_VScale	Usage and Description $ \begin{tabular}{ll} N_VScale(c, x, z); \\ Scales the N_Vector x by the scalar c and returns the result in z: \\ z_i = cx_i, \ i = 0, \dots, n-1. \\ N_VAbs(x, y); \\ Sets the components of the N_Vector y to be the absolute values of the components of the N_Vector x: \\ y_i =  x_i , \ i = 0, \dots, n-1. \\ \end{tabular} $
N_VAbs I	Scales the N_Vector x by the scalar c and returns the result in z: $z_i=cx_i,\ i=0,\dots,n-1.$ N_VAbs(x, y); Sets the components of the N_Vector y to be the absolute values of
1	Sets the components of the $N\_Vector$ y to be the absolute values of
N VTnv	
]	N_VInv(x, z); Sets the components of the N_Vector z to be the inverses of the components of the N_Vector x: $z_i = 1.0/x_i, i = 0, \ldots, n-1$ . This routine may not check for division by 0. It should be called only with an x which is guaranteed to have all nonzero components.
	N_VAddConst(x, b, z); Adds the scalar b to all components of x and returns the result in the N_Vector z: $z_i = x_i + b, \ i = 0, \dots, n-1$ .
	d = N_VDotProd(x, y); Returns the value of the ordinary dot product of x and y: $d = \sum_{i=0}^{n-1} x_i y_i$ .
	m = N_VMaxNorm(x); Returns the maximum norm of the N_Vector x: $m = \max_i  x_i $ .
	m = N_VWrmsNorm(x, w) Returns the weighted root-mean-square norm of the N_Vector x with weight vector w: $m = \sqrt{\left(\sum_{i=0}^{n-1} (x_i w_i)^2\right)/n}$ .
]	m = N_VWrmsNormMask(x, w, id); Returns the weighted root mean square norm of the N_Vector x with weight vector w built using only the elements of x corresponding to nonzero elements of the N_Vector id: $m = \sqrt{\left(\sum_{i=0}^{n-1} (x_i w_i \mathrm{sign}(id_i))^2\right)/n}.$
	m = N_VMin(x); Returns the smallest element of the N_Vector x: $m = \min_i x_i$ .
	m = N_VWL2Norm(x, w); Returns the weighted Euclidean $\ell_2$ norm of the N_Vector x with weight vector w: $m = \sqrt{\sum_{i=0}^{n-1} (x_i w_i)^2}$ .
	m = N_VL1Norm(x); Returns the $\ell_1$ norm of the N_Vector x: $m = \sum_{i=0}^{n-1}  x_i $ . continued on next page

continued from last page	
Name	Usage and Description
N_VCompare	N_VCompare(c, x, z); Compares the components of the N_Vector x to the scalar c and returns an N_Vector z such that: $z_i = 1.0$ if $ x_i  \ge c$ and $z_i = 0.0$ otherwise.
N_VInvTest	t = N_VInvTest(x, z); Sets the components of the N_Vector z to be the inverses of the components of the N_Vector x, with prior testing for zero values: $z_i = 1.0/x_i, i = 0, \ldots, n-1$ . This routine returns TRUE if all components of x are nonzero (successful inversion) and returns FALSE otherwise.
N_VConstrMask	t = N_VConstrMask(c, x, m); Performs the following constraint tests: $x_i > 0$ if $c_i = 2$ , $x_i \geq 0$ if $c_i = 1$ , $x_i \leq 0$ if $c_i = -1$ , $x_i < 0$ if $c_i = -2$ . There is no constraint on $x_i$ if $c_i = 0$ . This routine returns FALSE if any element failed the constraint test, TRUE if all passed. It also sets a mask vector m, with elements equal to 1.0 where the constraint test failed, and 0.0 where the test passed. This routine is used only for constraint checking.
N_VMinQuotient	$minq = N\_VMinQuotient(num, denom);$ This routine returns the minimum of the quotients obtained by termwise dividing $num_i$ by $denom_i$ . A zero element in $denom$ will be skipped. If no such quotients are found, then the large value BIG_REAL (defined in the header file sundialstypes.h) is returned.

