# User Documentation for KINSOL v2.8.0 (SUNDIALS v2.6.0)

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## Chapter 1

# Introduction

KINSOL is part of a software family called SUNDIALS: SUite of Nonlinear and DIfferential/ALgebraic equation Solvers [14]. This suite consists of CVODE, ARKODE, KINSOL, and IDA, and variants of these with sensitivity analysis capabilities.

KINSOL is a general-purpose nonlinear system solver based on Newton-Krylov solver technology. A fixed point iteration is also included with the release of KINSOL v.2.8.0.

## 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 the solution of partial differential equations [5]. 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, avoiding 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 in 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 methods and tolerances, and has a more modular design to provide flexibility for future enhancements.

At present, KINSOL contains four Krylov methods: the GMRES (Generalized Minimal RESidual) [20], FGMRES (Flexible Generalized Minimal RESidual) [19], Bi-CGStab (Bi-Conjugate Gradient Stabilized) [21], and TFQMR (Transpose-Free Quasi-Minimal Residual) [13] linear iterative methods. As Krylov methods, these require almost no matrix storage as compared to direct methods. However, the algorithms allow for a user-supplied preconditioner matrix, and for most problems preconditioning is essential for an efficient solution. For very large nonlinear algebraic systems, the Krylov methods are preferable over direct linear solver methods, and are often the only feasible choice. Among the three Krylov methods in KINSOL, we recommend GMRES as the best overall choice. However, users are encouraged to compare all three, especially if encountering convergence failures with GMRES. Bi-CGStab and TFQMR have an advantage in storage requirements, in that the number of workspace vectors they require is fixed, while that number for GMRES depends on the desired Krylov subspace size.

For the sake of completeness in functionality, direct linear system solvers are included in KINSOL. These include methods for both dense and banded linear systems, with Jacobians that are either user-supplied or generated internally by difference quotients. KINSOL also includes interfaces to the sparse direct solvers KLU [7, 1], and the threaded sparse direct solver, SuperLU\_MT [16, 9, 2].

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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 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 abstraction 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. SUNDIALS (and thus KINSOL) is supplied with serial, MPI-parallel, and both openMP and Pthreads thread-parallel NVECTOR implementations.

There are several motivations for choosing the C language for KINSOL. First, a general movement away from FORTRAN and toward C in scientific computing was 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.

## 1.2 Changes from previous versions

## Changes in v2.8.0

Two major additions were made to the globalization strategy options (KINSol argument strategy). One is fixed-point iteration, and the other is Picard iteration. Both can be accelerated by use of the Anderson acceleration method. See the relevant paragraphs in Chapter 2.

Three additions were made to the linear system solvers that are available for use with the KINSOL solver. First, in the serial case, an interface to the sparse direct solver KLU was added. Second, an interface to SuperLU\_MT, the multi-threaded version of SuperLU, was added as a thread-parallel sparse direct solver option, to be used with the serial version of the NVECTOR module. As part of these additions, a sparse matrix (CSC format) structure was added to KINSOL. Finally, a variation of GMRES called Flexible GMRES was added.

Otherwise, only relatively minor modifications were made to KINSOL:

In function KINStop, two return values were corrected to make the values of uu and fval consistent.

A bug involving initialization of mxnewtstep was fixed. The error affects the case of repeated user calls to KINSol with no intervening call to KINSetMaxNewtonStep.

A bug in the increments for difference quotient Jacobian approximations was fixed in function kinDlsBandDQJac.

In KINLapackBand, the line smu = MIN(N-1,mu+ml) was changed to smu = mu + ml to correct an illegal input error for DGBTRF/DGBTRS.

In order to avoid possible name conflicts, the mathematical macro and function names MIN, MAX, SQR, RAbs, RSqrt, RExp, RPowerI, and RPowerR were changed to SUNMIN, SUNMAX, SUNSQR, SUNRabs, SUNRsqrt, SUNRexp, SRpowerI, and SUNRpowerR, respectively. These names occur in both the solver and in various example programs.

In the FKINSOL module, an incorrect return value ier in FKINfunc was fixed.

In the FKINSOL optional input routines FKINSETIIN, FKINSETRIN, and FKINSETVIN, the optional fourth argument key\_length was removed, with hardcoded key string lengths passed to all strncmp tests.

In all FKINSOL examples, integer declarations were revised so that those which must match a C type long int are declared INTEGER\*8, and a comment was added about the type match. All other integer declarations are just INTEGER. Corresponding minor corrections were made to the user guide.

Two new NVECTOR modules have been added for thread-parallel computing environments — one for openMP, denoted NVECTOR\_OPENMP, and one for Pthreads, denoted NVECTOR\_PTHREADS.

With this version of SUNDIALS, support and documentation of the Autotools mode of installation is being dropped, in favor of the CMake mode, which is considered more widely portable.

## Changes in v2.7.0

One significant design change was made with this release: The problem size and its relatives, bandwidth parameters, related internal indices, pivot arrays, and the optional output lsflag have all been changed from type int to type long int, except for the problem size and bandwidths in user calls to routines specifying BLAS/LAPACK routines for the dense/band linear solvers. The function NewIntArray is replaced by a pair NewIntArray/NewLintArray, for int and long int arrays, respectively.

A large number of errors have been fixed. Three major logic bugs were fixed – involving updating the solution vector, updating the linesearch parameter, and a missing error return. Three minor errors were fixed – involving setting etachoice in the Matlab/KINSOL interface, a missing error case in KINPrintInfo, and avoiding an exponential overflow in the evaluation of omega. In each linear solver interface function, the linear solver memory is freed on an error return, and the \*\*Free function now includes a line setting to NULL the main memory pointer to the linear solver memory. In the installation files, we modified the treatment of the macro SUNDIALS\_USE\_GENERIC\_MATH, so that the parameter GENERIC\_MATH\_LIB is either defined (with no value) or not defined.

## Changes in v2.6.0

This release introduces a new linear solver module, based on Blas and Lapack for both dense and banded matrices.

The user interface has been further refined. Some of the API changes involve: (a) a reorganization of all linear solver modules into two families (besides the already present family of scaled preconditioned iterative linear solvers, the direct solvers, including the new Lapack-based ones, were also organized into a *direct* family); (b) maintaining a single pointer to user data, optionally specified through a Set-type function; (c) a general streamlining of the band-block-diagonal preconditioner module distributed with the solver.

## Changes in v2.5.0

The main changes in this release involve a rearrangement of the entire SUNDIALS source tree (see §3.1). At the user interface level, the main impact is in the mechanism of including SUNDIALS header files which must now include the relative path (e.g. #include <cvode/cvode.h>). Additional changes were made to the build system: all exported header files are now installed in separate subdirectories of the installation *include* directory.

The functions in the generic dense linear solver (sundials\_dense and sundials\_smalldense) were modified to work for rectangular  $m \times n$  matrices ( $m \le n$ ), while the factorization and solution functions were renamed to DenseGETRF/denGETRF and DenseGETRS/denGETRS, respectively. The factorization and solution functions in the generic band linear solver were renamed BandGBTRF and BandGBTRS, respectively.

## Changes in v2.4.0

KINSPBCG, KINSPTFQMR, KINDENSE, and KINBAND modules have been added to interface with the Scaled Preconditioned Bi-CGStab (SPBCG), Scaled Preconditioned Transpose-Free Quasi-Minimal Residual (SPTFQMR), DENSE, and BAND linear solver modules, respectively. (For details see Chapter 4.) Corresponding additions were made to the FORTRAN interface module FKINSOL. At the same time, function type names for Scaled Preconditioned Iterative Linear Solvers were added for the user-supplied Jacobian-times-vector and preconditioner setup and solve functions.

Regarding the FORTRAN interface module FKINSOL, optional inputs are now set using FKINSETIIN (integer inputs), FKINSETRIN (real inputs), and FKINSETVIN (vector inputs). Optional outputs are still obtained from the IOUT and ROUT arrays which are owned by the user and passed as arguments to FKINMALLOC.

The KINDENSE and KINBAND linear solver modules include support for nonlinear residual monitoring which can be used to control Jacobian updating.

4 Introduction

To reduce the possibility of conflicts, the names of all header files have been changed by adding unique prefixes (kinsol\_ and sundials\_). When using the default installation procedure, the header files are exported under various subdirectories of the target include directory. For more details see Appendix A.

## 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 system 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 4.

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 Get-type functions.

Installation of KINSOL (and all of SUNDIALS) has been completely redesigned and is now based on configure scripts.

## 1.3 Reading this User Guide

This user guide is a combination of general usage instructions and specific examples. We expect that some readers will want to concentrate on the general instructions, while others will refer mostly to the examples, and the organization is intended to accommodate both styles.

There are different possible levels of usage of KINSOL. The most casual user, with a small nonlinear system, can get by with reading all of Chapter 2, then Chapter 4 through §4.5.3 only, and looking at examples in [6]. In a different direction, a more expert user with a nonlinear system may want to (a) use a package preconditioner (§4.7), (b) supply his/her own Jacobian or preconditioner routines (§4.6), (c) supply a new NVECTOR module (Chapter 6), or even (d) supply a different linear solver module (§3.2 and Chapter 7).

The structure of this document is as follows:

- In Chapter 2, 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 (§3.1) and the software organization of the KINSOL solver (§3.2).
- Chapter 4 is the main usage document for KINSOL for C applications. It includes a complete description of the user interface for the solution of nonlinear algebraic systems.
- In Chapter 5, we describe FKINSOL, an interface module for the use of KINSOL with FORTRAN applications.

- Chapter 6 gives a brief overview of the generic NVECTOR module shared among the various components of SUNDIALS, and details on the four NVECTOR implementations provided with SUNDIALS: a serial implementation (§6.1), a distributed memory parallel implementation based on MPI (§6.2), and two thread-parallel implementations based on openMP (§6.3) and Pthreads (§6.4), respectively.
- Chapter 7 describes the interfaces to the linear solver modules, so that a user can provide his/her own such module.
- Chapter 8 describes in detail the generic linear solvers shared by all SUNDIALS solvers.
- Finally, in the appendices, we provide detailed instructions for the installation of KINSOL, within the structure of SUNDIALS (Appendix A), as well as a list of all the constants used for input to and output from KINSOL functions (Appendix B).

Finally, the reader should be aware of the following notational conventions in this user guide: program listings and identifiers (such as KINInit) 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. Usage and installation instructions that constitute important warnings are marked with a triangular symbol in the margin.



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

## Chapter 2

## Mathematical Considerations

KINSOL solves nonlinear algebraic systems in real N-space.

Using Newton's method, or the Picard iteration, one can solve

$$F(u) = 0, \quad F: \mathbf{R}^N \to \mathbf{R}^N,$$
 (2.1)

given an initial guess  $u_0$ . Using a fixed-point iteration, the convergence of which can be improved with Anderson acceleration, one can solve

$$G(u) = u, \quad G: \mathbf{R}^N \to \mathbf{R}^N,$$
 (2.2)

given an initial guess  $u_0$ .

#### **Basic Newton iteration**

Depending on the linear solver used, KINSOL can employ either an Inexact Newton method [4, 5, 8, 10, 15], or a Modified Newton method. At the highest level, KINSOL implements the following iteration scheme:

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

Here,  $u_n$  is the *n*th iterate to u, and J(u) = F'(u) is the system Jacobian. At each stage in the iteration process, a scalar multiple of the step  $\delta_n$ , is added to  $u_n$  to produce a new iterate,  $u_{n+1}$ . A test for convergence is made before the iteration continues.

#### Newton method variants

For solving the linear system given in step 2(a), KINSOL provides several choices, including the option of a user-supplied linear solver module. The linear solver modules distributed with SUNDIALS are organized in three families, a *direct* family comprising direct linear solvers for dense or banded matrices, a *sparse* family comprising direct linear solvers for matrices stored in compressed-sparse-column format, and a *spils* family comprising scaled preconditioned iterative (Krylov) linear solvers. The methods offered through these modules are as follows:

• dense direct solvers, using either an internal implementation or a Blas/Lapack implementation (serial or threaded vector modules only),

- band direct solvers, using either an internal implementation or a Blas/Lapack implementation (serial or threaded vector modules only),
- sparse direct solver interfaces, using either the KLU sparse solver library [7, 1], or the threadenabled SuperLU\_MT sparse solver library [16, 9, 2] (serial or threaded vector modules only) [Note that users will need to download and install the KLU or SuperLU\_MT packages independent of KINSOL],
- SPGMR, a scaled preconditioned GMRES (Generalized Minimal Residual method) solver without restarts,
- SPFGMR, a scaled preconditioned FGMRES (Flexible Generalized Minimal Residual method) solver without restarts,
- SPBCG, a scaled preconditioned Bi-CGStab (Bi-Conjugate Gradient Stable method) solver, or
- SPTFQMR, a scaled preconditioned TFQMR (Transpose-Free Quasi-Minimal Residual method) solver.

When using one of the direct linear solvers, the linear system in 2(a) is solved exactly, thus resulting in a Modified Newton method (the Jacobian matrix is normally out of date; see below<sup>1</sup>). Note that the direct linear solvers (dense, band, and sparse) can only be used with the serial and threaded vector representations.

On the other hand, when using any of the iterative linear solvers (GMRES, FGMRES, Bi-CGStab, or TFQMR), the linear system in 2(a) is solved only approximately, thus resulting in an Inexact Newton method. Here right preconditioning is available by way of the preconditioning setup and solve routines supplied by the user, in which case the iterative method is applied to the linear systems  $(JP^{-1})(P\delta) = -F$ , where P denotes the right preconditioning matrix.

## Jacobian information update strategy

In general, unless specified otherwise by the user, KINSOL strives to update Jacobian information (the actual system Jacobian J in the case of direct linear solvers, or the preconditioner matrix P in the case of iterative linear solvers) as infrequently as possible to balance the high costs of matrix operations against other costs. Specifically, these updates occur when:

- the problem is initialized,
- $\|\lambda \delta_{n-1}\|_{D_{u,\infty}} > 1.5$  (Inexact Newton only),
- mbset= 10 nonlinear iterations have passed since the last update,
- the linear solver failed recoverably with outdated Jacobian information,
- the global strategy failed with outdated Jacobian information, or
- $\|\lambda \delta_n\|_{D_{u,\infty}} < \text{STEPTOL}$  with outdated Jacobian information.

KINSOL allows, through optional solver inputs, changes to the above strategy. Indeed, the user can disable the initial Jacobian information evaluation or change the default value of mbset, the number of nonlinear iterations after which a Jacobian information update is enforced.

 $<sup>^{1}</sup>$ KINSOL allows the user to enforce a Jacobian evaluation at each iteration thus allowing for an Exact Newton iteration.

## Scaling

To address the case of ill-conditioned nonlinear systems, KINSOL allows prescribing scaling factors both for the solution vector and for the residual 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:

$$||z||_{D_u} = ||D_u z||_2, \quad ||z||_{D_F} = ||D_F z||_2, \quad ||z||_{D_u,\infty} = ||D_u z||_\infty, \quad \text{and} \quad ||z||_{D_F,\infty} = ||D_F z||_\infty$$
 (2.3)

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 the perturbations used for the default difference quotient approximations for Jacobian information; see (2.7) and (2.9) below.

### Globalization strategy

Two methods of applying a computed step  $\delta_n$  to the previously computed solution vector are implemented. The first and simplest is the standard 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 [10] 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.

KINSOL implements a backtracking algorithm to first find the value  $\lambda$  such that  $u_n + \lambda \delta_n$  satisfies the sufficient decrease condition (or  $\alpha$ -condition)

$$F(u_n + \lambda \delta_n) \leq F(u_n) + \alpha \nabla F(u_n) \lambda \delta_n$$

where  $\alpha = 10^{-4}$ . Although backtracking in itself guarantees that the step is not too small, KINSOL secondly relaxes  $\lambda$  to satisfy the so-called  $\beta$ -condition (equivalent to Wolfe's curvature condition):

$$F(u_n + \lambda \delta_n) \ge F(u_n) + \beta \nabla F(u_n) \lambda \delta_n$$
,

where  $\beta = 0.9$ . During this second phase,  $\lambda$  is allowed to vary in the interval  $[\lambda_{min}, \lambda_{max}]$  where

$$\lambda_{min} = \frac{\text{STEPTOL}}{\|\bar{\delta}_n\|_{\infty}}, \quad \bar{\delta}_n^j = \frac{\delta_n^j}{1/D_u^j + |u^j|},$$

and  $\lambda_{max}$  corresponds to the maximum feasible step size at the current iteration (typically  $\lambda_{max} = \text{STEPMAX}/\|\delta_n\|_{D_n}$ ). In the above expressions,  $v^j$  denotes the jth component of a vector v.

For more details, the reader is referred to [10].

#### Nonlinear iteration stopping criteria

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  $U^{1/3}$ . Here U is the machine unit roundoff. For the latter, the Newton method will terminate when the maximum scaled step is below a given tolerance

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

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.

#### Additional constraints

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 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.

## Residual monitoring for Modified Newton method

When using a Modified Newton method (i.e. when a direct linear solver is used), in addition to the strategy described above for the update of the Jacobian matrix, KINSOL also provides an optional nonlinear residual monitoring scheme to control when the system Jacobian is updated. Specifically, a Jacobian update will also occur when mbsetsub= 5 nonlinear iterations have passed since the last update and

$$||F(u_n)||_{D_E} > \omega ||F(u_m)||_{D_E}$$

where  $u_n$  is the current iterate and  $u_m$  is the iterate at the last Jacobian update. The scalar  $\omega$  is given by

$$\omega = \min\left(\omega_{min} e^{\max(0, \rho - 1)}, \omega_{max}\right), \qquad (2.4)$$

with  $\rho$  defined as

$$\rho = \frac{\|F(u_n)\|_{D_F}}{\text{FTOL}}, \qquad (2.5)$$

where FTOL is the input scalar tolerance discussed before. Optionally, a constant value  $\omega_{const}$  can be used for the parameter  $\omega$ .

The constants controlling the nonlinear residual monitoring algorithm can be changed from their default values through optional inputs to KINSOL. These include the parameters  $\omega_{min}$  and  $\omega_{max}$ , the constant value  $\omega_{const}$ , and the threshold mbsetsub.

### Stopping criteria for iterative linear solvers

When using an Inexact Newton method (i.e. when an iterative linear solver is used), the convergence of the overall nonlinear solver is intimately coupled with the accuracy with which the linear solver in 2(a) above is solved. KINSOL provides three options for stopping criteria for the linear system solver, including the two algorithms of Eisenstat and Walker [11]. More precisely, 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.

#### Constant $\eta$

$$\eta_n = constant,$$

with 0.1 as the default.

The default strategy 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 [11].

## Difference quotient Jacobian approximations

With the direct dense and band methods, the Jacobian may be supplied by a user routine, or approximated by difference quotients, at the user's option. In the latter case, we use the usual approximation

$$J^{ij} = [F^i(u + \sigma_j e^j) - F^i(u)]/\sigma_j.$$
 (2.6)

The increments  $\sigma_i$  are given by

$$\sigma_j = \sqrt{U} \max\left\{ |u^j|, 1/D_u^j \right\}. \tag{2.7}$$

In the dense case, this scheme requires N evaluations of F, one for each column of J. In the band case, the columns of J are computed in groups, by the Curtis-Powell-Reid algorithm, with the number of F evaluations equal to the bandwidth.

We note that with the sparse direct solvers, the Jacobian *must* be supplied by a user routine in compressed-sparse-column format, i.e. it is not approximated internally within KINSOL.

In the case of a Krylov method, Jacobian information is needed only as matrix-vector products Jv. If a routine for Jv is not supplied, these products are approximated by directional difference quotients as

$$J(u)v \approx [F(u+\sigma v) - F(u)]/\sigma, \qquad (2.8)$$

where u is the current approximation to a root of (2.1), and  $\sigma$  is a scalar. The choice of  $\sigma$  is taken from [5] and is given by

$$\sigma = \frac{\max\{|u^T v|, u_{typ}^T | v|\}}{\|v\|_2} \operatorname{sign}(u^T v) \sqrt{U}, \qquad (2.9)$$

where  $u_{typ}$  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). Convergence of the Newton method is maintained as long as the value of  $\sigma$  remains appropriately small, as shown in [4].

#### **Basic Fixed Point iteration**

The basic fixed-point iteration scheme implemented in KINSOL is given by:

- 1. Set  $u_0 = \text{an initial guess}$
- 2. For  $n = 0, 1, 2, \dots$  until convergence do:
  - (a) Set  $u_{n+1} = G(u_n)$ .
  - (b) Test for convergence.

Here,  $u_n$  is the *n*th iterate to u. At each stage in the iteration process, function G is applied to the current iterate to produce a new iterate,  $u_{n+1}$ . A test for convergence is made before the iteration continues.

For Picard iteration, as implemented in KINSOL, we consider a special form of the nonlinear function F, such that F(u) = Lu - N(u), where L is a constant nonsingular matrix and N is (in general) nonlinear. Then the fixed-point function G is defined as  $G(u) = u - L^{-1}F(u)$ . The Picard iteration is given by:

- 1. Set  $u_0 = \text{an initial guess}$
- 2. For  $n = 0, 1, 2, \dots$  until convergence do:
  - (a) Set  $u_{n+1} = G(u_n) = u_n L^{-1}F(u_n)$ .
  - (b) Test  $F(u_{n+1})$  for convergence.

Here,  $u_n$  is the *n*th iterate to u. Within each iteration, the Picard step is computed then added to  $u_n$  to produce the new iterate. Next, the nonlinear residual function is evaluated at the new iterate, and convergence is checked. Noting that  $L^{-1}N(u) = u - L^{-1}F(u)$ , the above iteration can be written in the same form as a Newton iteration except that here, L is in the role of the Jacobian. Within KINSOL, however, we leave this in a fixed-point form as above. For more information, see p. 182 of [18].

#### **Anderson Acceleration**

The Picard and fixed point methods can be significantly accelerated using Anderson's method [3, 22, 12, 17]. Anderson acceleration can be formulated as follows:

- 1. Set  $u_0 =$  an initial guess and  $m \ge 1$
- 2. Set  $u_1 = G(u_0)$
- 3. For  $n = 0, 1, 2, \dots$  until convergence do:
  - (a) Set  $m_n = \min\{m, n\}$
  - (b) Set  $F_n = (f_{n-m_n}, \dots, f_n)$ , where  $f_i = G(u_i) u_i$
  - (c) Determine  $\alpha^{(n)}=(\alpha_0^{(n)},\dots,\alpha_{m_n}^{(n)})$  that solves  $\min_{\alpha}\|F_n\alpha^T\|_2$  such that  $\sum_{i=0}^{m_n}\alpha_i=1$
  - (d) Set  $u_{n+1} = \sum_{i=0}^{m_n} \alpha_i^{(n)} G(u_{n-m_n+i})$
  - (e) Test for convergence

It has been implemented in KINSOL by turning the constrained linear least-squares problem in Step (c) into an unconstrained one leading to the algorithm given below:

- 1. Set  $u_0 =$  an initial guess and  $m \ge 1$
- 2. Set  $u_1 = G(u_0)$
- 3. For  $n = 0, 1, 2, \dots$  until convergence do:
  - (a) Set  $m_n = \min\{m, n\}$
  - (b) Set  $\Delta F_n = (\Delta f_{n-m_n}, \dots, \Delta f_{n-1})$ , where  $\Delta f_i = f_{i+1} f_i$  and  $f_i = G(u_i) u_i$
  - (c) Determine  $\gamma^{(n)} = (\gamma_0^{(n)}, \dots, \gamma_{m_n-1}^{(n)})$  that solves  $\min_{\gamma} \|f_n \Delta F_n \gamma^T\|_2$
  - (d) Set  $u_{n+1} = G(u_n) \sum_{i=0}^{m_n-1} \gamma_i^{(n)} \Delta g_{n-m_n+i}$  with  $\Delta g_i = G(u_{i+1}) G(u_i)$
  - (e) Test for convergence

The least-squares problem in (c) is solved by applying a QR factorization to  $\Delta F_n = Q_n R_n$  and solving  $R_n \gamma = Q_n^T f_n$ .

## Fixed-point - Anderson Acceleration Stopping Criterion

The default stopping criterion is

$$||G(u_{n+1}) - u_{n+1}||_{D_{F,\infty}} < \text{GTOL},$$

where  $D_F$  is a user-defined diagonal matrix that can be the identity or a scaling matrix chosen so that the components of  $D_F(G(u) - u)$  have roughly the same order of magnitude. Note that when using Anderson acceleration, convergence is checked after the acceleration is applied.

## Picard - Anderson Acceleration Stopping Criterion

The default stopping criterion is

$$||F(u_{n+1})||_{D_F,\infty} < \text{FTOL},$$

where  $D_F$  is a user-defined diagonal matrix that can be the identity or a scaling matrix chosen so that the components of  $D_F F(u)$  have roughly the same order of magnitude. Note that when using Anderson acceleration, convergence is checked after the acceleration is applied.

## Chapter 3

# **Code Organization**

## 3.1 SUNDIALS organization

The family of solvers referred to as SUNDIALS consists of the solvers CVODE and ARKODE (for ODE systems), KINSOL (for nonlinear algebraic systems), and IDA (for differential-algebraic systems). In addition, SUNDIALS also includes variants of CVODE and IDA with sensitivity analysis capabilities (using either forward or adjoint methods), called CVODES and IDAS, respectively.

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. 3.1). The following is a list of the solver packages presently available, and the basic functionality of each:

- CVODE, a solver for stiff and nonstiff ODE systems dy/dt = f(t,y) based on Adams and BDF methods:
- CVODES, a solver for stiff and nonstiff ODE systems with sensitivity analysis capabilities;
- ARKODE, a solver for ODE systems Mdy/dt = f(t, y) based on additive Runge-Kutta methods;
- IDA, a solver for differential-algebraic systems  $F(t, y, \dot{y}) = 0$  based on BDF methods;
- IDAS, a solver for differential-algebraic systems with sensitivity analysis capabilities;
- KINSOL, a solver for nonlinear algebraic systems F(u) = 0.

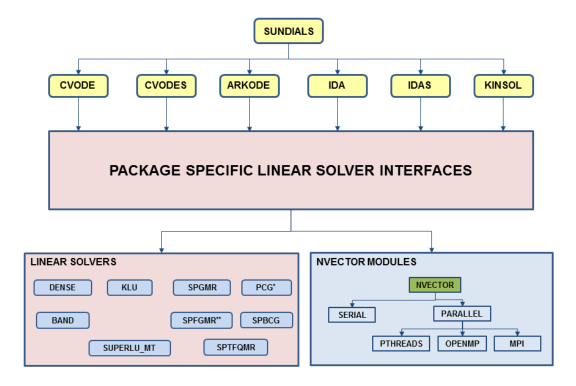
## 3.2 KINSOL organization

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.

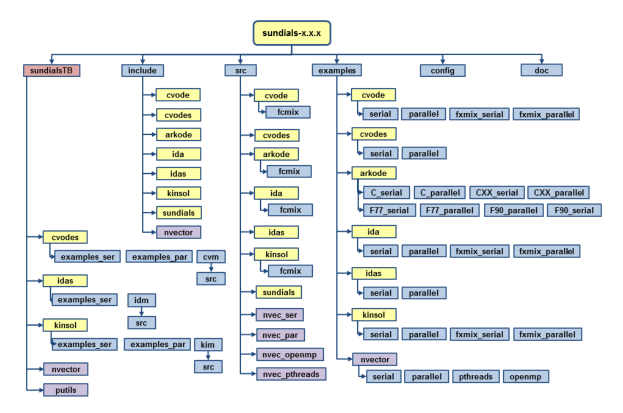
The overall organization of the KINSOL package is shown in Figure 3.2. The central solver module, implemented in the files kinsol.h, kinsol\_impl.h and kinsol.c, deals with the solution of a nonlinear algebraic system using either an Inexact Newton method or a line search method for the global strategy. 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, one of the linear system modules is specified, and is then invoked as needed.

