# User Documentation for CVODES v2.5.0 

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November 6, 2006


UCRL-SM-208111

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This research was supported under the auspices of the U.S. Department of Energy by the University of California, Lawrence Livermore National Laboratory under contract No. W-7405-Eng-48.

## Contents

List of Tables ..... vii
List of Figures ..... ix
1 Introduction ..... 1
1.1 Historical background ..... 1
1.2 Changes from previous versions ..... 2
1.3 Reading this user guide ..... 3
2 CVODES Installation Procedure ..... 5
2.1 Configuration options ..... 6
2.2 Configuration examples ..... 10
2.3 Installed libraries and exported header files ..... 11
2.4 Building SUNDIALS without the configure script ..... 11
3 Mathematical Considerations ..... 15
3.1 IVP solution ..... 15
3.2 Forward sensitivity analysis ..... 18
3.2.1 Forward sensitivity methods ..... 19
3.2.2 Selection of the absolute tolerances for sensitivity variables ..... 20
3.2.3 Evaluation of the sensitivity right-hand side ..... 21
3.3 Adjoint sensitivity analysis ..... 21
3.3.1 Checkpointing scheme ..... 23
3.4 Preconditioning ..... 24
3.5 BDF stability limit detection ..... 24
3.6 Rootfinding ..... 25
4 Code Organization ..... 27
4.1 SUNDIALS organization ..... 27
4.2 CVODES organization ..... 27
5 Using CVODES for IVP Solution ..... 31
5.1 Access to libraries and header files ..... 31
5.2 Data Types ..... 32
5.3 Header files ..... 32
5.4 A skeleton of the user's main program ..... 33
5.5 User-callable functions for IVP solution ..... 35
5.5.1 CVODES initialization and deallocation functions ..... 35
5.5.2 Advice on choice and use of tolerances ..... 36
5.5.3 Linear solver specification functions ..... 37
5.5.4 CVODE solver function ..... 40
5.5.5 Optional input functions ..... 41
5.5.5.1 Main solver optional input functions ..... 43
5.5.5.2 Dense linear solver ..... 48
5.5.5.3 Band linear solver ..... 49
5.5.5.4 SPILS linear solver ..... 49
5.5.6 Interpolated output function ..... 52
5.5.7 Optional output functions ..... 52
5.5.7.1 Main solver optional output functions ..... 52
5.5.7.2 Dense linear solver ..... 59
5.5.7.3 Band linear solver ..... 61
5.5.7.4 Diagonal linear solver ..... 62
5.5.7.5 SPILS linear solvers ..... 63
5.5.8 CVODES reinitialization function ..... 66
5.6 User-supplied functions ..... 67
5.6.1 ODE right-hand side ..... 67
5.6.2 Error message handler function ..... 68
5.6.3 Error weight function ..... 69
5.6.4 Jacobian information (direct method with dense Jacobian) ..... 69
5.6.5 Jacobian information (direct method with banded Jacobian) ..... 70
5.6.6 Jacobian information (matrix-vector product) ..... 71
5.6.7 Preconditioning (linear system solution) ..... 72
5.6.8 Preconditioning (Jacobian data) ..... 72
5.7 Integration of pure quadrature equations ..... 73
5.7.1 Quadrature initialization functions ..... 75
5.7.2 CVODE solver function ..... 76
5.7.3 Quadrature extraction functions ..... 76
5.7.4 Optional inputs for quadrature integration ..... 77
5.7.5 Optional outputs for quadrature integration ..... 78
5.7.6 User-supplied function for quadrature integration ..... 79
5.8 Rootfinding ..... 80
5.8.1 User-callable functions for rootfinding ..... 80
5.8.2 User-supplied function for rootfinding ..... 81
5.9 Preconditioner modules ..... 81
5.9.1 A serial banded preconditioner module ..... 81
5.9.2 A parallel band-block-diagonal preconditioner module ..... 85
6 Using CVODES for Forward Sensitivity Analysis ..... 93
6.1 A skeleton of the user's main program ..... 93
6.2 User-callable routines for forward sensitivity analysis ..... 95
6.2.1 Forward sensitivity initialization and deallocation functions ..... 95
6.2.2 CVODE solver function ..... 98
6.2.3 Forward sensitivity extraction functions ..... 98
6.2.4 Optional inputs for forward sensitivity analysis ..... 100
6.2.5 Optional outputs for forward sensitivity analysis ..... 103
6.3 User-supplied routines for forward sensitivity analysis ..... 107
6.3.1 Sensitivity equations right-hand side (all at once) ..... 107
6.3.2 Sensitivity equations right-hand side (one at a time) ..... 108
6.4 Note on using partial error control ..... 109
7 Using CVODES for Adjoint Sensitivity Analysis ..... 111
7.1 A skeleton of the user's main program ..... 111
7.2 User-callable functions for adjoint sensitivity analysis ..... 113
7.2.1 Adjoint sensitivity allocation and deallocation functions ..... 113
7.2.2 Forward integration function ..... 114
7.2.3 Backward problem initialization functions ..... 116
7.2.4 Linear solver initialization functions for backward problem ..... 117
7.2.5 Backward integration function ..... 118
7.2.6 Optional input functions for the backward problem ..... 119
7.2.6.1 Main solver optional input functions ..... 119
7.2.6.2 Dense linear solver ..... 119
7.2.6.3 Band linear solver ..... 120
7.2.6.4 SPILS linear solvers ..... 120
7.2.7 Optional output functions for the backward problem ..... 123
7.2.8 Backward integration of pure quadrature equations ..... 123
7.2.8.1 Backward quadrature initialization functions ..... 123
7.2.8.2 Backward quadrature extraction function ..... 124
7.2.8.3 Optional input/output functions for backward quadrature integration ..... 124
7.2.9 Optional output from the adjoint module ..... 124
7.2.9.1 Checkpoint information function ..... 124
7.2.9.2 Interpolation data ..... 125
7.2.9.3 Return flag name ..... 126
7.3 User-supplied functions for adjoint sensitivity analysis ..... 126
7.3.1 ODE right-hand side for the backward problem ..... 126
7.3.2 Quadrature right-hand side for the backward problem ..... 127
7.3.3 Jacobian information for the backward problem (direct method with dense Ja- cobian) ..... 128
7.3.4 Jacobian information for the backward problem (direct method with banded Jacobian) ..... 129
7.3.5 Jacobian information for the backward problem (matrix-vector product) ..... 130
7.3.6 Preconditioning for the backward problem (linear system solution) ..... 130
7.3.7 Preconditioning for the backward problem (Jacobian data) ..... 131
7.4 Using CVODES preconditioner modules for the backward problem ..... 132
7.4.1 Using the banded preconditioner CVBANDPRE ..... 132
7.4.2 Using the band-block-diagonal preconditioner CVBBDPRE ..... 134
7.4.2.1 Usage of CVBBDPRE for the backward problem ..... 134
7.4.2.2 User-supplied functions for CVBBDPRE ..... 137
8 Description of the NVECTOR module ..... 139
8.1 The NVECTOR_SERIAL implementation ..... 143
8.2 The NVECTOR_PARALLEL implementation ..... 145
8.3 NVECTOR functions used by CVODES ..... 147
9 Providing Alternate Linear Solver Modules ..... 149
9.1 Initialization function ..... 150
9.2 Setup function ..... 150
9.3 Solve function ..... 151
9.4 Memory deallocation function ..... 151
10 Generic Linear Solvers in SUNDIALS ..... 153
10.1 The DENSE module ..... 153
10.1.1 Type DenseMat ..... 154
10.1.2 Accessor Macros ..... 154
10.1.3 Functions ..... 154
10.1.4 Small Dense Matrix Functions ..... 155
10.2 The BAND module ..... 156
10.2.1 Type BandMat ..... 157
10.2.2 Accessor Macros ..... 157
10.2.3 Functions ..... 159
10.3 The SPGMR module ..... 159
10.3.1 Functions ..... 160
10.4 The SPBCG module ..... 160
10.4.1 Functions ..... 160
10.5 The SPTFQMR module ..... 161
10.5.1 Functions ..... 161
11 CVODES Constants ..... 163
11.1 CVODES input constants ..... 163
11.2 CVODES output constants ..... 164
Bibliography ..... 169
Index ..... 171

## List of Tables

2.1 SUNDIALS libraries and header files (names are relative to libdir for libraries and to includedir for header files) ..... 12
5.1 Optional inputs for CVODES, CVDENSE, CVBAND, and CVSPILS ..... 42
5.2 Optional outputs from CVODES, CVDENSE, CVBAND, CVDIAG, and CVSPILS ..... 53
6.1 Forward sensitivity optional inputs ..... 100
6.2 Forward sensitivity optional outputs ..... 103
8.1 Description of the NVECTOR operations ..... 141
8.2 List of vector functions usage by CVODES code modules ..... 148

## List of Figures

3.1 Illustration of the checkpointing algorithm for generation of the forward solution during the integration of the adjoint system. ..... 23
4.1 Organization of the SUNDIALS suite ..... 28
4.2 Overall structure of the CVODES package ..... 29
10.1 Diagram of the storage for a matrix of type BandMat ..... 158

## Chapter 1

## Introduction

CVODES [26] is part of a software family called sundials: SUite of Nonlinear and DIfferential/ALgebraic equation Solvers [16]. This suite consists of CVODE, KINSOL and IDA, and variants of these with sensitivity analysis capabilities. CVODES is a solver for stiff and nonstiff initial value problems (IVPs) for systems of ordinary differential equation (ODEs). In addition to solving stiff and nonstiff ODE systems, CVODES has sensitivity analysis capabilities, using either the forward or the adjoint methods.

### 1.1 Historical background

Fortran solvers for ODE initial value problems are widespread and heavily used. Two solvers that were previously written at LLNL are VODE [1] and VODPK [3]. VODE is a general-purpose solver that includes methods for both stiff and nonstiff systems, and in the stiff case uses direct methods (full or banded) for the solution of the linear systems that arise at each implicit step. Externally, VODE is very similar to the well known solver LSODE [24]. VODPK is a variant of VODE that uses a preconditioned Krylov (iterative) method, namely GMRES, for the solution of the linear systems. VODPK is a powerful tool for large stiff systems because it combines established methods for stiff integration, nonlinear iteration, and Krylov (linear) iteration with a problem-specific treatment of the dominant source of stiffness, in the form of the user-supplied preconditioner matrix [2]. The capabilities of both VODE and VODPK were combined in the C-language package CVODE [8, 9].

At present, CVODES contains three Krylov methods that can be used in conjuction with Newton iteration: the GMRES (Generalized Minimal RESidual) [25], Bi-CGStab (Bi-Conjugate Gradient Stabilized) [27], and TFQMR (Transpose-Free Quasi-Minimal Residual) linear iterative methods [11]. As Krylov methods, these require almost no matrix storage for solving the Newton equations 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 stiff ODE systems, the Krylov methods are preferable over direct linear solver methods, and are often the only feasible choice. Among the three Krylov methods in CVODES, we recommend GMRES as the best overall choice. However, users are encouraged to compare all three, especially if encountering convergence failures with GMRES. Bi-CGFStab 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.

In the process of translating the VODE and VODPK algorithms into C, the overall CVODE organization has changed considerably. One key feature of the CVODE organization is that the linear system solvers comprise a layer of code modules that is separated from the integration algorithm, thus allowing for easy modification and expansion of the linear solver array. A second key feature is a separate module devoted to vector operations; this facilitated the extension to multiprosessor environments with only a minimal impact on the rest of the solver, resulting in PVODE [5], the parallel variant of CVODE.

CVODES is written with a functionality that is a superset of that of the pair CVODE/PVODE. Sensitivity analysis capabilities, both forward and adjoint, have been added to the main integrator.

Enabling forward sensititivity computations in CVODES will result in the code integrating the socalled sensitivity equations simultaneously with the original IVP, yielding both the solution and its sensitivity with respect to parameters in the model. Adjoint sensitivity analysis, most useful when the gradients of relatively few functionals of the solution with respect to many parameters are sought, involves integration of the original IVP forward in time followed by the integration of the so-called adjoint equations backward in time. CVODES provides the infrastructure needed to integrate any final-condition ODE dependent on the solution of the original IVP (in particular the adjoint system).

Development of CVODES was concurrent with a redesign of the vector operations module across the sundials suite. The key feature of the new nVECTOR module is that it is written in terms of abstract vector operations with the actual vector functions attached by a particular implementation (such as serial or parallel) of NVECTOR. This allows writing the SUNDIALS solvers in a manner independent of the actual NVECTOR implementation (which can be user-supplied), as well as allowing more than one NVECTOR module to be linked into an executable file.

There were several motivations for choosing the C language for CVODE, and later for cvodes. First, a general movement away from Fortran and toward C in scientific computing was and still is apparent. Second, the pointer, structure, and dynamic memory allocation features in C are extremely useful in software of this complexity. Finally, we prefer C over C++ for CVODES 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 extended Fortran.

### 1.2 Changes from previous versions

## Changes in v2.5.0

The main changes in this release involve a rearrangement of the entire SUNDIALS source tree (see $\S 4.1$ ). 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 instaltion include directory.

In the adjoint solver module, the following two bugs were fixed: in CVodeF the solver was sometimes incorrectly taking an additional step before returning control to the user (in CV NORMAL mode) thus leading to a failure in the interpolated output function; in CVodeB, while searching for the current check point, the solver was sometimes reaching outside the integration interval resulting in a segmentation fault.

The functions in the generic dense linear solver (sundials_dense and sundials_smalldense) were modified to work for rectangular $m \times n$ matrices $(m \leq n)$, 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

CVSPBCG and CVSPTFQMR modules have been added to interface with the Scaled Preconditioned Bi-CGstab (SpBCG) and Scaled Preconditioned Transpose-Free Quasi-Minimal Residual (Sptfamr) linear solver modules, respectively (for details see Chapter 5). At the same time, function type names for Scaled Preconditioned Iterative Linear Solvers were added for the user-supplied Jacobian-timesvector and preconditioner setup and solve functions.

A new interpolation method was added to the CVODEA adjoint module. The function CVadjMalloc has an additional argument which can be used to select the desired interpolation scheme.

The deallocation functions now take as arguments the address of the respective memory block pointer.

To reduce the possibility of conflicts, the names of all header files have been changed by adding unique prefixes (cvodes_ 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 $\S 2$.

## Changes in v2.3.0

A minor bug was fixed in the interpolation functions of the adjoint CVODEA module.

## Changes in v2.2.0

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

## Changes in v2.1.2

A bug was fixed in the CVode function that was potentially leading to erroneous behaviour of the root finding procedure on the integration first step.

## Changes in v2.1.1

This CVODES release includes bug fixes related to forward sensitivity computations (possible loss of accuray on a BDF order increase and incorrect logic in testing user-supplied absolute tolerances). In addition, we have added the option of activating and deactivating forward sensitivity calculations on successive CVODES runs without memory allocation/deallocation.

Other changes in this minor sundials release affect the build system.

## Changes in v2.1.0

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, CVODES now provides a set of routines (with prefix CVodeSet) to change the default values for various quantities controlling the solver and a set of extraction routines (with prefix CVodeGet) 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 $\S 5.5 .5$ and $\S 5.5 .7$.

Additionally, the interfaces to several user-supplied routines (such as those providing Jacobians, preconditioner information, and sensitivity right hand sides) 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.

The rootfinding feature was added, whereby the roots of a set of given functions may be computed during the integration of the ODE system.

Installation of CVODES (and all of SUNDIALS) has been completely redesigned and is now based on a configure script.

### 1.3 Reading this user guide

This user guide is a combination of general usage instructions. Specific example programs are provided as a separate document. We expect that some readers will want to concentrate on the general instructions, while others will refer mostly to the examples.

There are different possible levels of usage of CVODES. The most casual user, with an IVP problem only, can get by with reading $\S 3.1$, then $\S 5$ through $\S 5.5 .4$ only, and looking at examples in [18]. In addition, to solve a forward sensitivity problem the user should read $\S 3.2$, followed by $\S 6$ through $\S 6.2 .3$ only, and look at examples in [18].

In a different direction, a more advanced user with an IVP problem may want to (a) use a package preconditioner (§5.9), (b) supply his/her own Jacobian or preconditioner routines (§5.6), (c) do multiple runs of problems of the same size (§5.5.8), (d) supply a new NVECTOR module (§8), or even (e)
supply a different linear solver module (§4.2). An advanced user with a forward sensitivity problem may also want to (a) provide his/her own sensitivity equations right-hand side routine (§6.3), (b) perform multiple runs with the same number of sensitivity parameters (§6.2.1), or (c) extract additional diagnostic information ( $\S 6.2 .3$ ). A user with an adjoint sensitivity problem needs to understand the IVP solution approach at the desired level and also go through $\S 3.3$ for a short mathematical description of the adjoint approach, $\S 7$ for the usage of the adjoint module in CVODES, and the examples in [18].

The structure of this document is as follows:

- In Chapter 2 we begin with instructions for the installation of CVODES, within the structure of SUNDIALS.
- In Chapter 3, we give short descriptions of the numerical methods implemented by CVODES for the solution of initial value problems for systems of ODEs, continue with an overview of the mathematical aspects of sensitivity analysis, both forward (§3.2) and adjoint (§3.3), and conclude with a description of stability limit detection (§3.5).
- The following chapter describes the structure of the sundials suite of solvers (§4.1) and the software organization of the CVODES solver (§4.2).
- Chapter 5 is the main usage document for CVODES for simulation applications. It includes a complete description of the user interface for the integration of ODE initial value problems. Readers that are not interested in using CVODES for sensitivity analysis can then skip the next two chapters.
- Chapter 6 describes the usage of CVODES for forward sensitivity analysis as an extension of its IVP integration capabilities. We begin with a skeleton of the user main program, with emphasis on the steps that are required in addition to those already described in Chapter 5. Following that we provide detailed descriptions of the user-callable interface routines specific to forward sensitivity analysis and of the additonal optional user-defined routines.
- Chapter 7 describes the usage of CVODES for adjoint sensitivity analysis. We begin by describing the CVODES checkpointing implementation for interpolation of the original IVP solution during integration of the adjoint system backward in time, and with an overview of a user's main program. Following that we provide complete descriptions of the user-callable interface routines for adjoint sensitivity analysis as well as descriptions of the required additional user-defined routines.
- Chapter 8 gives a brief overview of the generic NVECTOR module shared amongst the various components of SUNDIALS, as well as details on the two NVECTOR implementations provided with SUNDIALS: a serial implementation (§8.1) and a parallel implementation based on MPI (§8.2).
- Chapter 9 describes the specifications of linear solver modules as supplied by the user.
- Chapter 10 describes in detail the generic linear solvers shared by all SUNDIALS solvers.
- Finally, Chapter 11 lists the constants used for input to and output from CVODES.

Finally, the reader should be aware of the following notational conventions in this user guide: Program listings and identifiers (such as CVodeMalloc) within textual explanations appear in typewriter type style; fields in C structures (such as content) appear in italics; and packages or modules, such as CVDENSE, are written in all capitals. In the Index, page numbers that appear in bold indicate the main reference for that entry.

## Chapter 2

## CVODES Installation Procedure

The installation of CVODES is accomplished by installing the SUNDIALS suite as a whole, according to the instructions that follow. The same procedure applies whether or not the downloaded file contains solvers other than CVODES. ${ }^{1}$

The sundials suite (or individual solvers) are distributed as compressed archives (.tgz). The name of the distribution archive is of the form solver $-\mathrm{x} \cdot \mathrm{y} . \mathrm{z} . \mathrm{tgz}$, where solver is one of: sundials, cvode, cvodes, ida, 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.tgz
This will extract source files under a directory solver-x.y.z.
The installation procedure outlined below will work on commodity LINUX/UNIX systems without modification. However, users are still encouraged to carefully read the entire chapter before attempting to install the SUNDIALS suite, in case non-default choices are desired for compilers, compilation options, or the like. In lieu of reading the option list below, the user may invoke the configuration script with the help flag to view a complete listing of available options, which may be done by issuing

```
% ./configure --help
```

from within the directory created above.
In the remainder of this chapter, we make the following distinctions:

- srcdir
is the directory solver-x.y.z created above; i.e., the directory containing the SUNDIALS sources.
- builddir
is the directory under which SUNDIALS is built; i.e., the directory from within which the configure command is issued. Usually, this is the same as srcdir.
- instdir
is the directory under which the sundials exported header files and libraries will be installed. Typically, header files are exported under a directory instdir/include while libraries are installed under instdir/lib, with instdir specified with the --prefix flag to configure. See $\S 2.1$ for more details on the installation directories, including the special cases of the SUNDIALS examples and the SUNDIALSTB toolbox.

Note: The installation directory instdir should not be the same as the source directory srcdir.


[^0]The installation steps for sundials can be as simple as

```
% tar xzf solver-x.y.z.tgz
% cd solver-x.y.z
% ./configure
% make
% make install
```

in which case the SUNDIALS header files and libraries are installed under /usr/local/include and /usr/local/lib, respectively. Note that, by default, neither the example programs nor the SUNDIALSTB toolbox are built and installed.

If disk space is a priority, then to delete all temporary files created by building sundials, issue

```
% make clean
```

To prepare the sundials distribution for a new install (using, for example, different options and/or installation destinations), issue
\% make distclean

### 2.1 Configuration options

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

## General options

```
--prefix=PREFIX
```

Location for architecture-independent files.
Default: PREFIX=/usr/local
--exec-prefix=EPREFIX
Location for architecture-dependent files.
Default: EPREFIX=/usr/local
--includedir=DIR
Alternate location for installation of header files.
Default: DIR=PREFIX/include
--libdir=DIR
Alternate location for installation of libraries.
Default: DIR=EPREFIX/lib
--disable-solver
Although each existing solver module is built by default, support for a given solver can be explicitly disabled using this option. The valid values for solver are: cvode, cvodes, ida, and kinsol.
--enable-examples
Available example programs are not built by default. Use this option to enable compilation of all pertinent example programs. Upon completion of the make command, the example executables will be created under solver-specific subdirectories of builddir/examples:
builddir/examples/solver/serial: serial C examples
builddir/examples/solver/parallel : parallel C examples
builddir/examples/solver/fcmix_serial : serial FORTRAN examples
builddir/examples/solver/fcmix_parallel : parallel Fortran examples
Note: Some of these subdirectories may not exist depending upon the solver and/or the configuration options given.
--with-examples-instdir=EXINSTDIR
Alternate location for example executables and sample output files (valid only if examples are enabled). Note that installtion of example files can be completely disabled by issuing EXINSTDIR=no (in case building the examples is desired only as a test of the SUNDIALS libraries).
Default: DIR=EPREFIX/examples
--with-cppflags=ARG
Specify additional C preprocessor flags (e.g., ARG=-I<include_dir> if necessary header files are located in nonstandard locations).
--with-cflags=ARG
Specify additional C compilation flags.
--with-ldflags=ARG
Specify additional linker flags (e.g., ARG=-L<lib_dir> if required libraries are located in nonstandard locations).
--with-libs=ARG
Specify additional libraries to be used (e.g., ARG=-l<foo> to link with the library named libfoo.a or libfoo.so).
--with-precision=ARG
By default, SUNDIALS will define a real number (internally referred to as realtype) to be a double-precision floating-point numeric data type (double C-type); however, this option may be used to build sundials with realtype alternatively defined as a single-precision floating-point numeric data type (float C-type) if ARG=single, or as a long double C-type if
ARG=extended.
Default: ARG=double
Users should not build SUNDIALS with support for single-precision floating-point arithmetic on 32 - or 64 -bit systems. This will almost certainly result in unreliable numerical solutions. The configuration option --with-precision=single is intended for systems on which single-precision arithmetic involves at least 14 decimal digits.