## 6.1 The NVECTOR\_SERIAL implementation

The serial implementation of the NVECTOR module provided with SUNDIALS, NVECTOR\_SERIAL, defines the *content* field of N\_Vector to be a structure containing the length of the vector, a pointer to the beginning of a contiguous data array, and a boolean flag *own\_data* which specifies the ownership of *data*.

```
struct _N_VectorContent_Serial {
  long int length;
  booleantype own_data;
  realtype *data;
};
```

The following five macros are provided to access the content of an NVECTOR\_SERIAL vector. The suffix  $\mathcal{S}$  in the names denotes serial version.

#### • NV\_CONTENT\_S

This routine gives access to the contents of the serial vector N\_Vector.

The assignment  $v\_cont = NV\_CONTENT\_S(v)$  sets  $v\_cont$  to be a pointer to the serial  $N\_Vector$  content structure.

Implementation:

```
#define NV_CONTENT_S(v) ( (N_VectorContent_Serial)(v->content) )
```

#### • NV\_OWN\_DATA\_S, NV\_DATA\_S, NV\_LENGTH\_S

These macros give individual access to the parts of the content of a serial N\_Vector.

The assignment  $v_{data} = NV_DATA_S(v)$  sets  $v_{data}$  to be a pointer to the first component of the data for the  $N_V$ ctor v. The assignment  $NV_DATA_S(v) = v_{data}$  sets the component array of v to be  $v_{data}$  by storing the pointer  $v_{data}$ .

The assignment  $v_len = NV_LENGTH_S(v)$  sets  $v_len$  to be the length of v. On the other hand, the call  $NV_LENGTH_S(v) = len_v$  sets the length of v to be  $len_v$ .

Implementation:

```
#define NV_OWN_DATA_S(v) ( NV_CONTENT_S(v)->own_data )
#define NV_DATA_S(v) ( NV_CONTENT_S(v)->data )
#define NV_LENGTH_S(v) ( NV_CONTENT_S(v)->length )
```

#### • NV\_Ith\_S

This macro gives access to the individual components of the data array of an N\_Vector.

The assignment  $r = NV_i(v,i)$  sets r to be the value of the i-th component of v. The assignment  $NV_i(v,i) = r$  sets the value of the i-th component of v to be r.

Here i ranges from 0 to n-1 for a vector of length n.

Implementation:

```
#define NV_Ith_S(v,i) ( NV_DATA_S(v)[i] )
```

The NVECTOR\_SERIAL module defines serial implementations of all vector operations listed in Table 6.1 and provides the following user-callable routines:

#### • N\_VNew\_Serial

This function creates and allocates memory for a serial N\_Vector. Its only argument is the vector length.

N\_Vector N\_VNew\_Serial(long int vec\_length);

#### • N\_VNewEmpty\_Serial

This function creates a new serial N\_Vector with an empty (NULL) data array.

```
N_Vector N_VNewEmpty_Serial(long int vec_length);
```

#### • N\_VMake\_Serial

This function creates and allocates memory for a serial vector with user-provided data array. N\_Vector N\_VMake\_Serial(long int vec\_length, realtype \*v\_data);

#### • N\_VNewVectorArray\_Serial

This function creates an array of count serial vectors.

```
N_Vector *N_VNewVectorArray_Serial(int count, long int vec_length);
```

#### • N\_VNewVectorArrayEmpty\_Serial

This function creates an array of count serial vectors, each with an empty (NULL) data array. N\_Vector \*N\_VNewVectorArrayEmpty\_Serial(int count, long int vec\_length);

#### • N\_VDestroyVectorArray\_Serial

This function frees memory allocated for the array of count variables of type N\_Vector created with N\_VNewVectorArray\_Serial or with N\_VNewVectorArrayEmpty\_Serial.

```
void N_VDestroyVectorArray_Serial(N_Vector *vs, int count);
```

#### • N\_VPrint\_Serial

This function prints the content of a serial vector to stdout.

```
void N_VPrint_Serial(N_Vector v);
```

#### Notes

- When looping over the components of an N\_Vector v, it is more efficient to first obtain the component array via v\_data = NV\_DATA\_S(v) and then access v\_data[i] within the loop than it is to use NV\_Ith\_S(v,i) within the loop.
- I The NVECTOR\_SERIAL constructor functions N\_VNewEmpty\_Serial, N\_VMake\_Serial, and N\_VNewVectorArrayEmpty\_Serial set the field own\_data = FALSE. The functions N\_VDestroy\_Serial and N\_VDestroyVectorArray\_Serial will not attempt to free the pointer data for any N\_Vector with own\_data set to FALSE. In such a case, it is the user's responsibility to deallocate the data pointer.
- To maximize efficiency, vector operations in the NVECTOR\_SERIAL implementation that have more than one N\_Vector argument do not check for consistent internal representation of these vectors. It is the user's responsibility to ensure that such routines are called with N\_Vector arguments that were all created with the same internal representations.