At present, the package includes the following seven KINSOL linear algebra modules, organized into two families. The *direct* family of linear solvers provides solvers for the direct solution of linear systems with dense, banded, or sparse matrices and includes:

- KINDENSE: LU factorization and backsolving with dense matrices (using either an internal implementation or Blas/Lapack);
- KINBAND: LU factorization and backsolving with banded matrices (using either an internal implementation or Blas/Lapack);



- (a) High-level diagram (note that none of the Lapack-based linear solver modules are represented.)
  - \* only applies to ARKODE
  - \*\* only applies to ARKODE and KINSOL



(b) Directory structure of the source tree

Figure 3.1: Organization of the SUNDIALS suite

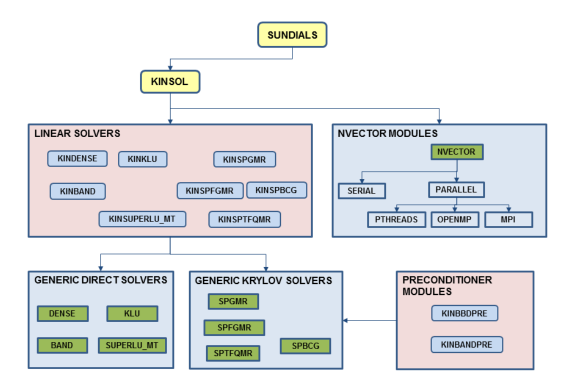


Figure 3.2: Overall structure diagram of the KINSOL package. Modules specific to KINSOL are distinguished by rounded boxes, while generic solver and auxiliary modules are in rectangular boxes. Grayed boxes refer to the encompassing SUNDIALS structure. Note that the direct linear solvers using Lapack implementations are not explicitly represented. Note also that the KLU and SuperLU\_MT support is through interfaces to packages. Users will need to download and compile those packages independently.

- KINKLU: LU factorization and backsolving with compressed-sparse-column (CSC) matrices using the KLU linear solver library [7, 1] (KLU to be downloaded and compiled by user independent of KINSOL);
- KINSUPERLUMT: LU factorization and backsolving with compressed-sparse-column (CSC) matrices using the threaded SuperLU\_MT linear solver library [16, 9, 2] (SuperLU\_MT to be downloaded and compiled by user independent of KINSOL).

The spils family of linear solvers providess scaled preconditioned iterative linear solvers and includes:

- KINSPGMR: scaled preconditioned GMRES method;
- KINSPBCG: scaled preconditioned Bi-CGStab method;
- KINSPTFQMR: scaled preconditioned TFQMR method.

The set of linear solver modules distributed with KINSOL is intended to be expanded in the future as new algorithms are developed. Note that users wishing to employ KLU or SuperLU\_MT will need to download and install these libraries independent of SUNDIALS. SUNDIALS provides only the interfaces between itself and these libraries.

In the case of the direct methods KINDENSE and KINBAND the package includes an algorithm for the approximation of the Jacobian by difference quotients, but the user also has the option of supplying the Jacobian (or an approximation to it) directly. When using the sparse direct linear solvers KINKLU and KINSUPERLUMT, the user must supply a routine for the Jacobian (or an approximation to it) in

16 Code Organization

CSC format, since standard difference quotient approximations do not leverage the inherent sparsity of the problem. In the case of the Krylov methods KINSPGMR, KINSPBCG and KINSPTFQMR, the package includes an algorithm for the approximation by difference quotients of the product between the Jacobian matrix and a vector of appropriate length. Again, the user has the option of providing a routine for this operation. For the Krylov methods, the preconditioning must be supplied by the user, in two phases: setup (preprocessing of Jacobian data) and solve.

Each 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 solution, 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.

These modules are also decomposed in another way. Each of the linear solver modules (KINDENSE, etc.) consists of an interface built on top of a generic linear system solver (DENSE etc.). The interface deals with the use of the particular method in the KINSOL context, whereas the generic solver is independent of the context. While some of the generic linear system solvers (DENSE, BAND, SPGMR, SPFGMR, SPBCG, and SPTFQMR) were written with SUNDIALS in mind, they are intended to be usable anywhere 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 for use with any of the Krylov iterative liear solvers. It works in conjunction with NVECTOR\_PARALLEL and generates a preconditioner that is a block-diagonal matrix with each block being a band matrix, as further described in §4.7.

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 multi-computer extension.

## Chapter 4

# Using KINSOL for C Applications

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 sample programs described in the companion document [6] 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.

Users with applications written in FORTRAN77 should see Chapter 5, which describes the FORTRAN/C interface module.

The user should be aware that not all linear solver modules are compatible with all NVECTOR implementations. For example, NVECTOR\_PARALLEL is not compatible with the direct dense, direct band or direct sparse linear solvers since these linear solver modules need to form the complete system Jacobian. The following KINSOL modules can only be used with NVECTOR\_SERIAL, NVECTOR\_OPENMP or NVECTOR\_PTHREADS: KINDENSE, KINBAND, KINKLU and KINSUPERLUMT. It is not recommended to use a threaded vector module with SuperLU\_MT unless it is the NVECTOR\_OPENMP module and SuperLU\_MT is also compiled with openMP. The preconditioner module KINBBDPRE can only be used with NVECTOR\_PARALLEL.

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 Appendix B.

## 4.1 Access to library and header files

At this point, it is assumed that the installation of KINSOL, following the procedure described in Appendix A, has been completed successfully.

Regardless of where the user's application program resides, its associated compilation and load commands must make reference to the appropriate locations for the library and header files required by KINSOL. The relevant library files are

- libdir/libsundials\_kinsol.lib.
- libdir/libsundials\_nvec\*.lib (one to four files),

where the file extension .lib is typically .so for shared libraries and .a for static libraries. The relevant header files are located in the subdirectories

- incdir/include
- incdir/include/kinsol
- incdir/include/sundials

The directories *libdir* and *incdir* are the install library and include directories, respectively. For a default installation, these are *builddir*/lib and *builddir*/include, respectively, where *builddir* was defined in Appendix A.

## 4.2 Data types

The sundials\_types.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  $\S A.1.2$  or  $\S ??$ ).

Additionally, based on the current precision, sundials\_types.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 that this usage is consistent with the typedef for realtype). 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 §??).

## 4.3 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 sundials\_types.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,
- nvector\_openmp.h, which defines the shared memory parallel openMP implementation,
- nvector\_pthreads.h, which defines the shared memory parallel Pthreads implementation.

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

Finally, a linear solver module header file is required. The header files corresponding to the various linear solver options in KINSOL are:

• kinsol\_dense.h, which is used with the dense direct linear solver;

- kinsol\_band.h, which is used with the band direct linear solver;
- kinsol\_lapack.h, which is used with Lapack implementations of dense or band direct linear solvers;
- kinsol\_klu.h, which is used with the KLU sparse direct linear solver;
- kinsol\_superlumt.h, which is used with the SuperLU\_MT threaded sparse direct linear solver;
- kinsol\_spgmr.h, which is used with the Krylov solver SPGMR;
- kinsol\_spfgmr.h, which is used with the Krylov solver SPFGMR;
- kinsol\_spbcgs.h, which is used with the Krylov solver SPBCG;
- kinsol\_sptfqmr.h, which is used with the Krylov solver SPTFQMR;

The header files for the dense and banded linear solvers (both internal and Lapack) include the file kinsol\_direct.h which defines common functions. This in turn includes a file (sundials\_direct.h) which defines the matrix type for these direct linear solvers (DlsMat), as well as various functions and macros acting on such matrices.

The header files for the KLU and SuperLU\_MT sparse linear solvers include the file kinsol\_sparse.h, which defines common functions. This in turn includes a file (sundials\_sparse.h) which defines the matrix type for these sparse direct linear solvers (SlsMat), as well as various functions and macros acting on such matrices.

The header files for the Krylov iterative solvers include kinsol\_spils.h which defined common functions and which in turn includes a header file (sundials\_iterative.h) which enumerates the kind of preconditioning and for the choices for the Gram-Schmidt process for SPGMR.

Other headers may be needed, according to the choice of preconditioner, etc. For example, in the kinFoodWeb\_kry\_p example (see [6]), preconditioning is done with a block-diagonal matrix. For this, even though the KINSPGMR linear solver is used, the header sundials\_dense.h is included for access to the underlying generic dense linear solver.

## 4.4 A skeleton of the user's main program

The following is a skeleton of the user's main program (or calling program) for the solution of a nonlinear problem. Some steps are independent of the NVECTOR implementation used; where this is not the case, usage specifications are given for the implementations provided with KINSOL: Steps marked [P] correspond to NVECTOR\_PARALLEL, steps marked [O] correspond to NVECTOR\_OPENMP, steps marked [T] correspond to NVECTOR\_PTHREADS, while steps marked [S] correspond to NVECTOR\_SERIAL.

#### 1. [P] Initialize MPI

Call 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], [O], [T] Set N, the problem size N.
- [O], [T] Set num\_threads, the number of threads to use within the threaded vector functions.
- [P] Set Nlocal, the local vector length (the sub-vector length for this process). Set N, the global vector length (the problem size N, and the sum of all the values of Nlocal). Set the active set of processes.

Note: The variables N and Nlocal should be of type long int. The variable num\_threads should be of type int.

## 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 = N_VMake_Serial(N, udata);
[O] y0 = N_VMake_OpenMP(N, num_threads, ydata);
[T] y0 = N_VMake_Pthreads(N, num_threads, ydata);
[P] u = N_VMake_Parallel(comm, Nlocal, N, udata);
Otherwise, make the call:
[S] u = N_VNew_Serial(N);
[O] y0 = N_VNew_OpenMP(N, num_threads);
[T] y0 = N_VNew_Pthreads(N, num_threads);
[P] u = N_VNew_Parallel(comm, Nlocal, N);
and load initial values into the structure defined by:
[S] NV_DATA_S(u)
[O] NV_DATA_OMP(y0)
[T] NV_DATA_PT(y0)
```

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. See §4.5.1 for details.

## 5. Set optional inputs

Call KINSet\* routines to change from their default values any optional inputs that control the behavior of KINSOL. See §4.5.4 for details.

## 6. Allocate internal memory

Call KINInit(...) to specify the problem defining function F, allocate internal memory for KINSOL, and initialize KINSOL. KINInit returns a flag to indicate success or an illegal argument value. See §4.5.1 for details.

## 7. Attach linear solver module

Initialize the linear solver module with one of the following calls (for details see §4.5.2).

```
[S], [O], [T] ier = KINDense(...);
[S], [O], [T] ier = KINBand(...);
[S], [O], [T] ier = KINLapackDense(...);
[S], [O], [T] ier = KINLapackBand(...);
[S], [O], [T] ier = KINKLU(...);
[S], [O], [T] ier = KINSuperLUMT(...);
ier = KINSpgmr(...);
```