## Options for Fortran support

--disable-fcmix
Using this option will disable all Fortran support. The fCVode, fkinsol, fida, and fnvector modules will not be built, regardless of availability.
--with-fflags=ARG
Specify additional Fortran compilation flags.
The configuration script will attempt to automatically determine the function name mangling scheme required by the specified Fortran compiler, but the following two options may be used to override the default behavior.
--with-f77underscore=ARG
This option pertains to the FCvode, fkinsol, fida, and fnvector Fortran-C interface modules and is used to specify the number of underscores to append to function names so Fortran routines can properly link with the associated SUNDIALS libraries. Valid values for ARG are: none, one and two.

Default: ARG=one
--with-f77case=ARG
Use this option to specify whether the external names of the FCVODE, FKINSOL, FIDA, and FNvector Fortran-C interface functions should be lowercase or uppercase so Fortran routines can properly link with the associated sundials libraries. Valid values for ARG are: lower and upper.

Default: ARG=lower

## Options for MPI support

The following configuration options are only applicable to the parallel SUNDIALS packages:
--disable-mpi
Using this option will completely disable MPI support.
--with-mpicc=ARG
--with-mpif77=ARG
By default, the configuration utility script will use the MPI compiler scripts named mpicc and mpif77 to compile the parallelized SUNDIALS subroutines; however, for reasons of compatibility, different executable names may be specified via the above options. Also, ARG=no can be used to disable the use of MPI compiler scripts, thus causing the serial C and Fortran compilers to be used to compile the parallelized sundials functions and examples.
--with-mpi-root=MPIDIR
This option may be used to specify which MPI implementation should be used. The sundiALS configuration script will automatically check under the subdirectories MPIDIR/include and MPIDIR/lib for the necessary header files and libraries. The subdirectory MPIDIR/bin will also be searched for the C and Fortran MPI compiler scripts, unless the user uses --with-mpicc=no or --with-mpif77=no.
--with-mpi-incdir=INCDIR
--with-mpi-libdir=LIBDIR
--with-mpi-libs=LIBS
These options may be used if the user would prefer not to use a preexisting MPI compiler script, but instead would rather use a serial complier and provide the flags necessary to compile the MPI-aware subroutines in SUNDIALS.

Often an MPI implementation will have unique library names and so it may be necessary to specify the appropriate libraries to use (e.g., LIBS=-lmpich).
Default: INCDIR=MPIDIR/include and LIBDIR=MPIDIR/lib
--with-mpi-flags=ARG
Specify additional MPI-specific flags.

## Options for library support

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

## --enable-shared

Using this particular option will result in both static and shared versions of the available SUNDIALS libraries being built if the system supports shared libraries. To build only shared libraries also specify --disable-static.
Note: The FCVODE, FKINsOL, and FIDA libraries can only be built as static libraries because they contain references to externally defined symbols, namely user-supplied Fortran subroutines. Although the Fortran interfaces to the serial and parallel implementations of the supplied NVECTOR module do not contain any unresolvable external symbols, the libraries are still built as static libraries for the purpose of consistency.

## Options for Matlab support

The following options are relevant only for configuring and building the sundialsTB Matlab toolbox:
--enable-sundialsTB
The sundialsTB Matlab toolbox is not built by default. Use this option to enable configuration and compilation of the mex files. Upon completion of the make command, the following mex files will be created:
builddir/sundialsTB/cvodes/cvm/cvm. mexext
builddir/sundialsTB/idas/idm/idm.mexext
builddir/sundialsTB/kinsol/kim/kim. mexext
where mexext is the platform-specific extension of mex files.
--with-sundialsTB-instdir=STBINSTDIR
Alternate location for the installed sundialsTB toolbox (valid only if sundialsTB is enabled). As for the example programs, installation of SUNDIALSTB can be completely disabled by issuing STBINSTDIR=no (in case building the toolbox is desired but its installtion will be done manually afterwards). Otherwise, all required sundialsTB files will be installed under the directory STBINSTDIR/sundialsTB.
Default: DIR=MATLAB/toolbox (see below for the definition of MATLAB).
--with-matlab=MATLAB
This option can be used to specify the location of the Matlab executable. The default is to search the path.
--with-mexopts=ARG
Specify the mex options file to be used.
Default: Standard Matlab mex options file.
--with-mexflags=ARG
Specify the mex compiler flags to be used.
Default: ARG=-0
--with-mexldadd=ARG
Specify additional mex linker flags.
Default: none

## Environment variables

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

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

## CFLAGS

## FFLAGS

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

### 2.2 Configuration examples

The following examples are meant to help demonstrate proper usage of the configure options.
To build sundials using the default C and Fortran compilers, and default mpicc and mpif77 parallel compilers, enable compilation of examples, build the Matlab mex files for sundialsTB, and install it under /home/myname/matlab/sundialsTB, use

```
% configure --prefix=/home/myname/sundials --enable-examples \
    --enable-sundialsTB --with-sundialsTB-instdir=/home/myname/matlab
```

To disable installation of the examples, use:

```
% configure --prefix=/home/myname/sundials \
    --enable-examples --with-examples-instdir=no \
    --enable-sundialsTB --with-sundialsTB-instdir=/home/myname/matlab
```

The following example builds sundials using gec as the serial C compiler, g 77 as the serial Fortran compiler, mpicc as the parallel C compiler, mpif77 as the parallel Fortran compiler, and appends the -g3 compilaton flag to the list of default flags:

```
% configure CC=gcc F77=g77 --with-cflags=-g3 --with-fflags=-g3 \
    --with-mpicc=/usr/apps/mpich/1.2.4/bin/mpicc \
    --with-mpif77=/usr/apps/mpich/1.2.4/bin/mpif77
```

The next example again builds sundials using gcc as the serial C compiler, but the --with-mpicc=no option explicitly disables the use of the corresponding MPI compiler script. In addition, since the --with-mpi-root option is given, the compilation flags $-I / u s r / a p p s / m p i c h / 1.2 .4 / i n c l u d e ~ a n d ~$ -L/usr/apps/mpich/1.2.4/lib are passed to gcc when compiling the MPI-enabled functions. The --disable-examples option explicitly disables the examples (which means a Fortran compiler is not required). The --with-mpi-libs option is required so that the configure script can check if gcc can link with the appropriate MPI library.

$$
\begin{aligned}
\% \text { configure } & \text { CC=gcc --disable-examples --with-mpicc=no \} } \\
{ } &{\text {--with-mpi-root=/usr/apps/mpich/1.2.4 \} } \\
{ } &{\text {--with-mpi-libs=-lmpich }}
\end{array}
\end{aligned}
$$

### 2.3 Installed libraries and exported header files

Using the standard 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 §2.1). 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 Table 2.1. The file extension .lib is typically .so for shared libraries and .a for static libraries (see Options for library support for additional details).

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_smalldense. h could be used in building a preconditioner).

### 2.4 Building SUNDIALS without the configure script

If the configure script cannot be used (e.g., when building sundials under Microsoft Windows without using Cygwin), or if the user prefers to own the build process (e.g., when SUNDIALS is incorporated into a larger project with its own build system), then the header and source files for a given module can be copied from the srcdir to some other location and compiled separately.

The following files are required to compile a sundials solver module:

- public header files located under srcdir/include/solver
- implementation header files and source files located under srcdir/src/solver
- (optional) Fortran/C interface files located under srcdir/src/solver/fcmix
- shared public header files located under srcdir/include/sundials
- shared source files located under srcdir/src/sundials
- (optional) NVECTOR_SERIAL header and source files located under srcdir/include/nvector and srcdir/src/nvec_ser
- (optional) NVECTOR_PARALLEL header and source files located under srcdir/include/nvector and srcdir/src/nvec_par
- configuration header file sundials_config.h (see below)

A sample header file that, appropriately modified, can be used as sundials_config.h (otherwise created automatically by the configure script) is provided below. The various preprocessor macros defined within sundials_config.h have the following uses:

- Precision of the SUNDIALS realtype type

Only one of the macros SUNDIALS_SINGLE_PRECISION, SUNDIALS_DOUBLE_PRECISION and SUNDIALS_EXTENDED_PRECISION should be defined to indicate if the SUNDIALS realtype type is an alias for float, double, or long double, respectively.

Table 2.1: SUNDIALS libraries and header files (names are relative to libdir for libraries and to includedir for header files)

| SHARED | Libraries | n/a |  |
| :---: | :---: | :---: | :---: |
|  | Header files | sundials/sundials_types.h sundials/sundials_config.h sundials/sunials_smalldense.h sundials/sundials_iterative.h sundials/sundials_spbcgs.h sundials/sundials_spgmr.h | sundials/sundials_math.h sundias/sundials_nvector.h sundials/sundials_dense.h sundials/sundials_band.h sundials/sundials_sptfqmr.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 |  |
| CVODE | Libraries | libsundials_cvode.lib | libsundials_fcvode.a |
|  | Header files | cvode/cvode.h cvode/cvode_dense.h cvode/cvode_diag.h cvode/cvode_bandpre.h cvode/cvode_spgmr.h cvode/cvode_sptfqmr.h | cvode/cvode_band.h cvode/cvode_spils.h cvode/cvode_bbdpre.h cvode/cvode_spbcgs.h cvode/cvode_impl.h |
| CVODES | Libraries | libsundials_cvodes.lib |  |
|  | Header files | cvodes/cvodes.h cvodes/cvodes_dense.h cvodes/cvodes_diag.h cvodes/cvodes_bandpre.h cvodes/cvodes_spgmr.h cvodes/cvodes_sptfqmr.h cvodes/cvodea_impl.h | cvodes/cvodes_band.h cvodes/cvodes_spils.h cvodes/cvodes_bbdpre.h cvodes/cvodes_spbcgs.h cvodes/cvodes_impl.h |
| IDA | Libraries | libsundials_ida.lib | libsundials_fida.a |
|  | Header files | ida/ida.h <br> ida/ida_dense.h <br> ida/ida_spils.h <br> ida/ida_spbcgs.h <br> ida/ida_bbdpre.h | ida/ida_band.h ida/ida_spgmr.h ida/ida_sptfqmr.h ida/ida_impl.h |
| KINSOL | Libraries | libsundials_kinsol.lib | libsundials_fkinsol.a |
|  | Header files | kinsol/kinsol.h <br> kinsol/kinsol_dense.h <br> kinsol/kinsol_spils.h <br> kinsol/kinsol_spbcgs.h <br> kinsol/kinsol_bbdpre.h | kinsol/kinsol_band.h kinsol/kinsol_spgmr.h kinsol/kinsol_sptfqmr.h kinsol/kinsol_impl.h |

- Use of generic math functions

If SUNDIALS_USE_GENERIC_MATH is defined, then the functions in sundials math. (h, c) will use the pow, sqrt, fabs, and exp functions from the standard math library (see math.h), regardless of the definition of realtype. Otherwise, if realtype is defined to be an alias for the float C-type, then SUNDIALS will use powf, sqrtf, fabsf, and expf. If realtype is instead defined to be a synonym for the long double C-type, then powl, sqrtl, fabsl, and expl will be used.

Note: Although the powf/powl, sqrtf/sqrtl, fabsf/fabsl, and expf/expl routines are not specified in the ANSI C standard, they are ISO C99 requirements. Consequently, these routines will only be used if available.

- Fortran name-mangling scheme

The macros given below are used to transform the C-language function names defined in the Fortran-C inteface modules in a manner consistent with the preferred Fortran compiler, thus allowing native C functions to be called from within a Fortran subroutine. The namemangling scheme can be specified either by appropriately defining the parameterized macros (using the stringization operator, \#\#, if necessary)

- F77_FUNC (name, NAME)
- F77_FUNC_(name,NAME)
or by defining one macro from each of the following lists:
- SUNDIALS_CASE_LOWER or SUNDIALS_CASE_UPPER
- SUNDIALS_UNDERSCORE_NONE, SUNDIALS_UNDERSCORE_ONE, or SUNDIALS_UNDERSCORE_TWO

For example, to specify that mangled C-language function names should be lowercase with one underscore appended include either

```
#define F77_FUNC(name,NAME) name ## _
#define F77_FUNC_(name,NAME) name ## _
```

or

```
#define SUNDIALS_CASE_LOWER 1
#define SUNDIALS_UNDERSCORE_ONE 1
```

in the sundials_config.h header file.

- Use of an MPI communicator other than MPI_COMM_WORLD in Fortran

If the macro SUNDIALS_MPI_COMM_F2C is defined, then the MPI implementation used to build SUNDIALS defines the type MPI_Fint and the function MPI_Comm_f2c, and it is possible to use MPI communicators other than MPI_COMM_WORLD with the FORTRAN-C interface modules.

```
/*
    * -------------------------------------------------------------------------
    * Copyright (c) 2005, The Regents of the University of California.
    * Produced at the Lawrence Livermore National Laboratory.
    * All rights reserved.
    * For details, see sundials/shared/LICENSE.
    *----------------------------------------------------------------------------
    * SUNDIALS configuration header file
    --------------------------------------------------------------------------
    */
/* Define SUNDIALS version number
    * ------------------------------- */
#define SUNDIALS_PACKAGE_VERSION "2.2.1"
/* Define precision of SUNDIALS data type 'realtype'
    * ------------------------------------------------------*/
/* Define SUNDIALS data type 'realtype' as 'double' */
#define SUNDIALS_DOUBLE_PRECISION 1
/* Define SUNDIALS data type 'realtype' as 'float' */
/* #define SUNDIALS_SINGLE_PRECISION 1 */
/* Define SUNDIALS data type 'realtype' as 'long double' */
/* #define SUNDIALS_EXTENDED_PRECISION 1 */
/* Use generic math functions
    * --------------------------- */
#define SUNDIALS_USE_GENERIC_MATH 1
/* FCMIX: Define Fortran name-mangling macro
    * ---------------------------------------------*/
#define F77_FUNC(name,NAME) name ## -
#define F77_FUNC_(name,NAME) name ## _
/* FCMIX: Define case of function names
    * ---------------------------------------*/
/* FCMIX: Make function names lowercase */
/* #define SUNDIALS_CASE_LOWER 1 */
/* FCMIX: Make function names uppercase */
/* #define SUNDIALS_CASE_UPPER 1 */
/* FCMIX: Define number of underscores to append to function names
    * ------------------------------------------------------------------------*/
/* FCMIX: Do NOT append any underscores to functions names */
/* #define SUNDIALS_UNDERSCORE_NONE 1 */
/* FCMIX: Append ONE underscore to function names */
/* #define SUNDIALS_UNDERSCORE_ONE 1 */
/* FCMIX: Append TWO underscores to function names */
/* #define SUNDIALS_UNDERSCORE_TWO 1 */
/* FNVECTOR: Allow user to specify different MPI communicator
    * ------------------------------------------------------------------*/
#define SUNDIALS_MPI_COMM_F2C 1
```


## Chapter 3

## Mathematical Considerations

CVODES solves ODE initial value problems (IVPs) in real $N$-space, which we write in the abstract form

$$
\begin{equation*}
\dot{y}=f(t, y), \quad y\left(t_{0}\right)=y_{0} \tag{3.1}
\end{equation*}
$$

where $y \in \mathbf{R}^{N}$. Here we use $\dot{y}$ to denote $d y / d t$. While we use $t$ to denote the independent variable, and usually this is time, it certainly need not be. CVODES solves both stiff and non-stiff systems. Roughly speaking, stiffness is characterized by the presence of at least one rapidly damped mode, whose time constant is small compared to the time scale of the solution itself.

Additionally, if (3.1) depends on some parameters $p \in \mathbf{R}^{N_{p}}$, i.e.

$$
\begin{align*}
& \dot{y}=f(t, y, p) \\
& y\left(t_{0}\right)=y_{0}(p) \tag{3.2}
\end{align*}
$$

CVODES can also compute first order derivative information, performing either forward sensitivity analysis or adjoint sensitivity analysis. In the first case, CVODES computes the sensitivities of the solution with respect to the parameters $p$, while in the second case, CVODES computes the gradient of a derived function with respect to the parameters $p$.

### 3.1 IVP solution

The methods used in CVODES are variable-order, variable-step multistep methods, based on formulas of the form

$$
\begin{equation*}
\sum_{i=0}^{K_{1}} \alpha_{n, i} y^{n-i}+h_{n} \sum_{i=0}^{K_{2}} \beta_{n, i} \dot{y}^{n-i}=0 \tag{3.3}
\end{equation*}
$$

Here the $y^{n}$ are computed approximations to $y\left(t_{n}\right)$, and $h_{n}=t_{n}-t_{n-1}$ is the step size. The user of CVODES must appropriately choose one of two multistep methods. For non-stiff problems, CVODES includes the Adams-Moulton formulas, characterized by $K_{1}=1$ and $K_{2}=q$ above, where the order $q$ varies between 1 and 12. For stiff problems, cvodes includes the Backward Differentiation Formulas (BDF) in so-called fixed-leading coefficient (FLC) form, given by $K_{1}=q$ and $K_{2}=0$, with order $q$ varying between 1 and 5 . The coefficients are uniquely determined by the method type, its order, the recent history of the step sizes, and the normalization $\alpha_{n, 0}=-1$. See [4] and [21].

For either choice of formula, the nonlinear system

$$
\begin{equation*}
G\left(y^{n}\right) \equiv y^{n}-h_{n} \beta_{n, 0} f\left(t_{n}, y^{n}\right)-a_{n}=0 \tag{3.4}
\end{equation*}
$$

where $a_{n} \equiv \sum_{i>0}\left(\alpha_{n, i} y^{n-i}+h_{n} \beta_{n, i} \dot{y}^{n-i}\right)$, must be solved (approximately) at each integration step. For this, CVODES offers the choice of either functional iteration, suitable only for non-stiff systems, and various versions of Newton iteration. Functional iteration, given by

$$
y^{n(m+1)}=h_{n} \beta_{n, 0} f\left(t_{n}, y^{n(m)}\right)+a_{n}
$$

involves evaluations of $f$ only. In contrast, Newton iteration requires the solution of linear systems

$$
\begin{equation*}
M\left[y^{n(m+1)}-y^{n(m)}\right]=-G\left(y^{n(m)}\right) \tag{3.5}
\end{equation*}
$$

in which

$$
\begin{equation*}
M \approx I-\gamma J, \quad J=\partial f / \partial y, \quad \text { and } \quad \gamma=h_{n} \beta_{n, 0} \tag{3.6}
\end{equation*}
$$

The initial guess for the iteration is a predicted value $y^{n(0)}$ computed explicitly from the available history data. For the Newton corrections, CVODES provides a choice of six methods:

- dense direct solver,
- band direct solver,
- diagonal approximate Jacobian solver,
- scaled preconditioned GMRES (Generalized Minimal Residual method) without restarts,
- scaled preconditioned Bi-CGStab (Bi-Conjugate Gradient Stable method), or
- scaled preconditioned TFQMR (Transpose-Free Quasi-Minimal Residual method).

For large stiff systems, where direct methods are not feasible, the combination of a BDF integrator and any of the preconditioned Krylov methods (SPGMR, SPBCG, or SPTFQMR) yields a powerful tool because it combines established methods for stiff integration, nonlinear iteration, and Krylov (linear) iteration with a problem-specific treatment of the dominant source of stiffness, in the form of the user-supplied preconditioner matrix [2]. Note that the direct linear solvers (dense and band) can only be used with serial vector representations.

In the process of controlling errors at various levels, CVODES uses a weighted root-mean-square norm, denoted $\|\cdot\|_{\text {WRMS }}$, for all error-like quantities. The multiplicative weights used are based on the current solution and on the relative and absolute tolerances input by the user, namely

$$
\begin{equation*}
W_{i}=1 /\left[\mathrm{RTOL} \cdot\left|y_{i}\right|+\mathrm{ATOL}_{i}\right] \tag{3.7}
\end{equation*}
$$

Because $1 / W_{i}$ represents a tolerance in the component $y_{i}$, a vector whose norm is 1 is regarded as "small." For brevity, we will usually drop the subscript WRMS on norms in what follows.

In the cases of a direct solver (dense, band, diagonal), the iteration is a modified Newton iteration since the iteration matrix $M$ is fixed throughout the nonlinear iterations. However, for any of the Krylov methods, it is an Inexact Newton iteration, in which $M$ is applied in a matrix-free manner, with matrix-vector products $J v$ obtained by either difference quotients or a user-supplied routine. The matrix $M$ (for the direct solvers) or preconditioner matrix $P$ (Krylov cases) is updated as infrequently as possible to balance the high costs of matrix operations against other costs. Specifically, this matrix update occurs when:

- starting the problem,
- more than 20 steps have been taken since the last update,
- the value $\bar{\gamma}$ of $\gamma$ at the last update satisfies $|\gamma / \bar{\gamma}-1|>0.3$,
- a non-fatal convergence failure just occurred, or
- an error test failure just occurred.

When forced by a convergence failure, an update of $M$ or $P$ may involve a reevaluation of $J$ (in $M$ ) or of Jacobian data (in $P$ ) if Jacobian error was the likely cause of the failure. More generally, the decision is made to reevaluate $J$ (or instruct the user to reevaluate Jacobian data in $P$ ) when:

- starting the problem,
- more than 50 steps have been taken since the last evaluation,
- a convergence failure occurred with an outdated matrix, and the value $\bar{\gamma}$ ( $\gamma$ at the last update) satisfies $|\gamma / \bar{\gamma}-1|<0.2$, or
- a convergence failure occurred that forced a reduction of the step size.

The stopping test for the Newton iteration is related to the subsequent local error test, with the goal of keeping the nonlinear iteration errors from interfering with local error control. As described below, the final computed value $y^{n(m)}$ will have to satisfy a local error test $\left\|y^{n(m)}-y^{n(0)}\right\| \leq \epsilon$. Letting $y^{n}$ denote the exact solution of (3.4), we want to ensure that the iteration error $y^{n}-y^{n(m)}$ is small relative to $\epsilon$, specifically that it is less than $0.1 \epsilon$. (The safety factor 0.1 can be changed by the user.) For this, we also estimate the linear convergence rate constant $R$ as follows. We initialize $R$ to 1 , and reset $R=1$ when $M$ or $P$ is updated. After computing a correction $\delta_{m}=y^{n(m)}-y^{n(m-1)}$, we update $R$ if $m>1$ as

$$
R \leftarrow \max \left\{0.3 R,\left\|\delta_{m}\right\| /\left\|\delta_{m-1}\right\|\right\}
$$

Now we use the estimate

$$
\left\|y^{n}-y^{n(m)}\right\| \approx\left\|y^{n(m+1)}-y^{n(m)}\right\| \approx R\left\|y^{n(m)}-y^{n(m-1)}\right\|=R\left\|\delta_{m}\right\|
$$

Therefore the convergence (stopping) test is

$$
R\left\|\delta_{m}\right\|<0.1 \epsilon
$$

We allow at most 3 iterations, but this limit can be changed by the user. We also declare the iteration diverged if any $\left\|\delta_{m}\right\| /\left\|\delta_{m-1}\right\|>2$ with $m>1$. If convergence fails with $J$ or $P$ current, we are forced to reduce the step size, and we replace $h_{n}$ by $h_{n} / 4$. The integration is halted after a preset number of convergence failures; the default value of this limit is 10 , but this can be changed by the user.