## 6.2 The NVECTOR\_PARALLEL implementation

The parallel implementation of the NVECTOR module provided with SUNDIALS, NVECTOR\_PARALLEL, defines the *content* field of N\_Vector to be a structure containing the global and local lengths of the vector, a pointer to the beginning of a contiguous local data array, an MPI communicator, an a boolean flag own\_data indicating ownership of the data array data.

```
struct _N_VectorContent_Parallel {
  long int local_length;
  long int global_length;
  booleantype own_data;
  realtype *data;
  MPI_Comm comm;
};
```

The following seven macros are provided to access the content of a NVECTOR\_PARALLEL vector. The suffix \_P in the names denotes parallel version.

#### • NV\_CONTENT\_P

This macro gives access to the contents of the parallel vector N\_Vector.

The assignment v\_cont = NV\_CONTENT\_P(v) sets v\_cont to be a pointer to the N\_Vector content structure of type struct \_N\_VectorParallelContent.

Implementation:

```
#define NV_CONTENT_P(v) ( (N_VectorContent_Parallel)(v->content) )
```

• NV\_OWN\_DATA\_P, NV\_DATA\_P, NV\_LOCLENGTH\_P, NV\_GLOBLENGTH\_P

These macros give individual access to the parts of the content of a parallel N\_Vector.

The assignment v\_data = NV\_DATA\_P(v) sets v\_data to be a pointer to the first component of the local data for the N\_Vector v. The assignment NV\_DATA\_P(v) = v\_data sets the component array of v to be v\_data by storing the pointer v\_data.

The assignment v\_llen = NV\_LOCLENGTH\_P(v) sets v\_llen to be the length of the local part of v. The call NV\_LENGTH\_P(v) = llen\_v sets the local length of v to be llen\_v.

The assignment  $v_glen = NV_GLOBLENGTH_P(v)$  sets  $v_glen$  to be the global length of the vector v. The call  $NV_GLOBLENGTH_P(v) = glen_v$  sets the global length of v to be  $glen_v$ .

Implementation:

```
#define NV_OWN_DATA_P(v) ( NV_CONTENT_P(v)->own_data )
#define NV_DATA_P(v) ( NV_CONTENT_P(v)->data )
#define NV_LOCLENGTH_P(v) ( NV_CONTENT_P(v)->local_length )
#define NV_GLOBLENGTH_P(v) ( NV_CONTENT_P(v)->global_length )
```

#### • NV\_COMM\_P

This macro provides access to the MPI communicator used by the NVECTOR\_PARALLEL vectors.

Implementation:

```
#define NV_COMM_P(v) ( NV_CONTENT_P(v)->comm )
```

#### • NV\_Ith\_P

This macro gives access to the individual components of the local data array of an N\_Vector.

The assignment  $r = NV_i(v,i)$  sets r to be the value of the i-th component of the local part of v. The assignment  $NV_i(v,i) = r$  sets the value of the i-th component of the local part of v to be r.

Here i ranges from 0 to n-1, where n is the local length.

Implementation:

```
#define NV_Ith_P(v,i) ( NV_DATA_P(v)[i] )
```

The NVECTOR\_PARALLEL module defines parallel implementations of all vector operations listed in Table 6.1 and provides the following user-callable routines:

#### • N\_VNew\_Parallel

This function creates and allocates memory for a parallel vector.

#### • N\_VNewEmpty\_Parallel

This function creates a new parallel N\_Vector with an empty (NULL) data array.

#### • N\_VMake\_Parallel

This function creates and allocates memory for a parallel vector with user-provided data array.

#### • N\_VNewVectorArray\_Parallel

This function creates an array of count parallel vectors.

• N\_VNewVectorArrayEmpty\_Parallel

This function creates an array of count parallel vectors, each with an empty (NULL) data array.