```
ier = KINSpbcg(...);
ier = KINSptfqmr(...);
```

#### 8. Set linear solver optional inputs

Call KIN\*Set\* functions from the selected linear solver module to change optional inputs specific to that linear solver. See §4.5.4 for details.

## 9. Solve problem

Call ier = KINSol(...) to solve the nonlinear problem for a given initial guess. See §4.5.3 for details.

### 10. Get optional outputs

Call KINGet\* and KIN\*Get\* functions to obtain optional output. See §4.5.5 for details.

## 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] N_VDestroy_Serial(u);
[O] N_VDestroy_OpenMP(y);
[T] N_VDestroy_Pthreads(y);
[P] N_VDestroy_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.

## 4.5 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 §4.5.4, 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 §4.4 for the correct order of these calls.

The return flag (when present) for each of these routines is a negative integer if an error occurred, and non-negative otherwise.

## 4.5.1 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, KINCreate returns a pointer to the newly created KINSOL memory block (of type void \*). If an error occurred, KINCreate prints an error message to stderr and returns NULL.

KINInit

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

Description The function KINInit specifies the problem-defining function, allocates internal mem-

ory, and initializes KINSOL.

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

func (KINSysFn) is the C function which computes the system function F (or G(u) for fixed-point iteration) in the nonlinear problem. This function has the form func(u, fval, user\_data). (For full details see §4.6.1.)

tmpl (N\_Vector) is any N\_Vector (e.g. the initial guess vector u) which is used as a template to create (by cloning) necessary vectors in kin\_mem.

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

KIN\_SUCCESS The call to KINInit 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 KINInit has an illegal value.

Notes If an error occurred, KINInit sends an error message to the error handler function.

KINFree

Call KINFree(&kin\_mem);

Description The function KINFree frees the memory allocated by a previous call to KINCreate.

Arguments The argument is the address of the pointer to the KINSOL memory block returned by

KINCreate (of type void \*).

Return value The function KINFree has no return value.

## 4.5.2 Linear solver specification functions

As previously explained, Newton and Picard iterations require the solution of linear systems of the form  $J\delta=-F$ . There are several KINSOL linear solvers currently available for this task: KINDENSE, KINBAND, KINKLU, KINSUPERLUMT, KINSPGMR, KINSPFGMR, KINSPBCG, and KINSPTFQMR.

The first two linear solvers are direct and derive their names from the type of approximation used for the Jacobian  $J = \partial F/\partial u$ ; KINDENSE and KINBAND work with dense and banded approximations to J, respectively. The SUNDIALS suite includes both internal implementations of these two linear solvers and interfaces to Lapack implementations. Together, these linear solvers are referred to as KINDLS (from Direct Linear Solvers).

The second two linear solvers are sparse direct solvers based on Gaussian elimination, and require user-supplied routines to construct the linear system matrix (in the case of Newton's method, this is the Jacobian  $J=\partial F/\partial u$ ) in compressed-sparse-column format. The SUNDIALS suite does not include internal implementations of these solver libraries, instead requiring compilation of SUNDIALS to link with existing installations of these libraries (if either is missing, SUNDIALS will install without the corresponding interface routines). Together, these linear solvers are referred to as KINSLS (from Sparse Linear Solvers).

The remaining KINSOL linear solvers — KINSPGMR, KINSPFGMR, KINSPBCG, and KINSPTFQMR — are Krylov iterative solvers, which use scaled preconditioned GMRES, scaled preconditioned Flexible GMRES, scaled preconditioned Bi-CGStab, and scaled preconditioned TFQMR, respectively. Together, they are referred to as KINSPILS (from Scaled Preconditioned Iterative Linear Solvers).

With any of the Krylov solvers, only right preconditioning is available. For specification of the preconditioner, see the Krylov solver sections within  $\S 4.5.4$  and  $\S 4.6$ . If preconditioning is done, user-supplied functions define the right preconditioner matrix P, which should approximate the system Jacobian matrix J.

To specify a KINSOL linear solver, after the call to KINCreate but before any calls to KINSO1, the user's program must call one of the functions KINDense/KINLapackDense, KINBand/KINLapackBand, KINKLU, KINSuperLUMT, KINSpgmr, KINSpfgmr, KINSpbcg, or KINSptfqmr, as documented below. The first argument passed to these functions is the KINSOL memory pointer returned by KINCreate. A call to one of these functions links the main KINSOL nonlinear solver to a linear solver and allows the user to specify parameters which are specific to a particular solver, such as the half-bandwidths in the KINBAND case. The use of each of the linear solvers involves certain constants and possibly some macros, that are likely to be needed in the user code. These are available in the corresponding header file associated with the linear solver, as specified below.

In each case, the linear solver module used by KINSOL is actually built on top of a generic linear system solver, which may be of interest in itself. These generic solvers, denoted DENSE, BAND, KLU, SUPERLUMT, SPGMR, SPFGMR, SPBCG, and SPTFQMR, are described separately in Chapter 8.

### KINDense

Call flag = KINDense(kin\_mem, N);

Description The function KINDense selects the KINDENSE linear solver and indicates the use of the

internal direct dense linear algebra functions.

The user's main program must include the kinsol\_dense.h header file.

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

N (long int) problem dimension.

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

KINDLS\_SUCCESS The KINDENSE initialization was successful.

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

KINDLS\_ILL\_INPUT The KINDENSE solver is not compatible with the current NVECTOR

module.

KINDLS\_MEM\_FAIL A memory allocation request failed.

Notes The KINDENSE linear solver is not compatible with all implementations of the NVECTOR

module. Of the NVECTOR modules provided with SUNDIALS, only NVECTOR\_SERIAL, NVECTOR\_OPENMP and NVECTOR\_PTHREADS are compatible, while NVECTOR\_PARALLEL

is not.

## KINLapackDense

Call flag = KINLapackDense(kin\_mem, N);

Description The function KINLapackDense selects the KINDENSE linear solver and indicates the use

of Lapack functions.

The user's main program must include the kinsol\_lapack.h header file.

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

N (int) problem dimension.

Return value The values of the returned flag (of type int) are identical to those of KINDense.

Notes Note that N is restricted to be of type int here, because of the corresponding type

restriction in the Lapack solvers.

## KINBand

Call flag = KINBand(kin\_mem, N, mupper, mlower);

Description The function KINBand selects the KINBAND linear solver and indicates the use of the

internal direct band linear algebra functions.

The user's main program must include the kinsol\_band.h header file.

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

N (long int) problem dimension.

mupper (long int) upper half-bandwidth of the problem Jacobian (or of the approx-

imation of it).

mlower (long int) lower half-bandwidth of the problem Jacobian (or of the approxi-

mation of it).

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

KINDLS\_SUCCESS The KINBAND initialization was successful.

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

KINDLS\_ILL\_INPUT The KINBAND solver is not compatible with the current NVECTOR

module, or one of the Jacobian half-bandwidths is outside its valid

range  $(0 \dots N-1)$ .

KINDLS\_MEM\_FAIL A memory allocation request failed.

Notes

The Kinband linear solver is not compatible with all implementations of the NVECTOR module. Of the NVECTOR modules provided with Sundials, only NVECTOR\_SERIAL, NVECTOR\_OPENMP and NVECTOR\_PTHREADS are compatible, while NVECTOR\_PARALLEL is not. The half-bandwidths are to be set so that the nonzero locations (i,j) in the banded (approximate) Jacobian satisfy  $-mlower \leq j-i \leq mupper$ .

## KINLapackBand

Call flag = KINLapackBand(kin\_mem, N, mupper, mlower);

Description The function KINLapackBand selects the KINBAND linear solver and indicates the use of

Lapack functions.

The user's main program must include the kinsol\_lapack.h header file.

 $\label{eq:arguments} \textbf{Arguments} \quad \textbf{The input arguments are identical to those of KINBand}, \ \textbf{except that N}, \ \textbf{mupper}, \ \textbf{and} \quad$ 

mlower are of type int here.

Return value The values of the returned flag (of type int) are identical to those of KINBand.

Notes Note that N, mupper, and mlower are restricted to be of type int here, because of the

corresponding type restriction in the Lapack solvers.

## KINKLU

Call flag = KINKLU(kin\_mem, N, NNZ);

Description The function KINKLU selects the KINKLU linear solver and indicates the use of sparse-

direct linear algebra functions.

The user's main program must include the kinsol\_klu.h header file.

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

N (int) problem dimension. NNZ (int) problem dimension.

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

KINSLS\_SUCCESS The KINKLU initialization was successful.

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

KINSLS\_ILL\_INPUT The KINKLU solver is not compatible with the current NVECTOR module.

KINSLS\_MEM\_FAIL A memory allocation request failed.

KINSLS\_PACKAGE\_FAIL A call to the KLU library returned a failure flag.

Notes

The KINKLU linear solver is not compatible with all implementations of the NVECTOR module. Of the NVECTOR modules provided with SUNDIALS, only NVECTOR\_SERIAL, NVECTOR\_OPENMP and NVECTOR\_PTHREADS are compatible, while NVECTOR\_PARALLEL is not.

## KINSuperLUMT

Call flag = KINSuperLUMT(kin\_mem, num\_threads, N, NNZ);

Description The function KINSuperLUMT selects the KINSUPERLUMT linear solver and indicates the

use of sparse-direct linear algebra functions.

The user's main program must include the kinsol\_superlumt.h header file.

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

num\_threads (int) the number of threads to use when factoring the linear systems.

Note that SuperLU\_MT is thread-parallel only in the factorization routine.

N (int) problem dimension.

NNZ (int) maximum number of nonzero entries in the system Jacobian.

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

KINSLS\_SUCCESS The KINSUPERLUMT initialization was successful.

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

KINSLS\_ILL\_INPUT The KINSUPERLUMT solver is not compatible with the current NVECTOR module.

KINSLS\_MEM\_FAIL A memory allocation request failed.

KINSLS\_PACKAGE\_FAIL A call to the SuperLU\_MT library returned a failure flag.

Notes

The KINSUPERLUMT linear solver is not compatible with all implementations of the NVECTOR module. Of the NVECTOR modules provided with SUNDIALS, only NVECTOR\_SERIAL, NVECTOR\_OPENMP and NVECTOR\_PTHREADS are compatible, while NVECTOR\_PARALLEL is not.

Performance will significantly degrade if the user applies the SuperLU\_MT package compiled with PThreads while using the NVECTOR\_OPENMP module. If a user wants to use a threaded vector kernel with this thread-parallel solver, then SuperLU\_MT should be compiled with openMP and the NVECTOR\_OPENMP module should be used. Also, note that the expected benefit of using the threaded vector kernel is minimal compared to the potential benefit of the threaded solver, unless very long (greater than 100,000 entries) vectors are used.

## KINSpgmr

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

Description The function KINSpgmr selects the KINSPGMR linear solver.

The user's main program must include the kinsol\_spgmr.h header file.

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 KINSPILS\_MAXL= 5.

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

KINSPILS\_SUCCESS The KINSPGMR initialization was successful.

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

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

KINSPILS\_MEM\_FAIL A memory allocation request failed.



KINSpfgmr

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

Description The function KINSpfgmr selects the KINSPFGMR linear solver.

The user's main program must include the kinsol\_spfgmr.h header file.

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 KINSPILS\_MAXL= 5.

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

KINSPILS\_SUCCESS The KINSPFGMR initialization was successful.

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

KINSPILS\_ILL\_INPUT The NVECTOR module used does not implement a required oper-

ation.

KINSPILS\_MEM\_FAIL A memory allocation request failed.

KINSpbcg

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

Description The function KINSpbcg selects the KINSPBCG linear solver.

The user's main program must include the kinsol\_spbcgs.h header file.

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 KINSPILS\_MAXL= 5.

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

KINSPILS\_SUCCESS The KINSPBCG initialization was successful.

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

KINSPILS\_ILL\_INPUT The NVECTOR module used does not implement a required oper-

ation.

KINSPILS\_MEM\_FAIL A memory allocation request failed.

KINSptfqmr

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

Description The function KINSptfqmr selects the KINSPTFQMR linear solver.

The user's main program must include the kinsol\_sptfqmr.h header file.

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 KINSPILS\_MAXL= 5.

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

KINSPILS\_SUCCESS The KINSPTFQMR initialization was successful.

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

KINSPILS\_ILL\_INPUT The NVECTOR module used does not implement a required oper-

ation.

KINSPILS\_MEM\_FAIL A memory allocation request failed.

## 4.5.3 KINSOL solver function

This is the central step in the solution process, the call to solve the nonlinear algebraic system.

KINSol

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

Description The function KINSol computes an approximate solution to 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 KINSol, but which upon return contains an approximate solution of the nonlinear system F(u) = 0.

strategy (int) strategy used to solve the nonlinear system. It must be of the following:

KIN\_NONE basic Newton iteration

KIN\_LINESEARCH Newton with globalization

KIN\_FP fixed-point iteration with Anderson Acceleration KIN\_PICARD Picard iteration with Anderson Acceleration

u\_scale (N\_Vector) vector containing diagonal elements of scaling matrix  $D_u$  for vector u chosen so that the components of  $D_u$ ·u (as a matrix multiplication) all have roughly the same magnitude when u is close to a root of F(u).

f\_scale (N\_Vector) vector containing diagonal elements of scaling matrix  $D_F$  for F(u) chosen so that the components of  $D_F \cdot F(u)$  (as a matrix multiplication) all have roughly the same magnitude when u is not too near a root of F(u). In the case of a fixed-point iteration, consider F(u) = G(u) - u.

Return value On return, KINSol returns the approximate solution in the vector u if successful. 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 §4.5.4 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 KINCreate.

#### 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 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  $||D_u p||_{L^2} > 0.99$  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 positive 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 solve the preconditioned linear system) encountered an unrecoverable error, or the linear solver routine (lsolve) encountered an error condition.

#### KIN\_SYSFUNC\_FAIL

The system function failed in an unrecoverable manner.

#### KIN\_FIRST\_SYSFUNC\_ERR

The system function failed recoverably at the first call.

#### KIN\_REPTD\_SYSFUNC\_ERR

The system function had repeated recoverable errors. No recovery is possible.

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.

#### 4.5.4 Optional input functions

There are numerous optional input parameters that control the behavior of the KINSOL solver. KINSOL provides functions that can be used to change these from their default values. Table 4.1 lists all optional input functions in KINSOL which are then described in detail in the remainder of this section, beginning with those for the main KINSOL solver and continuing with those for the linear solver modules. For the most casual use of KINSOL, the reader can skip to §4.6.

We note that, on error return, all of these functions also send an error message to the error handler function. We also note that all error return values are negative, so a test flag < 0 will catch any error.

#### 4.5.4.1 Main solver optional input functions

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

Table 4.1: Optional inputs for KINSOL, KINDENSE, KINSPARSE, and KINSPILS

Optional input	Function name	Default
KINSOL main solver		
Error handler function	KINSetErrHandlerFn	internal fn.
Pointer to an error file	KINSetErrFile	stderr
Info handler function	KINSetInfoHandlerFn	internal fn.
Pointer to an info file	KINSetInfoFile	stdout
Data for problem-defining function	KINSetUserData	NULL
Verbosity level of output	KINSetPrintLevel	0
Max. number of nonlinear iterations	KINSetNumMaxIters	200
No initial matrix setup	KINSetNoInitSetup	FALSE
No residual monitoring*	KINSetNoResMon	FALSE
Max. iterations without matrix setup	KINSetMaxSetupCalls	10
Max. iterations without residual check*	KINSetMaxSubSetupCalls	5
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
Values of $\omega_{min}$ and $\omega_{max}^*$	KINSetResMonParams	0.00001  and  0.9
Constant value of $\omega^*$	KINSetResMonConstValue	0.9
Lower bound on $\epsilon$	KINSetNoMinEps	FALSE
Max. scaled length of Newton step	KINSetMaxNewtonStep	$  1000  D_uu_0  _2$
Max. number of $\beta$ -condition failures	KINSetMaxBetaFails	10
Rel. error for D.Q. $Jv$	KINSetRelErrFunc	$\sqrt{\text{uround}}$
Function-norm stopping tolerance	KINSetFuncNormTol	$uround^{1/3}$
Scaled-step stopping tolerance	KINSetScaledSteptol	$uround^{2/3}$
Inequality constraints on solution	KINSetConstraints	NULL
Nonlinear system function	KINSetSysFunc	none
Anderson Acceleration subspace size	KINSetMAA	0
KINDLS linear solvers		
Dense Jacobian function	KINDlsSetDenseJacFn	DQ
Band Jacobian function	KINDlsSetBandJacFn	DQ
KINSLS linear solvers		
Sparse Jacobian function	KINSlsSetSparseJacFn	none
Sparse matrix ordering algorithm	KINKLUSetOrdering	1 for COLAMD
Sparse matrix ordering algorithm	KINSuperLUMTSetOrdering	3 for COLAMD
KINSPILS linear solvers		
Max. number of restarts**	KINSpilsSetMaxRestarts	0
Preconditioner functions and data	KINSpilsSetPreconditioner	NULL, NULL, NULL
Jacobian-times-vector function and data	KINSpilsSetJacTimesVecFn	internal DQ,
	_	NULL

<sup>\*</sup> Only for the KINDLS linear solvers \*\* Only for KINSPGMR and KINSPFGMR

#### KINSetErrFile

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

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

should be directed when the default KINSOL error handler function is used.

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 of NULL disables all future error message output (except for the case in which the KINSOL memory pointer is NULL). This use of KINSetErrFile is strongly

discouraged.

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.

#### KINSetErrHandlerFn

Call flag = KINSetErrHandlerFn(kin\_mem, ehfun, eh\_data);

Description The function KINSetErrHandlerFn specifies the optional user-defined function to be

used in handling error messages.

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

ehfun (KINErrHandlerFn) is the user's C error handler function (see §4.6.2).

eh\_data (void \*) pointer to user data passed to ehfun every time it is called.

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

KIN\_SUCCESS The function ehfun and data pointer eh\_data have been successfully set.

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

Notes The default internal error handler function directs error messages to the file specified

by the file pointer errfp (see KINSetErrFile above).

Error messages indicating that the KINSOL solver memory is NULL will always be directed

to stderr.

#### KINSetInfoFile

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

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

(non-error) 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 stdout.



#### KINSetInfoHandlerFn

Call flag = KINSetInfoHandlerFn(kin\_mem, ihfun, ih\_data);

Description The function KINSetInfoHandlerFn specifies the optional user-defined function to be

used in handling informative (non-error) messages.

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

ihfun (KINInfoHandlerFn) is the user's C information handler function (see §4.6.3).

ih\_data (void \*) pointer to user data passed to ihfun every time it is called.

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

KIN\_SUCCESS The function infun and data pointer in\_data have been successfully set.

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

Notes The default internal information handler function directs informative (non-error) mes-

sages to the file specified by the file pointer infofp (see KINSetInfoFile above).

#### 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\_NONE), 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\_NONE).

 $||F(u)||_{D_F,\infty}$  (for KIN\_NONE 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.

#### KINSetUserData

Call flag = KINSetUserData(kin\_mem, user\_data);

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

be passed to all user-supplied functions.

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

user\_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

If specified, the pointer to user\_data is passed to all user-supplied functions that have it as an argument. Otherwise, a NULL pointer is passed.

If user\_data is needed in user preconditioner functions, the call to KINSetUserData must be made *before* the call to specify the linear solver.



#### KINSetNumMaxIters

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

Description The function KINSetNumMaxIters specifies the maximum number of nonlinear iterations

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.

#### KINSetNoInitSetup

Call flag = KINSetNoInitSetup(kin\_mem, noInitSetup);

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

or Jacobian setup function should be made or not.

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

noInitSetup (booleantype) flag controlling whether an initial call to the precondi-

tioner or Jacobian setup function is made (pass FALSE) or not made (pass

TRUE).

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 noInitSetup is FALSE, meaning that an initial call to the precon-

ditioner or Jacobian setup function will be made.

A call to this function is useful when solving a sequence of problems, in which the final preconditioner or Jacobian value from one problem is to be used initially for the next

problem.

#### KINSetNoResMon

Call flag = KINSetNoResMon(kin\_mem, noNNIResMon);

Description The function KINSetNoResMon specifies whether or not the nonlinear residual monitoring

scheme is used to control Jacobian updating

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

noNNIResMon (booleantype) flag controlling whether residual monitoring is used (pass

FALSE) or not used (pass TRUE).

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

When using a direct solver, the default value for noNNIResMon is FALSE, meaning that the nonlinear residual will be monitored.

Residual monitoring is only available for use with the direct linear solver modules (meaning KINDENSE, KINBAND, KINKLU, and KINSUPERLUMT).

#### KINSetMaxSetupCalls

Call flag = KINSetMaxSetupCalls(kin\_mem, msbset);

Description The function KINSetMaxSetupCalls specifies the maximum number of nonlinear iterations that can be performed between calls to the preconditioner or Jacobian setup function

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

msbset (long int) maximum number of nonlinear iterations without a call to the preconditioner or Jacobian setup function. Pass 0 to indicate the default.

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 msbset was negative.

Notes The default value for msbset is MSBSET\_DEFAULT = 10.

#### KINSetMaxSubSetupCalls

Call flag = KINSetMaxSubSetupCalls(kin\_mem, msbsetsub);

Description The function KINSetMaxSubSetupCalls specifies the maximum number of nonlinear

iterations between checks by the residual monitoring algorithm.

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

msbsetsub (long int) maximum number of nonlinear iterations without checking the nonlinear residual. Pass 0 to indicate the default.

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 msbsetsub was negative.

Notes The default value for msbsetsub is MSBSET\_SUB\_DEFAULT = 5.

Residual monitoring is only available for use with the direct linear solver modules (meaning KINDENSE, KINBAND, KINKLU, and KINSUPERLUMT).

#### 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  $\eta$ . The value must be one of KIN\_ETACHOICE1, KIN\_ETACHOICE2, or KIN\_ETACONSTANT (see Chapter 2 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  $\eta$ . Pass 0.0 to indicate the default.

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 legal 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. Pass 0.0 to indicate the default. ealpha (realtype) value of the  $\alpha$  parameter. Pass 0.0 to indicate the default.

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 legal values are 0.0 < egamma < 1.0 and 1.0 < ealpha < 2.0.

#### KINSetResMonConstValue

Call flag = KINSetResMonConstValue(kin\_mem, omegaconst);

Description The function KINSetResMonConstValue specifies the constant value for  $\omega$  when using

residual monitoring.

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

omegaconst (realtype) constant value for  $\omega$ . Passing 0.0 results in using Eqn. (2.4).

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 omegaconst had an illegal value

Notes The default value for omegaconst is 0.9. The legal values are 0.0 < omegaconst < 1.0.

#### KINSetResMonParams

Call flag = KINSetResMonParams(kin\_mem, omegamin, omegamax);

Description The function KINSetResMonParams specifies the parameters  $\omega_{min}$  and  $\omega_{max}$  in the for-

mula (2.4) for  $\omega$ .

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

omegamin (realtype) value of the  $\omega_{min}$  parameter. Pass 0.0 to indicate the default. omegamax (realtype) value of the  $\omega_{max}$  parameter. Pass 0.0 to indicate the default.

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 omegamin or omegamax had an illegal value.

Notes The default values for omegamin and omegamax are 0.00001 and 0.9, respectively.

The legal values are  $0.0 < \mathtt{omegamin} < \mathtt{omegamax} < 1.0$ .

#### 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, meaning that a positive minimum value, equal

to 0.01\*fnormtol, is applied to  $\epsilon$ . (See KINSetFuncNormTol below.)

#### ${\tt KINSetMaxNewtonStep}$

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 ( $\geq 0.0$ ). Pass 0.0 to indicate the

default.

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 input value was negative.

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

#### KINSetMaxBetaFails

Call flag = KINSetMaxBetaFails(kin\_mem, mxnbcf);

Description The function KINSetMaxBetaFails specifies the maximum number of  $\beta$ -condition fail-

ures in the linesearch algorithm.

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

mxnbcf (realtype) maximum number of  $\beta$ -condition failures. Pass 0.0 to indicate the default.

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 mxnbcf was negative.

Notes The default value of mxnbcf is MXNBCF\_DEFAULT = 10.

#### 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) (relfunc  $\geq 0.0$ ). Pass 0.0 to indicate the

default.

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 relative error was negative.

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 ( $\geq 0.0$ ).

Pass 0.0 to indicate the default.

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 negative.

Notes The default value for fnormtol is (unit roundoff) $^{1/3}$ .

#### 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 ( $\geq 0.0$ ). Pass 0.0 to indicate the default.

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  $u_i \geq 0.0$ .

-1.0 then  $u_i$  will be constrained to be  $u_i \leq 0.0$ .

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

-2.0 then  $u_i$  will be constrained to be  $u_i < 0.0$ .

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) or G(u).

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

func (KINSysFn) user-supplied function that evaluates F(u) (or G(u) for fixed-point iteration).

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 argument func was NULL.

Notes The nonlinear system function is initially specified through KINInit. 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.

#### KINSetMAA

Call flag = KINSetMAA(kin\_mem, maa);

Description The function KINSetMAA specifies the size of the subspace used with Anderson acceleration in conjunction with Picard or fixed-point iteration.

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

maa (long int) subspace size for various methods. A value of 0 means no acceleration, while a positive value means acceleration will be done.

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 maa was negative.

Notes

This function sets the subspace size, which needs to be > 0 if Anderson Acceleration is to be used. It also allocates additional memory necessary for Anderson Acceleration.

The default value of maa is 0, indicating no acceleration. The value of maa should always be less than mxiter.

If the user calls the function KINSetNumMaxIters, that call should be made before the call to KINSetMAA, as the latter uses the value of mxiter.

#### 4.5.4.2 Dense direct linear solver optional input functions

The KINDENSE solver needs a function to compute a dense approximation to the Jacobian matrix J(u). This function must be of type KINDlsDenseJacFn. The user can supply his/her own dense Jacobian function, or use the default internal difference quotient approximation that comes with the KINDENSE solver. To specify a user-supplied Jacobian function djac, KINDENSE provides the function KINDlsSetDenseJacFn. The KINDENSE solver passes the pointer user\_data to the dense Jacobian function. This allows the user to create an arbitrary structure with relevant problem data and access it during the execution of the user-supplied Jacobian function, without using global data in the program. The pointer user\_data may be specified through KINSetUserData.

#### KINDlsSetDenseJacFn

Call flag = KINDlsSetDenseJacFn(kin\_mem, djac);

Description The function KINDlsSetDenseJacFn specifies the dense Jacobian approximation func-

tion to be used.

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

djac (KINDlsDenseJacFn) user-defined dense Jacobian approximation function.

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

KINDLS\_SUCCESS The optional value has been successfully set.

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

KINDLS\_LMEM\_NULL The KINDENSE linear solver has not been initialized.

Notes By default, KINDENSE uses an internal difference quotient function. If NULL is passed to

djac, this default function is used.

The function type KINDlsDenseJacFn is described in §4.6.4.

The KINBAND solver needs a function to compute a banded approximation to the Jacobian matrix J(u). This function must be of type KINDlsBandJacFn. The user can supply his/her own banded Jacobian approximation function, or use the default internal difference quotient approximation that comes with the KINBAND solver. To specify a user-supplied Jacobian function bjac KINBAND provides the function KINDlsSetBandJacFn. The KINBAND solver passes the pointer user\_data to the banded Jacobian approximation function. This allows the user to create an arbitrary structure with relevant problem data and access it during the execution of the user-supplied Jacobian function, without using global data in the program. The pointer user\_data may be specified through KINSetUserData.

#### KINDlsSetBandJacFn

Call flag = KINDlsSetBandJacFn(kin\_mem, bjac);

to be used.

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

bjac (KINDlsBandJacFn) user-defined banded Jacobian approximation function.

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

KINDLS\_SUCCESS The optional value has been successfully set.

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

KINDLS\_LMEM\_NULL The KINBAND linear solver has not been initialized.

Notes By default, KINBAND uses an internal difference quotient approximation. If NULL is passed to bjac, this default function is used.

The function type KINDlsBandJacFn is described in §4.6.5.

#### 4.5.4.3 Sparse linear solvers optional input functions

The KINKLU and KINSUPERLUMT solvers require a function to compute a compressed-sparse-column approximation to the Jacobian matrix J(u). This function must be of type KINSlsSparseJacFn. The user must supply a custom sparse Jacobian function since a difference-quotient approximation would not leverage the underlying sparse matrix structure of the problem. To specify a user-supplied Jacobian function sjac, KINKLU and KINSUPERLUMT provide the function KINSlsSetSparseJacFn. The KINKLU and KINSUPERLUMT solvers pass the pointer user\_data to the sparse Jacobian function. This mechanism allows the user to create an arbitrary structure with relevant problem data and access it during the execution of the user-supplied Jacobian function, without using global data in the program. The pointer user\_data may be specified through KINSetUserData.

#### KINSlsSetSparseJacFn

Call flag = KINSlsSetSparseJacFn(kin\_mem, sjac);

Description The function KINS1sSetSparseJacFn specifies the sparse Jacobian approximation func-

tion to be used.

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

sjac (KINSlsSparseJacFn) user-defined sparse Jacobian approximation function.

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

KINSLS\_SUCCESS The optional value has been successfully set.

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

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

Notes The function type KINSlsSparseJacFn is described in §4.6.6.

When using a spasre direct solver, there may be instances when the number of state variables does not change, but the number of nonzeroes in the Jacobian does change. For the KINKLU solver, we provide a reinitialization. This reinitialization routine reinitializes the Jacobian matrix memory for the new number of nnozeroes and sets flags for a new factorization (symbolic and numeric) to be conducted at the next solver setup call. This routine is useful in the cases where the number of nonzeroes has changed or if the structure of the linear system has changed which would require a new symbolic (and numeric factorization).

#### KINKLUReInit

Call flag = KINKLUReInit(kin\_mem, n, nnz, reinit\_type);

Description The function KINKLUReInit reinitializes Jacobian matrix memory and flags for new

symbolic and numeric KLU factorizations.

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

n (int) number of state variables in the system.

nnz (int) number of nonzeroes in the Jacobian matrix.

reinit\_type (int) type of reinitialization:

- 1 The Jacobian matrix will be destroyed and a new one will be allocated based on the nnz value passed to this call. New symbolic and numeric factorizations will be completed at the next solver setup.
- 2 Only symbolic and numeric factorizations will be completed. It is assumed that the Jacobian size has not exceeded the size of nnz given in the prior call to KINKLU.

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

 ${\tt KINSLS\_SUCCESS} \quad {\tt The \ reinitialization \ succeeded}.$ 

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

KINSLS\_LMEM\_NULL The KINKLU linear solver has not been initialized.

KINSLS\_ILL\_INPUT The given reinit\_type has an illegal value.

KINSLS\_MEM\_FAIL A memory allocation failed.

Notes The default value for reinit\_type is 2.

Both the KINKLU and KINSUPERLUMT solvers can apply reordering algorithms to minimize fill-in for the resulting sparse LU decomposition internal to the solver. The approximate minimal degree ordering for nonsymmetric matrices given by the COLAMD algorithm is the default algorithm used within both solvers, but alternate orderings may be chosen through one of the following two functions. The input values to these functions are the numeric values used in the respective packages, and the user-supplied value will be passed directly to the package.

#### KINKLUSetOrdering

Call flag = KINKLUSetOrdering(kin\_mem, ordering\_choice);

Description The function KINKLUSetOrdering specifies the ordering algorithm used by KINKLU for

reducing fill.

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

ordering\_choice (int) flag denoting algorithm choice:

O AMD

1 COLAMD

2 natural ordering

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

KINSLS\_SUCCESS The optional value has been successfully set.

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

KINSLS\_ILL\_INPUT The supplied value of ordering\_choice is illegal.

Notes The default ordering choice is 1 for COLAMD.

#### KINSuperLUMTSetOrdering

Call flag = KINSuperLUMTSetOrdering(kin\_mem, ordering\_choice);

Description The function KINSuperLUMTSetOrdering specifies the ordering algorithm used by KIN-

SUPERLUMT for reducing fill.

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

ordering\_choice (int) flag denoting algorithm choice:

0 natural ordering

1 minimal degree ordering on  $J^TJ$ 

2 minimal degree ordering on  $J^T + J$ 

3 COLAMD

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

KINSLS\_SUCCESS The optional value has been successfully set.

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

KINSLS\_ILL\_INPUT The supplied value of ordering\_choice is illegal.

Notes The default ordering choice is 3 for COLAMD.

#### 4.5.4.4 Iterative linear solvers optional input functions

If any preconditioning is to be done with one of the KINSPILS linear solvers, then the user must supply a preconditioner solve function psolve and specify its name in a call to KINSPILSEtPreconditioner.

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 §4.6. If used, the psetup function should also be specified in the call to KINSPILS solver passes the pointer user\_data received through KINSetUserData to the preconditioner psetup and psolve 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.

Ther KINSPILS solvers require 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-times-vector approximation function, or use the internal difference quotient approximation that comes with the KINSPILS solvers. A user-defined Jacobian-vector function must be of type KINSpilsJacTimesVecFn and can be specified through a call to KINSpilsSetJacTimesVecFn (see §4.6.7 for specification details). A KINSPILS solver passes the pointer user\_data received through KINSetUserData to the Jacobian-times-vector function jtimes each time it is called.

#### KINSpilsSetPreconditioner

Call flag = KINSpilsSetPreconditioner(kin\_mem, psetup, psolve);

Description The function KINSpilsSetPreconditioner specifies the preconditioner setup and solve

functions.

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

psetup (KINSpilsPrecSetupFn) user-defined preconditioner setup function. Pass NULL

if no setup operation is to be done.

psolve (KINSpilsPrecSolveFn) user-defined preconditioner solve function.

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

 ${\tt KINSPILS\_SUCCESS} \qquad {\tt The~optional~values~have~been~successfully~set}.$ 

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

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

The function type KINSpilsPrecSolveFn is described in §4.6.8. The function type

KINSpilsPrecSetupFn is described in §4.6.9.

## ${\tt KINSpilsSetJacTimesVecFn}$

Notes

Call flag = KINSpilsSetJacTimesVecFn(kin\_mem, jtimes);

Description The function KINSpilsSetJacTimesFn specifies the Jacobian-vector function to be used.

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

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

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

KINSPILS\_SUCCESS The optional value has been successfully set.

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

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

Notes

By default, the KINSPILS linear solvers use an internal difference quotient function KINSpilsDQJtimes. If NULL is passed as jtimes, this default function is used.

The function type KINSpilsJacTimesVecFn is described in §4.6.7.

#### KINSpilsSetMaxRestarts

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

Description The function KINSpilsSetMaxRestarts specifies the maximum number of times the

iterative linear solver can be restarted.

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

(int) maximum number of restarts (>0).

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

KINSPILS\_SUCCESS The optional value has been successfully set.

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

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

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

Notes The default value is 0 (meaning no restarts).

This option is available only for the KINSPGMR and KINSPFGMR linear solvers.



#### 4.5.5Optional output functions

KINSOL provides an extensive list of functions that can be used to obtain solver performance information. Table 4.2 lists all optional output functions in KINSOL, which are then described in detail in the remainder of this section, beginning with those for the main KINSOL solver and continuing with those for the linear solver modules. Where the name of an output from a linear solver module would otherwise conflict with the name of an optional output from the main solver, a suffix LS (for Linear Solver) has been added here (e.g., lenrwLS).

#### Main solver optional output functions 4.5.5.1

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.

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

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

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).

Table 4.2: Optional outputs from kinsol, kindles, kinsles, and kinspils

Optional output	Function name	
KINSOL main solver		
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	
KINDLS linear solvers		
Size of real and integer workspaces	KINDlsGetWorkSpace	
No. of Jacobian evaluations	KINDlsGetNumJacEvals	
No. of $F$ calls for D.Q. Jacobian evals.	KINDlsGetNumFuncEvals	
Last return from a KINDLS function	KINDlsGetLastFlag	
KINSLS linear solvers		
No. of Jacobian evaluations	KINSlsGetNumJacEvals	
Last return from a linear solver function	KINSlsGetLastFlag	
Name of constant associated with a return flag	KINSlsGetReturnFlagName	
KINSPILS linear solvers		
Size of real and integer workspaces	KINSpilsGetWorkSpace	
No. of linear iterations	KINSpilsGetNumLinIters	
No. of linear convergence failures	KINSpilsGetNumConvFails	
No. of preconditioner evaluations	KINSpilsGetNumPrecEvals	
No. of preconditioner solves	KINSpilsGetNumPrecSolves	
No. of Jacobian-vector product evaluations	KINSpilsGetNumJtimesEvals	
No. of $F$ calls for D.Q. Jacobian-vector evals.	KINSpilsGetNumFuncEvals	
Last return from a linear solver function	KINSpilsGetLastFlag	

#### 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.

#### 4.5.5.2 Dense direct linear solvers optional output functions

The following optional outputs are available from the KINDLS module: workspace requirements, number of calls to the Jacobian routine, number of calls to the system function routine for difference quotient Jacobian approximation, and last return value from a KINDLS function.

#### KINDlsGetWorkSpace

Call flag = KINDlsGetWorkSpace(kin\_mem, &lenrwLS, &leniwLS);

Description The function KINDlsGetWorkSpace returns the KINDENSE real and integer workspace

sizes.

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

lenrwLS (long int) the number of realtype values in the KINDLS workspace.

leniwLS (long int) the number of integer values in the KINDLS workspace.

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

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

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

KINDLS\_LMEM\_NULL The KINDENSE linear solver has not been initialized.

Notes For the KINDENSE linear soler, in terms of the problem size N, the actual size of the real workspace is  $N^2$  realtype words, and the actual size of the integer workspace is

N integer words.

For the KINBAND linear solver, in terms of the problem size N and Jacobian half-bandwidths, the actual size of the real workspace, in realtype words, is approximately (2 mupper + 3 mlower + 2) N, and the actual size of the integer workspace is N integer words.

#### KINDlsGetNumJacEvals

Call flag = KINDlsGetNumJacEvals(kin\_mem, &njevals);

Description The function KINDlsGetNumJacEvals returns the number of calls to the dense Jacobian

approximation function.

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

njevals (long int) the number of calls to the Jacobian function.

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

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

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

KINDLS\_LMEM\_NULL The KINDENSE linear solver has not been initialized.

#### KINDlsGetNumFuncEvals

Call flag = KINDlsGetNumFuncEvals(kin\_mem, &nfevalsLS);

Description The function KINDlsGetNumFuncEvals returns the number of calls to the user system

function used to compute the difference quotient approximation to the dense or banded

Jacobian.

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

nfevalsLS (long int) the number of calls to the user system function.

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

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

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

KINDLS\_LMEM\_NULL The KINDENSE or KINBAND linear solver has not been initialized.

Notes The value nfevalsLS is incremented only if the internal difference quotient function is

used.

#### KINDlsGetLastFlag

Call flag = KINDlsGetLastFlag(kin\_mem, &lsflag);

Description The function KINDlsGetLastFlag returns the last return value from a KINDENSE rou-

tine.

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

1sflag (long int) the value of the last return flag from a KINDENSE function.

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

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

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

KINDLS\_LMEM\_NULL The KINDENSE linear solver has not been initialized.

Notes If the KINDLS setup function failed (KINSol returned KIN\_LSETUP\_FAIL), then lsflag is

equal to the column index (numbered from one) at which a zero diagonal element was encountered during the LU factorization of the dense Jacobian matrix. For all other

failures, lsflag is negative.

#### 4.5.5.3 Sparse direct linear solvers optional output functions

The following optional outputs are available from the KINSLS module: number of calls to the Jacobian routine and last return value from a KINSLS function.

#### KINSlsGetNumJacEvals

Call flag = KINSlsGetNumJacEvals(kin\_mem, &njevals);

Description The function KINSlsGetNumJacEvals returns the number of calls to the sparse Jacobian

approximation function.

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

njevals (long int) the number of calls to the Jacobian function.

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

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

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

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

#### ${\tt KINSlsGetLastFlag}$

Call flag = KINSlsGetLastFlag(kin\_mem, &lsflag);

Description The function KINSlsGetLastFlag returns the last return value from a KINSLS routine.

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

lsflag (long int) the value of the last return flag from a KINSLS function.

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

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

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

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

Notes

#### KINSlsGetReturnFlagName

Description The function KINSlsGetReturnFlagName returns the name of the KINSLS constant cor-

responding to 1sflag.

Arguments The only argument, of type long int, is a return flag from a KINSLS function.

Return value The return value is a string containing the name of the corresponding constant.

#### 4.5.5.4 Iterative linear solvers optional output functions

The following optional outputs are available from the KINSPILS modules: workspace requirements, number of linear iterations, number of linear convergence failures, number of calls to the preconditioner setup and solve routines, number of calls to the Jacobian-vector product routine, number of calls to the system function routine for difference quotient Jacobian-vector product approximation, and last return value from a linear solver function.

#### KINSpilsGetWorkSpace

Call flag = KINSpilsGetWorkSpace(kin\_mem, &lenrwLS, &leniwLS);

Description The function KINSpilsGetWorkSpace returns the global sizes of the linear solver real

and integer workspaces.

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

lenrwLS (long int) the number of realtype values in the linear solver workspace.

leniwLS (long int) the number of integer values in the linear solver workspace.

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

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

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

KINSPILS\_LMEM\_NULL The 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, in realtype words, is roughly:  $(\max 1+3)*N+\max 1*(\max 1+4)+1$  for KINSPGMR,

 $(2\max 1+3)*N+\max 1*(\max 1+4)+1$  for KINSPEGMR,

7\*N for KINSPBCG, and 11\*N for KINSPTFQMR.

In a parallel setting, this value is global, summed over all processes.

#### KINSpilsGetNumLinIters

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

Description The function KINSpilsGetNumLinIters returns the cumulative number of linear itera-

tions.

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:

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

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

KINSPLIS\_LMEM\_NULL The linear solver module has not been initialized.

#### KINSpilsGetNumConvFails

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

Description The function KINSpilsGetNumConvFails returns the cumulative number of linear con-

vergence 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:

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

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

KINSPILS\_LMEM\_NULL The linear solver module has not been initialized.

#### KINSpilsGetNumPrecEvals

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

Description The function KINSpilsGetNumPrecEvals returns the number of preconditioner evalua-

tions, 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:

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

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

KINSPILS\_LMEM\_NULL The linear solver module has not been initialized.

#### KINSpilsGetNumPrecSolves

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

Description The function KINSpilsGetNumPrecSolves 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:

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

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

KINSPILS\_LMEM\_NULL The linear solver module has not been initialized.

#### KINSpilsGetNumJtimesEvals

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

Description The function KINSpilsGetNumJtimesEvals 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:

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

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

KINSPILS\_LMEM\_NULL The linear solver module has not been initialized.

#### KINSpilsGetNumFuncEvals

Call flag = KINSpilsGetNumFuncEvals(kin\_mem, &nfevalsLS);

Description The function KINSpilsGetNumFuncEvals returns the number of calls to the user system

function for difference quotient Jacobian-vector product approximations.

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

nfevalsLS (long int) the number of calls to the user system function.

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

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

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

KINSPILS\_LMEM\_NULL The linear solver module has not been initialized.

Notes The value nfevalsLS is incremented only if the default KINSpilsDQJtimes difference

quotient function is used.

#### KINSpilsGetLastFlag

Call flag = KINSpilsGetLastFlag(kin\_mem, &lsflag);

Description The function KINSpilsGetLastFlag returns the last return value from a KINSPILS rou-

tine.

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

lsflag (long int) the value of the last return flag from a KINSPILS function.

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

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

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

KINSPILS\_LMEM\_NULL The linear solver module has not been initialized.

Notes If the KINSPILS setup function failed (KINSOL returned KIN\_LSETUP\_FAIL), lsflag will be set to SPGMR\_PSET\_FAIL\_UNREC, SPFGMR\_PSET\_FAIL\_UNREC, SPBCG\_PSET\_FAIL\_UNREC,

or SPTFQMR\_PSET\_FAIL\_UNREC.

If the KINSPGMR solve function failed (KINSol returned KIN\_LSOLVE\_FAIL), lsflag contains the error return flag from SpgmrSolve and will be one of: SPGMR\_MEM\_NULL, indicating that the SPGMR memory is NULL; SPGMR\_ATIMES\_FAIL\_UNREC, indicating an unrecoverable failure in the Jacobian-times-vector 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  $\mathsf{SPGMR\_QRSOL\_FAIL}$ , indicating that the matrix R was found to be singular during the QR solve phase.

If the KINSPFGMR solve function failed (KINSol returned KIN\_LSOLVE\_FAIL), lsflag contains the error return flag from SpfgmrSolve and will be a similar value to one of the return codes for KINSPGMR.

If the KINSPBCG solve function failed (KINSol returned KIN\_LSOLVE\_FAIL), lsflag contains the error return flag from SpbcgSolve and will be one of: SPBCG\_MEM\_NULL, indicating that the SPBCG memory is NULL; SPBCG\_ATIMES\_FAIL\_UNREC, indicating an unrecoverable failure in the Jacobian-times-vector function; or SPBCG\_PSOLVE\_FAIL\_UNREC, indicating that the preconditioner solve function, psolve, failed unrecoverably.

If the KINSPTFQMR solve function failed (KINSol returned KIN\_LSOLVE\_FAIL), lsflag contains the error return flag from SptfqmrSolve and will be one of: SPTFQMR\_MEM\_NULL, indicating that the SPTFQMR memory is NULL; SPTFQMR\_ATIMES\_FAIL\_UNREC, indicating an unrecoverable failure in the J\*v function; or SPTFQMR\_PSOLVE\_FAIL\_UNREC, indicating that the preconditioner solve function, psolve, failed unrecoverably.

## 4.6 User-supplied functions

The user-supplied functions consist of one function defining the nonlinear system, (optionally) a function that handles error and warning messages, (optionally) a function that handles informational messages, (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 any of the Krylov iterative algorithms.

#### 4.6.1 Problem-defining function

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

### KINSysFn

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

Purpose This function computes F(u) (or G(u) for fixed-point iteration and Anderson accelera-

tion) 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).

user\_data is a pointer to user data, the pointer user\_data passed to KINSetUserData.

Return value A KINSysFn function should return 0 if successful, a positive value if a recoverable error occurred (in which case KINSOL will attempt to correct), or a negative value if it failed unrecoverably (in which case the solution process is halted and KIN\_SYSFUNC\_FAIL is

returned).

Notes Allocation of memory for fval is handled within KINSOL.

#### 4.6.2 Error message handler function

As an alternative to the default behavior of directing error and warning messages to the file pointed to by errfp (see KINSetErrFile), the user may provide a function of type KINErrHandlerFn to process any such messages. The function type KINErrHandlerFn is defined as follows:

#### KINErrHandlerFn

Definition typedef void (\*KINErrHandlerFn)(int error\_code, const char \*module, const char \*function, char \*msg, void \*eh\_data);

Purpose This function processes error and warning messages from KINSOL and its sub-modules.

Arguments error\_code is the error code.

module is the name of the KINSOL module reporting the error.

function is the name of the function in which the error occurred.

msg is the error message.

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

KINSetErrHandlerFn.

Return value A KINErrHandlerFn function has no return value.

Notes  $error\_code$  is negative for errors and positive (KIN\_WARNING) for warnings. If a function

that returns a pointer to memory encounters an error, it sets error\_code to 0.

#### 4.6.3 Informational message handler function

As an alternative to the default behavior of directing informational (meaning non-error) messages to the file pointed to by infofp (see KINSetInfoFile), the user may provide a function of type KINInfoHandlerFn to process any such messages. The function type KINInfoHandlerFn is defined as follows:

#### KINInfoHandlerFn

Definition typedef void (\*KINInfoHandlerFn)(const char \*module, const char \*function, char \*msg, void \*ih\_data);

Purpose This function processes informational messages from KINSOL and its sub-modules.

Arguments module is the name of the KINSOL module reporting the information.

function is the name of the function reporting the information.

msg is the message.

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

 ${\tt KINSetInfoHandlerFn}.$ 

Return value A KINInfoHandlerFn function has no return value.

## 4.6.4 Jacobian information (direct method with dense Jacobian)

If the direct linear solver with dense treatment of the Jacobian is used (KINDense or KINLapackDense is called in Step 7 of §4.4), the user may provide a function of type KINDlsDenseJacFn defined by

#### KINDlsDenseJacFn

Definition typedef int (\*KINDlsDenseJacFn)(long int N, N\_Vector u, N\_Vector fu, DlsMat J, void \*user\_data, N\_Vector tmp1, N\_Vector tmp2);

Purpose This function computes the dense Jacobian J(u) or an approximation to it.

Arguments N is the problem size.

u is the current (unscaled) iterate.

fu is the current value of the vector F(u).

J is the output approximate Jacobian matrix,  $J = \partial F/\partial u$ .

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

tmp1

tmp2 are pointers to memory allocated for variables of type N\_Vector which can be used by KINDenseJacFn as temporary storage or work space.

Return value A function of type KINDlsDenseJacFn should return 0 if successful or a non-zero value otherwise.

Notes

A user-supplied dense Jacobian function must load the N by N dense matrix J with an approximation to the Jacobian matrix J(u) at u. Only nonzero elements need to be loaded into J because J is set to the zero matrix before the call to the Jacobian function. The type of J is DlsMat.

The accessor macros DENSE\_ELEM and DENSE\_COL allow the user to read and write dense matrix elements without making explicit references to the underlying representation of the DlsMat type. DENSE\_ELEM(J, i, j) references the (i, j)-th element of the dense matrix J (i, j=0...N-1). This macro is for use in small problems in which efficiency of access is not a major concern. Thus, in terms of indices m and n running from 1 to N, the Jacobian element  $J_{m,n}$  can be loaded with the statement DENSE\_ELEM(J, m-1, n-1) =  $J_{m,n}$ . Alternatively, DENSE\_COL(J, j) returns a pointer to the storage for the jth column of J (j=0...N-1), and the elements of the jth column are then accessed via ordinary array indexing. Thus  $J_{m,n}$  can be loaded with the statements col\_n = DENSE\_COL(J, n-1); col\_n[m-1] =  $J_{m,n}$ . For large problems, it is more efficient to use DENSE\_COL than to use DENSE\_ELEM. Note that both of these macros number rows and columns starting from 0, not 1.

The DlsMat type and the accessor macros DENSE\_ELEM and DENSE\_COL are documented in  $\S 8.1.3$ .

If the user's KINDlsDenseJacFn function uses difference quotient approximations, it may need to access quantities not in the call list. To obtain these, the user will need to add the kin\_mem structure to their user\_data and use the KINGet\* functions described in §4.5.5.1. The unit roundoff can be accessed as UNIT\_ROUNDOFF defined in sundials\_types.h.

For the sake of uniformity, the argument N is of type long int, even in the case that the Lapack dense solver is to be used.

#### 4.6.5 Jacobian information (direct method with banded Jacobian)

If the direct linear solver with banded treatment of the Jacobian is used (KINBand or KINLapackBand is called in Step 7 of §4.4), the user may provide a function of type KINDlsBandJacFn defined by:

#### KINDlsBandJacFn