When a Krylov method is used to solve the linear system, its errors must also be controlled, and this also involves the local error test constant. The linear iteration error in the solution vector $\delta_{m}$ is approximated by the preconditioned residual vector. Thus to ensure (or attempt to ensure) that the linear iteration errors do not interfere with the nonlinear error and local integration error controls, we require that the norm of the preconditioned residual be less than $0.05 \cdot(0.1 \epsilon)$.

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

$$
J_{i j}=\left[f_{i}\left(t, y+\sigma_{j} e_{j}\right)-f_{i}(t, y)\right] / \sigma_{j}
$$

The increments $\sigma_{j}$ are given by

$$
\sigma_{j}=\max \left\{\sqrt{U}\left|y_{j}\right|, \sigma_{0} / W_{j}\right\}
$$

where $U$ is the unit roundoff, $\sigma_{0}$ is a dimensionless value, and $W_{j}$ is the error weight defined in (3.7). 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.

In the case of a Krylov method, preconditioning may be used on the left, on the right, or both, with user-supplied routines for the preconditioning setup and solve operations, and optionally also for the required matrix-vector products $J v$. If a routine for $J v$ is not supplied, these products are computed as

$$
\begin{equation*}
J v=[f(t, y+\sigma v)-f(t, y)] / \sigma \tag{3.8}
\end{equation*}
$$

The increment $\sigma$ is $1 /\|v\|$, so that $\sigma v$ has norm 1 .
A critical part of CVODES, that makes it an ODE "solver" rather than just an ODE method, is its control of local error. At every step, the local error is estimated and required to satisfy tolerance conditions, and the step is redone with reduced step size whenever that error test fails. As with any linear multistep method, the local truncation error LTE, at order $q$ and step size $h$, satisfies an asymptotic relation

$$
\mathrm{LTE}=C h^{q+1} y^{(q+1)}+O\left(h^{q+2}\right)
$$

for some constant $C$, under mild assumptions on the step sizes. A similar relation holds for the error in the predictor $y^{n(0)}$. These are combined to get a relation

$$
\mathrm{LTE}=C^{\prime}\left[y^{n}-y^{n(0)}\right]+O\left(h^{q+2}\right)
$$

The local error test is simply $\|$ LTE $\| \leq 1$. Using the above, it is performed on the predictor-corrector difference $\Delta_{n} \equiv y^{n(m)}-y^{n(0)}$ (with $y^{n(m)}$ the final iterate computed), and takes the form

$$
\left\|\Delta_{n}\right\| \leq \epsilon \equiv 1 /\left|C^{\prime}\right|
$$

If this test passes, the step is considered successful. If it fails, the step is rejected and a new step size $h^{\prime}$ is computed based on the asymptotic behavior of the local error, namely by the equation

$$
\left(h^{\prime} / h\right)^{q+1}\left\|\Delta_{n}\right\|=\epsilon / 6
$$

Here $1 / 6$ is a safety factor. A new attempt at the step is made, and the error test repeated. If it fails three times, the order $q$ is reset to 1 (if $q>1$ ), or the step is restarted from scratch (if $q=1$ ). The ratio $h^{\prime} / h$ is limited above to 0.2 after two error test failures, and limited below to 0.1 after three. After seven failures, CVODES returns to the user with a give-up message.

In addition to adjusting the step size to meet the local error test, CVODES periodically adjusts the order, with the goal of maximizing the step size. The integration starts out at order 1 , but the order is varied dynamically after that. The basic idea is to pick the order $q$ for which a polynomial of order $q$ best fits the discrete data involved in the multistep method. However, if either a convergence failure or an error test failure occurred on the step just completed, no change is made to the step size or order. At the current order $q$, selecting a new step size is done exactly as when the error test fails, giving a tentative step size ratio

$$
h^{\prime} / h=\left(\epsilon / 6\left\|\Delta_{n}\right\|\right)^{1 /(q+1)} \equiv \eta_{q}
$$

We consider changing order only after taking $q+1$ steps at order $q$, and then we consider only orders $q^{\prime}=q-1($ if $q>1)$ or $q^{\prime}=q+1($ if $q<5)$. The local truncation error at order $q^{\prime}$ is estimated using the history data. Then a tentative step size ratio is computed on the basis that this error, $\operatorname{LTE}\left(q^{\prime}\right)$, behaves asymptotically as $h^{q^{\prime}+1}$. With safety factors of $1 / 6$ and $1 / 10$ respectively, these ratios are:

$$
h^{\prime} / h=[1 / 6\|\operatorname{LTE}(q-1)\|]^{1 / q} \equiv \eta_{q-1}
$$

and

$$
h^{\prime} / h=[1 / 10\|\operatorname{LTE}(q+1)\|]^{1 /(q+2)} \equiv \eta_{q+1}
$$

The new order and step size are then set according to

$$
\eta=\max \left\{\eta_{q-1}, \eta_{q}, \eta_{q+1}\right\}, \quad h^{\prime}=\eta h
$$

with $q^{\prime}$ set to the index achieving the above maximum. However, if we find that $\eta<1.5$, we do not bother with the change. Also, $h^{\prime} / h$ is always limited to 10 , except on the first step, when it is limited to $10^{4}$.

The various algorithmic features of CVODES described above, as inherited from VODE and VODPK, are documented in $[1,3,15]$. They are also summarized in [16].

Normally, CVODES takes steps until a user-defined output value $t=t_{\text {out }}$ is overtaken, and then it computes $y\left(t_{\text {out }}\right)$ by interpolation. However, a "one step" mode option is available, where control returns to the calling program after each step. There are also options to force CVODES not to integrate past a given stopping point $t=t_{\text {stop }}$.

### 3.2 Forward sensitivity analysis

Typically, the governing equations of complex, large-scale models depend on various parameters, through the right-hand side vector and/or through the vector of initial conditions, as in (3.2). In
addition to numerically solving the ODEs, it may be desirable to determine the sensitivity of the results with respect to the model parameters. Such sensitivity information can be used to estimate which parameters are most influential in affecting the behavior of the simulation or to evaluate optimization gradients (in the setting of dynamic optimization, parameter estimation, optimal control, etc.).

The solution sensitivity with respect to the model parameter $p_{i}$ is defined as the vector $s_{i}(t)=$ $\partial y(t) / \partial p_{i}$ and satisfies the following forward sensitivity equations (or in short sensitivity equations):

$$
\begin{equation*}
\dot{s}_{i}=\frac{\partial f}{\partial y} s_{i}+\frac{\partial f}{\partial p_{i}}, \quad s_{i}\left(t_{0}\right)=\frac{\partial y_{0}(p)}{\partial p_{i}} \tag{3.9}
\end{equation*}
$$

obtained by applying the chain rule of differentiation to the original ODEs (3.2).
When performing forward sensitivity analysis, CVODES carries out the time integration of the combined system, (3.2) and (3.9), by viewing it as an ODE system of size $N\left(N_{s}+1\right)$, where $N_{s}$ is the number of model parameters $p_{i}$, with respect to the desired sensitivities $\left(N_{s} \leq N_{p}\right)$. However, major improvements in efficiency can be made by taking advantage of the special form of the sensitivity equations as linearizations of the original ODEs. In particular, for stiff systems, for which CVODES employs a Newton iteration, the original ODE system and all sensitivity systems share the same Jacobian matrix, and therefore the same iteration matrix $M$ in (3.6).

The sensitivity equations are solved with the same linear multistep formula that was selected for the original ODEs and, if Newton iteration was selected, the same linear solver is used in the correction phase for both state and sensitivity variables. In addition, CVODES offers the option of including (full error control) or excluding (partial error control) the sensitivity variables from the local error test.

### 3.2.1 Forward sensitivity methods

In what follows we briefly describe three methods that have been proposed for the solution of the combined ODE and sensitivity system for the vector $\hat{y}=\left[y, s_{1}, \ldots, s_{N_{s}}\right]$.

## - Staggered Direct

In this approach [7], the nonlinear system (3.4) is first solved and, once an acceptable numerical solution is obtained, the sensitivity variables at the new step are found by directly solving (3.9) after the (BDF or Adams) discretization is used to eliminate $\dot{s}_{i}$. Although the system matrix of the above linear system is based on exactly the same information as the matrix $M$ in (3.6), it must be updated and factored at every step of the integration, in contrast to $M$ which is updated only occasionally. For problems with many parameters (relative to the problem size), the staggered direct method can outperform the methods described below [22]. However, the computational cost associated with matrix updates and factorizations makes this method unattractive for problems with many more states than parameters (such as those arising from semidiscretization of PDEs) and is therefore not implemented in CVODES.

## - Simultaneous Corrector

In this method [23], the discretization is applied simultaneously to both the original equations (3.2) and the sensitivity systems (3.9) resulting in the following nonlinear system

$$
\hat{G}\left(\hat{y}_{n}\right) \equiv \hat{y}_{n}-h_{n} \beta_{n, 0} \hat{f}\left(t_{n}, \hat{y}_{n}\right)-\hat{a}_{n}=0
$$

where $\hat{f}=\left[f(t, y, p), \ldots,(\partial f / \partial y)(t, y, p) s_{i}+\left(\partial f / \partial p_{i}\right)(t, y, p), \ldots\right]$, and $\hat{a}_{n}$ is comprised of the terms in the discretization that depend on the solution at previous integration steps. This combined nonlinear system can be solved using a modified Newton method as in (3.5) by solving the corrector equation

$$
\begin{equation*}
\hat{M}\left[\hat{y}_{n(m+1)}-\hat{y}_{n(m)}\right]=-\hat{G}\left(\hat{y}_{n(m)}\right) \tag{3.10}
\end{equation*}
$$

at each iteration, where

$$
\hat{M}=\left[\begin{array}{ccccc}
M & & & & \\
-\gamma J_{1} & M & & & \\
-\gamma J_{2} & 0 & M & & \\
\vdots & \vdots & \ddots & \ddots & \\
-\gamma J_{N_{s}} & 0 & \ldots & 0 & M
\end{array}\right]
$$

$M$ is defined as in (3.6), and $J_{i}=(\partial / \partial y)\left[(\partial f / \partial y) s_{i}+\left(\partial f / \partial p_{i}\right)\right]$. It can be shown that 2-step quadratic convergence can be attained by using only the block-diagonal portion of $\hat{M}$ in the corrector equation (3.10). This results in a decoupling that allows the reuse of $M$ without additional matrix factorizations. However, the products $(\partial f / \partial y) s_{i}$ and the vectors $\partial f / \partial p_{i}$ must still be reevaluated at each step of the iterative process (3.10) to update the sensitivity portions of the residual $\hat{G}$.

## - Staggered corrector

In this approach [10], as in the staggered direct method, the nonlinear system (3.4) is solved first using the Newton iteration (3.5). Then a separate Newton iteration is used to solve the sensitivity system (3.9):

$$
\begin{align*}
M\left[s_{i}^{n(m+1)}-s_{i}^{n(m)}\right]= & \\
& -\left[s_{i}^{n(m)}-\gamma\left(\frac{\partial f}{\partial y}\left(t_{n}, y^{n}, p\right) s_{i}^{n(m)}+\frac{\partial f}{\partial p_{i}}\left(t_{n}, y^{n}, p\right)\right)-a_{i, n}\right], \tag{3.11}
\end{align*}
$$

where $a_{i, n}=\sum_{j>0}\left(\alpha_{n, j} s_{i}^{n-j}+h_{n} \beta_{n, j} \dot{s}_{i}^{n-j}\right)$. In other words, a modified Newton iteration is used to solve a linear system. In this approach, the vectors $\partial f / \partial p_{i}$ need be updated only once per integration step, after the state correction phase (3.5) has converged. Note also that Jacobian-related data can be reused at all iterations (3.11) to evaluate the products $(\partial f / \partial y) s_{i}$.

CVODES implements the simultaneous corrector method and two flavors of the staggered corrector method which differ only if the sensitivity variables are included in the error control test. In the full error control case, the first variant of the staggered corrector method requires the convergence of the iterations (3.11) for all $N_{s}$ sensitivity systems and then performs the error test on the sensitivity variables. The second variant of the method will perform the error test for each sensitivity vector $s_{i},\left(i=1,2, \ldots, N_{s}\right)$ individually, as they pass the convergence test. Differences in performance between the two variants may therefore be noticed whenever one of the sensitivity vectors $s_{i}$ fails a convergence or error test.

An important observation is that the staggered corrector method, combined with a Krylov linear solver, effectively results in a staggered direct method. Indeed, the Krylov solver requires only the action of the matrix $M$ on a vector and this can be provided with the current Jacobian information. Therefore, the modified Newton procedure (3.11) will theoretically converge after one iteration.

### 3.2.2 Selection of the absolute tolerances for sensitivity variables

If the sensitivities are included in the error test, CVODES provides an automated estimation of absolute tolerances for the sensitivity variables based on the absolute tolerance for the corresponding state variable. The relative tolerance for sensitivity variables is set to be the same as for the state variables. The selection of absolute tolerances for the sensitivity variables is based on the observation that the sensitivity vector $s_{i}$ will have units of $[y] /\left[p_{i}\right]$. With this, the absolute tolerance for the $j$-th component of the sensitivity vector $s_{i}$ is set to $\mathrm{ATOL}_{j} /\left|\bar{p}_{i}\right|$, where $\mathrm{ATOL}_{j}$ are the absolute tolerances for the state variables and $\bar{p}$ is a vector of scaling factors that are dimensionally consistent with the model parameters $p$ and give an indication of their order of magnitude. This choice of relative and absolute tolerances is equivalent to requiring that the weighted root-mean-square norm of the sensitivity vector $s_{i}$ with weights based on $s_{i}$ be the same as the weighted root-mean-square norm of the vector of scaled sensitivities $\bar{s}_{i}=\left|\bar{p}_{i}\right| s_{i}$ with weights based on the state variables (the scaled sensitivities $\bar{s}_{i}$ being
dimensionally consistent with the state variables). However, this choice of tolerances for the $s_{i}$ may be a poor one, and the user of CVODES can provide different values as an option.

### 3.2.3 Evaluation of the sensitivity right-hand side

There are several methods for evaluating the right-hand side of the sensitivity systems (3.9): analytic evaluation, automatic differentiation, complex-step approximation, and finite differences (or directional derivatives). CVODES provides all the software hooks for implementing interfaces to automatic differentiation ( AD ) or complex-step approximation; future versions will include a generic interface to AD-generated functions. At the present time, besides the option for analytical sensitivity righthand sides (user-provided), CVODES can evaluate these quantities using various finite difference-based approximations to evaluate the terms $(\partial f / \partial y) s_{i}$ and $\left(\partial f / \partial p_{i}\right)$, or using directional derivatives to evaluate $\left[(\partial f / \partial y) s_{i}+\left(\partial f / \partial p_{i}\right)\right]$. As is typical for finite differences, the proper choice of perturbations is a delicate matter. CVODES takes into account several problem-related features: the relative ODE error tolerance RTOL, the machine unit roundoff $U$, the scale factor $\bar{p}_{i}$, and the weighted root-mean-square norm of the sensitivity vector $s_{i}$.

Using central finite differences as an example, the two terms $(\partial f / \partial y) s_{i}$ and $\partial f / \partial p_{i}$ in the righthand side of (3.9) can be evaluated separately:

$$
\begin{gather*}
\frac{\partial f}{\partial y} s_{i} \approx \frac{f\left(t, y+\sigma_{y} s_{i}, p\right)-f\left(t, y-\sigma_{y} s_{i}, p\right)}{2 \sigma_{y}}  \tag{3.12}\\
\frac{\partial f}{\partial p_{i}} \approx \frac{f\left(t, y, p+\sigma_{i} e_{i}\right)-f\left(t, y, p-\sigma_{i} e_{i}\right)}{2 \sigma_{i}} \\
\sigma_{i}=\left|\bar{p}_{i}\right| \sqrt{\max (\operatorname{RTOL}, U)}, \quad \sigma_{y}=\frac{1}{\max \left(1 / \sigma_{i},\left\|s_{i}\right\|_{\mathrm{WRMS}} /\left|\bar{p}_{i}\right|\right)}
\end{gather*}
$$

simultaneously:

$$
\begin{gather*}
\frac{\partial f}{\partial y} s_{i}+\frac{\partial f}{\partial p_{i}} \approx \frac{f\left(t, y+\sigma s_{i}, p+\sigma e_{i}\right)-f\left(t, y-\sigma s_{i}, p-\sigma e_{i}\right)}{2 \sigma},  \tag{3.13}\\
\sigma=\min \left(\sigma_{i}, \sigma_{y}\right)
\end{gather*}
$$

or by adaptively switching between $(3.12)+\left(3.12^{\prime}\right)$ and (3.13), depending on the relative size of the estimated finite difference increments $\sigma_{i}$ and $\sigma_{y}$.

These procedures for choosing the perturbations $\left(\delta_{i}, \delta_{y}, \delta\right)$ and switching ( $\rho_{\max }$ ) between finite difference and directional derivative formulas have also been implemented for first-order formulas. Forward finite differences can be applied to $(\partial f / \partial y) s_{i}$ and $\partial f / \partial p_{i}$ separately, or the single directional derivative formula

$$
\frac{\partial f}{\partial y} s_{i}+\frac{\partial f}{\partial p_{i}} \approx \frac{f\left(t, y+\delta s_{i}, p+\delta e_{i}\right)-f(t, y, p)}{\delta}
$$

can be used. In CVODES, the default value of $\rho_{\max }=0$ indicates the use of the second-order centered directional derivative formula (3.13) exclusively. Otherwise, the magnitude of $\rho_{\max }$ and its sign (positive or negative) indicates whether this switching is done with regard to (centered or forward) finite differences, respectively.

### 3.3 Adjoint sensitivity analysis

In the forward sensitivity approach described in the previous section, obtaining sensitivities with respect to $N_{s}$ parameters is roughly equivalent to solving an ODE system of size $\left(1+N_{s}\right) N$. This can become prohibitively expensive, especially for large-scale problems, if sensitivities with respect to many parameters are desired. In this situation, the adjoint sensitivity method is a very attractive alternative, provided that we do not need the solution sensitivities $s_{i}$, but rather the gradients with
respect to model parameters of a relatively few derived functionals of the solution. In other words, if $y(t)$ is the solution of (3.2), we wish to evaluate the gradient $d G / d p$ of

$$
\begin{equation*}
G(p)=\int_{t_{0}}^{t_{\mathrm{f}}} g(t, y, p) d t \tag{3.14}
\end{equation*}
$$

or, alternatively, the gradient $d g / d p$ of the function $g(t, x, p)$ at time $t_{\mathrm{f}}$. The function $g$ must be smooth enough that $\partial g / \partial y$ and $\partial g / \partial p$ exist and are bounded.

In what follows, we only sketch the analysis for the sensitivity problem for both $G$ and $g$. For details on the derivation see [6]. Introducing a Lagrange multiplier $\lambda$, we form the augmented objective function

$$
\begin{equation*}
I(p)=G(p)-\int_{t_{0}}^{t_{1}} \lambda^{*}(\dot{y}-f(t, y, p)) d t \tag{3.15}
\end{equation*}
$$

where $*$ denotes the conjugate transpose. The gradient of $G$ with respect to $p$ is

$$
\begin{equation*}
\frac{d G}{d p}=\frac{d I}{d p}=\int_{t_{0}}^{t_{1}}\left(g_{p}+g_{y} s\right) d t-\int_{t_{0}}^{t_{1}} \lambda^{*}\left(\dot{s}-f_{y} s-f_{p}\right) d t \tag{3.16}
\end{equation*}
$$

where subscripts on functions such as $f$ or $g$ are used to denote partial derivatives and $s=\left[s_{1}, \ldots, s_{N_{s}}\right]$ is the matrix of solution sensitivities. Applying integration by parts to the term $\lambda^{*} \dot{s}$ and selecting $\lambda$ such that

$$
\begin{align*}
& \dot{\lambda}=-\left(\frac{\partial f}{\partial y}\right)^{*} \lambda-\left(\frac{\partial g}{\partial y}\right)^{*}  \tag{3.17}\\
& \lambda\left(t_{1}\right)=0
\end{align*}
$$

the gradient of $G$ with respect to $p$ is nothing but

$$
\begin{equation*}
\frac{d G}{d p}=\lambda^{*}\left(t_{0}\right) s\left(t_{0}\right)+\int_{t_{0}}^{t_{1}}\left(g_{p}+\lambda^{*} f_{p}\right) d t \tag{3.18}
\end{equation*}
$$

The gradient of $g\left(t_{1}, y, p\right)$ with respect to $p$ can be then obtained by using the Leibnitz differentiation rule. Indeed, from (3.14),

$$
\frac{d g}{d p}\left(t_{1}\right)=\frac{d}{d t_{1}} \frac{d G}{d p}
$$

and therefore, taking into account that $d G / d p$ in (3.18) depends on $t_{1}$ both through the upper integration limit and through $\lambda$ and that $\lambda\left(t_{1}\right)=0$,

$$
\begin{equation*}
\frac{d g}{d p}\left(t_{1}\right)=\mu^{*}\left(t_{0}\right) s\left(t_{0}\right)+g_{p}\left(t_{1}\right)+\int_{t_{0}}^{t_{1}} \mu^{*} f_{p} d t \tag{3.19}
\end{equation*}
$$

where $\mu$ is the sensitivity of $\lambda$ with respect to the final integration limit and thus satisfies the following equation, obtained by taking the total derivative with respect to $t_{1}$ of (3.17):

$$
\begin{align*}
& \dot{\mu}=-\left(\frac{\partial f}{\partial y}\right)^{*} \mu \\
& \mu\left(t_{1}\right)=\left(\frac{\partial g}{\partial y}\right)_{t=t_{1}}^{*} \tag{3.20}
\end{align*}
$$

The final condition on $\mu\left(t_{1}\right)$ follows from $(\partial \lambda / \partial t)+\left(\partial \lambda / \partial t_{1}\right)=0$ at $t_{1}$, and therefore, $\mu\left(t_{1}\right)=-\dot{\lambda}\left(t_{1}\right)$.
The first thing to notice about the adjoint system (3.17) is that there is no explicit specification of the parameters $p$; this implies that, once the solution $\lambda$ is found, the formula (3.18) can then be used to find the gradient of $G$ with respect to any of the parameters $p$. The same holds true for the system (3.20) and the formula (3.19) for gradients of $g\left(t_{1}, y, p\right)$. The second important remark is that the adjoint systems (3.17) and (3.20) are terminal value problems which depend on the solution $y(t)$ of the original IVP (3.2). Therefore, a procedure is needed for providing the states $y$ obtained during a forward integration phase of (3.2) to CVODES during the backward integration phase of (3.17) or (3.20). The approach adopted in CVODES, based on checkpointing, is described below.


Figure 3.1: Illustration of the checkpointing algorithm for generation of the forward solution during the integration of the adjoint system.

### 3.3.1 Checkpointing scheme

During the backward integration, the evaluation of the right-hand side of the adjoint system requires, at the current time, the states $y$ which were computed during the forward integration phase. Since CVODES implements variable-step integration formulas, it is unlikely that the states will be available at the desired time and so some form of interpolation is needed. The cVODES implementation being also variable-order, it is possible that during the forward integration phase the order may be reduced as low as first order, which means that there may be points in time where only $y$ and $\dot{y}$ are available. These requirements therefore limit the choices for possible interpolation schemes. CVODES implements two interpolation methods: a cubic Hermite interpolation algorithm and a variable-degree polynomial interpolation method which attempts to mimic the BDF interpolant for the forward integration.