• N\_VDestroyVectorArray\_Parallel

This function frees memory allocated for the array of count variables of type N\_Vector created with N\_VNewVectorArray\_Parallel or with N\_VNewVectorArray\_Empty\_Parallel.

```
void N_VDestroyVectorArray_Parallel(N_Vector *vs, int count);
```

• N\_VPrint\_Parallel

This function prints the content of a parallel vector to stdout.

```
void N_VPrint_Parallel(N_Vector v);
```

#### Notes

- When looping over the components of an N\_Vector v, it is more efficient to first obtain the local component array via v\_data = NV\_DATA\_P(v) and then access v\_data[i] within the loop than it is to use NV\_Ith\_P(v,i) within the loop.
- The NVECTOR\_PARALLEL constructor functions N\_VNewEmpty\_Parallel, N\_VMake\_Parallel, and N\_VNewVectorArrayEmpty\_Parallel set the field own\_data = FALSE. The functions N\_VDestroy\_Parallel and N\_VDestroyVectorArray\_Parallel will not attempt to free the pointer data for any N\_Vector with own\_data set to FALSE. In such a case, it is the user's responsibility to deallocate the data pointer.
- It is the user's responsability to ensure that such routines are called with N-Vector arguments that were all created with the same internal representations.

## 6.3 NVECTOR functions used by KINSOL

In Table 6.2 below, we list the vector functions in the NVECTOR module within the KINSOL package. The table also shows, for each function, which of the code modules uses the function. The KINSOL column shows function usage within the main solver module, the KINSPGMR column shows function usage within the linear solver, the KINBBDPRE column shows function usage within the band-block-diagonal preconditioner module, and the FKINSOL column shows function usage within the FKINSOL interface module.

There is one subtlety in the KINSPGMR column hidden by the table. The dot product function N\_VDotProd is called both within the implementation file kinspgmr.c for the KINSPGMR solver and within the implementation files spgmr.c and iterative.c for the generic SPGMR solver upon which the KINSPGMR solver is implemented.

At this point, we should emphasize that the KINSOL user does not need to know anything about the usage of vector functions by the KINSOL code modules in order to use KINSOL. The information is presented as an implementation detail for the interested reader.

The following vector operations listed in Table 6.1 are *not* used by KINSOL: N\_VAddConst, N\_VWrmsNorm, N\_VWrmsNormMask, N\_VCompare, N\_VCloneEmpty, and N\_VInvTest. Therefore a user-supplied NVECTOR module for KINSOL could omit these five functions.

Table 6.2: List of vector functions usage by KINSOL code modules

	KINSOL	KINSPGMR	KINBBDPRE	FKINSOL
N_VClone	<b>√</b>		<b>√</b>	<b>√</b>
N_VDestroy	<b>√</b>		<b>√</b>	<b>√</b>
N_VSpace	<b>√</b>			
N_VGetArrayPointer			<b>√</b>	<b>√</b>
N_VSetArrayPointer				<b>√</b>
N_VLinearSum	<b>√</b>	<b>√</b>		
N_VConst		<b>√</b>		
N_VProd	<b>√</b>	<b>√</b>		
N_VDiv	<b>√</b>			
N_VMinQuotient	<b>√</b>			
N_VScale	<b>√</b>	<b>√</b>	<b>√</b>	
N_VAbs	<b>√</b>			
N_VInv	<b>√</b>			
N_VDotProd		<b>√</b>		
N_VConstrMask	<b>√</b>			
N_VMaxNorm	<b>√</b>			
N_VL1Norm		<b>√</b>		
N_VWL2Norm	<b>√</b>	<b>√</b>		
N_VMin	<b>√</b>			

# Chapter 7

# Providing Alternate Linear Solver Modules

The central KINSOL module interfaces with the linear solver module by way of calls to four routines. These are denoted here by linit, lsetup, lsolve, and lfree. Briefly, their purposes are as follows:

- linit: initialize and allocate memory specific to the linear solver;
- lsetup: evaluate and preprocess the Jacobian or preconditioner;
- lsolve: solve the linear system;
- lfree: free the linear solver memory.

A linear solver module must also provide a user-callable specification routine (like that described in §5.4.2 for KINSPGMR) which will attach the above four routines to the main KINSOL memory block. Note that of the four interface routines, only the lsolve routine is required. The lfree routine must be provided only if the solver specification routine makes any memory allocation.

These four routines that interface between KINSOL and the linear solver module necessarily have fixed call sequences. Thus, a user wishing to implement another linear solver within the KINSOL package must adhere to this set of interfaces. The following is a complete description of the call list for each of these routines. Note that the call list of each routine includes a pointer to the main KINSOL memory block, by which the routine can access various data related to the KINSOL solution. The contents of this memory block are given in the file kinsol\_impl.h (but not reproduced here, for the sake of space).