```
Definition typedef int (*KINDlsBandJacFn)(long int N, long int mupper,
long int mlower, N_Vector u, N_Vector fu,
DlsMat J, void *user_data,
N_Vector tmp1, N_Vector tmp2);
```

Purpose This function computes the banded Jacobian J(u) or a banded approximation to it.

Arguments N is the problem size.

mlower

mupper are the lower and upper half-bandwidths of the Jacobian.

u is the current (unscaled) iterate.

fu is the current value of the vector F(u).

J is the output approximate Jacobian matrix,  $J = \partial F/\partial u$ .

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

tmp1

are pointers to memory allocated for variables of type N\_Vector which can be used by KINBandJacFn as temporary storage or work space.

Return value A function of type KINDlsBandJacFn should return 0 if successful or a non-zero value otherwise.

Notes

A user-supplied band Jacobian function must load the band matrix J of type DlsMat with the elements of the Jacobian J(u) at u. Only nonzero elements need to be loaded into J because J is preset to zero before the call to the Jacobian function.

The accessor macros BAND\_ELEM, BAND\_COL, and BAND\_COL\_ELEM allow the user to read and write band matrix elements without making specific references to the underlying representation of the DlsMat type. BAND\_ELEM(J, i, j) references the (i, j)th element of the band matrix J, counting from 0. This macro is for use in small problems in which efficiency of access is not a major concern. Thus, in terms of indices m and n running from 1 to N with (m,n) within the band defined by mupper and mlower, the Jacobian element  $J_{m,n}$  can be loaded with the statement BAND\_ELEM(J, m-1, n-1) =  $J_{m,n}$ . The elements within the band are those with -mupper  $\leq$  m-n  $\leq$  mlower. Alternatively, BAND\_COL(J, j) returns a pointer to the diagonal element of the jth column of J, and if we assign this address to realtype \*col\_j, then the ith element of the jth column is given by BAND\_COL\_ELEM(col\_j, i, j), counting from 0. Thus for (m,n) within the band,  $J_{m,n}$  can be loaded by setting col\_n = BAND\_COL(J, n-1); BAND\_COL\_ELEM(col\_n, m-1, n-1) =  $J_{m,n}$ . The elements of the jth column can also be accessed via ordinary array indexing, but this approach requires knowledge of the underlying storage for a band matrix of type DlsMat. The array coln can be indexed from -mupper to mlower. For large problems, it is more efficient to use the combination of BAND\_COL and BAND\_COL\_ELEM than to use the BAND\_ELEM. As in the dense case, these macros all number rows and columns starting from 0, not 1.

The DlsMat type and the accessor macros BAND\_ELEM, BAND\_COL, and BAND\_COL\_ELEM are documented in §8.1.4.

If the user's KINDlsBandJacFn function uses difference quotient approximations, it may need to access quantities not in the call list. To obtain these, the user will need to add the kin\_mem structure to their user\_data and use the KINGet\* functions described in §4.5.5.1. The unit roundoff can be accessed as UNIT\_ROUNDOFF defined in sundials\_types.h.

For the sake of uniformity, the arguments N, mlower, and mupper are of type long int, even in the case that the Lapack band solver is to be used.

## 4.6.6 Jacobian information (direct method with sparse Jacobian)

If the direct linear solver with sparse treatment of the Jacobian is used (KINKLU or KINSuperLUMT is called in Step 7 of §4.4), the user may provide a function of type KINSlsSparseJacFn defined by

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

tmp1

are pointers to memory allocated for variables of type N\_Vector which can tmp2 be used by KINSlsSparseJacFn as temporary storage or work space.

Return value A function of type KINSlsSparseJacFn should return 0 if successful or a non-zero value otherwise.

Notes

A user-supplied sparse Jacobian function must load the compressed-sparse-column matrix J with an approximation to the Jacobian matrix J(u) at the point (u). Storage for J already exists on entry to this function, although the user should ensure that sufficient space is allocated in J to hold the nonzero values to be set; if the existing space is insufficient the user may reallocate the data and row index arrays as needed. The type of J is SlsMat, and the amount of allocated space is available within the SlsMat structure as NNZ. The SlsMat type is further documented in the Section §8.2.

If the user's KINSlsSparseJacFn function uses difference quotient approximations to set the specific nonzero matrix entries, then it may need to access quantities not in the argument list. To obtain these, use the KINGet\* functions described in §4.5.5.1. The unit roundoff can be accessed as UNIT\_ROUNDOFF defined in sundials\_types.h.

#### 4.6.7 Jacobian information (matrix-vector product)

If one of the Krylov iterative linear solvers SPGMR, SPBCG, or SPTFQMR is selected (KINSp\* is called in step 7 of §4.4), the user may provide a jtimes function of type KINSpilsJacTimesVecFn to compute products Jv. If such a function is not supplied, the default is a difference quotient approximation of these products.

#### KINSpilsJacTimesVecFn

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

N\_Vector u, booleantype new\_u,

void \*user\_data);

This jtimes function computes the product Jv (or an approximation to it). Purpose

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

> T 77 is the computed output vector.

is the current value of the dependent variable vector.

is a flag, input from KINSOL and possibly reset by the user's jtimes function, new\_u

> indicating whether the iterate vector u has been updated since the last call to jtimes. This is useful if the jtimes function computes and saves Jacobian data that depends on u for use in computing J(u)v. The input value of new\_u is TRUE following an update by KINSOL, and in that case any saved Jacobian data depending on u should be recomputed. The jtimes routine should then set new\_u to FALSE, so that on subsequent calls to jtimes with the same u,

the saved data can be reused.

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

Return value The value to be returned by the Jacobian-times-vector function should be 0 if successful. If a recoverable failure occurred, the return value should be positive. In this case, KINSOL will attempt to correct by calling the preconditioner setup function. If this information is current, KINSOL halts. If the Jacobian-times-vector function encounters an unrecoverable error, it should return a negative value, prompting KINSOL to halt.

Notes

If a user-defined routine is not given, then an internal KINSPGMR function, using difference quotient approximations, is used.

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

#### 4.6.8 Preconditioning (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, approximating (at least crudely) the system Jacobian  $J = \partial F/\partial u$ . This function must be of type KINSpilsPrecSolveFn, defined as follows:

#### KINSpilsPrecSolveFn

Definition typedef int (\*KINSpilsPrecSolveFn)(N\_Vector u, N\_Vector uscale, N\_Vector fval, N\_Vector fscale, N\_Vector v, void \*user\_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, r. On

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

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

the function KINSetUserData.

tmp is a pointer to memory allocated for a variable of type 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.

Notes If the preconditioner solve function fails recoverably and if the preconditioner informa-

tion (set by the preconditioner setup function) is out of date, KINSOL attempts to correct by calling the setup function. If the preconditioner data is current, KINSOL halts.

#### 4.6.9 Preconditioning (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 KINSpilsPrecSetupFn, defined as follows:

#### ${\tt KINSpilsPrecSetupFn}$

Definition typedef int (\*KINSpilsPrecSetupFn)(N\_Vector u, N\_Vector uscale, N\_Vector fval, N\_Vector fscale, void \*user\_data, N\_Vector tmp1,

N\_Vector tmp2);

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

conditioner solve function.

Arguments The arguments of a KINSpilsPrecSetupFn 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.

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

the function KINSetUserData.

tmp1

are pointers to memory allocated for variables of type N\_Vector which can be used by KINSpilsPrecSetupFn 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, any other value resulting in halting the KINSOL solver.

The user-supplied preconditioner setup subroutine should compute the right preconditioner matrix P (stored in the memory block referenced by the user\_data pointer) used to form the scaled preconditioned linear system

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

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 user's KINSpilsPrecSetupFn function uses difference quotient approximations, it may need to access quantities not in the call list. To obtain these, the user will need to add the kin\_mem structure to their user\_data and use the KINGet\* functions described in §4.5.5.1. The unit roundoff can be accessed as UNIT\_ROUNDOFF defined in sundials\_types.h.

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

# 4.7 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 band structure. This structure is given by upper and lower half-bandwidths, mudq and mldq, defined as the number of non-zero diagonals above and below the main diagonal, respectively. However, from the resulting approximate Jacobain blocks, only a matrix of bandwidth mukeep + mlkeep +1 is retained.

Neither pair of parameters need 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 mukeep and mlkeep while keeping mudq and mldq at their true values, discards the elements outside the narrower band. Reducing both pairs has the additional effect of lumping the outer Jacobian elements into the computed elements within the band, and requires more caution and experimentation to see whether the lower cost of narrower band matrices offsets the loss of accuracy in the blocks.

Notes

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 user\_data as that passed by the user to KINSetUserData and passed to the user's function func, and neither function has a return value. The user is responsible for providing space (presumably within user\_data) for components of u that are communicated by Gcomm from the other processes, and that are then used by Gloc, which should not do any communication.

#### KINLocalFn

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

Purpose This Gloc 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.

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

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

Notes This function must assume that all interprocess communication of data needed to cal-

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

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

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

#### KINCommFn

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

Purpose This Gcomm function performs all interprocess communications necessary for the execu-

tion 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.

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

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 user\_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 KINBBDPrecInit (see below).

Besides the header files required for the solution of a nonlinear problem (see §4.3), 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  $\S4.4$  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. Attach iterative linear solver, one of:

```
flag = KINSpgmr(kin_mem, max1);
flag = KINSpfgmr(kin_mem, max1);
flag = KINSpbcg(kin_mem, max1);
flag = KINSptfqmr(kin_mem, max1);
```

#### 8. Initialize the KINBBDPRE preconditioner module

Specify the upper and lower half-bandwidth pairs (mudq, mldq) and (mukeep, mlkeep), and call

to allocate memory for and initialize the internal preconditoner data. The last two arguments of KINBBDPrecInit are the two user-supplied functions described above.

9. Set linear solver optional inputs

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

10. Solve problem

#### 11. Get optional output

Additional optional outputs associated with KINBBDPRE are available by way of two routines described below, KINBBDPrecGetWorkSpace and KINBBDPrecGetNumGfnEvals.

- 12. Deallocate memory for solution vector
- 13. Free solver memory
- 14. Finalize MPI

The user-callable function that initializes KINBBDPRE (step 8), is described in more detail below.

#### KINBBDPrecInit

```
Call flag = KINBBDPrecInit(kin_mem, Nlocal, mudq, mldq, mukeep, mlkeep, dq_rel_u, Gloc, Gcomm);
```

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

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

Nlocal (long int) local vector length.

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

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

mukeep (long int) upper half-bandwidth of the retained banded approximate Jacobian block.

mlkeep (long int) lower half-bandwidth of the retained banded approximate Jacobian block.

dq\_rel\_u (realtype) the relative increment in components of u used in the difference quotient approximations. The default is  $dq_rel_u = \sqrt{unit\ roundoff}$ , 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 The return value flag (of type int) is one of

KINSPILS\_SUCCESS The call to KINBBDPrecInit was successful.

KINSPILS\_MEM\_NULL The kin\_mem pointer was NULL.

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

KINSPILS\_LMEM\_NULL A KINSPILS linear solver was not attached.

KINSPILS\_ILL\_INPUT The supplied vector implementation was not compatible with block band preconditioner.

Notes

If one of the half-bandwidths mudq or mldq to be used in the difference-quotient calculation of the approximate Jacobian is negative or exceeds the value Nlocal-1, it is replaced with 0 or Nlocal-1 accordingly.

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

Also, the half-bandwidths mukeep and mlkeep of the retained banded approximate Jacobian block may be even smaller, to reduce storage and computation costs further.

For all four half-bandwidths, the values need not be the same for every process.

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

#### KINBBDPrecGetWorkSpace

Call flag = KINBBDPrecGetWorkSpace(kin\_mem, &lenrwBBDP, &leniwBBDP);

Description The function KINBBDPrecGetWorkSpace returns the local KINBBDPRE real and integer workspace sizes.

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

lenrwBBDP (long int) local number of realtype values in the 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:

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

KINSPILS\_MEM\_NULL The kin\_mem pointer was NULL.

KINSPILS\_PMEM\_NULL The KINBBDPRE preconditioner has not been initialized.

Notes

In terms of the local vector dimension Nlocal and  $smu = min(N_l-1, mukeep + mlkeep)$ , the actual size of the real workspace is  $(2 \, mlkeep + mukeep + smu + 2) \, Nlocal realtype$  words, and the actual size of the integer workspace is Nlocal integer words. These values are local to the current processor.

The workspaces referred to here exist in addition to those given by the corresponding KINSp\*GetWorkSpace function.

#### KINBBDPrecGetNumGfnEvals

Call flag = KINBBDPrecGetNumGfnEvals(kin\_mem, &ngevalsBBDP);

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

function due to the difference quotient approximation of the Jacobian blocks used within

KINBBDPRE's preconditioner setup function.

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

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

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

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

KINSPILS\_MEM\_NULL The kin\_mem pointer was NULL.

KINSPILS\_PMEM\_NULL The KINBBDPRE preconditioner has not been initialized.

In addition to the ngevalsBBDP Gloc evaluations, the costs associated with KINBBDPRE also include nlinsetups LU factorizations, nlinsetups calls to Gcomm, npsolves banded backsolve calls, and nfevalsLS right-hand side function evaluations, where nlinsetups is an optional KINSOL output and npsolves and nfevalsLS are linear solver optional outputs (see §4.5.5).

# Chapter 5

# FKINSOL, an Interface Module for FORTRAN Applications

The FKINSOL interface module is a package of C functions which support the use of the KINSOL solver, for the solution of 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 all supplied serial and the parallel NVECTOR implementations.

## 5.1 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. By default, those mapping definitions depend in turn on the C macro F77\_FUNC defined in the header file sundials\_config.h. The mapping defined by F77\_FUNC in turn transforms the C interface names to match the name-mangling approach used by the supplied Fortran compiler.

By "name-mangling", we mean that due to the case-independent nature of the FORTRAN language, FORTRAN compilers convert all subroutine and object names to use either all lower-case or all uppercase characters, and append either zero, one or two underscores as a prefix or suffix to the name. For example, the FORTRAN subroutine MyFunction() will be changed to one of myfunction, MYFUNCTION, myfunction..., MYFUNCTION., and so on, depending on the FORTRAN compiler used.

SUNDIALS determines this name-mangling scheme at configuration time (see Appendix A).

# 5.2 Fortran Data Types

Throughout this documentation, we will refer to data types according to their usage in C. The equivalent types to these may vary, depending on your computer architecture and on how SUNDIALS was compiled (see Appendix A). A FORTRAN user should first determine the equivalent types for their architecture and compiler, and then take care that all arguments passed through this FORTRAN/C interface are declared of the appropriate type.

Integers: SUNDIALS uses both int and long int types:

- int equivalent to an INTEGER or INTEGER\*4 in FORTRAN
- long int this will depend on the computer architecture:
  - 32-bit architecture equivalent to an INTEGER or INTEGER\*4 in FORTRAN
  - 64-bit architecture equivalent to an INTEGER\*8 in FORTRAN

Real numbers: As discussed in Appendix A, at compilation SUNDIALS allows the configuration option --with-precision, that accepts values of single, double or extended (the default is double). This choice dictates the size of a realtype variable. The corresponding FORTRAN types for these realtype sizes are:

- single equivalent to a REAL or REAL\*4 in FORTRAN
- double equivalent to a DOUBLE PRECISION or REAL\*8 in FORTRAN
- extended equivalent to a REAL\*16 in FORTRAN

#### 5.3 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 N\_VNewEmpty\_Serial.
  - FNVINITOMP (defined by NVECTOR\_OPENMP) interfaces to N\_VNewEmpty\_OpenMP.
  - FNVINITPTS (defined by NVECTOR\_PTHREADS) interfaces to N\_VNewEmpty\_Pthreads.
  - FNVINITP (defined by NVECTOR\_PARALLEL) interfaces to N\_VNewEmpty\_Parallel.
- Interface to the main KINSOL module
  - FKINMALLOC interfaces to KINCreate, KINSetUserData, and KINInit.
  - FKINSETIIN and FKINSETRIN interface to KINSet\* functions.
  - FKINSETVIN interfaces to KINSetConstraints.
  - FKINSOL interfaces to KINSol, KINGet\* functions, and to the optional output functions for the selected linear solver module.
  - FKINFREE interfaces to KINFree.
- Interface to the linear solver modules
  - FKINDENSE interfaces to KINDense.
  - FKINDENSESETJAC interfaces to KINDlsSetDenseJacFn.
  - FKINLAPACKDENSE interfaces to KINLapackDense.
  - FKINLAPACKDENSESETJAC interfaces to KINDlsSetDenseJacFn.
  - FKINBAND interfaces to KINBand.
  - FKINBANDSETJAC interfaces to KINDlsSetBandJacFn.
  - $-\ {\tt FKINLAPACKBAND}$  interfaces to  ${\tt KINLapackBand}.$
  - FKINLAPACKBANDSETJAC interfaces to KINDlsSetBandJacFn.
  - FKINKLU interfaces to KINKLU.
  - FKINKLUREINIT interfaces to KINKLUReInit.
  - FKINSUPERLUMT interfaces to KINSuperLUMT.
  - FKINSPGMR interfaces to KINSpgmr and SPGMR optional input functions.
  - FKINSPFGMR interfaces to KINSpfgmr and SPFGMR optional input functions.
  - FKINSPBCG interfaces to KINSpbcg and SPBCG optional input functions.
  - FKINSPTFQMR interfaces to KINSptfqmr and SPTFQMR optional input functions.
  - FKINSPILSSETJAC interfaces to KINSpilsSetJacTimesVecFn.
  - FKINSPILSSETPREC interfaces to KINSpilsSetPreconditioner.

The user-supplied functions,	each listed with th	e corresponding i	internal interface	function which
calls it (and its type within KINS	SOL), are as follows:			

FKINSOL routine	KINSOL function	KINSOL type of		
(FORTRAN, user-supplied)	(C, interface)	interface function		
FKFUN	FKINfunc	KINSysFn		
FKDJAC	FKINDenseJac	KINDlsDenseJacFn		
	FKINLapackDenseJac	KINDlsDenseJacFn		
FKBJAC	FKINBandJac	KINDlsBandJacFn		
	FKINLapackBandJac	KINDlsBandJacFn		
FKINSPJAC	FKINSparseJac	KINSlsSparseJacFn		
FKPSET	FKINPSet	KINSpilsPrecSetupFn		
FKPSOL	FKINPSol	KINSpilsPrecSolveFn		
FKJTIMES	FKINJtimes	KINSpilsJacTimesVecFn		

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.

# 5.4 Usage of the FKINSOL interface module

The usage of FKINSOL requires calls to a few different 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.

In the instructions below, steps marked [S] apply to the NVECTOR module NVECTOR\_SERIAL, steps marked [O] apply to NVECTOR\_OPENMP, steps marked [T] apply to NVECTOR\_PTHREADS, while steps marked [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, IER)
DIMENSION U(*), FVAL(*)
```

It must set the FVAL array to F(u), the system function, as a function of U = u. IER is an error return flag which should be set to 0 if successful, a positive value if a recoverable error occurred (in which case KINSOL will attempt to correct), or a negative value if it failed unrecoverably (in which case the solution process is halted).

# 2. NVECTOR module initialization

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

```
CALL FNVINITS (KEY, NEQ, IER)
```

where KEY is the solver id (KEY = 3 for KINSOL), NEQ is the size of vectors, and IER is a return completion flag which is 0 on success and -1 if a failure occurred.

[O] To initialize the NVECTOR\_OPENMP NVECTOR module, the user must make the following call:

where KEY is the solver id (KEY = 3 for KINSOL), NEQ is the size of vectors, NUMTHREADS is the number of threads, and IER is a return completion flag which is 0 on success and -1 if a failure occurred.

[T] To initialize the NVECTOR\_PTHREADS NVECTOR module, the user must make the following call:

CALL FNVINITPTS (KEY, NEQ, NUMTHREADS, IER)

where KEY is the solver id (KEY = 3 for KINSOL), NEQ is the size of vectors, NUMTHREADS is the number of threads, and IER is a return completion flag which is 0 on success and -1 if a failure occurred.

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

CALL FNVINITP (COMM, KEY, NLOCAL, NGLOBAL, IER)

in which the arguments are: COMM = MPI communicator, KEY = 3 for KINSOL, NLOCAL = the local size of vectors on this processor, and NGLOBAL = the system size (and the global size of all vectors, equal to the sum of all values of NLOCAL). The return flag IER is set to 0 on a successful return and to -1 otherwise.

NOTE: The integers NEQ, NLOCAL, and NGLOBAL should be declared so as to match C type long int.

If the header file sundials\_config.h defines SUNDIALS\_MPI\_COMM\_F2C to be 1 (meaning the MPI implementation used to build SUNDIALS includes the MPI\_Comm\_f2c function), then COMM can be any valid MPI communicator. Otherwise, MPI\_COMM\_WORLD will be used, so just pass an integer value as a placeholder.

# 3. Problem specification

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

# FKINMALLOC

Call CALL FKINMALLOC (IOUT, ROUT, IER)

Description This function specifies the optional output arrays, allocates internal memory, and

initializes KINSOL.

Arguments IOUT is an integer array for integer optional outputs.

ROUT is a real array for real optional outputs.

Return value IER is the return completion flag. Values are 0 for successful return and -1 other-

wise. See printed message for details in case of failure.

Notes The user integer data array IOUT must be declared as INTEGER\*4 or INTEGER\*8

according to the C type long int.

The optional outputs associated with the main KINSOL integrator are listed in Ta-

ble 5.2.