However, especially for large-scale problems and long integration intervals, the number and size of the vectors $y$ and $\dot{y}$ that would need to be stored make this approach computationally intractable. Thus, CVODES settles for a compromise between storage space and execution time by implementing a so-called checkpointing scheme. At the cost of at most one additional forward integration, this approach offers the best possible estimate of memory requirements for adjoint sensitivity analysis. To begin with, based on the problem size $N$ and the available memory, the user decides on the number $N_{d}$ of data pairs $(y, \dot{y})$ if cubic Hermite interpolation is selected, or on the number $N_{d}$ of $y$ vectors in the case of variable-degree polynomial interpolation that can be kept in memory for the purpose of interpolation. Then, during the first forward integration stage, after every $N_{d}$ integration steps a checkpoint is formed by saving enough information (either in memory or on disk) to allow for a hot restart, that is a restart which will exactly reproduce the forward integration. In order to avoid storing Jacobian-related data at each checkpoint, a reevaluation of the iteration matrix is forced before each checkpoint. At the end of this stage, we are left with $N_{c}$ checkpoints, including one at $t_{0}$. During the backward integration stage, the adjoint variables are integrated from $t_{1}$ to $t_{0}$ going from one checkpoint to the previous one. The backward integration from checkpoint $i+1$ to checkpoint $i$ is preceded by a forward integration from $i$ to $i+1$ during which $N_{d}$ the vectors $y$ (and, if necessary $\dot{y}$ ) are generated and stored in memory for interpolation ${ }^{1}$ (see Fig. 3.1).

This approach transfers the uncertainty in the number of integration steps in the forward integration phase to uncertainty in the final number of checkpoints. However, $N_{c}$ is much smaller than the number of steps taken during the forward integration, and there is no major penalty for writing/reading the checkpoint data to/from a temporary file. Note that, at the end of the first forward integration stage, interpolation data are available from the last checkpoint to the end of the interval of integration. If no checkpoints are necessary ( $N_{d}$ is larger than the number of integration steps taken

[^1]in the solution of (3.2)), the total cost of an adjoint sensitivity computation can be as low as one forward plus one backward integration. In addition, CVODES provides the capability of reusing a set of checkpoints for multiple backward integrations, thus allowing for efficient computation of gradients of several functionals (3.14).

Finally, we note that the adjoint sensitivity module in CVODES provides the necessary infrastructure to integrate backwards in time any ODE terminal value problem dependent on the solution of the IVP (3.2), including adjoint systems (3.17) or (3.20), as well as any other quadrature ODEs that may be needed in evaluating the integrals in (3.18) or (3.19). In particular, for ODE systems arising from semi-discretization of time-dependent PDEs, this feature allows for integration of either the discretized adjoint PDE system or the adjoint of the discretized PDE.

### 3.4 Preconditioning

When using a Newton method to solve the nonlinear system (3.4), CVODES makes repeated use of a linear solver to solve linear systems of the form $M x=-r$, where $x$ is a correction vector and $r$ is a residual vector. If this linear system solve is done with one of the scaled preconditioned iterative linear solvers, these solvers are rarely successful if used without preconditioning; it is generally necessary to precondition the system in order to obtain acceptable efficiency. A system $A x=b$ can be preconditioned on the left, as $\left(P^{-1} A\right) x=P^{-1} b$; on the right, as $\left(A P^{-1}\right) P x=b$; or on both sides, as $\left(P_{L}^{-1} A P_{R}^{-1}\right) P_{R} x=P_{L}^{-1} b$. The Krylov method is then applied to a system with the matrix $P^{-1} A$, or $A P^{-1}$, or $P_{L}^{-1} A P_{R}^{-1}$, instead of $A$. In order to improve the convergence of the Krylov iteration, the preconditioner matrix $P$, or the product $P_{L} P_{R}$ in the last case, should in some sense approximate the system matrix $A$. Yet at the same time, in order to be cost-effective, the matrix $P$, or matrices $P_{L}$ and $P_{R}$, should be reasonably efficient to evaluate and solve. Finding a good point in this tradeoff between rapid convergence and low cost can be very difficult. Good choices are often problem-dependent (for example, see [2] for an extensive study of preconditioners for reaction-transport systems).

The CVODES solver allow for preconditioning either side, or on both sides, although we know of no situation where preconditioning on both sides is clearly superior to preconditioning on one side only (with the product $P_{L} P_{R}$ ). Moreover, for a given preconditioner matrix, the merits of left vs. right preconditioning are unclear in general, and the user should experiment with both choices. Performance will differ because the inverse of the left preconditioner is included in the linear system residual whose norm is being tested in the Krylov algorithm. As a rule, however, if the preconditioner is the product of two matrices, we recommend that preconditioning be done either on the left only or the right only, rather than using one factor on each side.

Typical preconditioners used with CVODES are based on approximations to the Newton iteration matrix of the systems involved; in other words, $P \approx I-\gamma J$, where $\gamma$ is a scalar proportional to the integration step size $h$ and $J$ is the system Jacobian, i.e. $J=\partial f / \partial y$. Because the Krylov iteration occurs within a Newton iteration and further also within a time integration, and since each of these iterations has its own test for convergence, the preconditioner may use a very crude approximation, as long as it captures the dominant numerical feature(s) of the system. We have found that the combination of a preconditioner with the Newton-Krylov iteration, using even a fairly poor approximation to the Jacobian, can be surprisingly superior to using the same matrix without Krylov acceleration (i.e., a modified Newton iteration), as well as to using the Newton-Krylov method with no preconditioning.

### 3.5 BDF stability limit detection

CVODES includes an algorithm, STALD (STAbility Limit Detection), which provides protection against potentially unstable behavior of the BDF multistep integration methods in certain situations, as described below.

When the BDF option is selected, cvodes uses Backward Differentiation Formula methods of orders 1 to 5 . At order 1 or 2 , the BDF method is A-stable, meaning that for any complex constant
$\lambda$ in the open left half-plane the method is unconditionally stable (for any step size) for the standard scalar model problem $\dot{y}=\lambda y$. For an ODE system, this means that, roughly speaking, as long as all modes in the system are stable, the method is also stable for any choice of step size, at least in the sense of a local linear stability analysis.

At orders 3 to 5, the BDF methods are not A-stable, although they are stiffly stable. In each case, in order for the method to be stable at step size $h$ on the scalar model problem, the product $h \lambda$ must lie within a region of absolute stability. That region excludes a portion of the left half-plane that is concentrated near the imaginary axis. The size of that region of instability grows as the order increases from 3 to 5 . What this means is that, when running BDF at any of these orders, if an eigenvalue $\lambda$ of the system lies close enough to the imaginary axis, the step sizes $h$ for which the method is stable are limited (at least according to the linear stability theory) to a set that prevents $h \lambda$ from leaving the stability region. The meaning of close enough depends on the order. At order 3, the unstable region is much narrower than at order 5, so the potential for unstable behavior grows with order.

System eigenvalues that are likely to run into this instability are ones that correspond to weakly damped oscillations. A pure undamped oscillation corresponds to an eigenvalue on the imaginary axis. Problems with modes of that kind call for different considerations since the oscillation generally must be followed by the solver, but this requires step sizes ( $h \sim 1 / \nu$, where $\nu$ is the frequency) that are stable for BDF anyway. But for a weakly damped oscillatory mode, the oscillation in the solution is eventually damped to the noise level, and at that time it is important that the solver not be restricted to step sizes on the order of $1 / \nu$. It is in this situation that the new option may be of great value.

In terms of partial differential equations, the typical problems for which the stability limit detection option is appropriate are ODE systems resulting from semi-discretized PDEs (i.e., PDEs discretized in space) with advection and diffusion, but with advection dominating over diffusion. Diffusion alone produces pure decay modes, while advection tends to produce undamped oscillatory modes. A mix of the two with advection dominant will have weakly damped oscillatory modes.

The stald algorithm attempts to detect, in a direct manner, the presence of a stability region boundary that is limiting the step sizes in the presence of a weakly damped oscillation [13]. The algorithm supplements (but differs greatly from) the existing algorithms in CVODES for choosing step size and order based on estimated local truncation errors. The stald algorithm works directly with history data that is readily available in CVODES. If it concludes that the step size is in fact stabilitylimited, it dictates a reduction in the method order regardless of the outcome of the error-based algorithm. The STALD algorithm has been tested in combination with the vode solver on linear advection-dominated advection-diffusion problems [14], where it works well. The implementation in CVODES has been successfully tested on linear and nonlinear advection-diffusion problems, among others.

This stability limit detection option adds some computational overhead to the CVODES solution. (In timing tests, these overhead costs have ranged from $2 \%$ to $7 \%$ of the total, depending on the size and complexity of the problem, with lower relative costs for larger problems.) Therefore, it should be activated only when there is reasonable expectation of modes in the user's system for which it is appropriate. In particular, if a CVODES solution with this option turned off appears to take an inordinately large number of steps for orders between 3 and 5 for no apparent reason in terms of the solution time scale, then there is a good chance that step sizes are being limited by stability, and that turning on the option will improve efficiency.

### 3.6 Rootfinding

The CVODES solver has been augmented to include a rootfinding feature. This means that, while integrating the Initial Value Problem (3.1), CVODES can also find the roots of a set of user-defined functions $g_{i}(t, y)$ that depend both on $t$ and on the solution vector $y=y(t)$. The number of these root functions is arbitrary, and if more than one $g_{i}$ is found to have a root in any given interval, the various root locations are found and reported in the order that they occur on the $t$ axis, in the direction of integration.

Generally, this rootfinding feature finds only roots of odd multiplicity, corresponding to changes
in sign of $g_{i}(t, y(t))$, denoted $g_{i}(t)$ for short. If a user root function has a root of even multiplicity (no sign change), it will probably be missed by cVODES. If such a root is desired, the user should reformulate the root function so that it changes sign at the desired root.

The basic scheme used is to check for sign changes of any $g_{i}(t)$ over each time step taken, and then (when a sign change is found) to hone in on the root(s) with a modified secant method [12]. In addition, each time $g$ is computed, CVODES checks to see if $g_{i}(t)=0$ exactly, and if so it reports this as a root. However, if an exact zero of any $g_{i}$ is found at a point $t$, CVODES computes $g$ at $t+\delta$ for a small increment $\delta$, slightly further in the direction of integration, and if any $g_{i}(t+\delta)=0$ also, CVODES stops and reports an error. This way, each time CVODES takes a time step, it is guaranteed that the values of all $g_{i}$ are nonzero at some past value of $t$, beyond which a search for roots is to be done.

At any given time in the course of the time-stepping, after suitable checking and adjusting has been done, CVODEs has an interval $\left(t_{l o}, t_{h i}\right]$ in which roots of the $g_{i}(t)$ are to be sought, such that $t_{h i}$ is further ahead in the direction of integration, and all $g_{i}\left(t_{l o}\right) \neq 0$. The endpoint $t_{h i}$ is either $t_{n}$, the end of the time step last taken, or the next requested output time $t_{\text {out }}$ if this comes sooner. The endpoint $t_{l o}$ is either $t_{n-1}$, the last output time $t_{\text {out }}$ (if this occurred within the last step), or the last root location (if a root was just located within this step), possibly adjusted slightly toward $t_{n}$ if an exact zero was found. The algorithm checks $g_{i}$ at $t_{h i}$ for zeros and for sign changes in $\left(t_{l o}, t_{h i}\right)$. If no sign changes were found, then either a root is reported (if some $g_{i}\left(t_{h i}\right)=0$ ) or we proceed to the next time interval (starting at $t_{h i}$ ). If one or more sign changes were found, then a loop is entered to locate the root to within a rather tight tolerance, given by

$$
\tau=100 * U *\left(\left|t_{n}\right|+|h|\right) \quad(U=\text { unit roundoff })
$$

Whenever sign changes are seen in two or more root functions, the one deemed most likely to have its root occur first is the one with the largest value of $\left|g_{i}\left(t_{h i}\right)\right| /\left|g_{i}\left(t_{h i}\right)-g_{i}\left(t_{l o}\right)\right|$, corresponding to the closest to $t_{l o}$ of the secant method values. At each pass through the loop, a new value $t_{\text {mid }}$ is set, strictly within the search interval, and the values of $g_{i}\left(t_{m i d}\right)$ are checked. Then either $t_{l o}$ or $t_{h i}$ is reset to $t_{\text {mid }}$ according to which subinterval is found to include the sign change. If there is none in $\left(t_{l o}, t_{\text {mid }}\right)$ but some $g_{i}\left(t_{\text {mid }}\right)=0$, then that root is reported. The loop continues until $\left|t_{h i}-t_{l o}\right|<\tau$, and then the reported root location is $t_{h i}$.

In the loop to locate the root of $g_{i}(t)$, the formula for $t_{\text {mid }}$ is

$$
t_{m i d}=t_{h i}-\left(t_{h i}-t_{l o}\right) g_{i}\left(t_{h i}\right) /\left[g_{i}\left(t_{h i}\right)-\alpha g_{i}\left(t_{l o}\right)\right]
$$

where $\alpha$ is a weight parameter. On the first two passes through the loop, $\alpha$ is set to 1 , making $t_{\text {mid }}$ the secant method value. Thereafter, $\alpha$ is reset according to the side of the subinterval (low vs. high, i.e., toward $t_{l o}$ vs. toward $t_{h i}$ ) in which the sign change was found in the previous two passes. If the two sides were opposite, $\alpha$ is set to 1 . If the two sides were the same, $\alpha$ is halved (if on the low side) or doubled (if on the high side). The value of $t_{\text {mid }}$ is closer to $t_{l o}$ when $\alpha<1$ and closer to $t_{h i}$ when $\alpha>1$. If the above value of $t_{m i d}$ is within $\tau / 2$ of $t_{l o}$ or $t_{h i}$, it is adjusted inward, such that its fractional distance from the endpoint (relative to the interval size) is between .1 and .5 (. 5 being the midpoint), and the actual distance from the endpoint is at least $\tau / 2$.

## Chapter 4

## Code Organization

### 4.1 SUNDIALS organization

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

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

- CVODE, a solver for stiff and nonstiff ODEs $d y / d t=f(t, y)$;
- CVODES, a solver for stiff and nonstiff ODEs $d y / d t=f(t, y, p)$ with sensitivity analysis capabilities;
- KINSOL, a solver for nonlinear algebraic systems $F(u)=0$;
- IDA, a solver for differential-algebraic systems $F\left(t, y, y^{\prime}\right)=0$.


### 4.2 CVODES organization

The cVodes package is written in ANSI C. The following summarizes the basic structure of the package, although knowledge of this structure is not necessary for its use.

The overall organization of the CVODES package is shown in Figure 4.2. The basic elements of the structure are a module for the basic integration algorithm (including forward sensitivity analysis), a module for adjoint sensitivity analysis, and a set of modules for the solution of linear systems that arise in the case of a stiff system. The central integration module, implemented in the files cvodes.h, cvodes_impl.h, cvodes.c, and cvodea.c, deals with the evaluation of integration coefficients, the functional or Newton iteration process, estimation of local error, selection of step size and order, and interpolation to user output points, among other issues. Although this module contains logic for the basic Newton iteration algorithm, it has no knowledge of the method being 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 during the integration.

In addition, if forward sensitivity analysis is turned on, the main module will integrate the forward sensitivity equations simultaneously with the original IVP. The sensitivity variables may be included in the local error control mechanism of the main integrator. CVODES provides three different strategies for dealing with the correction stage for the sensitivity variables: CV_SIMULTANEOUS, CV_STAGGERED

(a) High-level diagram

(b) Directory structure of the source tree

Figure 4.1: Organization of the SUNDIALS suite


Figure 4.2: Overall structure of the CVODES package. Modules specific to CVODES are distinguished by rounded boxes, while generic solver and auxiliary modules are in rectangular boxes.
and CV_STAGGERED1 (see $\S 3.2$ and $\S 6.2 .1$ ). The CVODES package includes an algorithm for the approximation of the sensitivity equations right-hand sides by difference quotients, but the user has the option of supplying these right-hand sides directly.

The adjoint sensitivity module provides the infrastructure needed for the backward integration of any system of ODEs which depends on the solution of the original IVP, in particular the adjoint system and any quadratures required in evaluating the gradient of the objective functional. This module deals with the setup of the checkpoints, the interpolation of the forward solution during the backward integration, and the backward integration of the adjoint equations.

At present, the package includes the following six CVODES linear solver modules:

- CVDENSE: LU factorization and backsolving with dense matrices;
- CVBAND: LU factorization and backsolving with banded matrices;
- CVDIAG: an internally generated diagonal approximation to the Jacobian;
- CVSPGMR: scaled preconditioned GMRES method;
- CVSPBCG: scaled preconditioned Bi-CGStab method;
- CVSPTFQMR: scaled preconditioned TFQMR method.

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

In the case of the direct methods CVDENSE and CVBAND, 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. In the case of the Krylov iterative methods CVSPGMR, CVSPBCG, and CVSPTFQMR, the package includes an algorithm for the approximation of the product between the Jacobian matrix and a vector of appropriate length by difference quotients. Again, the user has the option of providing a routine for this operation. For the Krylov methods, the preconditioner must be supplied by the user in two parts: setup (preprocessing of Jacobian data) and solve. While there is no default choice of preconditioner analogous to the difference-quotient
approximation in the direct case, the references [2,3], together with the example programs included with CVODES, offer considerable assistance in building preconditioners.

Each CVODES linear solver module consists of four routines devoted to (1) memory allocation and initialization, (2) setup of the matrix data involved, (3) solution of the system, and (4) freeing of memory. The setup and solution phases are separate because the evaluation of Jacobians and preconditioners is done only periodically during the integration, and only as required to achieve convergence. The call list within the central CVODES module for each of the five 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 modules CVDEnse, CVBAND, CVSPGMR, CVSPBCG, and CVSPTFQMR is a set of interface routines built on top of a generic solver module, named DENSE, BAND, SPGMR, SPBCG, and SPTFQMR, respectively. The interfaces deal with the use of these methods in the cVODES context, whereas the generic solver is independent of the context. While the generic solvers here were generated with sundials in mind, our intention is that they be usable in other applications as general-purpose solvers. This separation also allows for any generic solver to be replaced by an improved version, with no necessity to revise the CVODES package elsewhere.

CVODES also provides two preconditioner modules for use with any of the Krylov iterative linear solvers. The first one, CVBANDPRE, is intended to be used with NVECTOR_SERIAL and provides banded difference-quotient Jacobian-based preconditioner and solver routines. The second preconditioner module, CVBBDPRE, works in conjunction with NVECTOR_PARALLEL and generates a preconditioner that is a block-diagonal matrix with each block being a band matrix.

All state information used by CVODES 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 CVODES 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 CVODES memory structure. The reentrancy of cVODES was motivated by the anticipated multicomputer extension, but is also essential in a uniprocessor setting where two or more problems are solved by intermixed calls to the package from within a single user program.

## Chapter 5

## Using CVODES for IVP Solution

This chapter is concerned with the use of CVODES for the solution of initial value problems (IVPs). The following sections treat the header files and the layout of the user's main program, and provide descriptions of the CVODES user-callable functions and user-supplied functions.

This usage is essentially equivalent to using cvode [19]. The listings of the sample programs in the companion document [17] may also be helpful. Those codes may be used as templates and are included in the CVODES package.

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 or direct band linear solvers since these linear solver modules need to form the complete system Jacobian. The following cVodes modules can only be used with NVECTOR_SERIAL: CVDENSE, CVBAND and CVBANDPRE. Also, the preconditioner module CVBBDPRE can only be used with NVECTOR_PARALLEL.

CVODES uses various constants for both input and output. These are defined as needed in this chapter, but for convenience are also listed separately in Chapter 11.

### 5.1 Access to libraries and header files

At this point, it is assumed that the installation of CVODES, following the procedure described in Chapter 2, 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 cvodes. The relevant library files are

- libdir/libsundials_cvodes.lib,
- libdir/libsundials_nvec*. lib (one or two 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/cvodes
- incdir/include/sundials

The directories libdir and incdir are the install libray and include directories. For a default installation, these are builddir/lib and builddir/include, respectively, where builddir was defined in Chapter 2.

Note that an application cannot link to both the CVODE and CVODES libraries because both contain user-callable functions with the same names (to ensure that CVODES is backward compatible with CVODE). Therefore, applications that contain both IVP problems and IVPs with sensitivity analysis, should use CVODES.

### 5.2 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 §2.1).

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 $\operatorname{RCONST}(1.0)$ automatically expands to 1.0 if realtype is double, to 1.0 F if realtype is float, or to 1.0 L if realtype is long double. SUNDIALS uses the RCONST macro internally to declare all of its floating-point constants.

A user program which uses the type realtype and the RCONST macro to handle floating-point constants is precision-independent except for any calls to precision-specific standard math library functions. (Our example programs use both realtype and RCONST.) Users can, however, use the type double, float, or long double in their code (assuming the typedef for realtype matches this choice). Thus, a previously existing piece of ANSI C code can use SUNDIALS without modifying the code to use realtype, so long as the sundials libraries use the correct precision (for details see §2.1).

### 5.3 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:

- cvodes.h, the main header file for CVODES, which defines the several types and various constants, and includes function prototypes.

Note that cvodes.h includes sundials_types.h, which defines the types realtype and booleantype and the constants FALSE and TRUE.

The calling program must also include an NVECTOR implementation header file (see $\S 8$ for details). For the two NVECTOR implementations that are included in the CVODES package, the corresponding header files are:

- nvector_serial.h, which defines the serial implementation NVECTOR_SERIAL;
- nvector_parallel.h, which defines the parallel (MPI) implementation, NVECTOR_PARALLEL.

Note that both these files in turn include the header file sundials_nvector.h which defines the abstract N_Vector data type.

Finally, if the user chooses Newton iteration for the solution of the nonlinear systems, then a linear solver module header file will be required. The header files corresponding to the various linear solvers availble for use with CVODES are:

- cvodes_dense.h, which is used with the dense direct linear solver in the context of CVODES. This in turn includes a header file (sundials_dense.h) which defines the DenseMat type and corresponding accessor macros;
- cvodes_band.h, which is used with the band direct linear solver in the context of CVODES. This in turn includes a header file (sundials_band.h) which defines the BandMat type and corresponding accessor macros;
- cvodes_diag.h, which is used with the diagonal linear solver in the context of CVODES;
- cvodes_spgmr.h, which is used with the Krylov solver SPGMR in the context of CVODES;
- cvodes_spbcgs.h, which is used with the Krylov solver SPBCG in the context of CVODES;
- cvodes_sptfqmr.h; which is used with the Krylov solver SPTFQMR in the context of CVODES;

The header files for the Krylov iterative solvers include cvodes spils.h which defined common fuunctions 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, depending upon the choice of preconditioner, etc. In one of the examples in [18], preconditioning is done with a block-diagonal matrix. For this, the header sundials_smalldense.h is included.

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

The following is a skeleton of the user's main program (or calling program) for the integration of an ODE IVP. Some steps are independent of the NVECTOR implementation used; where this is not the case, usage specifications are given for the two implementations provided with CVODES: steps marked with $[\mathbf{P}]$ correspond to NVECTOR_PARALLEL, while steps marked with $[\mathbf{S}]$ correspond to NVECTOR_SERIAL.

## 1. [P] Initialize MPI

Call MPI_Init (\&argc, \&argv) ; to initialize MPI if used by the user's program. Here argc and argv are the command line argument counter and array received by main, respectively.