**Initialization routine.** The type definition of limit is

linit

Definition int (\*linit)(KINMem kin\_mem);

Purpose The purpose of limit is to complete initializations for a specific linear solver, such as

counters and statistics.

Arguments kin\_mem is the KINSOL memory pointer of type KINMem.

Return value An linit function should return 0 if it has successfully initialized the KINSOL linear

solver and -1 otherwise.

Notes If an error does occur, an appropriate message should be sent to kin\_mem->kin\_errfp.

**Setup routine.** The type definition of lsetup is

lsetup

Definition int (\*lsetup)(KINMem kin\_mem);

Purpose The job of lsetup is to prepare the linear solver for subsequent calls to lsolve. It

may recompute Jacobian-related data if it deems necessary.

Arguments kin\_mem is the KINSOL memory pointer of type KINMem.

Return value The 1setup routine should return 0 if successful, a positive value for a recoverable

error, and a negative value for an unrecoverable error.

Solve routine. The type definition of lsolve is

lsolve

Definition int (\*lsolve)(KINMem kin\_mem, N\_Vector x,

N\_Vector b, realtype \*res\_norm);

Purpose The routine 1solve must solve the linear equation Jx = b, where  $J = \partial F/\partial u$  is

evaluated at the current iterate and the right-hand side vector b is input.

Arguments kin\_mem is the KINSOL memory pointer of type KINMem.

x is a vector set to an initial guess prior to calling lsolve. On return it should

contain the solution to Jx = b.

b is the right-hand side vector b, set to -F(u), evaluated at the current iterate.

res\_norm holds the value of the  $L_2$  norm of the residual vector upon return.

Return value lsolve returns a positive value for a recoverable error and a negative value for an

unrecoverable error. Success is indicated by a 0 return value.

Memory deallocation routine. The type definition of lfree is

lfree

Definition void (\*lfree)(KINMem kin\_mem);

Purpose The routine lfree should free any linear solver memory allocated by the linit routine.

Arguments kin\_mem is the KINSOL memory pointer of type KINMem.

Return value This routine has no return value.

Notes This routine is called once a problem has been completed and the linear solver is no

longer needed.

# Chapter 8

# Generic Linear Solvers in SUNDIALS

In this chapter, we describe two generic linear solver code modules that are included in SUNDIALS, but which are of potential use as generic packages in themselves, either in conjunction with the use of KINSOL or separately. These modules are:

- The DENSE matrix package, which includes functions for small dense matrices treated as simple array types.
- The SPGMR package, which includes a solver for the scaled preconditioned GMRES method.

The functions for small dense matrices are fully described here because we expect that they will be useful in the implementation of preconditioners used with the combination of KINSOL and the KINSPGMR solver.

#### 8.1 The DENSE module

#### 8.1.1 Type DenseMat

The type DenseMat is defined to be a pointer to a structure with a size and a data field:

```
typedef struct {
  long int size;
  realtype **data;
} *DenseMat;
```

The size field indicates the number of columns (which is the same as the number of rows) of a dense matrix, while the data field is a two dimensional array used for component storage. The elements of a dense matrix are stored columnwise (i.e columns are stored one on top of the other in memory). If A is of type DenseMat, then the (i,j)-th element of A (with  $0 \le i, j \le size-1$ ) is given by the expression (A->data)[j][i] or by the expression (A->data)[0][j\*size+i]. The macros below allow a user to efficiently access individual matrix elements without writing out explicit data structure references and without knowing too much about the underlying element storage. The only storage assumption needed is that elements are stored columnwise and that a pointer to the j-th column of elements can be obtained via the DENSE\_COL macro. Users should use these macros whenever possible.

#### 8.1.2 Accessor Macros

The following two macros are defined by the DENSE module to provide access to data in the DenseMat type:

#### • DENSE\_ELEM

```
Usage: DENSE_ELEM(A,i,j) = a_ij; or a_ij = DENSE_ELEM(A,i,j); 
DENSE_ELEM references the (i,j)-th element of the N \times N DenseMat A, 0 \le i, j \le N - 1.
```

• DENSE\_COL

```
Usage : col_j = DENSE_COL(A,j);
```

DENSE\_COL references the j-th column of the  $N \times N$  DenseMat A,  $0 \le j \le N-1$ . The type of the expression DENSE\_COL(A,j) is realtype \*. After the assignment in the usage above, col\_j may be treated as an array indexed from 0 to N-1. The (i, j)-th element of A is referenced by col\_j[i].