### 4. Set optional inputs

Call FKINSETIIN, FKINSETRIN, and/or FKINSETVIN, to set desired optional inputs, if any. See §5.5 for details.

# 5. Linear solver specification



The solution method in KINSOL involves the solution of linear systems related to the Jacobian of the nonlinear system. The user of FKINSOL must call a routine with a specific name to make the desired choice of linear solver.

# [S] Dense treatment of the linear system

To use the direct dense linear solver based on the internal KINSOL implementation, the user must make the call:

```
CALL FKINDENSE (NEQ, IER)
```

where NEQ is the size of the nonlinear system. The argument IER is an error return flag which is 0 for success, -1 if a memory allocation failure occurred, or -2 for illegal input.

Alternatively, to use the Lapack-based direct dense linear solver, the user must make the call:

```
CALL FKINLAPACKDENSE(NEQ, IER)
```

where the arguments have the same meanings as for FKINDENSE, except that here NEQ must be declared so as to match C type int.

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

```
SUBROUTINE FKDJAC (NEQ, U, FVAL, DJAC, WK1, WK2, IER) DIMENSION U(*), FVAL(*), DJAC(NEQ,*), WK1(*), WK2(*)
```

Typically this routine will use only NEQ, U, and DJAC. It must compute the Jacobian and store it columnwise in DJAC. The input arguments U and FVAL contain the current values of u and F(u), respectively. The vectors WK1 and WK2, of length NEQ, are provided as work space for use in FKDJAC. IER is an error return flag which should be set to 0 if successful, a positive value if a recoverable error occurred (in which case KINSOL will attempt to correct), or a negative value if FKDJAC failed unrecoverably (in which case the solution process is halted). NOTE: The argument NEQ has a type consistent with C type long int even in the case when the Lapack dense solver is to be used.

If the FKDJAC routine is provided, then, following the call to FKINDENSE, the user must make the call:

```
CALL FKINDENSESETJAC (FLAG, IER)
```

with  $\mathtt{FLAG} \neq 0$  to specify use of the user-supplied Jacobian approximation. The argument <code>IER</code> is an error return flag which is 0 for success or non-zero if an error occurred. If using the Lapack-based direct dense linear solver, the use of a Jacobian approximation supplied by the user is indicated through the call

```
CALL FKINLAPACKDENSESETJAC (FLAG, IER)
```

Optional outputs specific to the DENSE case are listed in Table 5.2.

## [S] Band treatment of the linear system

To use the direct band linear solver based on the internal KINSOL implementation, the user must make the call:

```
CALL FKINBAND (NEQ, MU, ML, IER)
```

The arguments are: MU, the upper half-bandwidth; ML, the lower half-bandwidth; and IER, an error return flag which is 0 for success, -1 if a memory allocation failure occurred, or -2 in case an input has an illegal value.

Alternatively, to use the Lapack-based direct band linear solver, the user must make the call:

```
CALL FKINLAPACKBAND (NEQ, MU, ML, IER)
```

where the arguments have the same meanings as for FKINBAND, except that here NEQ, MU, and ML must be declared so as to match C type int.

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

```
SUBROUTINE FKBJAC (NEQ, MU, ML, MDIM, U, FVAL, BJAC, WK1, WK2, IER) DIMENSION U(*), FVAL(*), BJAC(MDIM,*), WK1(*), WK2(*)
```

Typically this routine will use only NEQ, MU, ML, U, and BJAC. It must load the MDIM by N array BJAC with the Jacobian matrix at the current u in band form. Store in BJAC(k,j) the Jacobian element  $J_{i,j}$  with k=i-j+ MU +1 ( $k=1\cdots$  ML + MU + 1) and  $j=1\cdots N$ . The input arguments U and FVAL contain the current values of u, and F(u), respectively. The vectors WK1 and WK2 of length NEQ are provided as work space for use in FKBJAC. IER is an error return flag, which should be set to 0 if successful, a positive value if a recoverable error occurred (in which case KINSOL will attempt to correct), or a negative value if FKBJAC failed unrecoverably (in which case the solution process is halted). NOTE: The arguments NEQ, MU, ML, and MDIM have a type consistent with C type long int even in the case when the Lapack band solver is to be used.

If the FKBJAC routine is provided, then, following the call to FKINBAND, the user must make the call:

```
CALL FKINBANDSETJAC (FLAG, IER)
```

with  ${\tt FLAG} \neq 0$  to specify use of the user-supplied Jacobian approximation. The argument IER is an error return flag which is 0 for success or non-zero if an error occurred. If using the Lapack-based direct band linear solver, the use of a Jacobian approximation supplied by the user is indicated through the call

```
CALL FKINLAPACKNBANDSETJAC (FLAG, IER)
```

Optional outputs specific to the BAND case are listed in Table 5.2.

# [S] Sparse direct treatment of the linear system

To use the KLU sparse direct linear solver, the user must make the call:

```
CALL FKINKLU (NEQ, NNZ, ORDERING, IER)
```

where NEQ is the size of the nonlinear system, NNZ is the maximum number of nonzeros in the Jacobian matrix, and ORDERING is the matrix ordering desired with possible values from the KLU package (0 = AMD, 1 = COLAMD). The argument IER is an error return flag which is 0 for success or negative for an error.

The KINSOL KLU solver will reuse much of the factorization information from one nonlinear iteration to the next. If at any time the user wants to force a full refactorization or if the number of nonzeros in the Jacobian matrix changes, the user should make the call

```
CALL FKINKLUREINIT(NEQ, NNZ, REINIT_TYPE)
```

where NEQ is the size of the nonlinear system, NNZ is the maximum number of nonzeros in the Jacobian matrix, and REINIT\_TYPE is 1 or 2. For a value of 1, the matrix will be destroyed and a new one will be allocated with NNZ nonzeros. For a value of 2, only symbolic and numeric factorizations will be completed.

Alternatively, to use the SuperLUMT linear solver, the user must make the call:

```
CALL FKINSUPERLUMT (NEQ, NNZ, ORDERING, IER)
```

where the arguments have the same meanings as for FKINKLU, except that here possible values for ORDERING derive from the SUPERLUMT package and include: 0 for Natural ordering, 1 for Minimum degree on  $A^TA$ , 2 for Minimum degree on  $A^T+A$ , and 3 for COLAMD.

If the either of the sparse direct interface packages are used, then the user must supply the FKINSPJAC routine that computes a compressed-sparse-column approximation of the system Jacobian J = dF(y)/dy. If supplied, it must have the following form:

```
SUBROUTINE FKINSPJAC(Y, FY, N, NNZ, JDATA, JRVALS, & JCPTRS, WK1, WK2, IER)
```

Typically this routine will use only N, NNZ, JDATA, JRVALS and JCPTRS. It must load the N by N compressed sparse column matrix with storage for NNZ nonzeros, stored in the arrays JDATA (nonzero values), JRVALS (row indices for each nonzero), JCOLPTRS (indices for start of each column), with the Jacobian matrix at the current (y) in CSC form (see sundials\_sparse.h for more information). The arguments are Y, an array containing state variables, FY, an array containing residual values, N, the number of matrix rows/columns in the Jacobian, NNZ, allocated length of nonzero storage, JDATA, nonzero values in the Jacobian (of length NNZ), JRVALS, row indices for each nonzero in Jacobian (of length NNZ), JCPTRS, pointers to each Jacobian column in the two preceding arrays (of length N+1), WK\*, work arrays containing temporary workspace of same size as Y, and IER, error return code (0 if successful, ¿0 if a recoverable error occurred, or ¡0 if an unrecoverable error occurred.)

Optional outputs specific to the DENSE case are listed in Table 5.2.

# [S][P] SPGMR and SPFGMR treatment of the linear systems

For the Scaled Preconditioned GMRES or the Scaled Preconditioned Flexible GMRES solution of the linear systems, the user must make either the call

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

or the call

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

The arguments are as follows. MAXL is the maximum Krylov subspace dimension. MAXLRST is the maximum number of restarts. IER is an error return flag which is 0 to indicate success, -1 if a memory allocation failure occurred, or -2 to indicate an illegal input.

Optional outputs specific to the SPGMR and SPFGMR cases are listed in Table 5.2.

For descriptions of the relevant optional user-supplied routines, see User-supplied routines for SPGMR/SPFMGR/SPBCG/SPTFQMR below.

```
[S][P] SPBCG treatment of the linear systems
```

For the Scaled Preconditioned Bi-CGStab solution of the linear systems, the user must make the call

```
CALL FKINSPBCG (MAXL, IER)
```

Its arguments are the same as those with the same names for FKINSPGMR.

Optional outputs specific to the SPBCG case are listed in Table 5.2.

For descriptions of the relevant optional user-supplied routines, see below.

# [S][P] SPTFQMR treatment of the linear systems

For the Scaled Preconditioned Transpose-Free Quasi-Minimal Residual solution of the linear systems, the user must make the call

```
CALL FKINSPTFQMR (MAXL, IER)
```

Its arguments are the same as those with the same names for FKINSPGMR.

Optional outputs specific to the SPTFQMR case are listed in Table 5.2.

For descriptions of the relevant optional user-supplied routines, see below.

# [S][P] Functions used by SPGMR/SPFGMR/SPBCG/SPTFQMR

An optional user-supplied routine, FKINJTIMES (see below), can be provided for Jacobian-vector products. (Note that this routine is required if Picard iteration is being used.) If it is, then, following the call to FKINSPGMR, FKINSPFGMR, FKINSPBCG, or FKINSPTFQMR, the user must make the call:

```
CALL FKINSPILSSETJAC (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 is 0 for success or non-zero if an error occurred.

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

```
CALL FKINSPILSSETPREC (FLAG, IER)
```

with FLAG  $\neq 0$ . The return flag IER is 0 if successful, or negative if a memory error occurred. In addition, the user program must include preconditioner routines FKPSOL and FKPSET (see below).

# [S][P] User-supplied routines for SPGMR/SPFGMR/SPBCG/SPTFQMR

With treatment of the linear systems by any of the Krylov iterative solvers, there are three optional user-supplied routines — FKINJTIMES, FKPSOL, and FKPSET. The specifications for these routines are given below.

As an option when using the SPGMR, SPFGMR, SPBCG, or SPTFQMR linear solvers, 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 FKINJTIMES (V, FJV, NEWU, U, IER) DIMENSION V(*), FJV(*), U(*)
```

Typically this routine will use only U, V, and FJV. It must compute the product vector Jv, where the vector v is stored in V, and store the product in FJV. The input argument U contains the current value of u. On return, set IER = 0 if FKINJTIMES was successful, and nonzero otherwise. NEWU is a flag to indicate if U has been changed since the last call; if it has, then NEWU = 1, and FKINJTIMES should recompute any saved Jacobian data it uses and reset NEWU to 0. (See §4.6.7.)

If preconditioning is to be included, the following routine must be supplied, 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.

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

If the user calls FKINSPILSSETPREC, the routine FKPSET must be provided, even if it is not needed, and then it should return IER = 0.



## 6. 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 (0 specifies Inexact Newton, 1 indicates Newton with line search, 2 indicates Picard iteration, and 3 indicates Fixed Point iteration). 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. These values correspond to the KINSo1 returns (see §4.5.3 and §B.2). The values of the optional outputs are available in IOPT and ROPT (see Table 5.2).

# 7. Memory deallocation

To free the internal memory created by the call to FKINMALLOC, make the call

CALL FKINFREE

# 5.5 FKINSOL optional input and output

In order to keep the number of user-callable FKINSOL interface routines to a minimum, optional inputs to the KINSOL solver are passed through only three routines: FKINSETIIN for integer optional inputs, FKINSETRIN for real optional inputs, and FKINSETVIN for real vector (array) optional inputs. These functions should be called as follows:

```
CALL FKINSETIIN (KEY, IVAL, IER)
CALL FKINSETRIN (KEY, RVAL, IER)
CALL FKINSETVIN (KEY, VVAL, IER)
```

integer optional inputs rainseriin			
Key Optional input		Default value	
PRNT_LEVEL	Verbosity level of output	0	
MAX_NITER	Maximum no. of nonlinear iterations	200	
ETA_FORM	Form of $\eta$ coefficient	1 (KIN_ETACHOICE1)	
MAX_SETUPS	Maximum no. of iterations without prec. setup	10	
MAX_SP_SETUPS	Maximum no. of iterations without residual check	5	
NO_INIT_SETUP	No initial preconditioner setup	FALSE	
NO_MIN_EPS	Lower bound on $\epsilon$	FALSE	
NO_RES_MON	No residual monitoring	FALSE	

Table 5.1: Keys for setting FKINSOL optional inputs

# Real optional inputs (FKINSETRIN)

Key	Optional input	Default value
FNORM_TOL	Function-norm stopping tolerance	$uround^{1/3}$
SSTEP_TOL	Scaled-step stopping tolerance	$uround^{2/3}$
MAX_STEP	Max. scaled length of Newton step	$  1000  D_uu_0  _2$
RERR_FUNC	Relative error for F.D. $Jv$	$\sqrt{\text{uround}}$
ETA_CONST	Constant value of $\eta$	0.1
ETA_PARAMS	Values of $\gamma$ and $\alpha$	0.9 and 2.0
RMON_CONST	Constant value of $\omega$	0.9
RMON_PARAMS	Values of $\omega_{min}$ and $\omega_{max}$	0.00001 and 0.9

where KEY is a quoted string indicating which optional input is set, IVAL is the integer input value to be used, RVAL is the real input value to be used, and VVAL is the input real array to be used. IER is an integer return flag which is set to 0 on success and a negative value if a failure occurred. For the legal values of KEY in calls to FKINSETIIN and FKINSETRIN, see Table 5.1. The one legal value of KEY for FKINSETVIN is CONSTR\_VEC, for providing the array of inequality constraints to be imposed on the solution, if any. The integer IVAL should be declared in a manner consistent with C type long int.

The optional outputs from the KINSOL solver are accessed not through individual functions, but rather through a pair of arrays, IOUT (integer type) of dimension at least 15, and ROUT (real type) of dimension at least 2. These arrays are owned (and allocated) by the user and are passed as arguments to FKINMALLOC. Table 5.2 lists the entries in these two arrays and specifies the optional variable as well as the KINSOL function which is actually called to extract the optional output.

For more details on the optional inputs and outputs, see §4.5.4 and §4.5.5.

# 5.6 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 §4.7), 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 KINBBDPrecInit.
- FKINBBDOPT interfaces to KINBBDPRE optional output functions.

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):

Table 5.2: Description of the fkinsol optional output arrays  ${\tt IOUT}$  and  ${\tt ROUT}$ 

	Integer output array IOUT			
Index	Optional output			
	KINSOL main solver			
1	LENRW	KINGetWorkSpace		
2	LENIW	KINGetWorkSpace		
3	NNI	KINGetNumNonlinSolvIters		
4	NFE	KINGetNumFuncEvals		
5	NBCF	KINGetNumBetaCondFails		
6	NBKTRK	KINGetNumBacktrackOps		
	KINDENSE, KIN	NBAND linear solver		
7	LENRWLS	KINDlsGetWorkSpace		
8	LENIWLS	KINDlsGetWorkSpace		
9	$\mathtt{LS\_FLAG}$	KINDlsGetLastFlag		
10	NFELS	KINDlsGetNumFuncEvals		
11	NJE	KINDlsGetNumJacEvals		
KINSPARSE linea		E linear solver		
8	LS_FLAG	KINSlsGetLastFlag		
10	NJE	KINSlsGetNumJacEvals		
KINSPGI	MR, KINSPFGMR, KINS	PBCG, KINSPTFQMR linear solvers		
7	LENRWLS	KINSpilsGetWorkSpace		
8	LENIWLS	KINSpilsGetWorkSpace		
9	$LS\_FLAG$	KINSpilsGetLastFlag		
10	NFELS	KINSpilsGetNumFuncEvals		
11	NJTV	KINSpilsGetNumJacEvals		
12	NPE	KINSpilsGetNumPrecEvals		
13	NPS	KINSpilsGetNumPrecSolves		
14	NLI	KINSpilsGetNumLinIters		
15	NCFL	KINSpilsGetNumConvFails		

# Real output array $\mathtt{ROUT}$

Index	Optional output	KINSOL function
1	FNORM	KINGetFuncNorm
2	SSTEP	KINGetStepLength

FKINBBD routine	KINSOL function	KINSOL type of		
(FORTRAN, user-supplied)	(C, interface)	interface function		
FKLOCFN	FKINgloc	KINLocalFn		
FKCOMMF	FKINgcomm	KINCommFn		
FKJTIMES	FKINJtimes	KINSpilsJacTimesVecFn		

As with the rest of the FKINSOL routines, the names of all user-supplied routines here are fixed, in order to maximize portability for the resulting mixed-language program. Additionally, based on flags discussed above in §5.3, the names of the user-supplied routines are mapped to actual values through a series of definitions in the header file fkinbbd.h.

The following is a summary of the usage of this module. Steps that are unchanged from the main program described in §5.4 are grayed-out.

- 1. Nonlinear system function specification
- 2. NVECTOR module initialization
- 3. Problem specification
- 4. Set optional inputs

# 5. Linear solver specification

First, specify one of the KINSPILS iterative linear solvers, by calling one of FKINSPGMR, FKINSPFGRM, FKINSPBCG, or FKINSPTFQMR.

To initialize the KINBBDPRE preconditioner, make the following call:

The arguments are as follows. NLOCAL is the local size of vectors for this process. MUDQ and MLDQ 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. MU and ML are the upper and lower half-bandwidths of the band matrix that is retained as an approximation of the local Jacobian block; these may be smaller than MUDQ and MLDQ. 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.

Optionally, to specify that SPGMR, SPFGMR, SPBCG, or SPTFQMR should use the supplied FKJTIMES, make the call

```
CALL FKINSPILSSETJAC (FLAG, IER)
```

with FLAG  $\neq 0$ . (See step 5 in §5.4).

6. Problem solution

# 7. KINBBDPRE **Optional outputs**

Optional outputs specific to the SPGMR, SPFGMR, SPBCG, or SPTFQMR solver are listed in Table 5.2. To obtain the optional outputs associated with the KINBBDPRE module, make the following call:

```
CALL FKINBBDOPT (LENRBBD, LENIBBD, NGEBBD)
```

The arguments should be consistent with C type long int. Their returned values are as follows: LENRBBD is the length of real preconditioner work space, in realtype words. LENIBBD is the length of integer preconditioner work space, in integer words. These sizes are local to the current process. NGEBBD is the cumulative number of G(u) evaluations (calls to FKLOCFN) so far.

# 8. Memory deallocation

(The memory allocated for the FKINBBD module is deallocated automatically by FKINFREE.)

# 9. User-supplied routines

The following two routines must be supplied for use with the KINBBDPRE module:

```
SUBROUTINE FKLOCFN (NLOC, ULOC, GLOC, IER)
DIMENSION ULOC(*), GLOC(*)
```

This routine is to evaluate the function G(u) approximating F (possibly identical to F), in terms of the array ULOC (of length NLOC), which is the sub-vector of u local to this processor. The resulting (local) sub-vector is to be stored in the array GLOC. IER is an error return flag which should be set to 0 if successful, a positive value if a recoverable error occurred (in which case KINSOL will attempt to correct), or a negative value if FKLOCFN failed unrecoverably (in which case the solution process is halted).

```
SUBROUTINE FKCOMMFN (NLOC, ULOC, IER) DIMENSION ULOC(*)
```

This routine is to perform the inter-processor communication necessary for the FKLOCFN routine. Each call to FKCOMMFN is preceded by a call to the system function routine FKFUN with the same argument ULOC. IER is an error return flag which should be set to 0 if successful, a positive value if a recoverable error occurred (in which case KINSOL will attempt to correct), or a negative value if FKCOMMFN failed recoverably (in which case the solution process is halted).

The subroutine FKCOMMFN must be supplied even if it is not needed and must return IER = 0.

Optionally, the user can supply a routine FKINJTIMES for the evaluation of Jacobian-vector products, as described above in step 5 in §5.4. Note that this routine is required if using Picard iteration.



# 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 four provided within SUNDIALS — a serial implementation and three parallel implementations. The generic operations are described below. In the sections following, the implementations provided with SUNDIALS are described.

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);
  void
              (*nvspace)(N_Vector, long int *, long int *);
              (*nvgetarraypointer)(N_Vector);
  realtype*
              (*nvsetarraypointer)(realtype *, N_Vector);
  void
              (*nvlinearsum)(realtype, N_Vector, realtype, N_Vector, N_Vector);
  void
              (*nvconst)(realtype, N_Vector);
  void
  void
              (*nvprod)(N_Vector, N_Vector, N_Vector);
              (*nvdiv)(N_Vector, N_Vector, N_Vector);
  void
              (*nvscale)(realtype, N_Vector, N_Vector);
  void
  void
              (*nvabs)(N_Vector, N_Vector);
              (*nvinv)(N_Vector, N_Vector);
  void
  void
              (*nvaddconst)(N_Vector, realtype, N_Vector);
              (*nvdotprod)(N_Vector, N_Vector);
  realtype
  realtype
              (*nvmaxnorm)(N_Vector);
  realtype
              (*nvwrmsnorm)(N_Vector, N_Vector);
```

```
realtype (*nvwrmsnormmask)(N_Vector, N_Vector);
realtype (*nvwin)(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 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  $N_{\text{-}}Vector$  can be destroyed by calling  $N_{\text{-}}VDestroyVectorArray$ , 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 data.</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. This function is advisory only, for use in determining a user's total space requirements; it could be a dummy function in a user-supplied NVECTOR module if that information is not of interest.
N_VGetArrayPointer	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 (serial) linear solvers, the sparse linear solvers (serial and threaded), and in the interfaces to the banded (serial) and band-block-diagonal (parallel) preconditioner modules provided with SUNDIALS.
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 (serial) linear solver, hence need not exist in a user-supplied NVECTOR module for a parallel environment.
N_VLinearSum	N_VLinearSum(a, x, b, y, z); Performs the operation $z = ax + by$ , where a and b are realtype 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 realtype c: $z_i=c,\ i=0,\dots,n-1.$
1	continued on next page

Name	Usage and Description
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: $z_i = x_i y_i$ , $i = 0,, n-1$ .
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: $z_i = x_i/y_i$ , $i = 0, \ldots, n-1$ . The $y_i$ may not be tested for 0 values. It should only be called with a y that is guaranteed to have all nonzero components.
N_VScale	N_VScale(c, x, z); Scales the N_Vector x by the realtype scalar c and returns the result in z: $z_i = cx_i, i = 0, \ldots, n-1$ .
N_VAbs	N_VAbs(x, z); Sets the components of the N_Vector z to be the absolute values of the components of the N_Vector x: $y_i =  x_i , i = 0, \ldots, n-1$ .
N_VInv	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	N_VAddConst(x, b, z); Adds the realtype scalar b to all components of x and returns the result in the N_Vector z: $z_i = x_i + b, \ i = 0, \ldots, n-1$ .
N_VDotProd	d = N_VDotProd(x, y); Returns the value of the ordinary dot product of x and y: $d = \sum_{i=0}^{n-1} x_i y_i$ .
N_VMaxNorm	$m = N_VMaxNorm(x);$ Returns the maximum norm of the N_Vector x: $m = \max_i  x_i .$
N_VWrmsNorm	$m = N_{VWrmsNorm}(x, w)$ Returns the weighted root-mean-square norm of the $N_{Vector}(x, w)$
N_VWrmsNormMask	realtype weight vector w: $m = \sqrt{\left(\sum_{i=0}^{n-1}(x_iw_i)^2\right)/n}$ . $\mathbf{m} = \mathbf{N}_{\text{VWrmsNormMask}}(\mathbf{x}, \mathbf{w}, \text{id});$ Returns the weighted root mean square norm of the N_Vector $\mathbf{x}$ with realtype weight vector $\mathbf{w}$ built using only the elements of $\mathbf{x}$ corresponding to nonzero elements of the N_Vector $\mathbf{id}$ : $m = \sqrt{\left(\sum_{i=0}^{n-1}(x_iw_i\text{sign}(id_i))^2\right)/n}.$
N_VMin	$m = \sqrt{\sum_{i=0}^{n} (x_i w_i \operatorname{sign}(\alpha u_i))}$ / $m = N_{\text{L}} \operatorname{VMin}(\mathbf{x})$ ;  Returns the smallest element of the $N_{\text{L}} \operatorname{Vector} \mathbf{x}$ : $m = \min_{i} x_i$ . $continued on next page$

continued from last pag	e
Name	Usage and Description
N_VWL2Norm	m = N_VWL2Norm(x, w); Returns the weighted Euclidean $\ell_2$ norm of the N_Vector x with realtype weight vector w: $m = \sqrt{\sum_{i=0}^{n-1} (x_i w_i)^2}$ .
N_VL1Norm	m = N_VL1Norm(x); Returns the $\ell_1$ norm of the N_Vector x: $m = \sum_{i=0}^{n-1}  x_i $ .
N_VCompare	N_VCompare(c, x, z); Compares the components of the N_Vector x to the realtype 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 a boolean assigned to 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 a boolean assigned to FALSE if any element failed the constraint test and assigned to 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 <sub>i</sub> by denom <sub>i</sub> . 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 sundials_types.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 \_S in the names denotes the 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  $N_{content}$  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  $N_Vector v$ . 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  $r = NV_ih_s(v,i)$  sets r to be the value of the i-th component of v. The assignment  $NV_ih_s(v,i) = r$  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. Their names are obtained from those in Table 6.1 by appending the suffix \_Serial. The module NVECTOR\_SERIAL provides the following additional 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\_VCloneVectorArray\_Serial

This function creates (by cloning) an array of count serial vectors.

```
N_Vector *N_VCloneVectorArray_Serial(int count, N_Vector w);
```

# • N\_VCloneEmptyVectorArray\_Serial

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

```
N_Vector *N_VCloneEmptyVectorArray_Serial(int count, N_Vector w);
```

• N\_VDestroyVectorArray\_Serial

This function frees memory allocated for the array of count variables of type N\_Vector created with N\_VCloneVectorArray\_Serial or with N\_VCloneEmptyVectorArray\_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.
- N\_VNewEmpty\_Serial, N\_VMake\_Serial, and N\_VCloneEmptyVectorArray\_Serial set the field own\_data = FALSE. 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 NVECTOR\_PARALLEL implementation of the NVECTOR module provided with SUNDIALS is based on MPI. It 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 the distributed memory 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  $N_V$ ctor 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 Tth P

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

The assignment  $r = NV_{i,i}$  sets r to be the value of the i-th component of the local part of v. The assignment  $NV_{i,i} = r$  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 Their names are obtained from those in Table 6.1 by appending the suffix \_Parallel. The module NVECTOR\_PARALLEL provides the following additional 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\_VCloneVectorArray\_Parallel

This function creates (by cloning) an array of count parallel vectors.

```
N_Vector *N_VCloneVectorArray_Parallel(int count, N_Vector w);
```

• N\_VCloneEmptyVectorArray\_Parallel

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

```
N_Vector *N_VCloneEmptyVectorArray_Parallel(int count, N_Vector w);
```

• N\_VDestroyVectorArray\_Parallel

This function frees memory allocated for the array of count variables of type N\_Vector created with N\_VCloneVectorArray\_Parallel or with N\_VCloneEmptyVectorArray\_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.
- N\_VNewEmpty\_Parallel, N\_VMake\_Parallel, and N\_VCloneEmptyVectorArray\_Parallel set the field own\_data = FALSE. 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.
- To maximize efficiency, vector operations in the NVECTOR\_PARALLEL 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.

# 6.3 The NVECTOR\_OPENMP implementation

In situations where a user has a multi-core processing unit capable of running multiple parallel threads with shared memory, SUNDIALS provides an implementation of NVECTOR using OpenMP, called NVECTOR\_OPENMP, and an implementation using pThreads, called NVECTOR\_PTHREADS. Testing has shown that vectors should be of length at least 100,000 before the overhead associated with creating and using the threads is made up by the parallelism in the vector calculations.