## 2. Set problem dimensions

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

## 3. Set vector of initial values

To set the vector y0 of initial values, use the appropriate functions defined by a particular NVECTOR implementation. If a realtype array ydata containing the initial values of $y$ already exists, then make the call:

```
[S] y0 = N_VMake_Serial(N, ydata);
[P] y0 = N_VMake_Parallel(comm, Nlocal, N, ydata);
```

Otherwise, make the call:

```
[S] y0 = N_VNew_Serial(N);
[P] y0 = N_VNew_Parallel(comm, Nlocal, N);
```

and load initial values into the structure defined by:
[S] NV_DATA_S (y0)
[P] NV_DATA_P (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 cvodes object

Call cvode_mem = CVodeCreate (lmm, iter) ; to create the CVODES memory block and to specify the solution method (linear multistep method and nonlinear solver iteration type). CVodeCreate returns a pointer to the CVODES memory structure. See $\S 5.5 .1$ for details.

## 5. Allocate internal memory

Call CVodeMalloc (. . ) ; to provide required problem specifications, allocate internal memory for CVODES, and initialize CVODES. CVodeMalloc returns a flag, the value of which indicates either success or an illegal argument value. See $\S 5.5 .1$ for details.

## 6. Set optional inputs

Call CVodeSet* functions to change any optional inputs that control the behavior of CVODES from their default values. See §5.5.5.1 for details.

## 7. Attach linear solver module

If Newton iteration is chosen, initialize the linear solver module with one of the following calls (for details see §5.5.3):
[S] ier = CVDense(...);
[S] ier = CVBand(...);

```
ier = CVDiag(...);
```

ier $=$ CVSpgmr (...);
ier $=\operatorname{CVSpbcg}(. .$.$) ;$
ier $=$ CVSptfqmr (...);

## 8. Set linear solver optional inputs

Call CV*Set* functions from the selected linear solver module to change optional inputs specific to that linear solver. See $\S 5.5 .5$ for details.

## 9. Specify rootfinding problem

Optionally, call CVodeRootInit to initialize a rootfinding problem to be solved during the integration of the ODE system. See §5.8.1 for details.

## 10. Advance solution in time

For each point at which output is desired, call ier = CVode (cvode_mem, tout, yout, \&tret, itask) ; Set itask to specify the return mode. The vector y (which can be the same as the vector y0 above) will contain $y(t)$. See $\S 5.5 .4$ for details.

## 11. Get optional outputs

Call CV*Get* functions to obtain optional output. See $\S 5.5 .7$ and $\S 5.8 .1$ for details.

## 12. Deallocate memory for solution vector

Upon completion of the integration, deallocate memory for the vector y by calling the destructor function defined by the NVECTOR implementation:
[S] N_VDestroy_Serial (y) ;
[P] N_VDestroy_Parallel (y);

## 13. Free solver memory

Call CVodeFree(\&cvode_mem) ; to free the memory allocated for CVODES.

## 14. [P] Finalize MPI <br> Call MPI_Finalize() ; to terminate MPI.

### 5.5 User-callable functions for IVP solution

This section describes the CVODES functions that are called by the user to setup and then solve an IVP. Some of these are required. However, starting with $\S 5.5 .5$, the functions listed involve optional inputs/outputs or restarting, and those paragraphs may be skipped for a casual use of CVODES. In any case, refer to $\S 5.4$ for the correct order of these calls. Calls related to rootfinding are described in §5.8.

### 5.5.1 CVODES 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 IVP solution is complete, as it frees the CVODES memory block created and allocated by the first two calls.

## CVodeCreate

Call cvode_mem = CVodeCreate(lmm, iter);
Description The function CVodeCreate instantiates a CVODES solver object and specifies the solution method.

Arguments lmm (int) specifies the linear multistep method and may be one of two possible values: CV_ADAMS or CV_BDF.
iter (int) specifies the type of nonlinear solver iteration and may be either CV_NEWTON or CV_FUNCTIONAL.
The recommended choices for (lmm, iter) are (CV_ADAMS, CV_FUNCTIONAL) for nonstiff problems and (CV_BDF, CV_NEWTON) for stiff problems.
Return value If successful, CVodeCreate returns a pointer to the newly created CVODES memory block (of type void *). If an error occurred, CVodeCreate prints an error message to stderr and returns NULL.

| CVodeMalloc |
| :--- | :--- |
| Call flag $=$ CVodeMalloc(cvode_mem, $f, ~ t 0, ~ y 0, ~ i t o l, ~ r e l t o l, ~ a b s t o l) ; ~$ |

Description The function CVodeMalloc provides required problem and solution specifications, allocates internal memory, and initializes CVODES.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block returned by CVodeCreate. $\mathrm{f} \quad$ (CVRhsFn) is the C function which computes $f$ in the ODE. This function has the form $f(t, y$, ydot, f_data) (for full details see §5.6.1).
t0 (realtype) is the initial value of $t$.
y0 (N_Vector) is the initial value of $y$.
itol (int) is one of CV_SS, CV_SV, or CV_WF. Here itol $=$ SS indicates scalar relative error tolerance and scalar absolute error tolerance, while itol $=$ CV_SV indicates scalar relative error tolerance and vector absolute error tolerance. The latter choice is important when the absolute error tolerance needs to be different for each component of the ODE. If itol $=C V \_W F$, the arguments reltol and abstol are ignored and the user is expected to provide a function to evaluate the error weight vector $W$, replacing (3.7). See CVodeSetEwtFn in §5.5.5.1.
reltol (realtype) is the relative error tolerance.
abstol $(\operatorname{void} *)$ is a pointer to the absolute error tolerance. If itol $=C V \_S S$, abstol must be a pointer to a realtype variable. If itol $=$ CV_SV, abstol must be an N_Vector variable.

Return value The return value flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeMalloc was successful.
CV_MEM_NULL The CVODES memory block was not initialized through a previous call to CVodeCreate.
CV_MEM_FAIL A memory allocation request has failed.
CV_ILL_INPUT An input argument to CVodeMalloc has an illegal value.
Notes See also §5.5.2 for advice on tolerances.
The tolerance values in reltol and abstol may be changed between calls to CVode (see CVodeSetTolerances in §5.5.5.1).
It is the user's responsibility to provide compatible itol and abstol arguments.
If an error occurred, CVodeMalloc also sends an error message to the error handler function.

## CVodeFree

Call CVodeFree (\&cvode_mem) ;
Description The function CVodeFree frees the memory allocated by a previous call to CVodeMalloc.
Arguments The argument is the pointer to the CVODES memory block (of type void $*$ ).
Return value The function CVodeFree has no return value.

### 5.5.2 Advice on choice and use of tolerances

General advice on choice of tolerances. For many users, the appropriate choices for tolerance values in reltol and abstol are a concern. The following pieces of advice are relevant.
(1) The scalar relative tolerance reltol is to be set to control relative errors. So reltol $=1.0 \mathrm{E}-4$ means that errors are controlled to $.01 \%$. We do not recommend using reltol larger than $1.0 \mathrm{E}-3$. On the other hand, reltol should not be so small that it is comparable to the unit roundoff of the machine arithmetic (generally around $1.0 \mathrm{E}-15$ ).
(2) The absolute tolerances abstol (whether scalar or vector) need to be set to control absolute errors when any components of the solution vector y may be so small that pure relative error control is meaningless. For example, if y[i] starts at some nonzero value, but in time decays to zero, then pure relative error control on y [i] makes no sense (and is overly costly) after y[i] is below some noise level. Then abstol (if scalar) or abstol[i] (if a vector) needs to be set to that noise level. If the different components have different noise levels, then abstol should be a vector. See the example cvsdenx in the cVode package, and the discussion of it in the cVode Examples document [17]. In that problem, the three components vary betwen 0 and 1 , and have different noise levels; hence the abstol vector. It is impossible to give any general advice on abstol values, because the appropriate noise levels are completely problem-dependent. The user or modeler hopefully has some idea as to what those noise levels are.
(3) Finally, it is important to pick all the tolerance values conservately, because they control the error committed on each individual time step. The final (global) errors are some sort of accumulation of those per-step errors. A good rule of thumb is to reduce the tolerances by a factor of .01 from the actual desired limits on errors. So if you want $.01 \%$ accuracy (globally), a good choice is reltol $=$ $1.0 \mathrm{E}-6$. But in any case, it is a good idea to do a few experiments with the tolerances to see how the computed solution values vary as tolerances are reduced.

Advice on controlling unphysical negative values. In many applications, some components in the true solution are always positive or non-negative, though at times very small. In the numerical
solution, however, small negative (hence unphysical) values can then occur. In most cases, these values are harmless, and simply need to be controlled, not eliminated. The following pieces of advice are relevant.
(1) The way to control the size of unwanted negative computed values is with tighter absolute tolerances. Again this requires some knowledge of the noise level of these components, which may or may not be different for different components. Some experimentation may be needed.
(2) If output plots or tables are being generated, and it is important to avoid having negative numbers appear there (for the sake of avoiding a long explanation of them, if nothing else), then eliminate them, but only in the context of the output medium. Then the internal values carried by the solver are unaffected. Remember that a small negative value in y returned by CVODES, with magnitude comparable to abstol or less, is equivalent to zero as far as the computation is concerned.
(3) The user's right-hand side routine $f$ should never change a negative value in the solution vector $y$ to a non-negative value, as a "solution" to this problem. This can cause instability. If the $f$ routine cannot tolerate a zero or negative value (e.g. because there is a square root or log of it), then the offending value should be changed to zero or a tiny positive number in a temporary variable (not in the input y vector) for the purposes of computing $f(t, y)$.

### 5.5.3 Linear solver specification functions

As previously explained, Newton iteration requires the solution of linear systems of the form (3.5). There are six CVODES linear solvers currently available for this task: CVDENSE, CVBAND, CVDIAG, CVSPGMR, CVSPBCG, and CVSPTFQMR. The first three are direct solvers and their names indicate the type of approximation used for the Jacobian $J=\partial f / \partial y$; CVDENSE, CVBAND, and CVDIAG work with dense, banded, and diagonal approximations to $J$, respectively. The last three cvodes linear solvers - CVSPGMR, CVSPBCG, and CVSPTFQMR - are Krylov iterative solvers, which use scaled preconditioned GMRES, scaled preconditioned Bi-CGStab, and scaled preconditioned TFQMR, respectively. Together, they are refered to as CVSPILS (from scaled preconditioned iterative linear solvers).

With any of the Krylov methods, preconditioning can be done on the left only, on the right only, on both the left and the right, or not at all. For the specification of a preconditioner, see the iterative linear solver sections in $\S 5.5 .5$ and $\S 5.6$.

If preconditioning is done, user-supplied functions define left and right preconditioner matrices $P_{1}$ and $P_{2}$ (either of which could be the identity matrix), such that the product $P_{1} P_{2}$ approximates the Newton matrix $M=I-\gamma J$ of (3.6).

To specify a CVODES linear solver, after the call to CVodeCreate but before any calls to CVode, the user's program must call one of the functions CVDense, CVBand, CVDiag, CVSpgmr, CVSpbcg, or CVSptfqmr, as documented below. The first argument passed to these functions is the CVODES memory pointer returned by CVodeCreate. A call to one of these functions links the main CVODES integrator 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 CVBAND 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 except the diagonal approximation case CVDIAG, the linear solver module used by CVODES is actually built on top of a generic linear system solver, which may be of interest in itself. These generic solvers, denoted DENSE, BAND, SPGMR, SPBCG, and SPTFQMR, are described separately in Chapter 10.

CVDense
Call flag = CVDense(cvode_mem, N);
Description
The function CVDense selects the CVDENSE linear solver.
The user's main program must include the cvodes_dense. h header file.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
$\mathrm{N} \quad$ (long int) problem dimension.

Return value The return value flag (of type int) is one of:
CVDENSE_SUCCESS The CVDENSE initialization was successful.
CVDENSE_MEM_NULL The cvode_mem pointer is NULL.
CVDENSE_ILL_INPUT The CVDENSE solver is not compatible with the current NVECTOR module.
CVDENSE_MEM_FAIL A memory allocation request failed.
Notes The CVDENSE linear solver may not be compatible with a particular implementation of the nVECTOR module. Of the two nVECTOR modules provided with sundials, only NVECTOR_SERIAL is compatible.

CVBand
Call
flag = CVBand(cvode_mem, N, mupper, mlower);
Description The function CVBand selects the CVBAND linear solver.
The user's main program must include the cvodes_band.h header file.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
$\mathrm{N} \quad$ (long int) problem dimension.
mupper (long int) upper half-bandwidth of the problem Jacobian (or of the approximation of it).
mlower (long int) lower half-bandwidth of the problem Jacobian (or of the approximation of it).
Return value The return value flag (of type int) is one of:
CVBAND_SUCCESS The CVBAND initialization was successful.
CVBAND_MEM_NULL The cvode_mem pointer is NULL.
CVBAND_ILL_INPUT The CVBAND solver is not compatible with the current NVECTOR module, or one of the Jacobian half-bandwidths is outside of its valid range $(0 \ldots \mathrm{~N}-1)$.
CVBAND_MEM_FAIL A memory allocation request failed.
Notes The CVBAND linear solver may not be compatible with a particular implementation of the NVECTOR module. Of the two NVECTOR modules provided with sundials, only NVECTOR_SERIAL is compatible. The half-bandwidths are to be set such that the nonzero locations $(i, j)$ in the banded (approximate) Jacobian satisfy -mlower $\leq j-i \leq$ mupper.

CVDiag
Call flag = CVDiag (cvode_mem);
Description The function CVDiag selects the CVDIAG linear solver.
The user's main function must include the cvodes_diag. h header file.
Arguments cvodemem (void $*$ ) pointer to the CVODES memory block.
Return value The return value flag (of type int) is one of:
CVDIAG_SUCCESS The CVDIAG initialization was successful.
CVDIAG_MEM_NULL The cvode_mem pointer is NULL.
CVDIAG_ILL_INPUT The CVDIAG solver is not compatible with the current NVECTOR module.

CVDIAG_MEM_FAIL A memory allocation request failed.

Notes The CVDIAG solver is the simplest of all of the current cVodes linear solvers. The CVDIAG solver uses an approximate diagonal Jacobian formed by way of a difference quotient. The user does not have the option of supplying a function to compute an approximate diagonal Jacobian.

CVSpgmr
Call
flag = CVSpgmr (cvode_mem, pretype, maxl);
Description
The function CVSpgmr selects the CVSPGMR linear solver.
The user's main function must include the cvodes_spgmr.h header file.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
pretype (int) specifies the preconditioning type and must be one of: PREC_NONE, PREC_LEFT, PREC_RIGHT, or PREC_BOTH.
$\operatorname{maxl} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.

Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The CVSPGMR initialization was successful.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
Notes The CVSPGMR solver uses a scaled preconditioned GMRES iterative method to solve the linear system (3.5).

CVSpbcg
Call
Description
flag = CVSpbcg(cvode_mem, pretype, maxl);
The function CVSpbcg selects the CVSPBCG linear solver.
The user's main function must include the cvodes_spbcgs.h header file.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
pretype (int) specifies the preconditioning type and must be one of: PREC_NONE, PREC_LEFT, PREC_RIGHT, or PREC_BOTH.
$\operatorname{maxl} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The CVSPBCG initialization was successful.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
Notes The cVSPBCG solver uses a scaled preconditioned Bi-CGStab iterative method to solve the linear system (3.5).

CVSptfqmr
Call
Description
flag = CVSptfqmr (cvode_mem, pretype, maxl);
The function CVSptfqmr selects the CVSPTFQMR linear solver.
The user's main function must include the cvodes_sptfqmr.h header file.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
pretype (int) specifies the preconditioning type and must be one of: PREC_NONE, PREC_LEFT, PREC_RIGHT, or PREC_BOTH.
$\operatorname{maxl} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The CVSPTFQMR initialization was successful.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
Notes The CVSPTFQMR solver uses a scaled preconditioned TFQMR iterative method to solve the linear system (3.5).

### 5.5.4 CVODE solver function

This is the central step in the solution process - the call to perform the integration of the IVP.

## CVode

Call
flag $=$ CVode(cvode_mem, tout, yout, tret, itask);
Description The function CVode integrates the ODE over an interval in $t$.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
tout (realtype) the next time at which a computed solution is desired.
yout (N_Vector) the computed solution vector.
tret (realtype *) the time reached by the solver.
itask (int) a flag indicating the job of the solver for the next user step. The CV_NORMAL option causes the solver to take internal steps until it has reached or just passed the user-specified tout parameter. The solver then interpolates in order to return an approximate value of $y$ (tout). The CV_ONE_STEP option tells the solver to take just one internal step and then return the solution at the point reached by that step. The CV_NORMAL_TSTOP and CV_ONE_STEP_TSTOP modes are similar to CV_NORMAL and CV_ONE_STEP, respectively, except that the integration never proceeds past the value tstop (specified through the function CVodeSetStopTime).
Return value On return, CVode returns a vector yout and a corresponding independent variable value $t=*$ tret, such that yout is the computed value of $y(t)$.
In CV_NORMAL mode (with no errors), *tret will be equal to tout and yout $=y$ (tout).
The return value flag (of type int) will be one of the following:
CV_SUCCESS CVode succeeded and no roots were found.
CV_TSTOP_RETURN CVode succeeded by reaching the stopping point specified through the optional input function CVodeSetStopTime (see §5.5.5.1).
CV_ROOT_RETURN CVode succeeded and found one or more roots. If nrtfn $>1$, call CVodeGetRootInfo to see which $g_{i}$ were found to have a root. See $\S 5.8$ for more information.
CV_MEM_NULL The cvode mem argument was NULL.
CV_NO_MALLOC The CVODES memory was not allocated by a call to CVodeMalloc.
CV_ILL_INPUT One of the inputs to CVode is illegal. This includes the situation where a root of one of the root functions was found both at a point $t$ and also very near $t$. It also includes the situation where a component of the error weight vector becomes negative during internal time-stepping. The CV_ILL_INPUT flag will also be returned if the
linear solver initialization function (called by the user after calling CVodeCreate) failed to set the linear solver-specific lsolve field in cvode mem. In any case, the user should see the error message for details.
CV_TOO_CLOSE The initial time $t_{0}$ and the final time $t_{\text {out }}$ are too close to each other and the user did not specify an initial step size.
CV_TOO_MUCH_WORK The solver took mxstep internal steps but still could not reach tout. The default value for mxstep is MXSTEP DEFAULT $=500$.
CV_TOO_MUCH_ACC The solver could not satisfy the accuracy demanded by the user for some internal step.
CV_ERR_FAILURE Either error test failures occurred too many times (MXNEF = 7) during one internal time step, or with $|h|=h_{\text {min }}$.
CV_CONV_FAILURE Either convergence test failures occurred too many times (MXNCF = 10) during one internal time step, or with $|h|=h_{\text {min }}$.

CV_LINIT_FAIL The linear solver's initialization function failed.
CV_LSETUP FAIL The linear solver's setup function failed in an unrecoverable manner.
CV_LSOLVE_FAIL The linear solver's solve function failed in an unrecoverable manner.
CV_RHSFUNC_FAIL The right-hand side function failed in an unrecoverable manner.
CV_FIRST_RHSFUNC_FAIL The right-hand side function had a recoverable error at the first call.
CV_REPTD_RHSFUNC_ERR Convergence tests occurred too many times due to repeated recoverable errors in the right-hand side function. The CV_REPTD_RHSFUNC_ERR will also be returned if the right-hand side function had repeated recoverable errors during the estimation of an initial step size.
CV_UNREC_RHSFUNC_ERR The right-hand function had a recoverable error, but no recovery was possible. This failure mode is rare, as it can occur only if the right-hand side function fails recoverably after an error test failed while at order one.
CV_RTFUNC_FAIL The rootfinding function failed.
Notes The vector yout can occupy the same space as the y0 vector of initial conditions that was passed to CVodeMalloc.
In the CV_ONE_STEP mode, tout is only used on the first call to get the direction and a rough scale of the independent variable.
All failure return values are negative and so the test ier < 0 will trap all CVode failures.
On any error return in which one or more internal steps were taken by CVode, the returned values of tret and yout correspond to the farthest point reached in the integration. On all other error returns, tret and yout are left unchanged from the previous CVode return.

### 5.5.5 Optional input functions

CVODES provides an extensive set of functions that can be used to change from their default values various optional input parameters that control the behavior of the CVODES solver. Table 5.1 lists all optional input functions in CVODES which are then described in detail in the remainder of this section, begining with those for the main CVODES solver and continuing with those for the linear solver modules. Note that the diagonal linear solver module has no optional inputs. For the most casual use of CVODES, the reader can skip to §5.6.

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

Table 5.1: Optional inputs for CVODES, CVDENSE, CVBAND, and CVSPILS

| Optional input | Function name | Default |
| :---: | :---: | :---: |
| CVODE main solver |  |  |
| Error handler function <br> Pointer to an error file <br> Data for right-hand side function <br> Maximum order for BDF method <br> Maximum order for Adams method <br> Maximum no. of internal steps before $t_{\text {out }}$ <br> Maximum no. of warnings for $t_{n}+h=t_{n}$ <br> Flag to activate stability limit detection <br> Initial step size <br> Minimum absolute step size <br> Maximum absolute step size <br> Value of $t_{\text {stop }}$ <br> Maximum no. of error test failures <br> Maximum no. of nonlinear iterations <br> Maximum no. of convergence failures <br> Coefficient in the nonlinear convergence test <br> Nonlinear iteration type <br> Integration tolerances <br> Ewt compuation function | CVodeSetErrHandlerFn <br> CVodeSetErrFile <br> CVodeSetFdata <br> CVodeSetMaxOrd <br> CVodeSetMaxOrd <br> CVodeSetMaxNumSteps <br> CVodeSetMaxHnilWarns <br> CVodeSetStabLimDet <br> CVodeSetInitStep <br> CVodeSetMinStep <br> CVodeSetMaxStep <br> CVodeSetStopTime <br> CVodeSetMaxErrTestFails <br> CVodeSetMaxNonlinIters <br> CVodeSetMaxConvFails <br> CVodeSetNonlinConvCoef <br> CVodeSetIterType <br> CVodeSetTolerances <br> CVodeSetEwtFn | internal fn. <br> stderr <br> NULL <br> 5 <br> 12 <br> 500 <br> 10 <br> FALSE <br> estimated <br> 0.0 <br> $\infty$ <br> undefined <br> 7 <br> 3 <br> 10 <br> 0.1 <br> none <br> none <br> internal fn. |
| CVDENSE linear solver |  |  |
| Dense Jacobian function and data | CVDenseSetJacFn | $\begin{aligned} & \text { internal DQ, } \\ & \text { NULL } \end{aligned}$ |
| CVBAND linear solver |  |  |
| Band Jacobian function and data | CVBandSetJacFn | $\begin{aligned} & \text { internal DQ, } \\ & \text { NULL } \end{aligned}$ |
| CVSPILS linear solvers |  |  |
| Preconditioner functions and data Jacobian-times-vector function and data <br> Preconditioning type <br> Ratio between linear and nonlinear tolerances Type of Gram-Schmidt orthogonalization ${ }^{(a)}$ Maximum Krylov subspace size ${ }^{(b)}$ | CVSpilsSetPreconditioner CVSpilsSetJacTimesVecFn <br> CVSpilsSetPrecType <br> CVSpilsSetDelt <br> CVSpilsSetGSType <br> CVSpilsSetMaxl | all NULL <br> internal DQ, <br> NULL <br> none <br> 0.05 <br> classical GS 5 |

(a) Only for CVSPGMR
(b) Only for CVSPBCG and CVSPTFQMR

### 5.5.5.1 Main solver optional input functions

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

## CVodeSetErrHandlerFn

Call flag = CVodeSetErrHandlerFn(cvode_mem, ehfun, eh_data);
Description The function CVodeSetErrHandlerFn specifies the optional user-defined function to be used in handling error messages.
Arguments cvode mem (void $*$ ) pointer to the CVODES memory block.
ehfun (CVErrHandlerFn) is the C error handler function (see §5.6.2).
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
CV_SUCCESS The function ehfun and data pointer eh_data have been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.
Notes The default internal error handler function directs error messages to the file specified by the file pointer errfp (see CVodeSetErrFile below).
Error messages indicating that the CVODES solver memory is NULL will always be directed to stderr.