#### 8.1.3 Functions

The following functions for DenseMat matrices are available in the DENSE package. For full details, see the header file dense.h.

- DenseAllocMat: allocation of a DenseMat matrix;
- DenseAllocPiv: allocation of a pivot array for use with DenseFactor/DenseBacksolve;
- DenseFactor: LU factorization with partial pivoting;
- DenseBacksolve: solution of Ax = b using LU factorization;
- DenseZero: load a matrix with zeros;
- DenseCopy: copy one matrix to another;
- DenseScale: scale a matrix by a scalar;
- DenseAddI: increment a matrix by the identity matrix;
- DenseFreeMat: free memory for a DenseMat matrix;
- DenseFreePiv: free memory for a pivot array;
- DensePrint: print a DenseMat matrix to standard output.

#### 8.1.4 Small Dense Matrix Functions

The following functions for small dense matrices are available in the DENSE package:

#### • denalloc

denalloc(n) allocates storage for an n by n dense matrix. It returns a pointer to the newly allocated storage if successful. If the memory request cannot be satisfied, then denalloc returns NULL. The underlying type of the dense matrix returned is realtype\*\*. If we allocate a dense matrix realtype\*\* a by a = denalloc(n), then a[j][i] references the (i,j)-th element of the matrix a,  $0 \le i$ ,  $j \le n-1$ , and a[j] is a pointer to the first element in the j-th column of a. The location a[0] contains a pointer to  $n^2$  contiguous locations which contain the elements of a.

#### ullet denallocpiv

denallocpiv(n) allocates an array of n integers. It returns a pointer to the first element in the array if successful. It returns NULL if the memory request could not be satisfied.

#### • gefa

gefa(a,n,p) factors the n by n dense matrix a. It overwrites the elements of a with its LU factors and keeps track of the pivot rows chosen in the pivot array p.

A successful LU factorization leaves the matrix **a** and the pivot array **p** with the following information:

- 1. p[k] contains the row number of the pivot element chosen at the beginning of elimination step k, k = 0, 1, ..., n-1.
- 2. If the unique LU factorization of a is given by Pa = LU, where P is a permutation matrix, L is a lower triangular matrix with all 1's on the diagonal, and U is an upper triangular matrix, then the upper triangular part of a (including its diagonal) contains U and the strictly lower triangular part of a contains the multipliers, I L.

gefa returns 0 if successful. Otherwise it encountered a zero diagonal element during the factorization. In this case it returns the column index (numbered from one) at which it encountered the zero.

#### • gesl

gesl(a,n,p,b) solves the n by n linear system ax = b. It assumes that a has been LU-factored and the pivot array p has been set by a successful call to gefa(a,n,p). The solution x is written into the b array.

#### • denzero

denzero(a,n) sets all the elements of the n by n dense matrix a to be 0.0;

#### • dencopy

dencopy(a,b,n) copies the n by n dense matrix a into the n by n dense matrix b;

#### • denscale

denscale(c,a,n) scales every element in the n by n dense matrix a by c;

#### • denaddI

denaddI(a,n) increments the n by n dense matrix a by the identity matrix;

#### denfreepiv

denfreepiv(p) frees the pivot array p allocated by denallocpiv;

#### • denfree

denfree(a) frees the dense matrix a allocated by denalloc;

#### • denprint

denprint(a,n) prints the n by n dense matrix a to standard output as it would normally appear on paper. It is intended as a debugging tool with small values of n. The elements are printed using the %g option. A blank line is printed before and after the matrix.

#### 8.2 The SPGMR Module

The SPGMR package, in the files spgmr.h and spgmr.c, includes an implementation of the scaled preconditioned GMRES method. A separate code module, iterative.h and iterative.c, contains auxiliary functions that support SPGMR, and also other Krylov solvers to be added later. For full details, including usage instructions, see the files spgmr.h and iterative.h.

Functions. The following functions are available in the SPGMR package:

- SpgmrMalloc: allocation of memory for SpgmrSolve;
- SpgmrSolve: solution of Ax = b by the SPGMR method;
- SpgmrFree: free memory allocated by SpgmrMalloc.

The following functions are available in the support package iterative.h and iterative.c:

- ModifiedGS: performs modified Gram-Schmidt procedure;
- ClassicalGS: performs classical Gram-Schmidt procedure;
- QRfact: performs QR factorization of Hessenberg matrix;
- QRsol: solves a least squares problem with a Hessenberg matrix factored by QRfact.