The OpenMP NVECTOR implementation provided with SUNDIALS, NVECTOR\_OPENMP, 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, a boolean flag own\_data which specifies the ownership of data, and the number of threads. Operations on the vector are threaded using OpenMP.





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

The following six macros are provided to access the content of an NVECTOR\_OPENMP vector. The suffix \_OMP in the names denotes the OpenMP version.

# • NV\_CONTENT\_OMP

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

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

Implementation:

```
#define NV_CONTENT_OMP(v) ( (N_VectorContent_OpenMP)(v->content) )
```

• NV\_OWN\_DATA\_OMP, NV\_DATA\_OMP, NV\_LENGTH\_OMP, NV\_NUM\_THREADS\_OMP

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

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

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

The assignment v\_num\_threads = NV\_NUM\_THREADS\_OMP(v) sets v\_num\_threads to be the number of threads from v. On the other hand, the call NV\_NUM\_THREADS\_OMP(v) = num\_threads\_v sets the number of threads for v to be num\_threads\_v.

Implementation:

```
#define NV_OWN_DATA_OMP(v) ( NV_CONTENT_OMP(v)->own_data )
#define NV_DATA_OMP(v) ( NV_CONTENT_OMP(v)->data )
#define NV_LENGTH_OMP(v) ( NV_CONTENT_OMP(v)->length )
#define NV_NUM_THREADS_OMP(v) ( NV_CONTENT_OMP(v)->num_threads )
```

## • NV\_Ith\_OMP

This macro gives access to the individual components of the data array of an N-Vector.

The assignment  $r = NV_{in}(v,i)$  sets r to be the value of the i-th component of v. The assignment  $NV_{in}(v,i) = r$  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_OMP(v,i) ( NV_DATA_OMP(v)[i] )
```

The NVECTOR\_OPENMP module defines OpenMP implementations of all vector operations listed in Table 6.1. Their names are obtained from those in Table 6.1 by appending the suffix \_OpenMP. The module NVECTOR\_OPENMP provides the following additional user-callable routines:

# • N\_VNew\_OpenMP

This function creates and allocates memory for a OpenMP N\_Vector. Arguments are the vector length and number of threads.

```
N_Vector N_VNew_OpenMP(long int vec_length, int num_threads);
```

# • N\_VNewEmpty\_OpenMP

This function creates a new OpenMP N\_Vector with an empty (NULL) data array. N\_Vector N\_VNewEmpty\_OpenMP(long int vec\_length, int num\_threads);

• N\_VMake\_OpenMP

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

• N\_VCloneVectorArray\_OpenMP

This function creates (by cloning) an array of count OpenMP vectors.

N\_Vector \*N\_VCloneVectorArray\_OpenMP(int count, N\_Vector w);

• N\_VCloneEmptyVectorArray\_OpenMP

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

N\_Vector \*N\_VCloneEmptyVectorArray\_OpenMP(int count, N\_Vector w);

• N\_VDestroyVectorArray\_OpenMP

This function frees memory allocated for the array of count variables of type N\_Vector created with N\_VCloneVectorArray\_OpenMP or with N\_VCloneEmptyVectorArray\_OpenMP.

void N\_VDestroyVectorArray\_OpenMP(N\_Vector \*vs, int count);

• N\_VPrint\_OpenMP

This function prints the content of a OpenMP vector to stdout. void N\_VPrint\_OpenMP(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\_OMP(v) and then access v\_data[i] within the loop than it is to use NV\_Ith\_OMP(v,i) within the loop.
- N\_VNewEmpty\_OpenMP, N\_VMake\_OpenMP, and N\_VCloneEmptyVectorArray\_OpenMP set the field  $own\_data = FALSE$ . N\_VDestroy\_OpenMP and N\_VDestroyVectorArray\_OpenMP 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\_OPENMP 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.

# 6.4 The NVECTOR\_PTHREADS implementation

In situations where a user has a multi-core processing unit capable of running multiple parallel threads with shared memory, SUNDIALS provides an implementation of NVECTOR using OpenMP, called NVECTOR\_OPENMP, and an implementation using pThreads, called NVECTOR\_PTHREADS. Testing has shown that vectors should be of length at least 100,000 before the overhead associated with creating and using the threads is made up by the parallelism in the vector calculations.

The Pthreads NVECTOR implementation provided with SUNDIALS, NVECTOR\_PTHREADS, 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, a boolean flag own\_data which specifies the ownership of data, and the number of threads. Operations on the vector are threaded using POSIX threads (Pthreads).





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

The following six macros are provided to access the content of an NVECTOR\_PTHREADS vector. The suffix \_PT in the names denotes the Pthreads version.

### NV\_CONTENT\_PT

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

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

Implementation:

```
#define NV_CONTENT_PT(v) ( (N_VectorContent_Pthreads)(v->content) )
```

• NV\_OWN\_DATA\_PT, NV\_DATA\_PT, NV\_LENGTH\_PT, NV\_NUM\_THREADS\_PT

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

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

The assignment v\_len = NV\_LENGTH\_PT(v) sets v\_len to be the length of v. On the other hand, the call NV\_LENGTH\_PT(v) = len\_v sets the length of v to be len\_v.

The assignment v\_num\_threads = NV\_NUM\_THREADS\_PT(v) sets v\_num\_threads to be the number of threads from v. On the other hand, the call NV\_NUM\_THREADS\_PT(v) = num\_threads\_v sets the number of threads for v to be num\_threads\_v.

Implementation:

```
#define NV_OWN_DATA_PT(v) ( NV_CONTENT_PT(v)->own_data )
#define NV_DATA_PT(v) ( NV_CONTENT_PT(v)->data )
#define NV_LENGTH_PT(v) ( NV_CONTENT_PT(v)->length )
#define NV_NUM_THREADS_PT(v) ( NV_CONTENT_PT(v)->num_threads )
```

## • NV\_Ith\_PT

This macro gives access to the individual components of the data array of an N-Vector.

The assignment  $r = NV_Ith_PT(v,i)$  sets r to be the value of the i-th component of v. The assignment  $NV_Ith_PT(v,i) = r$  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_PT(v,i) ( NV_DATA_PT(v)[i] )
```

The NVECTOR\_PTHREADS module defines Pthreads implementations of all vector operations listed in Table 6.1. Their names are obtained from those in Table 6.1 by appending the suffix \_Pthreads. The module NVECTOR\_PTHREADS provides the following additional user-callable routines:

#### • N\_VNew\_Pthreads

This function creates and allocates memory for a Pthreads N\_Vector. Arguments are the vector length and number of threads.

```
N_Vector N_VNew_Pthreads(long int vec_length, int num_threads);
```

# • N\_VNewEmpty\_Pthreads

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

N\_Vector N\_VNewEmpty\_Pthreads(long int vec\_length, int num\_threads);

# • N\_VMake\_Pthreads

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

N\_Vector N\_VMake\_Pthreads(long int vec\_length, realtype \*v\_data, int num\_threads);

# • N\_VCloneVectorArray\_Pthreads

This function creates (by cloning) an array of count Pthreads vectors.

N\_Vector \*N\_VCloneVectorArray\_Pthreads(int count, N\_Vector w);

# • N\_VCloneEmptyVectorArray\_Pthreads

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

N\_Vector \*N\_VCloneEmptyVectorArray\_Pthreads(int count, N\_Vector w);

# • N\_VDestroyVectorArray\_Pthreads

This function frees memory allocated for the array of count variables of type N\_Vector created with N\_VCloneVectorArray\_Pthreads or with N\_VCloneEmptyVectorArray\_Pthreads.

void N\_VDestroyVectorArray\_Pthreads(N\_Vector \*vs, int count);

# • N\_VPrint\_Pthreads

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

```
void N_VPrint_Pthreads(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\_PT(v) and then access v\_data[i] within the loop than it is to use NV\_Ith\_PT(v,i) within the loop.
- N\_VNewEmpty\_Pthreads, N\_VMake\_Pthreads, and N\_VCloneEmptyVectorArray\_Pthreads set the field own\_data = FALSE. N\_VDestroy\_Pthreads and N\_VDestroyVectorArray\_Pthreads 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\_PTHREADS 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.





# 6.5 NVECTOR Examples

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# 6.6 NVECTOR functions used by KINSOL

In Table 6.2 below, we list the vector functions in the NVECTOR module used 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, while the remaining five columns show function usage within each of the seven KINSOL linear solvers, the KINBBDPRE preconditioner module, and the FKINSOL module. Here KINDLS stands for KINDENSE and KINBAND; KINSPILS stands for KINSPGMR, KINSPFGMR, KINSPBCG, and KINSPTFQMR; and KINSLS stands for KINKLU and KINSUPERLUMT.

There is one subtlety in the KINSPILS column hidden by the table, explained here for the case of the KINSPGMR module. The N\_VDotProd function is called both within the interface file kinsol\_spgmr.c and within the implementation files sundials\_spgmr.c and sundials\_iterative.c for the generic SPGMR solver upon which the KINSPGMR solver is built. Also, although N\_VDiv and N\_VProd are not called within the interface file kinsol\_spgmr.c, they are called within the implementation file sundials\_spgmr.c, and so are required by the KINSPGMR solver module. Analogous statements apply to the KINSPFGMR, KINSPBCG and KINSPTFQMR modules, except that the latter two do not use sundials\_iterative.c. This issue does not arise for the direct KINSOL linear solvers because the generic DENSE and BAND solvers (used in the implementation of KINDENSE and KINBAND) do not make calls to any vector functions.

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.

	KINSOL	KINDLS	KINSPILS	KINSLS	KINBBDPRE	FKINSOL
$N_{-}VClone$	<b>√</b>		<b>√</b>		<b>√</b>	
$N_{-}VCloneEmpty$						<b>√</b>
$N_{-}VDestroy$	<b>√</b>		<b>√</b>		<b>√</b>	<b>√</b>
N_VSpace	<b>√</b>					
N_VGetArrayPointer		<b>√</b>		<b>√</b>	<b>√</b>	<b>√</b>
N_VSetArrayPointer		<b>√</b>				<b>√</b>
N_VLinearSum	<b>√</b>	<b>√</b>	<b>√</b>			
$N_{-}VConst$			<b>√</b>			
N_VProd	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>		
N_VDiv	<b>√</b>		<b>√</b>			
N_VScale	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>	
N_VAbs	<b>√</b>					
N_VInv	<b>√</b>					
$N_{-}VDotProd$	<b>√</b>	<b>√</b>	<b>√</b>	<b>√</b>		
N_VMaxNorm	<b>√</b>					
$N_{-}VMin$	<b>√</b>					
N_VWL2Norm	<b>√</b>		<b>√</b>			
N_VL1Norm			<b>√</b>			
N_VConstrMask	<b>√</b>					
$N_{-}VMinQuotient$	<b>√</b>					

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

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

# Chapter 7

# Providing Alternate Linear Solver Modules

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

- linit: initialize 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 function** (like that described in §4.5.2) which will attach the above four functions to the main KINSOL memory block. The KINSOL memory block is a structure defined in the header file kinsol\_impl.h. A pointer to such a structure is defined as the type KINMem. The four fields in a KINMem structure that must point to the linear solver's functions are kin\_linit, kin\_lsetup, kin\_lsolve, and kin\_lfree, respectively. Note that of the four interface functions, only the lsolve function is required. The lfree function must be provided only if the solver specification function makes any memory allocation. For any of the functions that are *not* provided, the corresponding field should be set to NULL. The linear solver specification function must also set the value of the field kin\_setupNonNull in the KINSOL memory block — to TRUE if lsetup is used, or FALSE otherwise.

Typically, the linear solver will require a block of memory specific to the solver, and a principal function of the specification function is to allocate that memory block, and initialize it. Then the field kin\_lmem in the KINSOL memory block is available to attach a pointer to that linear solver memory. This block can then be used to facilitate the exchange of data between the four interface functions.

If the linear solver involves adjustable parameters, the specification function should set the default values of those. User-callable functions may be defined that could, optionally, override the default parameter values.

We encourage the use of performance counters in connection with the various operations involved with the linear solver. Such counters would be members of the linear solver memory block, would be initialized in the linit function, and would be incremented by the lsetup and lsolve functions. Then, user-callable functions would be needed to obtain the values of these counters.

For consistency with the existing KINSOL linear solver modules, we recommend that the return value of the specification function be 0 for a successful return, and a negative value if an error occurs. Possible error conditions include: the pointer to the main KINSOL memory block is NULL, an input is illegal, the NVECTOR implementation is not compatible, or a memory allocation fails.

These four functions, which 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 functions. Note that the call list of each function includes a pointer to the main KINSOL memory block, by which the function 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).

# 7.1 Initialization function

The type definition of linit 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. It should also set pointers to data blocks that will later be passed to functions associated with the linear solver. The linit function is called once

only, at the start of the problem, by KINSol.

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 a negative value otherwise.

# 7.2 Setup function

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, in the solution of linear systems Ax=b. The exact nature of this system depends on the input strategy to KINSol. In the cases strategy = KIN\_NONE or KIN\_LINESEARCH, A is the Jacobian  $J=\partial F/\partial u$ . If strategy = KIN\_PICARD, A is the approximate Jacobian matrix L. If strategy = KIN\_FP, linear systems do not arise.

The lsetup function may call a user-supplied function, or a function within the linear solver module, to compute Jacobian-related data that is required by the linear solver. It may also preprocess that data as needed for lsolve, which may involve calling a generic function (such as for LU factorization). This data may be intended either for direct use (in a direct linear solver) or for use in a preconditioner (in a preconditioned iterative linear solver).

The 1setup function is not called at every Newton iteration, but only as frequently as the solver determines that it is appropriate to perform the setup task. In this way, Jacobian-related data generated by 1setup is expected to be used over a number of Newton iterations.

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

Return value The 1setup function should return 0 if successful and a non-zero value otherwise.

Notes

The current values of u and F(u) can be accessed by lsetup through the fields kin\_uu and kin\_fval (respectively) in kin\_mem. If needed, the scaling vectors u\_scale and f\_scale can be accessed by lsetup through the fields kin\_uscale and kin\_fscale (respectively) in kin\_mem.

7.3 Solve function 91

# 7.3 Solve function

The type definition of lsolve is

lsolve

Definition int (\*lsolve)(KINMem kin\_mem, N\_Vector x, N\_Vector b, realtype \*sJpnorm, realtype \*sFdotJp);

Purpose

The lsolve function must solve the linear system Ax = b, where A is either the Jacobian  $J = \partial F/\partial u$  (evaluated at the current iterate), or the approximate Jacobian, L, in the case of Picard iteration. 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.

sJpnorm is a pointer to a real scalar to be computed by 1solve. The returned value sJpnorm should be equal to  $||D_F Jp||_2$ , the scaled  $L_2$  norm of the product Jp, where p (= x) is the computed solution of the linear system Jp = b, and the scaling is that given by  $D_F$ . This value is not needed in all cases. See below.

sFdotJp is a pointer to a real scalar to be computed by lsolve. The returned value sFdotJp should be equal to  $(D_FF) \cdot (D_FJp)$ , the dot product of the scaled F vector and the scaled vector Jp, where p = x is the computed solution of the linear system Jp = b, and the scaling is that given by  $D_F$ . This value is not needed in all cases. See below.

Return value lsolve should return 0 if successful. If an error occurs and recovery could be possible by calling the lsetup function again, then it should return a positive value. Otherwise, lsolve should return a negative value.

Notes

The current values of u and F(u) can be accessed by lsolve through the fields kin\_uu and kin\_fval (respectively) in kin\_mem, and the scaling vectors u\_scale and f\_scale can be accessed through the fields kin\_uscale and kin\_fscale (respectively) in kin\_mem.

In the case of a direct solver, sJpnorm can be ignored, and sFdotJp can be computed with lines of the form

```
N_VProd(b, f_scale, b);
N_VProd(b, f_scale, b);
*sFdotJp = N_VDotProd(fval, b);
```

in which Jp is taken to be equal to the input right-hand side b, and f\_scale and fval (=F(u)) are taken from kin\_mem.

In the case of an iterative solver, the two terms, sJpnorm and sFdotJp, can be computed with lines of the form

```
ret = KINSpilsAtimes(kin_mem, x, b);
*sJpnorm = N_VWL2Norm(b, f_scale);
N_VProd(b, f_scale, b);
N_VProd(b, f_scale, b);
*sFdotJp = N_VDotProd(fval, b);
```

following the computation of the solution vector  $\mathbf{x}$ , in which  $\mathbf{f}$ \_scale and  $\mathbf{fval} \ (= F(u))$  are taken from  $\mathtt{kin}$ \_mem.

The values sFdotJp and sFdotJp need not be set in all cases, and so for maximum efficiency, the lsolve function could do these calculations conditionally, depending on the value of the input strategy to KINSol, and the choice (given by etachoice) of Forcing Term in the Krylov iteration stopping test (see KINSetEtaForm). The precise conditions are as follows: First, if strategy is KIN\_FP, neither of these quantities need to be computed. In the other cases, if the linear solver is iterative

and etachoice = KIN\_ETACHOICE1 (the default) then both sFdotJp and sFdotJp must be set. If strategy is KIN\_LINESEARCH, then sFdotJp must be set, regardless of the linear solver type.

The values of strategy and etachoice are available from the fields kin\_global\_strategy and kin\_etaflag (respectively) in kin\_mem.

# 7.4 Memory deallocation function

The type definition of lfree is

lfree

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

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

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

Return value The lfree function has no return value.

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

longer needed.

# Chapter 8

# General Use Linear Solver Components in SUNDIALS

In this chapter, we describe seven 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 SUNDIALS or separately.

These generic modules in SUNDIALS are organized in three families, the *dls* family, which includes direct linear solvers appropriate for sequential computations; the *sls* family, which includes sparse matrices; and the *spils* family, which includes scaled preconditioned iterative (Krylov) linear solvers. The solvers in each family share common data structures and functions.

The dls family contains the following two generic linear solvers:

- The DENSE package, a linear solver for dense matrices either specified through a matrix type (defined below) or as simple arrays.
- The BAND package, a linear solver for banded matrices either specified through a matrix type (defined below) or as simple arrays.

Note that this family also includes the Blas/Lapack linear solvers (dense and band) available to the SUNDIALS solvers, but these are not discussed here.

The sls family contains a sparse matrix package and interfaces between it and two sparse direct solver packages:

- The KLU package, a linear solver for compressed-sparse-column matrices, [1, 7].
- The SUPERLUMT package, a threaded linear solver for compressed-sparse-column matrices, [2, 16, 9].

The *spils* family contains the following generic linear solvers:

- The SPGMR package, a solver for the scaled preconditioned GMRES method.
- The SPFGMR package, a solver for the scaled preconditioned Flexible GMRES method.
- The SPBCG package, a solver for the scaled preconditioned Bi-CGStab method.
- The SPTFQMR package, a solver for the scaled preconditioned TFQMR method.

For reasons related to installation, the names of the files involved in these packages begin with the prefix sundials. But despite this, each of the *dls* and *spils* solvers is in fact generic, in that it is usable completely independently of SUNDIALS.

For the sake of space, the functions for the dense and band modules that work with a matrix type and the functions in the SPGMR, SPFGMR, SPBCG, and SPTFQMR modules are only summarized briefly, since they are less likely to be of direct use in connection with a SUNDIALS solver. However, the

functions for dense matrices treated as simple arrays and sparse matrices are fully described, because we expect that they will be useful in the implementation of preconditioners used with the combination of one of the SUNDIALS solvers and one of the *spils* linear solvers.

# 8.1 The DLS modules: DENSE and BAND

The files comprising the DENSE generic linear solver, and their locations in the SUNDIALS *srcdir*, are as follows:

- header files (located in *srcdir*/include/sundials) sundials\_direct.h, sundials\_dense.h, sundials\_types.h, sundials\_math.h, sundials\_config.h
- source files (located in *srcdir*/src/sundials) sundials\_direct.c, sundials\_dense.c, sundials\_math.c

The files comprising the BAND generic linear solver are as follows:

- header files (located in *srcdir*/include/sundials) sundials\_direct.h, sundials\_band.h, sundials\_types.h, sundials\_math.h, sundials\_config.h
- source files (located in *srcdir*/src/sundials) sundials\_direct.c, sundials\_band.c, sundials\_math.c

Only two of the preprocessing directives in the header file sundials\_config.h are relevant to the DENSE and BAND packages by themselves (see §?? for details):

• (required) definition of the precision of the SUNDIALS type realtype. One of the following lines must be present:

```
#define SUNDIALS_DOUBLE_PRECISION 1
#define SUNDIALS_SINGLE_PRECISION 1
#define SUNDIALS_EXTENDED_PRECISION 1
```

• (optional) use of generic math functions: #define SUNDIALS\_USE\_GENERIC\_MATH 1

The sundials\_types.h header file defines the SUNDIALS realtype and booleantype types and the macro RCONST, while the sundials\_math.h header file is needed for the macros SUNMIN and SUNMAX, and the function SUNRabs.

The files listed above for either module can be extracted from the SUNDIALS *srcdir* and compiled by themselves into a separate library or into a larger user code.

# 8.1.1 Type DlsMat

The type DlsMat, defined in sundials\_direct.h is a pointer to a structure defining a generic matrix, and is used with all linear solvers in the *dls* family:

```
typedef struct _DlsMat {
  int type;
  long int M;
  long int N;
  long int ldim;
  long int mu;
  long int ml;
  long int s_mu;
  realtype *data;
  long int ldata;
  realtype **cols;
} *DlsMat;
```

For the DENSE module, the relevant fields of this structure are as follows. Note that a dense matrix of type DlsMat need not be square.

```
\mathbf{type} - SUNDIALS_DENSE (=1)
```

M - number of rows

N - number of columns

**ldim** - leading dimension ( $ldim \ge M$ )

data - pointer to a contiguous block of realtype variables

**ldata** - length of the data array (= ldim·N). The (i,j)-th element of a dense matrix A of type DlsMat (with  $0 \le i < M$  and  $0 \le j < N$ ) is given by the expression (A->data)[0][j\*M+i]

cols - array of pointers. cols[j] points to the first element of the j-th column of the matrix in the array data. The (i,j)-th element of a dense matrix A of type DlsMat (with  $0 \le i < M$  and  $0 \le j < N$ ) is given by the expression (A->cols)[j][i]

For the BAND module, the relevant fields of this structure are as follows (see Figure 8.1 for a diagram of the underlying data representation in a banded matrix of type DlsMat). Note that only square band matrices are allowed.

```
type - SUNDIALS_BAND (=2)
```

M - number of rows

N - number of columns (N = M)

 $\mathbf{mu}$  - upper half-bandwidth,  $0 \leq \mathbf{mu} < \min(\mathbf{M}, \mathbf{N})$ 

 $\mathbf{ml}$  - lower half-bandwidth,  $0 \le \mathtt{ml} < \min(\mathtt{M}, \mathtt{N})$ 

 $s\_mu$  - storage upper bandwidth,  $mu \le s\_mu < N$ . The LU decomposition routine writes the LU factors into the storage for A. The upper triangular factor U, however, may have an upper bandwidth as big as min(N-1,mu+ml) because of partial pivoting. The  $s\_mu$  field holds the upper half-bandwidth allocated for A.

ldim - leading dimension (ldim ≥ s\_mu)

data - pointer to a contiguous block of realtype variables. The elements of a banded matrix of type DlsMat are stored columnwise (i.e. columns are stored one on top of the other in memory). Only elements within the specified half-bandwidths are stored. data is a pointer to ldata contiguous locations which hold the elements within the band of A.

 $ldata - length of the data array (= ldim \cdot (s_mu + ml + 1))$ 

cols - array of pointers. cols[j] is a pointer to the uppermost element within the band in the j-th column. This pointer may be treated as an array indexed from  $s_mu-mu$  (to access the uppermost element within the band in the j-th column) to  $s_mu+ml$  (to access the lowest element within the band in the j-th column). Indices from 0 to  $s_mu-mu-1$  give access to extra storage elements required by the LU decomposition function. Finally,  $cols[j][i-j+s_mu]$  is the (i,j)-th element,  $j-mu \le i \le j+ml$ .

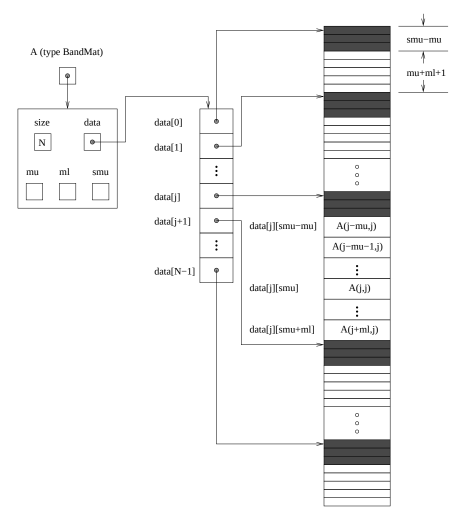


Figure 8.1: Diagram of the storage for a banded matrix of type DlsMat. Here A is an  $N \times N$  band matrix of type DlsMat with upper and lower half-bandwidths mu and ml, respectively. The rows and columns of A are numbered from 0 to N-1 and the (i,j)-th element of A is denoted A(i,j). The greyed out areas of the underlying component storage are used by the BandGBTRF and BandGBTRS routines.

# 8.1.2 Accessor macros for the DLS modules

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 or BAND\_COL macros. Users should use these macros whenever possible.

The following two macros are defined by the DENSE module to provide access to data in the DlsMat 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 M \times N DlsMat A, 0 \le i < M, 0 \le j < N.
```

#### • DENSE\_COL

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

DENSE\_COL references the j-th column of the  $M \times N$  DlsMat A,  $0 \le j < N$ . 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 M-1. The (i, j)-th element of A is referenced by col\_j[i].

The following three macros are defined by the BAND module to provide access to data in the DlsMat type:

#### • BAND\_ELEM

```
Usage : BAND_ELEM(A,i,j) = a_ij; or a_ij = BAND_ELEM(A,i,j); 
BAND_ELEM references the (i,j)-th element of the N \times N band matrix A, where 0 \le i, j \le N-1. The location (i,j) should further satisfy j-(A->mu) \le i \le j+(A->m1).
```

# • BAND\_COL

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

BAND\_COL references the diagonal element of the j-th column of the  $N \times N$  band matrix A,  $0 \le j \le N-1$ . The type of the expression BAND\_COL(A,j) is realtype \*. The pointer returned by the call BAND\_COL(A,j) can be treated as an array which is indexed from -(A-mu) to (A-ml).

### • BAND\_COL\_ELEM

```
Usage : BAND_COL_ELEM(col_j,i,j) = a_ij; or a_ij = BAND_COL_ELEM(col_j,i,j);
```

This macro references the (i,j)-th entry of the band matrix A when used in conjunction with BAND\_COL to reference the j-th column through col\_j. The index (i,j) should satisfy  $j-(A->mu) \le i \le j+(A->m1)$ .

# 8.1.3 Functions in the DENSE module

The DENSE module defines two sets of functions with corresponding names. The first set contains functions (with names starting with a capital letter) that act on dense matrices of type DlsMat. The second set contains functions (with names starting with a lower case letter) that act on matrices represented as simple arrays.

The following functions for DlsMat dense matrices are available in the DENSE package. For full details, see the header files sundials\_direct.h and sundials\_dense.h.