CVodeSetErrFile
Call flag = CVodeSetErrFile(cvode_mem, errfp);
Description The function CVodeSetErrFile specifies a pointer to the file where all CVODES messages should be directed in case the default CVODES error handler function is used.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. errfp (FILE $*)$ pointer to output file.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_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 CVODES memory pointer is NULL).
If CVodeSetErrFile is to be called, it should be called before any other optional input functions, in order to take effect for any later error message.

## CVodeSetFdata

Call flag = CVodeSetFdata(cvode_mem, f_data);
Description The function CVodeSetFdata specifies the user-defined data block $f$ data to be passed to the user-supplied right-hand side function $f$, and attaches it to the main CVODES memory block.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. f_data (void *) pointer to the user-defined data block.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes If $f$ _data is not specified, a NULL pointer is passed to the $f$ function.

| CVodeSetMax |  |
| :---: | :---: |
| Call | flag = CVodeSetMaxOrder(cvode mem, maxord); |
| Description | The function CVodeSetMaxOrder specifies the maximum order of the linear multistep method. |
| Arguments | cvode_mem (void *) pointer to the CVODES memory block. maxord (int) value of the maximum method order. |
| Return value | The return value flag (of type int) is one of CV_SUCCESS The optional value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL. CV_ILL_INPUT The specified value maxord is negative or larger than its previous value. |
| Notes | The default value is ADAMS_Q MAX = 12 for the Adams-Moulton method and BDF_Q MAX $=5$ for the BDF method. Since maxord affects the memory requirements for the internal CVODES memory block, its value cannot be increased past its previous value. |

CVodeSetMaxNumSteps
Call flag = CVodeSetMaxNumSteps (cvode_mem, mxsteps);

Description The function CVodeSetMaxNumSteps specifies the maximum number of steps to be taken by the solver in its attempt to reach the next output time.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
mxsteps (long int) maximum allowed number of steps.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_ILL_INPUT mxsteps is non-positive.
Notes Passing mxsteps $=0$ results in CVODES using the default value (500).

## CVodeSetMaxHnilWarns

Call flag = CVodeSetMaxHnilWarns (cvode_mem, mxhnil);
Description The function CVodeSetMaxHnilWarns specifies the maximum number of messages issued by the solver warning that $t+h=t$ on the next internal step.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
mxhnil (int) maximum number of warning messages
Return value The return value flag (of type int) is one of CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode mem pointer is NULL.
Notes The default value is 10. A negative value for mxhnil indicates that no warning messages should be issued.

## CVodeSetStabLimDet

Call
flag = CVodeSetstabLimDet(cvode_mem, stldet);
Description The function CVodeSetStabLimDet indicates if the BDF stability limit detection algorithm should be used. See $\S 3.5$ for further details.

Arguments cvode_mem (void *) pointer to the CVODES memory block.
stldet (booleantype) flag controlling stability limit detection (TRUE $=$ on; FALSE $=\mathrm{off})$.

Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_ILL_INPUT The linear multistep method is not set to CV_BDF.
Notes The default value is FALSE. If stldet $=$ TRUE when BDF is used and the method order is greater than or equal to 3 , then an internal function, CVsldet, is called to detect a possible stability limit. If such a limit is detected, then the order is reduced.

```
CVodeSetInitStep
Call flag = CVodeSetInitStep(cvode_mem, hin);
Description The function CVodeSetInitStep specifies the initial step size.
Arguments cvodemem (void \(*\) ) pointer to the CVODES memory block. hin (realtype) value of the initial step size.
```

Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes By default, CVODES estimates the initial step size to be the solution $h$ of the equation $\left\|0.5 h^{2} \ddot{y}\right\|_{\mathrm{WRMS}}=1$, where $\ddot{y}$ is an estimated second derivative of the solution at to.
CVodeSetMinStep
Call flag $=$ CVodeSetMinStep(cvode_mem, hmin);

Description The function CVodeSetMinStep specifies a lower bound on the magnitude of the step size.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
hmin (realtype) minimum absolute value of the step size.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_ILL_INPUT Either hmin is nonpositive or it exceeds the maximum allowable step size.
Notes The default value is 0.0 .
CVodeSetMaxStep
Call flag $=$ CVodeSetMaxStep(cvode_mem, hmax);

Description The function CVodeSetMaxStep specifies an upper bound on the magnitude of the step size.
Arguments cvode mem (void $*$ ) pointer to the CVODES memory block. hmax (realtype) maximum absolute value of the step size.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_ILL_INPUT Either hmax is nonpositive or it is smaller than the minimum allowable step size.
Notes Pass hmax $=0$ to obtain the default value $\infty$.
CVodeSetStopTime
Call flag $=$ CVodeSetStopTime (cvode_mem, tstop);

Description The function CVodeSetStopTime specifies the value of the independent variable $t$ past which the solution is not to proceed.
Arguments cvode mem (void $*$ ) pointer to the CVODES memory block.
tstop (realtype) value of the independent variable past which the solution should not proceed.

Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes The default value is $\infty$.

## CVodeSetMaxErrTestFails

Call flag = CVodeSetMaxErrTestFails(cvode_mem, maxnef);
Description The function CVodeSetMaxErrTestFails specifies the maximum number of error test failures permitted in attempting one step.

Arguments cvode_mem (void *) pointer to the CVODES memory block.
maxnef (int) maximum number of error test failures allowed on one step.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes $\quad$ The default value is 7 .
CVodeSetMaxNonlinIters
Call flag = CVodeSetMaxNonlinIters (cvode_mem, maxcor);

Description The function CVodeSetMaxNonlinIters specifies the maximum number of nonlinear solver iterations permitted per step.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
maxcor (int) maximum number of nonlinear solver iterations allowed per step.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes The default value is 3 .

## CVodeSetMaxConvFails

Call flag = CVodeSetMaxConvFails (cvode_mem, maxncf);
Description The function CVodeSetMaxConvFails specifies the maximum number of nonlinear solver convergence failures permitted during one step.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. maxncf (int) maximum number of allowable nonlinear solver convergence failures per step.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.
Notes The default value is 10 .

```
CVodeSetNonlinConvCoef
Call flag = CVodeSetNonlinConvCoef(cvode_mem, nlscoef);
```

Description The function CVodeSetNonlinConvCoef specifies the safety factor used in the nonlinear convergence test (see §3.1).
Arguments cvode mem (void $*$ ) pointer to the CVODES memory block.
nlscoef (realtype) coefficient in nonlinear convergence test.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes $\quad$ The default value is 0.1 .

| CVodeSetIterType |
| :--- | :--- |
| Call flag $=$ CVodeSetIterType(cvode_mem, iter); |

Description The function CVodeSetIterType resets the nonlinear solver iteration type to iter.
Arguments cvode mem (void $*$ ) pointer to the CVODES memory block.
iter (int) specifies the type of nonlinear solver iteration and may be either CV_NEWTON or CV_FUNCTIONAL.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_ILL_INPUT The iter value passed is neither CV_NEWTON nor CV FUNCTIONAL.
Notes The nonlinear solver iteration type is initially specified in the call to CVodeCreate (see §5.5.1). This function call is needed only if iter is being changed from its value in the prior call to CVodeCreate.

## CVodeSetTolerances

Call flag = CVodeSetTolerances(cvode_mem, itol, reltol, abstol);
Description The function CVodeSetTolerances resets the integration tolerances.
Arguments cvode mem (void $*$ ) pointer to the CVODES memory block.
itol (int) is either CV_SS or CV_SV, where itol = CV_SS indicates scalar relative error tolerance and scalar absolute error tolerance, while itol=CV_SV indicates scalar relative error tolerance and vector absolute error tolerance. The latter choice is important when the absolute error tolerance needs to be different for each component of the ODE.
reltol (realtype) the relative error tolerance.
abstol $(\operatorname{void} *)$ is a pointer to the absolute error tolerance. If itol=CV_SS, abstol must be a pointer to a realtype variable. If itol=CV_SV, abstol must be an N_Vector variable.
Return value The return value flag (of type int) is one of
CV_SUCCESS The tolerances have been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_ILL_INPUT An input argument has an illegal value.
Notes The integration tolerances are initially specified in the call to CVodeMalloc (see §5.5.1). This function call is needed only if the tolerances are being changed from their values beween successiv calls to CVode.

It is the user's responsibility to provide compatible itol and abstol arguments.
It is illegal to call CVodeSetTolerances before a call to CVodeMalloc.

## CVodeSetEwtFn

Call flag = CVodeSetEwtFn(cvode_mem, efun, e_data);
Description The function CVodeSetEwtFn specifies the user-defined function to be used in computing the error weight vector $W$, which is normally defined by Eq.(3.7).
Arguments cvode_mem (void *) pointer to the CVODES memory block.
efun (CVEwtFn) is the C function which defines the ewt vector (see §5.6.3).
e_data (void $*$ ) pointer to user data passed to efun every time it is called.
Return value The return value flag (of type int) is one of
CV_SUCCESS The function fun and data pointer e_data have been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.

Notes This function can be called between successive calls to CVode.
If not needed, pass NULL for edata.
It is illegal to call CVodeSetEwtFn before a call to CVodeMalloc.

### 5.5.5.2 Dense linear solver

The cVDENSE solver needs a function to compute a dense approximation to the Jacobian matrix $J(t, y)$. This function must be of type CVDenseJacFn. The user can supply his/her own dense Jacobian function, or use the default difference quotient function CVDenseDQJac that comes with the CVDENSE solver. To specify a user-supplied Jacobian function djac and associated user data jac_data, CVDENSE provides the function CVDenseSetJacFn. The CVDEnse solver passes the pointer jac_data to its 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 jac_data may be identical to f_data, if the latter was specified through CVodeSetFdata.

```
CVDenseSetJacFn
Call flag = CVDenseSetJacFn(cvode_mem, djac, jac_data);
```

Description The function CVDenseSetJacFn specifies the dense Jacobian approximation function to be used and the pointer to user data.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
djac (CVDenseJacFn) user-defined dense Jacobian approximation function.
jac_data (void $*$ ) pointer to the user-defined data structure.
Return value The return value flag (of type int) is one of
CVDENSE_SUCCESS The optional value has been successfully set.
CVDENSE_MEM_NULL The cvode_mem pointer is NULL.
CVDENSE_LMEM_NULL The CVDENSE linear solver has not been initialized.
Notes By default, CVDENSE uses the difference quotient function CVDenseDQJac. If NULL is passed to djac, this default function is used.

The function type CVDenseJacFn is described in §5.6.4.

### 5.5.5.3 Band linear solver

The CVBAND solver needs a function to compute a banded approximation to the Jacobian matrix $J(t, y)$. This function must be of type CVBandJacFn. The user can supply his/her own banded Jacobian approximation function, or use the default difference quotient function CVBandDQJac that comes with the CVBAND solver. To specify a user-supplied Jacobian function bjac and associated user data jac_data, CVBAND provides the function CVBandSetJacFn. The CVBAND solver passes the pointer jac_data to its 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 jac_data may be identical to f_data, if the latter was specified through CVodeSetFdata.

| CVBandSetJacFn |
| :--- |
| Call flag $=$ CVBandSetJacFn(cvode_mem, bjac, jac_data); |

Description The function CVBandSetJacFn specifies the banded Jacobian approximation function to be used and the pointer to user data.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
bjac (CVBandJacFn) user-defined banded Jacobian approximation function.
jac_data (void $*$ ) pointer to the user-defined data structure.
Return value The return value flag (of type int) is one of
CVBAND_SUCCESS The optional value has been successfully set.
CVBAND_MEM_NULL The cvode_mem pointer is NULL.
CVBAND_LMEM_NULL The CVBAND linear solver has not been initialized.
Notes By default, CVBAND uses the difference quotient function CVBandDQJac. If NULL is passed to bjac, this default function is used.
The function type CVBandJacFn is described in §5.6.5.

### 5.5.5.4 SPILS linear solver

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

The evaluation and preprocessing of any Jacobian-related data needed by the user's preconditioner solve function is done in the optional user-supplied function psetup. Both of these functions are fully specified in §5.6. If used, the psetup function should also be specified in the call to CVSpilsSetPreconditioner. Optionally, a CVSPILS solver passes the pointer p_data received through CVSpilsSetPreconditioner 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. The pointer p_data may be identical to f_data, if the latter was specified through CVodeSetFdata.

Ther CVSPILS solvers require a function to compute an approximation to the product between the Jacobian matrix $J(t, y)$ and a vector $v$. The user can supply his/her own Jacobian-times-vector approximation function, or use the difference quotient function CVSpilsDQJtimes that comes with the CVSPILS solvers. A user-defined Jacobian-vector function must be of type CVSpilsJacTimesVecFn and can be specified through a call to CVSpilsSetJacTimesVecFn (see $\S 5.6 .6$ for specification details). As with the preconditioner user data structure p_data, the user can also specify, in the call to CVSpilsSetJacTimesVecFn, a pointer to a user-defined data structure, jac_data, which the CVSPILS solver passes to the Jacobian-times-vector function jtimes each time it is called. The pointer jac_data may be identical to p_data and/or f_data.

CVSpilsSetPreconditioner
Call flag = CVSpilsSetPreconditioner (cvode_mem, psetup, psolve, p_data);

Description The function CVSpilsSetPreconditioner specifies the preconditioner setup and solve functions and the pointer to user data.
Arguments cvodemem (void $*$ ) pointer to the CVODES memory block.
psetup (CVSpilsPrecSetupFn) user-defined preconditioner setup function.
psolve (CVSpilsPrecSolveFn) user-defined preconditioner solve function.
p_data (void $*$ ) pointer to the user-defined data structure.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
Notes The function type CVSpilsPrecSolveFn is described in §5.6.7. The function type CVSpilsPrecSetupFn is described in §5.6.8.

CVSpilsSetJacTimesVecFn
Call flag = CVSpilsSetJacTimesVecFn(cvode_mem, jtimes, jac_data);
Description The function CVSpilsSetJacTimesFn specifies the Jacobian-vector function to be used and the pointer to user data.

Arguments cvode mem (void *) pointer to the CVODES memory block.
jtimes (CVSpilsJacTimesVecFn) user-defined Jacobian-vector product function. jac_data (void $*$ ) pointer to the user-defined data structure.
Return value The return value flag (of type int) is one of CVSPILS_SUCCESS The optional value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
Notes By default, the CVSPILS linear solvers use an internal difference quotient function CVSpilsDQJtimes. If NULL is passed to jtimes, this default function is used.

The function type CVSpilsJacTimesVecFn is described in §5.6.6.

CVSpilsSetPrecType
Call flag = CVSpilsSetPrecType (cvode_mem, pretype);
Description The function CVSpilsSetPrecType resets the type of preconditioning to be used.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
pretype (int) specifies the type of preconditioning and must be one of: PREC_NONE,
PREC_LEFT, PREC_RIGHT, or PREC_BOTH.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
Notes The preconditioning type is initially set in the call to the linear solver's specification function (see $\S 5.5 .3$ ). This function call is needed only if pretype is being changed from its original value.
CVSpilsSetGSType
Call flag $=$ CVSpilsSetGSType (cvode_mem, gstype);

Description The function CVSpilsSetGSType specifies the Gram-Schmidt orthogonalization to be used with the CVSPGMR solver (one of the enumeration constants MODIFIED_GS or CLASSICAL_GS). These correspond to using modified Gram-Schmidt and classical GramSchmidt, respectively.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
gstype (int) type of Gram-Schmidt orthogonalization.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
CVSPILS_ILL_INPUT The Gram-Schmidt orthogonalization type gstype is not valid.
Notes The default value is MODIFIED_GS.
This option is available only for the CVSPGMR linear solver.


| CVSpilsSetDelt |
| :--- |
| Call flag $=$ CVSpilsSetDelt (cvode_mem, delt); |

Description The function CVSpilsSetDelt specifies the factor by which the Krylov linear solver's convergence test constant is reduced from the Newton iteration test constant.

Arguments cvode mem (void $*$ ) pointer to the CVODES memory block.
delt (realtype)
Return value The return value flag (of type int) is one of CVSPILS_SUCCESS The optional value has been successfully set. CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
CVSPILS_ILL_INPUT The factor delt is negative.
Notes The default value is 0.05 .
Passing a value delt $=0.0$ also indicates using the default value.

CVSpilsSetMaxl
Call flag = CVSpilsSetMaxl(cv_mem, maxl);
Description The function CVSpilsSetMaxl resets maximum Krylov subspace dimension for the BiCGStab or TFQMR methods.
Arguments cv_mem (void *) pointer to the CVODES memory block.
$\operatorname{maxl}$ (int) maximum dimension of the Krylov subspace.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional value has been successfuly set.
CVSPILS_MEM_NULL The cv_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
Notes The maximum subspace dimension is initially specified in the call to the linear solver specification function (see $\S 5.5 .3$ ). This function call is needed only if maxl is being changed from its previous value.
This option is available only for the CVSPBCG and CVSPTFQMR linear solvers.


### 5.5.6 Interpolated output function

An optional function CVodeGetDky is available to obtain additional output values. This function should only be called after a successful return from CVode as it provides interpolated values either of $y$ or of its derivatives (up to the current order of the integration method) interpolated to any value of $t$ in the last internal step taken by CVODES.

The call to the CVodeGetDky function has the following form:

| CVodeGetDky |  |
| :---: | :---: |
| Call | flag = CVodeGetDky (cvode mem, t, k, dky) |
| Description | The function CVodeGetDky computes the $k$-th derivative of the function y at time t , i.e. $d^{(k)} y / d t^{(k)}(t)$, where $t_{n}-h_{u} \leq \mathrm{t} \leq t_{n}, t_{n}$ denotes the current internal time reached, and $h_{u}$ is the last internal step size successfully used by the solver. The user may request k $=0,1, \ldots, q_{u}$, where $q_{u}$ is the current order. |
| Arguments | ```cvode_mem (void *) pointer to the CVODES memory block. t (realtype) the value of the independent variable at which the derivative is to be evaluated.``` |
|  | k (int) the derivative order requested. |
|  | dky (N_Vector) vector containing the derivative. This vector must be allocated by the user. |
| Return value | The return value flag (of type int) is one of |
|  | CV_SUCCESS CVodeGetDky succeeded. |
|  | CV_BAD_K k is not in the range $0,1, \ldots, q_{u}$. |
|  | CV_BAD_T $\quad \mathrm{t}$ is not in the interval $\left[t_{n}-h_{u}, t_{n}\right]$. |
|  | CV_BAD_DKY The dky argument was NULL. |
|  | CV_MEM_nULL The cvode mem argument was NULL. |
| Notes | It is only legal to call the function CVodeGetDky after a successful return from CVode. See CVodeGetCurrentTime, CVodeGetLastOrder, and CVodeGetLastStep in the next section for access to $t_{n}, q_{u}$, and $h_{u}$, respectively. |

### 5.5.7 Optional output functions

CVODES provides an extensive set of functions that can be used to obtain solver performance information. Table 5.2 lists all optional output functions in CVODES, which are then described in detail in the remainder of this section, begining with those for the main CVODES 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).

### 5.5.7.1 Main solver optional output functions

CVODES 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, solver performance statistics, as well as additional data from the CVODES memory block (a suggested tolerance scaling factor, the error weight vector, and the vector of estimated local errors). Functions are also provided to extract statistics related to the performance of the CVODES nonlinear solver used. As a convenience, additional information extraction functions provide the optional outputs in groups. These optional output functions are described next.

Table 5.2: Optional outputs from CVODES, CVDENSE, CVBAND, CVDIAG, and CVSPILS

| Optional output | Function name |
| :---: | :---: |
| CVODES main solver |  |
| Size of CVODES real and integer workspaces | CVodeGetWorkSpace |
| Cumulative number of internal steps | CVodeGetNumSteps |
| No. of calls to r.h.s. function | CVodeGetNumRhsEvals |
| No. of calls to linear solver setup function | CVodeGetNumLinSolvSetups |
| No. of local error test failures that have occurred | CVodeGetNumErrTestFails |
| Order used during the last step | CVodeGetLastOrder |
| Order to be attempted on the next step | CVodeGetCurrentOrder |
| No. of order reductions due to stability limit detection | CVodeGetNumStabLimOrderReds |
| Actual initial step size used | CVodeGetActualInitStep |
| Step size used for the last step | CVodeGetLastStep |
| Step size to be attempted on the next step | CVodeGetCurrentStep |
| Current internal time reached by the solver | CVodeGetCurrentTime |
| Suggested factor for tolerance scaling | CVodeGetTolScaleFactor |
| Error weight vector for state variables | CVodeGetErrWeights |
| Estimated local error vector | CVodeGetEstLocalErrors |
| No. of nonlinear solver iterations | CVodeGetNumNonlinSolvIters |
| No. of nonlinear convergence failures | CVodeGetNumNonlinSolvConvFails |
| All CVODES integrator statistics | CVodeGetIntegratorStats |
| CVODES nonlinear solver statistics | CVodeGetNonlinSolvStats |
| Array showing roots found | CvodeGetRootInfo |
| No. of calls to user root function | CVodeGetNumGEvals |
| Name of constant associated with a return flag | CVodeGetReturnFlagName |
| CVDENSE linear solver |  |
| Size of CVDENSE real and integer workspaces | CVDenseGetWorkSpace |
| No. of Jacobian evaluations | CVDenseGetNumJacEvals |
| No. of r.h.s. calls for finite diff. Jacobian evals. | CVDenseGetNumRhsEvals |
| Last return from a CVDENSE function | CVDenseGetLastFlag |
| Name of constant associated with a return flag | CVDenseGetReturnFlagName |
| CVBAND linear solver |  |
| Size of CVBAND real and integer workspaces | CVBandGetWorkSpace |
| No. of Jacobian evaluations | CVBandGetNumJacEvals |
| No. of r.h.s. calls for finite diff. Jacobian evals. | CVBandGetNumRhsEvals |
| Last return from a CVBAND function | CVBandGetLastFlag |
| Name of constant associated with a return flag | CVBandGetReturnFlagName |
| CVDIAG linear solver |  |
| Size of CVDIAG real and integer workspaces | CVDiagGetWorkSpace |
| No. of r.h.s. calls for finite diff. Jacobian evals. | CVDiagGetNumRhsEvals |
| Last return from a CVDIAG function | CVDiagGetLastFlag |
| Name of constant associated with a return flag | CVDiagGetReturnFlagName |
| CVSPILS linear solvers |  |
| Size of real and integer workspaces | CVSpilsGetWorkSpace |
| No. of linear iterations | CVSpilsGetNumLinIters |
| No. of linear convergence failures | CVSpilsGetNumConvFails |
| No. of preconditioner evaluations | CVSpilsGetNumPrecEvals |
| No. of preconditioner solves | CVSpilsGetNumPrecSolves |
| No. of Jacobian-vector product evaluations | CVSpilsGetNumJtimesEvals |
| No. of r.h.s. calls for finite diff. Jacobian-vector evals. | CVSpilsGetNumRhsEvals |
| Last return from a linear solver function | CVSpilsGetLastFlag |
| Name of constant associated with a return flag | CVSpilsGetReturnFlagName |

## CVodeGetWorkSpace

Call flag = CVodeGetWorkSpace(cvode_mem, \&lenrw, \&leniw);
Description The function CVodeGetWorkSpace returns the CVODES real and integer workspace sizes.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
lenrw (long int) the number of realtype values in the CVODES workspace.
leniw (long int) the number of integer values in the CVODES workspace.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional output values have been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.
Notes In terms of the problem size $N$, the maximum method order maxord, and the number nrtfn of root functions (see §5.8), the actual size of the real workspace, in realtype words, is given by the following:

- base value: lenrw $=96+(\operatorname{maxord}+5) * N_{r}+3 * \operatorname{nrtfn} ;$
- if itol $=$ CV_SV: lenrw $=$ lenrw $+N_{r}$;
where $N_{r}$ is the number of real words in one N_Vector $(\approx N)$.
The size of the integer workspace (without distinction between int and long int words) is given by:
- base value: leniw $=40+($ maxord +5$) * N_{i}+\operatorname{nrtfn} ;$
- if itol $=$ CV_SV: leniw $=$ leniw $+N_{i}$;
where $N_{i}$ is the number of integer words in one N_Vector ( $=1$ for NVECTOR_SERIAL and $2 *$ npes for NVECTOR_PARALLEL and npes processors).