# Chapter 9

## KINSOL Constants

Below we list all input and output constants used by the main solver and linear solver modules, together with their numerical values and a short description of their meaning.

## 9.1 KINSOL input constants

#### KINSOL main solver module

KIN\_ETACHOICE1 1 Use Eisenstat and Walker Choice 1 for  $\eta$ . KIN\_ETACHOICE2 2 Use Eisenstat and Walker Choice 2 for  $\eta$ .

KIN\_ETACONSTANT 3 Use constant value for  $\eta$ .

KIN\_INEXACT\_NEWTON 1 Use inexact Newton globalization.

KIN\_LINESEARCH 2 Use line search globalization.

#### Iterative linear solver module

PREC\_NONE 0 No preconditioning

PREC\_RIGHT 2 Preconditioning on the right.

MODIFIED\_GS 1 Use modified Gram-Schmidt procedure.

CLASSICAL\_GS 2 Use classical Gram-Schmidt procedure.

## 9.2 KINSOL output constants

#### KINSOL main solver module

KIN\_SUCCESS 0 Successful function return.

KIN\_INITIAL\_GUESS\_OK 1 The initial user-supplied guess already satisfies the stopping

criterion.

KIN\_STEP\_LT\_STPTOL 2 The stopping tolerance on scaled step length was satisfied.

KIN\_MEM\_NULL -1 The cvode\_mem argument was NULL.
KIN\_ILL\_INPUT -2 One of the function inputs is illegal.

KIN\_NO\_MALLOC -3 The KINSOL memory was not allocated by a call to KINMalloc.

KIN\_MEM\_FAIL -4 A memory allocation failed.

KIN\_LINESEARCH\_NONCONV -5 The line search algorithm was unable to find an iterate suffi-

ciently distinct from the current iterate.

KIN\_MAXITER\_REACHED -6 The maximum number of nonlinear iterations has been

reached.

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KIN_MXNEWT_5X_EXCEEDED	-7	Five consecutive steps have been taken that satisfy a scaled
WIN I INCOCADOU DOCATI	0	step length test.
KIN_LINESEARCH_BCFAIL	-8	The line search algorithm was unable to satisfy the $\beta$ -condition
		for nbcfails iterations.
KIN_LINSOLV_NO_RECOVERY	-9	The user-supplied routine preconditioner slve function failed
		recoverably, but the preconditioner is already current.
KIN_LINIT_FAIL	-10	The linear solver's initialization function failed.
KIN_LSETUP_FAIL	-11	The linear solver's setup function failed in an unrecoverable
		manner.
KIN_LSOLVE_FAIL	-12	The linear solver's solve function failed in an unrecoverable
		manner.
KIN_PDATA_NULL	-13	The preconditioner module has not been initialized.
		The preconditioner integrate new new seem integrated.

#### ${\tt KINSPGMR} \ \textbf{linear solver module}$

KINSPGMR_SUCCESS	0	Successful function return.
KINSPGMR_MEM_NULL	-1	The cvode_mem argument was NULL.
KINSPGMR_LMEM_NULL	-2	The KINSPGMR linear solver has not been initialized.
KINSPGMR_ILL_INPUT	-3	The KINSPGMR solver is not compatible with the current
		NVECTOR module.
KINSPGMR_MEM_FAIL	-4	A memory allocation request failed.

PGMR generic linear solver module			
SPGMR_SUCCESS	0	Converged.	
SPGMR_RES_REDUCED	1	No convergence, but the residual norm was reduced.	
SPGMR_CONV_FAIL	2	Failure to converge.	
SPGMR_QRFACT_FAIL	3	A singular matrix was found during the QR factorization.	
SPGMR_PSOLVE_FAIL_REC	4	The preconditioner solve function failed recoverably.	
SPGMR_MEM_NULL	-1	The SPGMR memory is NULL	
SPGMR_ATIMES_FAIL	-2	The Jacobian tims vector function failed.	
SPGMR_PSOLVE_FAIL_UNREC	-3	The preconditioner solve function failed unrecoverably.	
SPGMR_GS_FAIL	-4	Failure in the Gram-Schmidt procedure.	
SPGMR_QRSOL_FAIL	-5	The matrix $R$ was found to be singular during the QR solve	
		phase.	

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