- NewDenseMat: allocation of a DlsMat dense matrix;
- DestroyMat: free memory for a DlsMat matrix;

- PrintMat: print a DlsMat matrix to standard output.
- NewLintArray: allocation of an array of long int integers for use as pivots with DenseGETRF and DenseGETRS:
- NewIntArray: allocation of an array of int integers for use as pivots with the Lapack dense solvers;
- NewRealArray: allocation of an array of realtype for use as right-hand side with DenseGETRS;
- DestroyArray: free memory for an array;
- SetToZero: load a matrix with zeros;
- AddIdentity: increment a square matrix by the identity matrix;
- DenseCopy: copy one matrix to another;
- DenseScale: scale a matrix by a scalar;
- DenseGETRF: LU factorization with partial pivoting;
- DenseGETRS: solution of Ax = b using LU factorization (for square matrices A);
- DensePOTRF: Cholesky factorization of a real symmetric positive matrix;
- DensePOTRS: solution of Ax = b using the Cholesky factorization of A;
- DenseGEQRF: QR factorization of an  $m \times n$  matrix, with  $m \ge n$ ;
- DenseORMQR: compute the product w = Qv, with Q calculated using DenseGEQRF;
- DenseMatvec: compute the product y = Ax, for an M by N matrix A;

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

# newDenseMat

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

# • destroyMat

destroyMat(a) frees the dense matrix a allocated by newDenseMat;

# newLintArray

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

### • newIntArray

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

# • newRealArray

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

### • destroyArray

destroyArray(p) frees the array p allocated by newLintArray, newIntArray, or newRealArray;

### • denseCopy

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

#### • denseScale

denseScale(c,a,m,n) scales every element in the m by n dense matrix a by the scalar c;

### • denseAddIdentity

denseAddIdentity(a,n) increments the square n by n dense matrix a by the identity matrix  $I_n$ ;

#### • denseGETRF

denseGETRF(a,m,n,p) factors the m by n dense matrix a, using Gaussian elimination with row pivoting. 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 an m by n lower trapezoidal matrix with all diagonal elements equal to 1, and U is an n by n upper triangular matrix, then the upper triangular part of a (including its diagonal) contains U and the strictly lower trapezoidal part of a contains the multipliers, I L. If a is square, L is a unit lower triangular matrix.

denseGETRF returns 0 if successful. Otherwise it encountered a zero diagonal element during the factorization, indicating that the matrix **a** does not have full column rank. In this case it returns the column index (numbered from one) at which it encountered the zero.

### • denseGETRS

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

### • densePOTRF

densePOTRF(a,m) calculates the Cholesky decomposition of the m by m dense matrix a, assumed to be symmetric positive definite. Only the lower triangle of a is accessed and overwritten with the Cholesky factor.

### • densePOTRS

densePOTRS(a,m,b) solves the m by m linear system ax = b. It assumes that the Cholesky factorization of a has been calculated in the lower triangular part of a by a successful call to densePOTRF(a,m).

### • denseGEQRF

denseGEQRF(a,m,n,beta,wrk) calculates the QR decomposition of the m by n matrix a  $(m \ge n)$  using Householder reflections. On exit, the elements on and above the diagonal of a contain the n by n upper triangular matrix R; the elements below the diagonal, with the array beta, represent the orthogonal matrix Q as a product of elementary reflectors. The real array wrk, of length m, must be provided as temporary workspace.

### • denseORMQR

denseORMQR(a,m,n,beta,v,w,wrk) calculates the product w = Qv for a given vector v of length n, where the orthogonal matrix Q is encoded in the m by n matrix a and the vector beta of length n, after a successful call to denseGEQRF(a,m,n,beta,wrk). The real array wrk, of length m, must be provided as temporary workspace.

### • denseMatvec

denseMatvec(a,x,y,m,n) calculates the product y = ax for a given vector x of length n, and m by n matrix a.

### 8.1.4 Functions in the BAND module

The BAND module defines two sets of functions with corresponding names. The first set contains functions (with names starting with a capital letter) that act on band matrices of type DlsMat. The second set contains functions (with names starting with a lower case letter) that act on matrices represented as simple arrays.

The following functions for DlsMat banded matrices are available in the BAND package. For full details, see the header files sundials\_direct.h and sundials\_band.h.

- NewBandMat: allocation of a DlsMat band matrix;
- DestroyMat: free memory for a DlsMat matrix;
- PrintMat: print a DlsMat matrix to standard output.
- NewLintArray: allocation of an array of int integers for use as pivots with BandGBRF and BandGBRS:
- NewIntArray: allocation of an array of int integers for use as pivots with the Lapack band solvers;
- NewRealArray: allocation of an array of realtype for use as right-hand side with BandGBRS;
- DestroyArray: free memory for an array;
- SetToZero: load a matrix with zeros;
- AddIdentity: increment a square matrix by the identity matrix;
- BandCopy: copy one matrix to another;
- BandScale: scale a matrix by a scalar;
- BandGBTRF: LU factorization with partial pivoting;
- BandGBTRS: solution of Ax = b using LU factorization;
- BandMatvec: compute the product y = Ax, for a square band matrix A;

The following functions for small band matrices are available in the BAND package:

### newBandMat

newBandMat(n, smu, ml) allocates storage for an n by n band matrix with lower half-bandwidth ml.

### • destroyMat

destroyMat(a) frees the band matrix a allocated by newBandMat;

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### • newLintArray

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

### • newIntArray

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

### • newRealArray

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

### • destroyArray

destroyArray(p) frees the array p allocated by newLintArray, newIntArray, or newRealArray;

### bandCopy

bandCopy(a,b,n,a\_smu, b\_smu,copymu, copyml) copies the n by n band matrix a into the n by n band matrix b;

#### • bandScale

bandScale(c,a,n,mu,ml,smu) scales every element in the n by n band matrix a by c;

### • bandAddIdentity

bandAddIdentity(a,n,smu) increments the n by n band matrix a by the identity matrix;

### • bandGETRF

bandGETRF(a,n,mu,ml,smu,p) factors the n by n band matrix a, using Gaussian elimination with row pivoting. It overwrites the elements of a with its LU factors and keeps track of the pivot rows chosen in the pivot array p.

### • bandGETRS

bandGETRS(a,n,smu,ml,p,b) solves the n by n linear system ax = b. It assumes that a (of size  $n \times n$ ) has been LU-factored and the pivot array p has been set by a successful call to bandGETRF(a,n,mu,ml,smu,p). The solution x is written into the b array.

### bandMatvec

bandMatvec(a,x,y,n,mu,ml,smu) calculates the product y = ax for a given vector x of length n, and n by n band matrix a.

### 8.2 The SLS module

SUNDIALS provides a compressed-sparse-column matrix type and sparse matrix support functions. In addition, SUNDIALS provides interfaces to the publically available KLU and SuperLU\_MT sparse direct solver packages. The files comprising the SLS matrix module, used in the KLU and SUPERLUMT linear solver packages, and their locations in the SUNDIALS *srcdir*, are as follows:

- header files (located in srcdir/include/sundials) sundials\_sparse.h, sundials\_klu\_impl.h, sundials\_superlumt\_impl.h, sundials\_types.h, sundials\_math.h, sundials\_config.h
- source files (located in srcdir/src/sundials) sundials\_sparse.c, sundials\_math.c

Only two of the preprocessing directives in the header file sundials\_config.h are relevant to the SLS package by itself (see §?? for details):

• (required) definition of the precision of the SUNDIALS type realtype. One of the following lines must be present:

```
#define SUNDIALS_DOUBLE_PRECISION 1
#define SUNDIALS_SINGLE_PRECISION 1
#define SUNDIALS_EXTENDED_PRECISION 1
```

• (optional) use of generic math functions: #define SUNDIALS\_USE\_GENERIC\_MATH 1

The sundials\_types.h header file defines the SUNDIALS realtype and booleantype types and the macro RCONST, while the sundials\_math.h header file is needed for the macros SUNMIN and SUNMAX, and the function SUNRabs.

### 8.2.1 Type SlsMat

The type SlsMat, defined in sundials\_sparse.h is a pointer to a structure defining a generic compressed-sparse-column matrix, and is used with all linear solvers in the sls family:

```
typedef struct _SlsMat {
  int M;
  int N;
  int NNZ;
  realtype *data;
  int *rowvals;
  int *colptrs;
} *SlsMat;
```

The fields of this structure are as follows (see Figure 8.2 for a diagram of the underlying compressed-sparse-column representation in a sparse matrix of type SlsMat). Note that a sparse matrix of type SlsMat need not be square.

M - number of rows

 ${f N}$  - number of columns

NNZ - maximum number of nonzero entries in the matrix (allocated length of data and rowvals arrays)

data - pointer to a contiguous block of realtype variables (of length NNZ), containing the values of the nonzero entries in the matrix

rowvals - pointer to a contiguous block of int variables (of length NNZ), containing the row indices of each nonzero entry held in data

colptrs - pointer to a contiguous block of int variables (of length N+1). Each entry provides the index of the first column entry into the data and rowvals arrays, e.g. if colptr[3]=7, then the first nonzero entry in the fourth column of the matrix is located in data[7], and is located in row rowvals[7] of the matrix. The last entry contains the total number of nonzero values in the matrix and hence points one past the end of the active data in the data and rowvals arrays.

For example, the  $5 \times 4$  matrix

$$\left[\begin{array}{ccccc}
0 & 3 & 1 & 0 \\
3 & 0 & 0 & 2 \\
0 & 7 & 0 & 0 \\
1 & 0 & 0 & 9 \\
0 & 0 & 0 & 5
\end{array}\right]$$

could be stored in a SlsMat structure as either

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```
M = 5;
N = 4;
NNZ = 8;
data = {3.0, 1.0, 3.0, 7.0, 1.0, 2.0, 9.0, 5.0};
rowvals = {1, 3, 0, 2, 0, 1, 3, 4};
colptrs = {0, 2, 4, 5, 8};

or

M = 5;
N = 4;
NNZ = 10;
data = {3.0, 1.0, 3.0, 7.0, 1.0, 2.0, 9.0, 5.0, *, *};
rowvals = {1, 3, 0, 2, 0, 1, 3, 4, *, *};
colptrs = {0, 2, 4, 5, 8};
```

where the first has no unused space, and the second has additional storage (the entries marked with \* may contain any values). Note in both cases that the final value in colptrs is 8. The work associated with operations on the sparse matrix is proportional to this value and so one should use the best understanding of the number of nonzeroes here.

### 8.2.2 Functions in the SLS module

The SLS module defines functions that act on sparse matrices of type SlsMat. For full details, see the header file sundials\_sparse.h.

### • NewSparseMat

NewSparseMat(M, N, NNZ) allocates storage for an M by N sparse matrix, with storage for up to NNZ nonzero entries.

### • SlsConvertDls

SlsConvertDls(A) converts a dense or band matrix A of type DlsMat into a new sparse matrix of type SlsMat by retaining only the nonzero values of the matrix A.

### • DestroySparseMat

DestroySparseMat(A) frees the memory for a sparse matrix A allocated by either NewSparseMat or SlsConvertDls.

• SlsSetToZero(A) zeros out the SlsMat matrix A. The storage for A is left unchanged.

### • CopySparseMat

CopySparseMat(A, B) copies the SlsMat A into the SlsMat B. It is assumed that the matrices have the same row/column dimensions. If B has insufficient storage to hold all the nonzero entries of A, the data and row index arrays in B are reallocated to match those in A.

### • ScaleSparseMat

ScaleSparseMat(c, A) scales every element in the SlsMat A by the realtype scalar c.

### • AddIdentitySparseMat

AddIdentitySparseMat(A) increments the SlsMat A by the identity matrix. If A is not square, only the existing diagonal values are incremented. Resizes the data and rowvals arrays of A to allow for new nonzero entries on the diagonal.

### • SlsAddMat

SlsAddMat(A, B) adds two SlsMat matrices A and B, placing the result back in A. Resizes the data and rowvals arrays of A upon completion to exactly match the nonzero storage for the result. Upon successful completion, the return value is zero; otherwise -1 is returned.

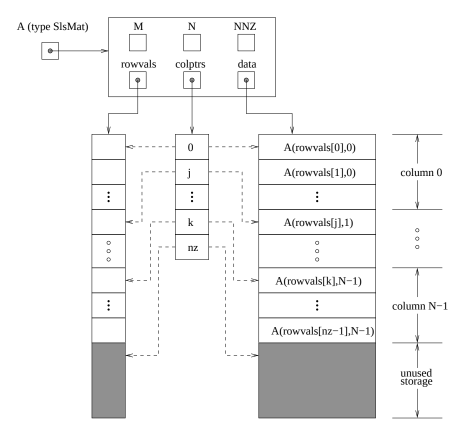


Figure 8.2: Diagram of the storage for a compressed-sparse-column matrix of type SlsMat. Here A is an  $M \times N$  sparse matrix of type SlsMat with storage for up to NNZ nonzero entries (the allocated length of both data and rowvals). The entries in rowvals may assume values from 0 to M-1, corresponding to the row index (zero-based) of each nonzero value. The entries in data contain the values of the nonzero entries, with the row i, column j entry of A (again, zero-based) denoted as A(i,j). The colptrs array contains N+1 entries; the first N denote the starting index of each column within the rowvals and data arrays, while the final entry points one past the final nonzero entry. Here, although NNZ values are allocated, only nz are actually filled in; the greyed-out portions of data and rowvals indicate extra allocated space.

### • ReallocSparseMat

ReallocSparseMat(A) eliminates unused storage in the SlsMat A by resizing the internal data and rowvals arrays to contain exactly colptrs[N] values.

### • SlsMatvec

SlsMatvec(A, x, y) computes the sparse matrix-vector product, y = Ax. If the SlsMat A is a sparse matrix of dimension  $M \times N$ , then it is assumed that x is a realtype array of length N, and y is a realtype array of length M. Upon successful completion, the return value is zero; otherwise -1 is returned.

### • PrintSparseMat

PrintSparseMat(A) Prints the SlsMat matrix A to standard output.

### 8.2.3 The KLU solver

KLU is a sparse matrix factorization and solver library written by Tim Davis [1, 7]. KLU has a symbolic factorization routine that computes the permutation of the linear system matrix to block triangular form and the permutations that will pre-order the diagonal blocks (the only ones that need to be factored) to reduce fill-in (using AMD, COLAMD, CHOLAMD, natural, or an ordering given by the user). Note that SUNDIALS uses the COLAMD ordering by default with KLU.

KLU breaks the factorization into two separate parts. The first is a symbolic factorization and the second is a numeric factorization that returns the factored matrix along with final pivot information. KLU also has a refactor routine that can be called instead of the numeric factorization. This routine will reuse the pivot information. This routine also returns diagnostic information that a user can examine to determine if numerical stability is being lost and a full numerical factorization should be done instead of the refactor.

The KLU interface in SUNDIALS will perform the symbolic factorization once. It then calls the numerical factorization once and will call the refactor routine until estimates of the numerical conditioning suggest a new factorization should be completed. The KLU interface also has a ReInit routine that can be used to force a full refactorization at the next solver setup call.

In order to use the SUNDIALS interface to KLU, it is assumed that KLU has been installed on the system prior to installation of SUNDIALS, and that SUNDIALS has been configured appropriately to link with KLU (see Appendix A for details).

Designed for serial calculations only, KLU is supported for calculations employing SUNDIALS' serial or shared-memory parallel NVECTOR modules (see Sections 6.1, 6.3 and 6.4 for details).

### 8.2.4 The SUPERLUMT solver

SUPERLUMT is a threaded sparse matrix factorization and solver library written by X. Sherry Li [2, 16, 9]. The package performs matrix factorization using threads to enhance efficiency in shared memory parallel environments. It should be noted that threads are only used in the factorization step.

In order to use the SUNDIALS interface to SUPERLUMT, it is assumed that SUPERLUMT has been installed on the system prior to installation of SUNDIALS, and that SUNDIALS has been configured appropriately to link with SUPERLUMT (see Appendix A for details).

Designed for serial and threaded calculations only, SUPERLUMT is supported for calculations employing SUNDIALS' serial or shared-memory parallel NVECTOR modules (see Sections 6.1, 6.3 and 6.4 for details).

# 8.3 The SPILS modules: SPGMR, SPFGMR, SPBCG, and SPTFQMR

The *spils* modules contain implementations of some of the most commonly use scaled preconditioned Krylov solvers. A linear solver module from the *spils* family can only be used in conjunction with any

NVECTOR implementation library.

### 8.3.1 The SPGMR module

The SPGMR package, in the files sundials\_spgmr.h and sundials\_spgmr.c, includes an implementation of the scaled preconditioned GMRES method. A separate code module, implemented in sundials\_iterative.(h,c), contains auxiliary functions that support SPGMR, as well as the other Krylov solvers in SUNDIALS (SPFGMR, SPBCG, and SPTFQMR). For full details, including usage instructions, see the header files sundials\_spgmr.h and sundials\_iterative.h.

The files comprising the SPGMR generic linear solver, and their locations in the SUNDIALS *srcdir*, are as follows:

- header files (located in srcdir/include/sundials)
   sundials\_spgmr.h, sundials\_iterative.h, sundials\_nvector.h,
   sundials\_types.h, sundials\_math.h, sundials\_config.h
- source files (located in *srcdir*/src/sundials) sundials\_spgmr.c, sundials\_iterative.c, sundials\_nvector.c

Only two of the preprocessing directives in the header file sundials\_config.h are required to use the SPGMR package by itself (see §?? for details):

• (required) definition of the precision of the SUNDIALS type realtype. One of the following lines must be present:

```
#define SUNDIALS_DOUBLE_PRECISION 1
#define SUNDIALS_SINGLE_PRECISION 1
#define SUNDIALS_EXTENDED_PRECISION 1
```

• (optional) use of generic math functions: #define SUNDIALS\_USE\_GENERIC\_MATH 1

The sundials\_types.h header file defines the SUNDIALS realtype and booleantype types and the macro RCONST, while the sundials\_math.h header file is needed for the macros SUNMIN, SUNMAX, and SUNSQR, and the functions SUNRabs and SUNRagrt.

The generic NVECTOR files, sundials\_nvector.(h,c) are needed for the definition of the generic N\_Vector type and functions. The NVECTOR functions used by the SPGMR module are: N\_VDotProd, N\_VLinearSum, N\_VScale, N\_VProd, N\_VDiv, N\_VConst, N\_VClone, N\_VCloneVectorArray, N\_VDestroy, and N\_VDestroyVectorArray.

The nine files listed above can be extracted from the SUNDIALS *srcdir* and compiled by themselves into an SPGMR library or into a larger user code.

The following functions are available in the SPGMR package:

- $\bullet \ \, {\tt SpgmrMalloc:} \ \, {\tt allocation} \ \, {\tt of} \ \, {\tt memory} \ \, {\tt for} \ \, {\tt SpgmrSolve};$
- SpgmrSolve: solution of Ax = b by the SPGMR method;
- SpgmrFree: free memory allocated by SpgmrMalloc.

The following functions are available in the support package sundials\_iterative.(h,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.

### 8.3.2 The SPFGMR module

The SPFGMR package, in the files sundials\_spfgmr.h and sundials\_spfgmr.c, includes an implementation of the scaled preconditioned Flexible GMRES method. For full details, including usage instructions, see the file sundials\_spfgmr.h.

The files needed to use the SPFGMR module by itself are the same as for the SPGMR module, but with sundials\_spfgmr.(h,c) in place of sundials\_spgmr.(h,c).

The following functions are available in the SPFGMR package:

- SpfgmrMalloc: allocation of memory for SpfgmrSolve;
- SpfgmrSolve: solution of Ax = b by the SPFGMR method;
- SpfgmrFree: free memory allocated by SpfgmrMalloc.

### 8.3.3 The SPBCG module

The SPBCG package, in the files sundials\_spbcgs.h and sundials\_spbcgs.c, includes an implementation of the scaled preconditioned Bi-CGStab method. For full details, including usage instructions, see the file sundials\_spbcgs.h.

The files needed to use the SPBCG module by itself are the same as for the SPGMR module, but with sundials\_spbcgs.(h,c) in place of sundials\_spgmr.(h,c).

The following functions are available in the SPBCG package:

- SpbcgMalloc: allocation of memory for SpbcgSolve;
- SpbcgSolve: solution of Ax = b by the SPBCG method;
- SpbcgFree: free memory allocated by SpbcgMalloc.

### 8.3.4 The SPTFQMR module

The SPTFQMR package, in the files sundials\_sptfqmr.h and sundials\_sptfqmr.c, includes an implementation of the scaled preconditioned TFQMR method. For full details, including usage instructions, see the file sundials\_sptfqmr.h.

The files needed to use the SPTFQMR module by itself are the same as for the SPGMR module, but with sundials\_sptfqmr.(h,c) in place of sundials\_spgmr.(h,c).

The following functions are available in the SPTFQMR package:

- SptfqmrMalloc: allocation of memory for SptfqmrSolve;
- SptfqmrSolve: solution of Ax = b by the SPTFQMR method;
- SptfqmrFree: free memory allocated by SptfqmrMalloc.

## Appendix A

## SUNDIALS Package Installation Procedure

The installation of any SUNDIALS package 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 one or all solvers in SUNDIALS.

The SUNDIALS suite (or individual solvers) are distributed as compressed archives (.tar.gz). The name of the distribution archive is of the form solver-x.y.z.tar.gz, where solver is one of: sundials, cvode, cvodes, ida, idas, or kinsol, and x.y.z represents the version number (of the SUNDIALS suite or of the individual solver). To begin the installation, first uncompress and expand the sources, by issuing

% tar xzf solver-x.y.z.tar.gz

This will extract source files under a directory *solver*-x.y.z.

Starting with version 2.6.0 of SUNDIALS, CMake is the only supported method of installation. The explanations on the installation procedure begins with a few common observations:

• The remainder of this chapter will follow these conventions:

srcdir is the directory solver-x.y.z created above; i.e., the directory containing the SUNDIALS sources.

builddir is the (temporary) directory under which SUNDIALS is built.

installdir is the directory under which the SUNDIALS exported header files and libraries will be installed. Typically, header files are exported under a directory installdir/include while libraries are installed under installdir/lib, with installdir specified at configuration time.

- For sundials CMake-based installation, in-source builds are prohibited; in other words, the build directory *builddir* can **not** be the same as *srcdir* and such an attempt will lead to an error. This prevents "polluting" the source tree and allows efficient builds for different configurations and/or options.
- The installation directory *installdir* can **not** be the same as the source directory *srcdir*.
- By default, only the libraries and header files are exported to the installation directory installdir. If enabled by the user (with the appropriate toggle for CMake), the examples distributed with SUNDIALS will be built together with the solver libraries but the installation step will result in exporting (by default in a subdirectory of the installation directory) the example sources and sample outputs together with automatically generated configuration files that reference the installed SUNDIALS headers and libraries. As such, these configuration files for the SUNDIALS examples can be used as "templates" for your own problems. CMake installs CMakeLists.txt files and also (as an option available only under Unix/Linux) makefiles. Note this installation



approach also allows the option of building the SUNDIALS examples without having to install them. (This can be used as a sanity check for the freshly built libraries.)

• Even if generation of shared libraries is enabled, only static libraries are created for the FCMIX modules. (Because of the use of fixed names for the Fortran user-provided subroutines, FCMIX shared libraries would result in "undefined symbol" errors at link time.)

### A.1 CMake-based installation

CMake-based installation provides a platform-independent build system. CMake can generate Unix and Linux Makefiles, as well as KDevelop, Visual Studio, and (Apple) XCode project files from the same configuration file. In addition, CMake also provides a GUI front end and which allows an interactive build and installation process.

The SUNDIALS build process requires CMake version 2.8.1 or higher and a working compiler. On Unix-like operating systems, it also requires Make (and curses, including its development libraries, for the GUI front end to CMake, ccmake), while on Windows it requires Visual Studio. While many Linux distributions offer CMake, the version included is probably out of date. Many new CMake features have been added recently, and you should download the latest version from http://www.cmake.org. Build instructions for CMake (only necessary for Unix-like systems) can be found on the CMake website. Once CMake is installed, Linux/Unix users will be able to use ccmake, while Windows users will be able to use CMakeSetup.

As previously noted, when using CMake to configure, build and install SUNDIALS, it is always required to use a separate build directory. While in-source builds are possible, they are explicitly prohibited by the SUNDIALS CMake scripts (one of the reasons being that, unlike autotools, CMake does not provide a make distclean procedure and it is therefore difficult to clean-up the source tree after an in-source build). By ensuring a separate build directory, it is an easy task for the user to clean-up all traces of the build by simply removing the build directory. CMake does generate a make clean which will remove files generated by the compiler and linker.

### A.1.1 Configuring, building, and installing on Unix-like systems

The default CMake configuration will build all included solvers and associated examples and will build static and shared libraries. The *installdir* defaults to /usr/local and can be changed by setting the CMAKE\_INSTALL\_PREFIX variable. Support for FORTRAN and all other options are disabled.

CMake can be used from the command line with the cmake command, or from a curses-based GUI by using the ccmake command. Examples for using both methods will be presented. For the examples shown it is assumed that there is a top level SUNDIALS directory with appropriate source, build and install directories:

```
% mkdir (...)sundials/installdir
% mkdir (...)sundials/builddir
% cd (...)sundials/builddir
```

### Building with the GUI

Using CMake with the GUI follows this general process:

- Select and modify values, run configure (c key)
- New values are denoted with an asterisk
- To set a variable, move the cursor to the variable and press enter
  - If it is a boolean (ON/OFF) it will toggle the value
  - If it is string or file, it will allow editing of the string

- For file and directories, the <tab> key can be used to complete
- Repeat until all values are set as desired and the generate option is available (g key)
- Some variables (advanced variables) are not visible right away
- To see advanced variables, toggle to advanced mode (t key)
- To search for a variable press / key, and to repeat the search, press the n key

To build the default configuration using the GUI, from the builddir enter the ccmake command and point to the sourcedir:

### % ccmake ../sourcedir

The default configuration screen is shown in Figure A.1.

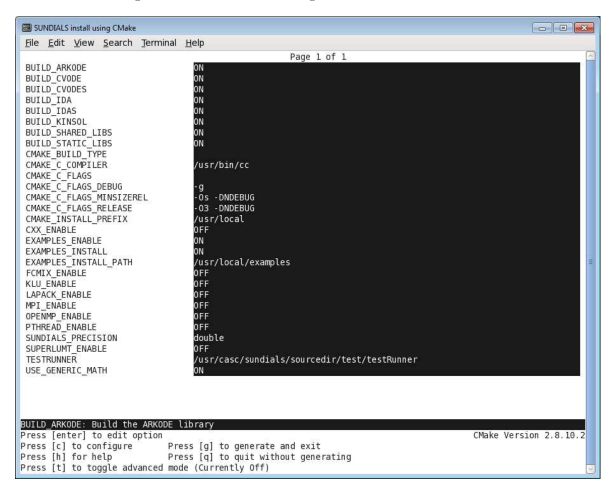


Figure A.1: Default configuration screen. Note: Initial screen is empty. To get this default configuration, press 'c' repeatedly (accepting default values denoted with asterisk) until the 'g' option is available.

The default *installdir* for both SUNDIALS and corresponding examples can be changed by setting the CMAKE\_INSTALL\_PREFIX and the EXAMPLES\_INSTALL\_PATH as shown in figure A.2.

Pressing the (g key) will generate makefiles including all dependencies and all rules to build SUN-DIALS on this system. Back at the command prompt, you can now run:

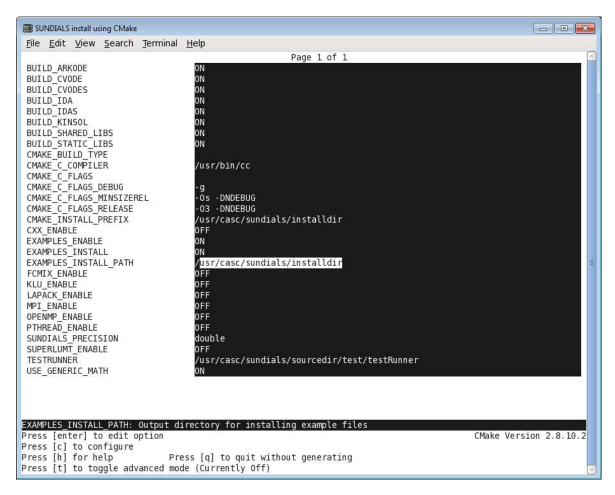


Figure A.2: Changing the *installdir* for SUNDIALS and corresponding examples

To install SUNDIALS in the installation directory specified in the configuration, simply run:

% make install

### Building from the command line

Using CMake from the command line is simply a matter of specifying CMake variable settings with the cmake command. The following will build the default configuration:

```
% cmake -DCMAKE_INSTALL_PREFIX=/home/myname/sundials/installdir \
> -DEXAMPLES_INSTALL_PATH=/home/myname/sundials/installdir/examples \
> ../sourcedir
% make
% make test
```

### A.1.2 Configuration options (Unix/Linux)

A complete list of all available options for a CMake-based SUNDIALS configuration is provide below. Note that the default values shown are for a typical configuration on a Linux system and are provided as illustration only.