For the default value of maxord, with no rootfinding, and with itol $\neq \mathrm{CV}$ _SV, these lengths are given roughly by:

- For the Adams method: lenrw $=96+17 N$ and leniw $=57$
- For the BDF method: lenrw $=96+10 N$ and leniw $=50$

Note that additional memory is allocated if quadratures and/or forward sensitivity integration is enabled. See $\S 5.7 .1$ and $\S 6.2 .1$ for more details.

## CVodeGetNumSteps <br> Call flag = CVodeGetNumSteps (cvode_mem, \&nsteps);

Description The function CVodeGetNumSteps returns the cumulative number of internal steps taken by the solver (total so far).
Arguments cvode_mem (void *) pointer to the CVODES memory block.
nsteps (long int) number of steps taken by CVODES.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetNumRhsEvals

Call flag = CVodeGetNumRhsEvals(cvode_mem, \&nfevals);
Description The function CVodeGetNumRhsEvals returns the number of calls to the user's right-hand side function.

Arguments cvode_mem (void *) pointer to the CVODES memory block. nfevals (long int) number of calls to the user's $f$ function.

Return value The return value flag (of type int) is one of CV_SUCCESS The optional output value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.
Notes The nfevals value returned by CVodeGetNumRhsEvals does not account for calls made to $f$ by a linear solver or preconditioner module.

## CVodeGetNumLinSolvSetups

Call flag = CVodeGetNumLinSolvSetups(cvode_mem, \&nlinsetups);
Description The function CVodeGetNumLinSolvSetups returns the number of calls made to the linear solver's setup function.

Arguments cvode_mem (void *) pointer to the CVODES memory block. nlinsetups (long int) number of calls made to the linear solver setup function.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional output value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetNumErrTestFails

Call flag = CVodeGetNumErrTestFails(cvode_mem, \&netfails);
Description The function CVodeGetNumErrTestFails returns the number of local error test failures that have occurred.
Arguments cvodemem (void $*$ ) pointer to the CVODES memory block. netfails (long int) number of error test failures.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

CVodeGetLastOrder
Call flag = CVodeGetLastOrder (cvode_mem, \&qlast);
Description The function CVodeGetLastOrder returns the integration method order used during the last internal step.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. qlast (int) method order used on the last internal step.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional output value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetCurrentOrder

Call flag = CVodeGetCurrentOrder (cvode_mem, \&qcur);
Description The function CVodeGetCurrentOrder returns the integration method order to be used on the next internal step.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
qcur (int) method order to be used on the next internal step.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetLastStep

Call flag = CVodeGetLastStep(cvode_mem, \&hlast);
Description The function CVodeGetLastStep returns the integration step size taken on the last internal step.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
hlast (realtype) step size taken on the last internal step.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetCurrentStep <br> Call flag = CVodeGetCurrentStep(cvode_mem, \&hcur);

Description The function CVodeGetCurrentStep returns the integration step size to be attempted on the next internal step.

Arguments cvode_mem (void *) pointer to the CVODES memory block. hcur (realtype) step size to be attempted on the next internal step.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional output value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.

CVodeGetActualInitStep
Call flag = CVodeGetActualInitStep(cvode_mem, \&hinused);
Description The function CVodeGetActualInitStep returns the value of the integration step size used on the first step.
Arguments cvodemem (void *) pointer to the CVODES memory block. hinused (realtype) actual value of initial step size.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional output value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.
Notes Even if the value of the initial integration step size was specified by the user through a call to CVodeSetInitStep, this value might have been changed by CVODES to ensure that the step size is within the prescribed bounds ( $h_{\min } \leq h_{0} \leq h_{\max }$ ), or to satisfy the local error test condition.

## CVodeGetCurrentTime

Call flag = CVodeGetCurrentTime(cvode_mem, \&tcur);
Description The function CVodeGetCurrentTime returns the current internal time reached by the solver.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
tcur (realtype) current internal time reached.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetNumStabLimOrderReds

Call flag = CVodeGetNumStabLimOrderReds(cvode_mem, \&nslred);
Description The function CVodeGetNumStabLimOrderReds returns the number of order reductions dictated by the BDF stability limit detection algorithm (see §3.5).
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. nslred (long int) number of order reductions due to stability limit detection.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes If the stability limit detection algorithm was not initialized through a call to CVodeSetStabLimDet, then nslred $=0$.

## CVodeGetTolScaleFactor

Call flag = CVodeGetTolScaleFactor (cvode_mem, \&tolsfac);
Description The function CVodeGetTolScaleFactor returns a suggested factor by which the user's tolerances should be scaled when too much accuracy has been requested for some internal step.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
tolsfac (realtype) suggested scaling factor for user-supplied tolerances.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetErrWeights

Call flag = CVodeGetErrWeights(cvode_mem, eweight);
Description The function CVodeGetErrWeights returns the solution error weights at the current time. These are the reciprocals of the $W_{i}$ given by (3.7).
Arguments cvode_mem (void *) pointer to the CVODES memory block.
eweight (N_Vector) solution error weights at the current time.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional output value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL.
Notes The user must allocate memory for eweight.

## CVodeGetEstLocalErrors

Call flag = CVodeGetEstLocalErrors (cvode_mem, ele);
Description The function CVodeGetEstLocalErrors returns the vector of estimated local errors.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
ale (N_Vector) estimated local errors.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
The user must allocate memory for ale.
The ale vector, together with the eweight vector from CVodeGetErrWeights, can be used to determine how the various components of the system contributed to the estimated local error test. Specifically, that error test uses the RMS norm of a vector whose components are the products of the components of the two vectors. Thus, for example, if there were recent error test failures, the components causing the failures are those with largest values for the products, denoted loosely as eweight[i]*ele[i].

```
CVodeGetIntegratorStats
Call flag = CVodeGetIntegratorStats(cvode_mem, &nsteps, &nfevals,
    &nlinsetups, &netfails, &qlast, &qcur,
    &hinused, &hlast, &hcur, &tcur);
```

Description The function CVodeGetIntegratorStats returns the CVODES integrator statistics as a group.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
nsteps (long int) number of steps taken by CVODES.
nfevals (long int) number of calls to the user's $f$ function.
nlinsetups (long int) number of calls made to the linear solver setup function.
netfails (long int) number of error test failures.
qlast (int) method order used on the last internal step.
qcur (int) method order to be used on the next internal step.
hinused (realtype) actual value of initial step size.
hlast (realtype) step size taken on the last internal step.
hour (realtype) step size to be attempted on the next internal step.
tour (realtype) current internal time reached.
Return value The return value flag (of type int) is one of
CV_SUCCESS the optional output values have been successfully set.
CV_MEM_NULL the cvode_mem pointer is NULL.

## CVodeGetNumNonlinSolvIters

Call flag = CVodeGetNumNonlinSolvIters(cvode_mem, \&nniters);
Description The function CVodeGetNumNonlinSolvIters returns the number of nonlinear (functional or Newton) iterations performed.

Arguments cvode_mem (void *) pointer to the CVODES memory block. nniters (long int) number of nonlinear iterations performed.
Return value The return value flag (of type int) is one of CV_SUCCESS The optional output values have been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

```
CVodeGetNumNonlinSolvConvFails
Call flag = CVodeGetNumNonlinSolvConvFails(cvode_mem, &nncfails);
```

Description The function CVodeGetNumNonlinSolvConvFails returns the number of nonlinear convergence failures that have occurred.
Arguments cvodemem (void $*$ ) pointer to the CVODES memory block.
nncfails (long int) number of nonlinear convergence failures.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetNonlinSolvStats

Call flag = CVodeGetNonlinSolvStats(cvode_mem, \&nniters, \&nncfails);
Description The function CVodeGetNonlinSolvStats returns the CVODES nonlinear solver statistics as a group.
Arguments cvode mem (void $*$ ) pointer to the CVODES memory block.
nniters (long int) number of nonlinear iterations performed.
nncfails (long int) number of nonlinear convergence failures.
Return value The return value flag (of type int) is one of
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

## CVodeGetReturnFlagName

Call name = CVodeGetReturnFlagName (flag);
Description The function CVodeGetReturnFlagName returns the name of the CVODE constant corresponding to flag.

Arguments The only argument, of type int is a return flag from a CVODE function.
Return value The return value is a string containing the name of the corresponding constant.

### 5.5.7.2 Dense linear solver

The following optional outputs are available from the CVDENSE module: workspace requirements, number of calls to the Jacobian routine, number of calls to the right-hand side routine for finitedifference Jacobian approximation, and last return value from a CVDENSE function.

```
CVDenseGetWorkSpace
Call flag = CVDenseGetWorkSpace(cvode_mem, &lenrwLS, &leniwLS);
```

Description The function CVDenseGetWorkSpace returns the CVDENSE real and integer workspace sizes.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
lenrwLS (long int) the number of realtype values in the CVDENSE workspace.
leniwLS (long int) the number of integer values in the CVDENSE workspace.
Return value The return value flag (of type int) is one of
CVDENSE_SUCCESS The optional output values have been successfully set.
CVDENSE_MEM_NULL The cvode_mem pointer is NULL.
CVDENSE_LMEM_NULL The CVDENSE linear solver has not been initialized.

Notes In terms of the problem size $N$, the actual size of the real workspace is $2 N^{2}$ realtype words, and the actual size of the integer workspace is $N$ integer words.

## CVDenseGetNumJacEvals

Call flag = CVDenseGetNumJacEvals(cvode.mem, \&njevals);
Description The function CVDenseGetNumJacEvals returns the number of calls made to the dense Jacobian approximation function.

Arguments cvode mem (void $*$ ) pointer to the CVODES 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
CVDENSE_SUCCESS The optional output value has been successfully set.
CVDENSE_MEM_NULL The cvode_mem pointer is NULL.
CVDENSE_LMEM_NULL The CVDENSE linear solver has not been initialized.

CVDenseGetNumRhsEvals
Call flag = CVDenseGetNumRhsEvals(cvode_mem, \&nfevalsLS);
Description The function CVDenseGetNumRhsEvals returns the number of calls made to the usersupplied right-hand side function due to the finite difference dense Jacobian approximation.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
nfevalsLS (long int) the number of calls made to the user-supplied right-hand side function.

Return value The return value flag (of type int) is one of
CVDENSE_SUCCESS The optional output value has been successfully set.
CVDENSE_MEM_NULL The cvode_mem pointer is NULL.
CVDENSE_LMEM_NULL The CVDENSE linear solver has not been initialized.
Notes The value nfevalsLS is incremented only if the default CVDenseDQJac difference quotient function is used.

CVDenseGetLastFlag
Call flag = CVDenseGetLastFlag(cvode_mem, \&lsflag);
Description The function CVDenseGetLastFlag returns the last return value from a CVDENSE routine.
Arguments cvodemem (void $*$ ) pointer to the CVODES memory block. lsflag (int) the value of the last return flag from a CVDENSE function.
Return value The return value flag (of type int) is one of CVDENSE_SUCCESS The optional output value has been successfully set.
CVDENSE_MEM_NULL The cvode_mem pointer is NULL.
CVDENSE_LMEM_NULL The CVDENSE linear solver has not been initialized.
Notes If the CVDENSE setup function failed (CVode returned CV_LSETUP_FAIL), then the value of lsflag corresponds to the column index (numbered from one) of a diagonal element with value zero that was encountered during the LU factorization of the dense Jacobian matrix.

```
CVDenseGetReturnFlagName
Call name = CVDenseGetReturnFlagName(flag);
```

Description The function CVDenseGetReturnFlagName returns the name of the CVDENSE constant corresponding to flag.
Arguments The only argument, of type int is a return flag from a CVDENSE function.
Return value The return value is a string containing the name of the corresponding constant.

### 5.5.7.3 Band linear solver

The following optional outputs are available from the CVBAND module: workspace requirements, number of calls to the Jacobian routine, number of calls to the right-hand side routine for finitedifference Jacobian approximation, and last return value from a CVBAND function.

CVBandGetWorkSpace
Call flag = CVBandGetWorkSpace(cvode_mem, \&lenrwLS, \&leniwLS);
Description The function CVBandGetWorkSpace returns the CVBAND real and integer workspace sizes.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
lenrwLS (long int) the number of realtype values in the CVBAND workspace.
leniwLS (long int) the number of integer values in the CVBAND workspace.
Return value The return value flag (of type int) is one of
CVBAND_SUCCESS The optional output values have been successfully set.
CVBAND_MEM_NULL The cvode_mem pointer is NULL.
CVBAND_LMEM_NULL The CVBAND linear solver has not been initialized.
Notes In terms of the problem size $N$ and Jacobian half-bandwidths, the actual size of the real workspace is ( 2 mupper +3 mlower +2 ) $N$ realtype words, and the actual size of the integer workspace is $N$ integer words.

## CVBandGetNumJacEvals

Call flag = CVBandGetNumJacEvals (cvode_mem, \&njevals);
Description The function CVBandGetNumJacEvals returns the number of calls made to the banded Jacobian approximation function.
Arguments cvode_mem (void *) pointer to the CVODES 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
CVBAND_SUCCESS The optional output value has been successfully set.
CVBAND_MEM_NULL The cvode_mem pointer is NULL.
CVBAND_LMEM_NULL The CVBAND linear solver has not been initialized.

CVBandGetNumRhsEvals
Call flag = CVBandGetNumRhsEvals(cvode_mem, \&nfevalsLS);
Description The function CVBandGetNumRhsEvals returns the number of calls made to the usersupplied right-hand side function due to the finite difference banded Jacobian approximation.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
nfevalsLS (long int) the number of calls made to the user-supplied right-hand side function.
Return value The return value flag (of type int) is one of
CVBAND_SUCCESS The optional output value has been successfully set.
CVBAND_MEM_NULL The cvode_mem pointer is NULL.
CVBAND_LMEM_NULL The CVBAND linear solver has not been initialized.
Notes The value nfevalsLS is incremented only if the default CVBandDQJac difference quotient function is used.

CVBandGetLastFlag
Call flag = CVBandGetLastFlag (cvode_mem, \&lsflag);
Description The function CVBandGetLastFlag returns the value of the last return flag from a CVBAND routine.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
lsflag (int) the value of the last return flag from a CVBAND function.
Return value The return value flag (of type int) is one of CVBAND_SUCCESS The optional output value has been successfully set.
CVBAND_MEM_NULL The cvode_mem pointer is NULL.
CVBAND_LMEM_NULL The CVBAND linear solver has not been initialized.
Notes If the CVBAND setup function failed (CVode returned CV_LSETUP_FAIL), the value of lsflag corresponds to the column index (numbered from one) of a diagonal element with value zero that was encountered during the LU factorization of the banded Jacobian matrix.

CVBandGetReturnFlagName
Call name = CVBandGetReturnFlagName(flag);
Description The function CVBandGetReturnFlagName returns the name of the CVBAND constant corresponding to flag.
Arguments The only argument, of type int is a return flag from a CVBAND function.
Return value The return value is a string containing the name of the corresponding constant.

### 5.5.7.4 Diagonal linear solver

The following optional outputs are available from the CVDIAG module: workspace requirements, number of calls to the right-hand side routine for finite-difference Jacobian approximation, and last return value from a CVDIAG function.

## CVDiagGetWorkSpace

Call flag = CVDiagGetWorkSpace (cvode_mem, \&lenrwLS, \&leniwLS);
Description The function CVDiagGetWorkSpace returns the CVDIAG real and integer workspace sizes.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
lenrwLS (long int) the number of realtype values in the CVDIAG workspace.
leniwLS (long int) the number of integer values in the CVDIAG workspace.
Return value The return value flag (of type int) is one of
CVDIAG_SUCCESS The optional output valus have been successfully set.
CVDIAG_MEM_NULL The cvode_mem pointer is NULL.

CVDIAG_LMEM_NULL The CVDIAG linear solver has not been initialized.
Notes In terms of the problem size $N$, the actual size of the real workspace is roughly $3 N$ realtype words.

| CVDiagGetNumRhsEvals |
| :--- |
| Call flag = CVDiagGetNumRhsEvals (cvode_mem, \&nfevalsLS); |

Description The function CVDiagGetNumRhsEvals returns the number of calls made to the usersupplied right-hand side function due to the finite difference Jacobian approximation.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
nfevalsLS (long int) the number of calls made to the user-supplied right-hand side function.
Return value The return value flag (of type int) is one of
CVDIAG_SUCCESS The optional output value has been successfully set.
CVDIAG_MEM_NULL The cvode_mem pointer is NULL.
CVDIAG_LMEM_NULL The CVDIAG linear solver has not been initialized.
Notes The number of diagonal approximate Jacobians formed is equal to the number of calls made to the linear solver setup function (see CVodeGetNumLinSolvSetups).

## CVDiagGetLastFlag

Call flag = CVDiagGetLastFlag (cvode_mem, \&lsflag);
Description The function CVDiagGetLastFlag returns the last return value from a CVDIAG routine.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
lsflag (int) the value of the last return flag from a CVDIAG function.
Return value The return value flag (of type int) is one of CVDIAG_SUCCESS The optional output value has been successfully set.
CVDIAG_MEM_NULL The cvode_mem pointer is NULL. CVDIAG_LMEM_NULL The CVDIAG linear solver has not been initialized.
Notes If the CVDIAG setup function failed (CVode returned CV_LSETUP FAIL), the value of lsflag is equal to CVDIAG_INV_FAIL, indicating that a diagonal element with value zero was encountered. The same value is also returned if the CVDIAG solve function failed (CVode returned CV_LSOLVE_FAIL).

CVDiagGetReturnFlagName
Call name = CVDiagGetReturnFlagName (flag);
Description The function CVDiagGetReturnFlagName returns the name of the CVDIAG constant corresponding to flag.
Arguments The only argument, of type int is a return flag from a CVDIAG function.
Return value The return value is a string containing the name of the corresponding constant.

### 5.5.7.5 SPILS linear solvers

The following optional outputs are available from the CVSPILS 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 right-hand side routine for finite-difference Jacobian-vector product approximation, and last return value from a linear solver function.
CVSpilsGetWorkSpace
Call flag = CVSpilsGetWorkSpace(cvode_mem, \&lenrwLS, \&leniwLS);

Description The function CVSpilsGetWorkSpace returns the global sizes of the CVSPGMR real and integer workspaces.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
lenrwLS (long int) the number of realtype values in the CVSPILS workspace.
leniwLS (long int) the number of integer values in the CVSPILS workspace.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional output value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
Notes In terms of the problem size $N$ and maximum subspace size maxl, the actual size of the real workspace is roughly:
$(\operatorname{maxl}+5) * N+\operatorname{maxl} *(\operatorname{maxl}+4)+1$ realtype words for CVSPGMR, $9 * N$ realtype words for CVSPBCG, and $11 * N$ realtype words for IDASPTFQMR.
In a parallel setting, the above values are global - summed over all processors.

## CVSpilsGetNumLinIters

Call flag = CVSpilsGetNumLinIters(cvode_mem, \&nliters);
Description The function CVSpilsGetNumLinIters returns the cumulative number of linear iterations.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
nliters (long int) the current number of linear iterations.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional output value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.

CVSpilsGetNumConvFails
Call flag = CVSpilsGetNumConvFails(cvode_mem, \&nlcfails);
Description The function CVSpilsGetNumConvFails returns the cumulative number of linear convergence failures.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
nlcfails (long int) the current number of linear convergence failures.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional output value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.

## CVSpilsGetNumPrecEvals

Call flag = CVSpilsGetNumPrecEvals(cvode_mem, \&npevals);
Description The function CVSpilsGetNumPrecEvals returns the number of preconditioner evaluations, i.e., the number of calls made to psetup with jok = FALSE.

Arguments cvode mem (void $*$ ) pointer to the CVODES memory block. npevals (long int) the current number of calls to psetup.

Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional output value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.

## CVSpilsGetNumPrecSolves

Call flag = CVSpilsGetNumPrecSolves(cvode_mem, \&npsolves);
Description The function CVSpilsGetNumPrecSolves returns the cumulative number of calls made to the preconditioner solve function, psolve.

Arguments cvode_mem (void *) pointer to the CVODES memory block. npsolves (long int) the current number of calls to psolve.

Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional output value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.

CVSpilsGetNumJtimesEvals
Call flag = CVSpilsGetNumJtimesEvals(cvode_mem, \&njvevals);
Description The function CVSpilsGetNumJtimesEvals returns the cumulative number made to the Jacobian-vector function, jtimes.

Arguments cvode mem (void $*$ ) pointer to the CVODES memory block. njvevals (long int) the current number of calls to jtimes.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional output value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.

CVSpilsGetNumRhsEvals
Call flag = CVSpilsGetNumRhsEvals(cvode_mem, \&nfevalsLS);
Description The function CVSpilsGetNumRhsEvals returns the number of calls to the user righthand side function for finite difference Jacobian-vector product approximation.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. nfevalsLS (long int) the number of calls to the user right-hand side function.

Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional output value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
Notes The value nfevalsLS is incremented only if the default CVSpilsDQJtimes difference quotient function is used.


Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The optional output value has been successfully set.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_LMEM_NULL The CVSPILS linear solver has not been initialized.
Notes If the CVSPILS setup function failed (CVode returned CV_LSETUP_FAIL), lsflag will be SPGMR_PSET_FAIL_UNREC, SPBCG_PSET_FAIL_UNREC, or SPTFQMR_PSET FAIL_UNREC.
If the CVSPGMR solve function failed (CVode returned CV_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 SPGMR_QRSOL_FAIL, indicating that the matrix $R$ was found to be singular during the QR solve phase.
If the CVSPBCG solve function failed (CVode returned CV LSOLVEFFAIL), 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 CVSPTFQMR solve function failed (CVode returned CV_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 Jacobian-times-vector function; or SPTFQMR_PSOLVE_FAIL_UNREC, indicating that the preconditioner solve function psolve failed unrecoverably.

## CVSpilsGetReturnFlagName

Call name = CVSpilsGetReturnFlagName (flag);
Description The function CVSpilsGetReturnFlagName returns the name of the CVSPILS constant corresponding to flag.
Arguments The only argument, of type int is a return flag from a cVSPILS function.
Return value The return value is a string containing the name of the corresponding constant.