```
 \begin{array}{c} {\tt BUILD\_ARKODE} \ - \ {\tt Build} \ \ {\tt the} \ \ {\tt ARKODE} \ \ {\tt library} \\ {\tt Default:} \ \ {\tt ON} \end{array}
```

BUILD\_CVODE - Build the CVODE library

Default: ON

BUILD\_CVODES - Build the CVODES library

Default: ON

BUILD\_IDA - Build the IDA library

Default: ON

BUILD\_IDAS - Build the IDAS library

Default: ON

BUILD\_KINSOL - Build the KINSOL library

Default: ON

BUILD\_SHARED\_LIBS - Build shared libraries

Default: OFF

BUILD\_STATIC\_LIBS - Build static libraries

Default: ON

CMAKE\_BUILD\_TYPE - Choose the type of build, options are: None (CMAKE\_C\_FLAGS used) Debug

 $Release\ RelWithDebInfo\ MinSizeRel$ 

Default:

CMAKE\_C\_COMPILER - C compiler

Default: /usr/bin/cc

CMAKE\_C\_FLAGS - Flags for C compiler

Default:

CMAKE\_C\_FLAGS\_DEBUG - Flags used by the compiler during debug builds

Default: -g

CMAKE\_C\_FLAGS\_MINSIZEREL - Flags used by the compiler during release minsize builds

Default: -Os -DNDEBUG

CMAKE\_C\_FLAGS\_RELEASE - Flags used by the compiler during release builds

Default: -O3 -DNDEBUG

CMAKE\_Fortran\_COMPILER - Fortran compiler

Default: /usr/bin/gfortran

Note: Fortran support (and all related options) are triggered only if either Fortran-C support is enabled (FCMIX\_ENABLE is ON) or Blas/Lapack support is enabled (LAPACK\_ENABLE is ON).

CMAKE\_Fortran\_FLAGS - Flags for Fortran compiler

Default:

 ${\tt CMAKE\_Fortran\_FLAGS\_DEBUG~- Flags~used~by~the~compiler~during~debug~builds}$ 

Default:

 ${\tt CMAKE\_Fortran\_FLAGS\_MINSIZEREL~Flags~used~by~the~compiler~during~release~minsize~builds}$ 

Default:

CMAKE\_Fortran\_FLAGS\_RELEASE - Flags used by the compiler during release builds

Default:

CMAKE\_INSTALL\_PREFIX - Install path prefix, prepended onto install directories

Default: /usr/local

Note: The user must have write access to the location specified through this option. Exported SUNDIALS header files and libraries will be installed under subdirectories include and lib of CMAKE\_INSTALL\_PREFIX, respectively.

### EXAMPLES\_ENABLE - Build the SUNDIALS examples

Default: ON

### EXAMPLES\_INSTALL - Install example files

Default: ON

Note: This option is triggered only if building example programs is enabled (EXAMPLES\_ENABLE ON). If the user requires installation of example programs then the sources and sample output files for all SUNDIALS modules that are currently enabled will be exported to the directory specified by EXAMPLES\_INSTALL\_PATH. A CMake configuration script will also be automatically generated and exported to the same directory. Additionally, if the configuration is done under a Unix-like system, makefiles for the compilation of the example programs (using the installed SUNDIALS libraries) will be automatically generated and exported to the directory specified by EXAMPLES\_INSTALL\_PATH.

### EXAMPLES\_INSTALL\_PATH - Output directory for installing example files

Default: /usr/local/examples

Note: The actual default value for this option will an examples subdirectory created under CMAKE\_INSTALL\_PREFIX.

### FCMIX\_ENABLE - Enable Fortran-C support

Default: OFF

### KLU\_ENABLE - Enable KLU support

Default: OFF

### LAPACK\_ENABLE - Enable Lapack support

Default: OFF

Note: Setting this option to ON will trigger the two additional options see below.

### LAPACK\_LIBRARIES - Lapack (and Blas) libraries

Default: /usr/lib/liblapack.so;/usr/lib/libblas.so

Note: CMake will search for these libraries in your LD\_LIBRARY\_PATH prior to searching default

system paths.

### MPI\_ENABLE - Enable MPI support

Default: OFF

Note: Setting this option to ON will trigger several additional options related to MPI.

### MPI\_MPICC - mpicc program

Default:

### MPI\_RUN\_COMMAND - Specify run command for MPI

Default: mpirun

Note: This can either be set to mpirun for OpenMPI or srun if jobs are managed by SLURM - Simple Linux Utility for Resource Management as exists on LLNL's high performance computing clusters.

### MPI\_MPIF77 - mpif77 program

Default:

Note: This option is triggered only if using MPI compiler scripts (MPI\_USE\_MPISCRIPTS is ON) and Fortran-C support is enabled (FCMIx\_ENABLE is ON).

### OPENMP\_ENABLE - Enable OpenMP support

Default: OFF

Turn on support for the OpenMP based nvector.

```
PTHREAD_ENABLE - Enable Pthreads support
     Default: OFF
     Turn on support for the Pthreads based nvector.
SUNDIALS_PRECISION - Precision used in SUNDIALS, options are: double, single or extended
     Default: double
SUPERLUMT_ENABLE - Enable SUPERLU_MT support
     Default: OFF
TESTRUNNER - Location of testRunner script
     Default: sourcedir/testRunner
USE_GENERIC_MATH - Use generic (stdc) math libraries
     Default: ON
```

#### A.1.3 Configuration examples

The following examples will help demonstrate usage of the CMake configure options.

To configure SUNDIALS using the default C and Fortran compilers, and default mpic and mpif77 parallel compilers, enable compilation of examples, and install libraries, headers, and example sources under subdirectories of /home/myname/sundials/, use:

```
% cmake \
> -DCMAKE_INSTALL_PREFIX=/home/myname/sundials/installdir \
> -DEXAMPLES_INSTALL_PATH=/home/myname/sundials/installdir/examples \
> -DMPI_ENABLE=ON \
> -DFCMIX_ENABLE=ON \
> /home/myname/sundials/sourcedir
% make install
```

To disable installation of the examples, use:

```
% cmake \
> -DCMAKE_INSTALL_PREFIX=/home/myname/sundials/installdir \
> -DEXAMPLES_INSTALL_PATH=/home/myname/sundials/installdir/examples \
> -DMPI_ENABLE=ON \
> -DFCMIX_ENABLE=ON \
> -DEXAMPLES_INSTALL=OFF \
> /home/myname/sundials/sourcedir
% make install
```

### Working with external Libraries

The SUNDIALS Suite contains many options to enable implementation flexibility when developing solutions. The following are some notes addressing specific configurations when using the supported third party libraries.

### Building with LAPACK and BLAS

To enable LAPACK and BLAS libraries, set the LAPACK\_ENABLE option to ON. If the directory containing the LAPACK and BLAS libraries is in the LD\_LIBRARY\_PATH environment variable, CMake will set the LAPACK\_LIBRARIES variable accordingly, otherwise CMake will attemp to find the LAPACK

and BLAS libraries in standard system locations. To explicitly tell CMake what libraries to use, the LAPACK\_LIBRARIES varible can be set to the desired libraries. Example:

```
% cmake \
> -DCMAKE_INSTALL_PREFIX=/home/myname/sundials/installdir \
> -DEXAMPLES_INSTALL_PATH=/home/myname/sundials/installdir/examples \
> -DLAPACK_LIBRARIES=/mypath/lib/liblapack.so;/mypath/lib/libblas.so \
> /home/myname/sundials/sourcedir
%
    make install
%
```

### Building with KLU

The KLU libraries are part of SuiteSparse, a suite of sparse matrix software, available from the Texas A&M University website: http://faculty.cse.tamu.edu/davis/suitesparse.html SUNDIALS has been tested with SuiteSparse version 4.2.1. To enable KLU, set KLU\_ENABLE to ON, set KLU\_INCLUDE\_DIR to the include path of the KLU installation and set KLU\_LIBRARY\_DIR to the lib path of the KLU installation. The CMake configure will result in populating the variables: AMD\_LIBRARY, AMD\_LIBRARY\_DIR, BTF\_LIBRARY, BTF\_LIBRARY\_DIR, COLAMD\_LIBRARY, COLAMD\_LIBRARY\_DIR, and KLU\_LIBRARY

### Building with SuperLU\_MT

The SuperLU\_MT libraries are available for download from the Lawrence Berkeley National Laboratory website: http://crd-legacy.lbl.gov/~xiaoye/SuperLU/#superlu\_mt. SUNDIALS has been tested with SuperLU\_MT version 2.4. To enable SuperLU\_MT, set SUPERLUMT\_ENABLE to ON, set SUPERLUMT\_INCLUDE\_DIR to the SRC path of the SuperLU\_MT installation and set SUPERLUMT\_LIBRARY\_DIR to the lib path of the SuperLU\_MT installation. Also, the SUPERLUMT\_THREAD\_TYPE must be set to either Pthread or OpenMP.



Do not mix thread types when building SUNDIALS solvers. If threading is enabled for SUNDIALS by having either OPENMP\_ENABLE or PTHREAD\_ENABLE set to ON then SuperLU\_MT should be set to use the same threading type.

## A.2 Building and Running Examples

Each of the SUNDIALS solvers is distributed with a set of examples demonstrating basic usage. To build and install the examples, set both EXAMPLES\_ENABLE and EXAMPLES\_INSTALL to ON. Specify the installation path for the examples with the variable EXAMPLES\_INSTALL\_PATH. CMake will generate CMakeLists.txt configuration files (and Makefile files if on Linux/Unix) that reference the *installed* SUNDIALS headers and libraries.

From within the installed example directory, run CMake (either with the GUI or command line) to compile the example code. The resulting output from running the example can be compared with example output bundled in the SUNDIALS distribution.



NOTE: There will likely differences in the output due to machine architecture, compiler versions, use of third party libraries etc.

These examples with the installed configuration files can be used as "templates" for user developed solutions.

## A.3 Configuring, building, and installing on Windows

Use CMakeSetup from the CMake install location. Make sure to select the appropriate source and the build directory. Also, make sure to pick the appropriate generator (on Visual Studio 6, pick the

Visual Studio 6 generator). Some CMake versions will ask you to select the generator the first time you press Configure instead of having a drop-down menu in the main dialog.

CMake will now create Visual Studio project files. You should now be able to open the SUNDIALS project (or workspace) file. Make sure to select the appropriate build type (Debug, Release, ...). To build SUNDIALS, simply build the ALL\_BUILD target. To install SUNDIALS, simply run the INSTALL target within the build system.

### A.4 Installed libraries and exported header files

Using the CMake SUNDIALS build system, the command

% make install

will install the libraries under *libdir* and the public header files under *includedir*. The default values for these directories are *instdir*/lib and *instdir*/include, respectively, but can be changed using the configure script options --prefix, --exec-prefix, --includedir and --libdir (see the appropriate CMake options). For example, a global installation of SUNDIALS on a \*NIX system could be accomplished using

% configure --prefix=/opt/sundials-2.1.1

Although all installed libraries reside under *libdir*, the public header files are further organized into subdirectories under *includedir*.

The installed libraries and exported header files are listed for reference in Tables A.1 and A.2. The file extension .lib is typically .so for shared libraries and .a for static libraries. Note that, in the Tables, names are relative to libdir for libraries and to includedir for header files.

A typical user program need not explicitly include any of the shared SUNDIALS header files from under the *includedir*/sundials directory since they are explicitly included by the appropriate solver header files (e.g., cvode\_dense.h includes sundials\_dense.h). However, it is both legal and safe to do so (e.g., the functions declared in sundials\_dense.h could be used in building a preconditioner).

Table A.1: SUNDIALS libraries and header files

SHARED	Libraries	n/a	
	Header files	sundials/sundials_config.h sundials/sundials_math.h	sundials/sundials_types.h
		sundials/sundials_nvector.h	sundials/sundials_fnvector.h
		sundials/sundials_direct.h	sundials/sundials_lapack.h
		sundials/sundials_dense.h	sundials/sundials_band.h
		sundials/sundials_sparse.h	,
		sundials/sundials_iterative.h	sundials/sundials_spgmr.h
		sundials/sundials_spbcgs.h	sundials/sundials_sptfqmr.h
		sundials/sundials_pcg.h	sundials/sundials_spfgmr.h
NVECTOR_SERIAL	Libraries	libsundials_nvecserial.lib	libsundials_fnvecserial.a
	Header files	nvector/nvector_serial.h	
NVECTOR_PARALLEL	Libraries	libsundials_nvecparallel.lib	libsundials_fnvecparallel.a
	Header files	nvector/nvector_parallel.h	
NVECTOR_OPENMP	Libraries	$libsundials\_nvecopenmp.lib$	libsundials_fnvecopenmp.a
	Header files	nvector/nvector_openmp.h	
NVECTOR_PTHREADS	Libraries	$libsundials\_nvecpthreads.lib$	$lib sundials\_fnvecpth reads. a$
	Header files	nvector/nvector_pthreads.h	
CVODE	Libraries	$libsundials\_cvode.lib$	libsundials_fcvode.a
	Header files	cvode/cvode.h	$cvode/cvode\_impl.h$
		cvode/cvode_direct.h	$cvode/cvode\_lapack.h$
		cvode/cvode_dense.h	$cvode/cvode\_band.h$
		cvode/cvode_diag.h	
		cvode/cvode_sparse.h	$cvode/cvode\_klu.h$
		cvode/cvode_superlumt.h	
		cvode/cvode_spils.h	cvode/cvode_spgmr.h
		cvode/cvode_sptfqmr.h	cvode/cvode_spbcgs.h
		cvode/cvode_bandpre.h	cvode/cvode_bbdpre.h
CVODES	Libraries	libsundials_cvodes.lib	
	Header files	cvodes/cvodes.h	cvodes/cvodes_impl.h
		cvodes/cvodes_direct.h	cvodes/cvodes_lapack.h
		cvodes/cvodes_dense.h	$cvodes/cvodes\_band.h$
		cvodes/cvodes_diag.h	1 / 1 11 1
		cvodes/cvodes_sparse.h	cvodes/cvodes_klu.h
		cvodes/cvodes_superlumt.h	1 / 1
		cvodes/cvodes_spils.h	cvodes/cvodes_spgmr.h
		cvodes/cvodes_sptfqmr.h	cvodes/cvodes_spbcgs.h
	T '1 '	cvodes/cvodes_bandpre.h	cvodes/cvodes_bbdpre.h
ARKODE	Libraries	libsundials_arkode.lib	libsundials_farkode.a
	Header files	arkode/arkode.h	arkode/arkode_impl.h
		arkode/arkode_direct.h	arkode/arkode_lapack.h
		arkode/arkode_dense.h	arkode/arkode_band.h
		arkode/arkode_sparse.h	arkode/arkode_klu.h
		arkode/arkode_superlumt.h	arkoda/arkoda anama h
		arkode/arkode_spils.h	arkode/arkode_spgmr.h
		arkode/arkode_sptfqmr.h arkode/arkode_pcg.h	arkode/arkode_spbcgs.h arkode/arkode_spfgmr.h
		arkode/arkode_pcg.n arkode/arkode_bandpre.h	arkode/arkode_spigmr.n arkode/arkode_bbdpre.h
		arkode/arkode_bandpre.h	arkode/arkode_bbdpre.ii

Table A.2: SUNDIALS libraries and header files (cont.)

IDA	Libraries	libsundials_ida.lib	libsundials_fida.a
	Header files	ida/ida.h	ida/ida_impl.h
		ida/ida_direct.h	ida/ida_lapack.h
		ida/ida_dense.h	ida/ida_band.h
		ida/ida_sparse.h	ida/ida_klu.h
		ida/ida_superlumt.h	,
		ida/ida_spils.h	ida/ida_spgmr.h
		ida/ida_spbcgs.h	ida/ida_sptfqmr.h
		ida/ida_bbdpre.h	,
IDAS	Libraries	libsundials_idas.lib	
	Header files	idas/idas.h	idas/idas_impl.h
		idas/idas_direct.h	idas/idas_lapack.h
		idas/idas_dense.h	idas/idas_band.h
		idas/idas_sparse.h	idas/idas_klu.h
		idas/idas_superlumt.h	
		idas/idas_spils.h	$idas/idas\_spgmr.h$
		$idas/idas\_spbcgs.h$	$idas/idas\_sptfqmr.h$
		idas/idas_bbdpre.h	
KINSOL	Libraries	libsundials_kinsol.lib	libsundials_fkinsol.a
	Header files	kinsol/kinsol.h	kinsol/kinsol_impl.h
		kinsol/kinsol_direct.h	kinsol/kinsol_lapack.h
		kinsol/kinsol_dense.h	kinsol/kinsol_band.h
		kinsol/kinsol_sparse.h	kinsol/kinsol_klu.h
		kinsol/kinsol_superlumt.h	
		kinsol/kinsol_spils.h	kinsol/kinsol_spgmr.h
		kinsol/kinsol_spbcgs.h	kinsol/kinsol_sptfqmr.h
		kinsol/kinsol_bbdpre.h	kinsol/kinsol_spfgmr.h

## Appendix B

## 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.

## **B.1** KINSOL input constants

KINSOL main solver module			
KIN_ETACHOICE1 KIN_ETACHOICE2 KIN_ETACONSTANT KIN_NONE KIN_LINESEARCH	1 2 3 0 1	Use Eisenstat and Walker Choice 1 for $\eta$ . Use Eisenstat and Walker Choice 2 for $\eta$ . Use constant value for $\eta$ . Use inexact Newton globalization. Use linesearch globalization.	
	Ite	rative linear solver module	
PREC_NONE PREC_RIGHT MODIFIED_GS CLASSICAL_GS	0 2 1 2	No preconditioning Preconditioning on the right. Use modified Gram-Schmidt procedure. Use classical Gram-Schmidt procedure.	

## B.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_WARNING	99	A non-fatal warning. The solver will continue.	
KIN_MEM_NULL	-1	The kin_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 linesearch algorithm was unable to find an iterate suffi- ciently distinct from the current iterate.	

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KIN_MAXITER_REACHED	-6	The maximum number of nonlinear iterations has been reached.
KIN_MXNEWT_5X_EXCEEDED	-7	Five consecutive steps have been taken that satisfy a scaled
KIN_LINESEARCH_BCFAIL	-8	step length test. The linesearch algorithm was unable to satisfy the $\beta$ - condition for nbcfails iterations.
KIN_LINSOLV_NO_RECOVERY	-9	The user-supplied routine preconditioner sive 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_SYSFUNC_FAIL	-13	The system function failed in an unrecoverable manner.
KIN_FIRST_SYSFUNC_ERR	-14	The system function failed recoverably at the first call.
KIN_REPTD_SYSFUNC_ERR	-15	The system function had repeated recoverable errors.
	KI	NDLS linear solver module
KINDLS_SUCCESS	0	Successful function return.
KINDLS_MEM_NULL	-1	The kin_mem argument was NULL.
KINDLS_LMEM_NULL	-2	The KINDLS linear solver has not been initialized.
KINDLS_ILL_INPUT	-3	The KINDLS solver is not compatible with the current NVEC-
		TOR module.
KINDLS_MEM_FAIL	-4	A memory allocation request failed.
KINDLS_JACFUNC_UNRECVR	-5	The Jacobian function failed in an unrecoverable manner.
KINDLS_JACFUNC_RECVR	-6	The Jacobian function had a recoverable error.
	KI	NSLS linear solver module
KINSLS_SUCCESS	0	Successful function return.
KINSLS_MEM_NULL	0 -1	The kin_mem argument was NULL.
KINSLS_MEM_NULL KINSLS_LMEM_NULL	0 -1 -2	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.
KINSLS_MEM_NULL	0 -1	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVEC-
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT	0 -1 -2 -3	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT KINSLS_MEM_FAIL	0 -1 -2 -3	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT	0 -1 -2 -3	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT KINSLS_MEM_FAIL KINSLS_JAC_NOSET	0 -1 -2 -3 -4 -5	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT KINSLS_MEM_FAIL KINSLS_JAC_NOSET KINSLS_PACKAGE_FAIL	0 -1 -2 -3 -4 -5	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT  KINSLS_MEM_FAIL KINSLS_JAC_NOSET  KINSLS_PACKAGE_FAIL KINSLS_JACFUNC_UNRECVR	0 -1 -2 -3 -4 -5 -6 -7	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.  The Jacobian function failed in an unrecoverable manner.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT KINSLS_MEM_FAIL KINSLS_JAC_NOSET KINSLS_PACKAGE_FAIL	0 -1 -2 -3 -4 -5	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT  KINSLS_MEM_FAIL KINSLS_JAC_NOSET  KINSLS_PACKAGE_FAIL KINSLS_JACFUNC_UNRECVR	0 -1 -2 -3 -4 -5 -6 -7 -8	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.  The Jacobian function failed in an unrecoverable manner.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT  KINSLS_MEM_FAIL KINSLS_JAC_NOSET  KINSLS_PACKAGE_FAIL KINSLS_JACFUNC_UNRECVR KINSLS_JACFUNC_RECVR	0 -1 -2 -3 -4 -5 -6 -7 -8	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.  The Jacobian function failed in an unrecoverable manner.  The Jacobian function had a recoverable error.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT  KINSLS_MEM_FAIL KINSLS_JAC_NOSET  KINSLS_PACKAGE_FAIL KINSLS_JACFUNC_UNRECVR KINSLS_JACFUNC_RECVR	0 -1 -2 -3 -4 -5 -6 -7 -8 KIN	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.  The Jacobian function failed in an unrecoverable manner.  The Jacobian function had a recoverable error.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT  KINSLS_MEM_FAIL KINSLS_JAC_NOSET  KINSLS_PACKAGE_FAIL KINSLS_JACFUNC_UNRECVR KINSLS_JACFUNC_RECVR  KINSPILS_SUCCESS KINSPILS_MEM_NULL	0 -1 -2 -3 -4 -5 -6 -7 -8 KIN	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.  The Jacobian function failed in an unrecoverable manner.  The Jacobian function had a recoverable error.  SPILS linear solver modules  Successful function return.  The kin_mem argument was NULL.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT  KINSLS_MEM_FAIL KINSLS_JAC_NOSET  KINSLS_JACFUNC_UNRECVR KINSLS_JACFUNC_RECVR  KINSLS_JACFUNC_RECVR  KINSPILS_SUCCESS KINSPILS_MEM_NULL KINSPILS_LMEM_NULL	0 -1 -2 -3 -4 -5 -6 -7 -8 KIN:	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.  The Jacobian function failed in an unrecoverable manner.  The Jacobian function had a recoverable error.  SPILS linear solver modules  Successful function return.  The kin_mem argument was NULL.  The KINSPILS linear solver has not been initialized.
KINSLS_MEM_NULL KINSLS_LMEM_NULL KINSLS_ILL_INPUT  KINSLS_MEM_FAIL KINSLS_JAC_NOSET  KINSLS_PACKAGE_FAIL KINSLS_JACFUNC_UNRECVR KINSLS_JACFUNC_RECVR  KINSPILS_SUCCESS KINSPILS_MEM_NULL	0 -1 -2 -3 -4 -5 -6 -7 -8 KIN	The kin_mem argument was NULL.  The KINSLS linear solver has not been initialized.  The KINSLS solver is not compatible with the current NVECTOR module or other input is invalid.  A memory allocation request failed.  The Jacobian evaluation routine was not been set before the linear solver setup routine was called.  An external package call return a failure error code.  The Jacobian function failed in an unrecoverable manner.  The Jacobian function had a recoverable error.  SPILS linear solver modules  Successful function return.  The kin_mem argument was NULL.

KINSPILS_MEM_FAIL	-4	A memory allocation request failed.
KINSPILS_PMEM_NULL	-5	The preconditioner module has not be

The preconditioner module has not been initialized. -5

### SPGMR 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_ATIMES_FAIL_REC	5	The Jacobian-times-vector function failed recoverably.
SPGMR_PSET_FAIL_REC	6	The preconditioner setup routine failed recoverably.
SPGMR_MEM_NULL	-1	The SPGMR memory is NULL
SPGMR_ATIMES_FAIL_UNREC	-2	The Jacobian-times-vector function failed unrecoverably.
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.
SPGMR_PSET_FAIL_UNREC	-6	The preconditioner setup routine failed unrecoverably.

### SPFGMR generic linear solver module (only available in KINSOL and ARKODE)

0	Converged.
1	No convergence, but the residual norm was reduced.
$^2$	Failure to converge.
3	A singular matrix was found during the QR factorization.
4	The preconditioner solve function failed recoverably.
5	The Jacobian-times-vector function failed recoverably.
6	The preconditioner setup routine failed recoverably.
-1	The SPFGMR memory is NULL
-2	The Jacobian-times-vector function failed unrecoverably.
-3	The preconditioner solve function failed unrecoverably.
-4	Failure in the Gram-Schmidt procedure.
-5	The matrix $R$ was found to be singular during the QR solve
	phase.
-6	The preconditioner setup routine failed unrecoverably.
	1 2 3 4 5 6 -1 -2 -3 -4 -5

### SPBCG generic linear solver module

SPBCG_SUCCESS	0	Converged.
SPBCG_RES_REDUCED	1	No convergence, but the residual norm was reduced.
SPBCG_CONV_FAIL	2	Failure to converge.
SPBCG_PSOLVE_FAIL_REC	3	The preconditioner solve function failed recoverably.
SPBCG_ATIMES_FAIL_REC	4	The Jacobian-times-vector function failed recoverably.
SPBCG_PSET_FAIL_REC	5	The preconditioner setup routine failed recoverably.
SPBCG_MEM_NULL	-1	The SPBCG memory is NULL
SPBCG_ATIMES_FAIL_UNREC	-2	The Jacobian-times-vector function failed unrecoverably.
SPBCG_PSOLVE_FAIL_UNREC	-3	The preconditioner solve function failed unrecoverably.

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SPBCG\_PSET\_FAIL\_UNREC -4 The preconditioner setup routine failed unrecoverably.

### SPTFQMR generic linear solver module

SPTFQMR_SUCCESS	0	Converged.
SPTFQMR_RES_REDUCED	1	No convergence, but the residual norm was reduced.
SPTFQMR_CONV_FAIL	2	Failure to converge.
SPTFQMR_PSOLVE_FAIL_REC	3	The preconditioner solve function failed recoverably.
SPTFQMR_ATIMES_FAIL_REC	4	The Jacobian-times-vector function failed recoverably.
SPTFQMR_PSET_FAIL_REC	5	The preconditioner setup routine failed recoverably.
SPTFQMR_MEM_NULL	-1	The SPTFQMR memory is NULL
SPTFQMR_ATIMES_FAIL_UNREC	-2	The Jacobian-times-vector function failed.
SPTFQMR_PSOLVE_FAIL_UNREC	-3	The preconditioner solve function failed unrecoverably.
SPTFQMR_PSET_FAIL_UNREC	-4	The preconditioner setup routine failed unrecoverably.

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