### 5.5.8 CVODES reinitialization function

The function CVodeReInit reinitializes the main CVODES solver for the solution of a problem, where a prior call to CVodeMalloc has been made. The new problem must have the same size as the previous one. CVodeReInit performs the same input checking and initializations that CVodeMalloc does, but does no memory allocation as it assumes that the existing internal memory is sufficient for the new problem.

The use of CVodeReInit requires that the maximum method order, denoted by maxord, be no larger for the new problem than for the previous problem. This condition is automatically fulfilled if the multistep method parameter 1 mm is unchanged (or changed from CV_ADAMS to CV BDF ) and the default value for maxord is specified.

If there are changes to the linear solver specifications, make the appropriate CV*Set* calls, as described in $\S 5.5 .3$


Return value The return flag flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeReInit was successful.
CV_MEM_NULL The CVODES memory block was not initialized through a previous call to CVodeCreate.
CV_NO_MALLOC Memory space for the Cvodes memory block was not allocated through a previous call to CVodeMalloc.
CV_ILL_INPUT An input argument to CVodeReInit has an illegal value.
Notes If an error occurred, CVodeReInit also sends an error message to the error handler function.

It is the user's responsibility to provide compatible itol and abstol arguments.

### 5.6 User-supplied functions

The user-supplied functions consist of one function defining the ODE, (optionally) a function that handles error and warning messages, (optionally) a function that provides the error weight vector, (optionally) a function that provides Jacobian-related information for the linear solver (if Newton iteration is chosen), and (optionally) one or two functions that define the preconditioner for use in any of the Krylov iterative algorithms.

### 5.6.1 ODE right-hand side

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

Definition typedef int (*CVRhsFn) (realtype t, N_Vector y, N_Vector ydot, void *f_data) ;
Purpose This function computes the ODE right-hand side for a given value of the independent variable $t$ and state vector $y$.

Arguments $t \quad$ is the current value of the independent variable.
$\mathrm{y} \quad$ is the current value of the dependent variable vector, $y(t)$.
ydot is the output vector $f(t, y)$.
f_data is the f_data pointer passed to CVodeSetFdata.
Return value A CVRhsFn should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverably (in which case the integration is halted and CV_RHSFUNC_FAIL is returned).
Notes Allocation of memory for ydot is handled within CVODES.
For efficiency considerations, the right-hand side function is not evaluated at the converged solution of the nonlinear solver. Therefore, a recoverable error in CVRhsFn at that point cannot be corrected (as it will occur when the right-hand side function is called the first time during the following integration step and a successful step cannot be undone). However, if the use program also includes quadrature integration, the state variables can be checked for legality in the call to CVQuadRhsFn which is called at the converged solution of the nonlinear system and therefore CVODES can be flagged to attempt to recover from such a situation. Also, if sensitivity analysis is performed with one of the staggered methods, the ODE right-hand side function is called at the converged solution of the nonlinear system and a recoverable error at that point will be captured and CVODES will try to correct it.
There are two other situations in which recovery is not possible even if the right-hand side function returns a recoverable error flag. This include the situation when this occurrs at the very first call to the CVRhsFn (in which case CVODES returns CV_FIRST_RHSFUNC_ERR) or if a recoverable error is reported when CVRhsFn is called after an error test failure, while the linear multistep method order is equal to 1 (in which case CVODES returns CV_UNREC_RHSFUNC_ERR).

### 5.6.2 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 CVSetErrFile), the user may provide a function of type CVErrHandlerFn to process any such messages. The function type CVErrHandlerFn is defined as follows:

## CVErrHandlerFn

Definition typedef void (*CVErrHandlerFn)(int error_code,
const char *module, const char *function,
char *msg, void *eh_data);

Purpose This function processes error and warning messages from CVODES and its sub-modules.
Arguments error_code is the error code.
module is the name of the CVODES module reporting the error.
function is the name of the function in which the error occurred.
$\mathrm{msg} \quad$ is the error message.
eh_data is a pointer to user data, the same as the eh_data parameter passed to CVodeSetErrHandlerFn.

Return value A CVErrHandlerFn function has no return value.
Notes error_code is negative for errors and positive (CV_WARNING) for warnings. If a function returning a pointer to memory (e.g. CVBBDPrecAlloc) encounters an error, it sets error_code to 0 before returning NULL.

### 5.6.3 Error weight function

As an alternative to providing the relative and absolute tolerances, the user may provide a function of type CVEwtFn to compute a vector ewt containing the weights in the WRMS norm $\|v\|_{\text {WRMS }}=$ $\sqrt{(1 / N) \sum_{1}^{N}\left(W_{i} \cdot v_{i}\right)^{2}}$. The function type CVEwtFn is defined as follows:

## CVEwtFn

Definition typedef int (*CVEwtFn) (N_Vector y, N_Vector dwt, void *e_data);
Purpose This function computes the WRMS error weights for the vector $y$.
Arguments y is the value of the vector for which the WRMS norm must be computed.
ewt is the output vector containing the error weights.
e_data is the e_data pointer passed to CVodeSetEwtFn.
Return value $A$ CVEwtFn function type must return 0 if it successfully set the error weights and -1 otherwise. In case of failure, a message is printed and the integration stops.
Notes Allocation of memory for ewt is handled within CVODES.
The error weight vector must have all components positive. It is the user's responsiblity to perform this test and return -1 if it is not satisfied.

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

If the direct linear solver with dense treatment of the Jacobian is used (i.e., CVDense is called in Step 7 of $\S 5.4$ ), the user may provide a function of type CVDenseJacFn defined by:


Return value $A$ CVDenseJacFn should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct, while CVDENSE sets last_flag on CVDENSE_JACFUNC_RECVR), or a negative value if it failed unrecoverable (in which case the integration is halted, CVode returns CV_LSETUP_FAIL and CVDENSE sets last_flag on CVDENSE_JACFUNC_UNRECVR).
Notes A user-supplied dense Jacobian function must load the $N$ by $N$ dense matrix $J$ with an approximation to the Jacobian matrix $J$ at the point ( $\mathrm{t}, \mathrm{y}$ ). 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 DenseMat.

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 DenseMat type. DENSE_ELEM (J, i, $j$ ) references the ( $i, j$ )-th element of the dense $\operatorname{matrix} \mathrm{J}(\mathrm{i}, \mathrm{j}=0 \ldots N-1)$. This macro is meant for small problems for which efficiency of access is not a major concern. Thus, in terms of the indices $m$ and $n$ ranging from 1 to $N$, the Jacobian element $J_{m, n}$ can be set using the statement DENSE_ELEM (J, m-1, $\mathrm{n}-1)=J_{m, n}$. Alternatively, DENSE_COL ( $\mathrm{J}, \mathrm{j}$ ) returns a pointer to the first element of the j -th column of $\mathrm{J}(\mathrm{j}=0 \ldots N-1)$, and the elements of the j -th column can then be accessed using ordinary array indexing. Consequently, $J_{m, n}$ can be loaded using 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 .

The DenseMat type and accessor macros DENSE_ELEM and DENSE_COL are documented in $\S 10.1$.

If the user's CVDenseJacFn function uses difference quotient approximations, then it may need to access quantities not in the argument list. These include the current step size, the error weights, etc. To obtain these, use the CVodeGet* functions described in $\S 5.5 .7 .1$. The unit roundoff can be accessed as UNIT_ROUNDOFF defined in sundials_types.h.

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

If the direct linear solver with banded treatment of the Jacobian is used (i.e. CVBand is called in Step 7 of $\S 5.4$ ), the user may provide a function of type CVBandJacFn defined as follows:

| CVBandJacFn |  |
| :---: | :---: |
| Definition | typedef int (*CVBandJacFn)(long int N, long int mupper, <br> long int mlower, BandMat J, realtype t, <br> N_Vector y, N_Vector fy, void *jac_data, <br> N_Vector tmp1, N_Vector tmp2, N_Vector tmp3); |
| Purpose | This function computes the banded Jacobian $J=\partial f / \partial y$ (or a banded approximation to it). |
| Arguments | $\mathrm{N} \quad$ is the problem size. |
|  | mlower |
|  | mupper are the lower and upper half-bandwidths of the Jacobian. |
|  | is the output Jacobian matrix. |
|  | is the current value of the independent variable. |
|  | is the current value of the dependent variable vector, namely the predicted value of $y(t)$. |
|  | fy is the current value of the vector $f(t, y)$. |
|  | jac_data is the jac_data pointer passed to CVBandSetJacFn. |
|  | tmp1 |
|  | tmp2 |
|  | tmp3 are pointers to memory allocated for variables of type N_Vector which can be used by CVBandJacFn as temporary storage or work space. |
| Return value | A CVBandJacFn should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct, while CVBAND sets last_flag on CVBAND_JACFUNC_RECVR), or a negative value if it failed unrecoverably (in which case the integration is halted, CVode returns CV LSETUP FAIL and CVBAND sets last_flag on CVBAND_JACFUNC_UNRECVR). |

Notes A user-supplied band Jacobian function must load the band matrix J of type BandMat with the elements of the Jacobian $J(t, y)$ at the point ( $\mathrm{t}, \mathrm{y}$ ). Only nonzero elements need to be loaded into J because J is initialized to the zero matrix 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 BandMat type. BAND_ELEM (J, i, j) references the (i, j)-th element of the band matrix J , counting from 0 . This macro is meant for use in small problems for which efficiency of access is not a major concern. Thus, in terms of the indices $m$ and $n$ ranging from 1 to $N$ with $(m, n)$ within the band defined by mupper and mlower, the Jacobian element $J_{m, n}$ can be loaded using the statement BAND_ELEM (J, $\mathrm{m}-1, \mathrm{n}-1)=J_{m, n}$. The elements within the band are those with -mupper $\leq \mathrm{m}-\mathrm{n} \leq$ mlower. Alternatively, BAND_COL ( $\mathrm{J}, \mathrm{j}$ ) returns a pointer to the diagonal element of the $j$-th column of $J$, and if we assign this address to realtype $* \operatorname{col} l_{-}$, then the i-th element of the $j$-th 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 , $\mathrm{n}-1)$; BAND_COL_ELEM (col_n, $\mathrm{m}-1, \mathrm{n}-1)=J_{m, n}$. The elements of the j -th column can also be accessed via ordinary array indexing, but this approach requires knowledge of the underlying storage for a band matrix of type BandMat. The array coln can be indexed from -mupper to mlower. For large problems, it is more efficient to use BAND_COL and BAND_COL_ELEM than to use the BAND ELEM macro. As in the dense case, these macros all number rows and columns starting from 0 .

The BandMat type and the accessor macros BAND_ELEM, BAND_COL and BAND_COL_ELEM are documented in $\S 10.2$.
If the user's CVBandJacFn function uses difference quotient approximations, then it may need to access quantities not in the argument list. These include the current step size, the error weights, etc. To obtain these, use the CVodeGet* functions described in §5.5.7.1. The unit roundoff can be accessed as UNIT_ROUNDOFF defined in sundials_types.h.

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

If one of the Krylov iterative linear solvers SPGMR, SPBCG, or SPTFQMR is selected (CVSp* is called in step 7 of $\S 5.4$ ), the user may provide a function of type CVSpilsJacTimesVecFn in the following form:


Return value The value to be returned by the Jacobian-vector product function should be 0 if successful. Any other return value will result in an unrecoverable error of the SPGMR generic solver, in which case the integration is halted.
Notes If the user's CVSpilsJacTimesVecFn function uses difference quotient approximations, it may need to access quantities not in the argument list. These include the current step size, the error weights, etc. To obtain these, use the CVodeGet* functions described in §5.5.7.1. The unit roundoff can be accessed as UNIT_ROUNDOFF defined in sundials_types.h.

### 5.6.7 Preconditioning (linear system solution)

If preconditioning is used, then the user must provide a C function to solve the linear system $P z=$ $r$, where $P$ may be either a left or right preconditioner matrix. This function must be of type CVSpilsPrecSolveFn, defined as follows:

```
CVSpilsPrecSolveFn
```

Definition typedef int (*CVSpilsPrecSolveFn) (realtype t, N_Vector y, N_Vector fy, N_Vector $r, N_{-}$Vector $z$, realtype gamma, realtype delta, int lr, void *p_data, N_Vector tmp);

Purpose This function solves the preconditioned system $P z=r$.
Arguments $t \quad$ is the current value of the independent variable.
$y \quad$ is the current value of the dependent variable vector.
fy is the current value of the vector $f(t, y)$.
$r \quad$ is the right-hand side vector of the linear system.
$z \quad$ is the computed output vector.
gamma is the scalar $\gamma$ appearing in the Newton matrix given by $M=I-\gamma J$.
delta is an input tolerance to be used if an iterative method is employed in the solution. In that case, the residual vector Res $=r-P z$ of the system should be made less than delta in the weighted $l_{2}$ norm, i.e., $\sqrt{\sum_{i}\left(R e s_{i} \cdot e w t_{i}\right)^{2}}<$ delta. To obtain the N_Vector ewt call CVodeGetErrWeights (see §5.5.7.1).
$\operatorname{lr} \quad$ is an input flag indicating whether the preconditioner solve function is to use the left preconditioner $(\operatorname{lr}=1)$ or the right preconditioner $(\operatorname{lr}=2)$;
p_data is the p_data pointer passed to CVSp*SetPreconditioner.
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 (in which case the step will be retried), or negative for an unrecoverable error (in which case the integration is halted).

### 5.6.8 Preconditioning (Jacobian data)

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

Definition typedef int (*CVSpilsPrecSetupFn) (realtype t, N_Vector y, N_Vector fy, booleantype jok, booleantype *jcurPtr, realtype gamma, void *p_data, N_Vector tmp1, N_Vector tmp2, N_Vector tmp3);
Purpose This function preprocesses and/or evaluates Jacobian-related data needed by the preconditioner.
Arguments The arguments of a CVSpilsPrecSetupFn are as follows:
$t \quad$ is the current value of the independent variable.
y is the current value of the dependent variable vector, namely the predicted value of $y(t)$.
fy $\quad$ is the current value of the vector $f(t, y)$.
jok is an input flag indicating whether the Jacobian-related data needs to be updated. The jok argument provides for the reuse of Jacobian data in the preconditioner solve function. jok = FALSE means that the Jacobian-related data must be recomputed from scratch. jok = TRUE means that the Jacobian data, if saved from the previous call to this function, can be reused (with the current value of gamma). A call with jok = TRUE can only occur after a call with jok = FALSE.
jcurPtr is a pointer to a flag which should be set to TRUE if Jacobian data was recomputed, or set to FALSE if Jacobian data was not recomputed, but saved data was still reused.
gamma is the scalar $\gamma$ appearing in the Newton matrix $M=I-\gamma P$.
p_data is the p_data pointer passed to CVSp*SetPreconditioner.
tmp1
tmp2
tmp3 are pointers to memory allocated for variables of type N_Vector which can be used by CVSpilsPrecSetupFn as temporary storage or work space.
Return value The value to be returned by the preconditioner setup function is a flag indicating whether it was successful. This value should be 0 if successful, positive for a recoverable error (in which case the step will be retried), or negative for an unrecoverable error (in which case the integration is halted).
Notes The operations performed by this function might include forming a crude approximate Jacobian, and performing an LU factorization of the resulting approximation to $M=$ $I-\gamma J$.
Each call to the preconditioner setup function is preceded by a call to the CVRhsFn user function with the same ( $t, y$ ) arguments. Thus, the preconditioner setup function can use any auxiliary data that is computed and saved during the evaluation of the ODE right-hand side.
This function is not called in advance of every call to the preconditioner solve function, but rather is called only as often as needed to achieve convergence in the Newton iteration.
If the user's CVSpilsPrecSetupFn function uses difference quotient approximations, it may need to access quantities not in the call list. These include the current step size, the error weights, etc. To obtain these, use the CVodeGet* functions described in §5.5.7.1. The unit roundoff can be accessed as UNIT_ROUNDOFF defined in sundials_types.h.

### 5.7 Integration of pure quadrature equations

If the system of ODEs contains pure quadratures, it is more efficient to treat them separately by excluding them from the nonlinear solution stage. To do this, begin by excluding the quadrature
variables from the vector $y$ and the quadrature equations from within $f$. The following is an overview of the sequence of calls in a user's main program in this situation. Steps that are unchanged from the skeleton program presented in $\S 5.4$ are grayed out.

1. [P] Initialize MPI

## 2. Set problem dimensions

$[\mathbf{S}]$ Set N to the problem size $N$ (excluding quadrature variables), and Nq to the number of quadrature variables.
$[\mathbf{P}]$ Set Nlocal to the local vector length (excluding quadrature variables), and Nqlocal to the local number of quadrature variables.
3. Set vector of initial values
4. Create CVODES object
5. Allocate internal memory
6. Set optional inputs
7. Attach linear solver module
8. Set linear solver optional inputs

## 9. Set vector of initial values for quadrature variables

Typically, the quadrature variables should be initialized to 0 .

## 10. Initialize quadrature integration

Call CVodeQuadMalloc to specify the quadrature equation right-hand side function and to allocate internal memory related to quadrature integration. See §5.7.1 for details.

## 11. Set optional inputs for quadrature integration

Call CVodeSetQuadFdata to specify user data required for the evaluation of the quadrature equation right-hand side. Call CVodeSetQuadErrCon to indicate whether or not quadrature variables shoule be used in the step size control mechanism, and to specify the integration tolerances for quadrature variables. See §5.7.4 for details.
12. Advance solution in time

## 13. Extract quadrature variables

Call CVodeGetQuad to obtain the values of the quadrature variables at the current time. See §5.7.3 for details.
14. Get optional outputs

## 15. Get quadrature optional outputs

Call CVodeGetQuad* functions to obtain optional output related to the integration of quadratures. See §5.7.5 for details.
16. Deallocate memory for solution vector and for the vector of quadrature variables
17. Free solver memory
18. [P] Finalize MPI

CVodeQuadMalloc can be called and quadrature-related optional inputs (step 11 above) can be set, anywhere between steps 4 and 12 .

### 5.7.1 Quadrature initialization functions

The function CVodeQuadMalloc activates integration of quadrature equations and allocates internal memory related to these calculations. The form of the call to this function is as follows:

## CVodeQuadMalloc

Call flag = CVodeQuadMalloc (cvode_mem, fQ, yQO);
Description The function CVodeQuadMalloc provides required problem specifications, allocates internal memory, and initializes quadrature integration.
Arguments cvode_mem (void *) pointer to the CVODES memory block returned by CVodeCreate.
fQ (CVQuadRhsFn) is the C function which computes $f_{Q}$, the right-hand side of the quadrature equations. This function has the form $f Q(t, y, y Q d o t$, fQ_data) (for full details see §5.7.6).
yQ0 (N_Vector) is the initial value of $y_{Q}$.
Return value The return value flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeQuadMalloc was successful.
CV_MEM_NULL The CVODES memory was not initialized by a prior call to CVodeCreate. CV_MEM_FAIL A memory allocation request failed.
Notes If an error occured, CVodeQuadMalloc also sends an error message to the error handler function.
In terms of the number of quadrature variables $N_{q}$ and maximum method order maxord, the size of the real workspace is increased by:

- Base value: lenrw $=$ lenrw $+(\operatorname{maxord}+5) N_{q}$
- With itolQ = CV_SV (see CVodeSetQuadErrCon): lenrw $=$ lenrw $+N_{q}$
the size of the integer workspace is increased by:
- Base value: leniw $=$ leniw $+(\operatorname{maxord}+5) N_{q}$
- With itolQ $=$ CV_SV: leniw $=$ leniw $+N_{q}$

The function CVodeQuadReInit, useful during the solution of a sequence of problems of same size, reinitializes the quadrature related internal memory and must follow a call to CVodeQuadMalloc (and maybe a call to CVodeReInit). The number Nq of quadratures is assumed to be unchanged from the prior call to CVodeQuadMalloc. The call to the CVodeQuadReInit function has the form:

| CVodeQuadReInit |
| :--- |
| Call flag $=$ CVodeQuadReInit (cvode_mem, fQ, yQ0); |

Description The function CVodeQuadReInit provides required problem specifications and reinitializes the quadrature integration.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
$\mathrm{fQ} \quad$ (CVQuadRhsFn) is the C function which computes $f_{Q}$, the right-hand side of the quadrature equations.
yQO (N_Vector) is the initial value of $y_{Q}$.
Return value The return value flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeReInit was successful.
CV_MEM_NULL The CVODES memory was not initialized by a prior call to CVodeCreate.
CV_NO_QUAD Memory space for the quadrature integration was not allocated by a prior call to CVodeQuadMalloc.
Notes If an error occured, CVodeQuadReInit also sends an error message to the error handler function.

### 5.7.2 CVODE solver function

Even if quadrature integration was enabled, the call to the main solver function CVode is exactly the same as in §5.5.4. However, in this case the return value flag can also be one of the following:
CV_QRHSFUNC_FAIL The quadrature right-hand side function failed in an unrecoverable manner.
CV_FIRST_QRHSFUNC_FAIL The quadrature right-hand side function failed at the first call.
CV_REPTD_QRHSFUNC_ERR Convergence tests occurred too many times due to repeated recoverable errors in the quadrature right-hand side function. The CV_REPTD_RHSFUNC_ERR will also be returned if the quadrature right-hand side function had repeated recoverable errors during the estimation of an initial step size (assuming the quadrature variables are included in the error tests).
CV_UNREC_RHSFUNC_ERR The quadrature right-hand function had a recoverable error, but no recovery was possible. This failure mode is rare, as it can occur only if the quadrature right-hand side function fails recoverably after an error test failed while at order one.

### 5.7.3 Quadrature extraction functions

If quadrature integration has been initialized by a call to CVodeQuadMalloc, or reinitialized by a call to CVodeQuadReInit, then CVODES computes both a solution and quadratures at time t . However, CVode will still return only the solution $y$ in $y$. Solution quadratures can be obtained using the following function:

| CVodeGetQuad |
| :--- | :--- |
| Call flag $=$ CVodeGetQuad (cvode_mem, $t, y Q) ; ~$ |

Description The function CVodeGetQuad returns the quadrature solution vector after a successful return from CVode.

Arguments cvode_mem (void $*$ ) pointer to the memory previously allocated by CVodeMalloc.
t (realtype) the time at which quadrature information is requested. The time $t$ must fall within the interval defined by the last successful step taken by cVodes.
yQ (N_Vector) the computed quadrature vector.
Return value The return value flag of CVodeGetQuad is one of:
CV_SUCCESS CVodeGetQuad was successful.
CV_MEM_NULL cvode_mem was NULL.
CV_NO_QUAD Quadrature integration was not initialized.
CV_BAD_DKY yQ is NULL.
CV_BAD_T The time $t$ is not in the allowed range.
Notes In case of an error return, an error message is also sent to the error handler function.
The function CVodeGetQuadDky computes the k -th derivatives of the interpolating polynomials for the quadrature variables at time t . This function is called by CVodeGetQuad with $\mathrm{k}=0$, but may also be called directly by the user.
CVodeGetQuadDky
Call flag $=$ CVodeGetQuadDky (cvode_mem, t, k, dkyQ);

Description The function CVodeGetQuadDky returns derivatives of the quadrature solution vector after a successful return from CVode.

Arguments cvode_mem (void *) pointer to the memory previously allocated by CVodeMalloc.
t (realtype) the time at which quadrature information is requested. The time $t$ must fall within the interval defined by the last successful step taken by CVODES.
$\mathrm{k} \quad$ (int) order of the requested derivative.
dkyQ
(N_Vector) the vector containing the derivative. This vector must be allocated by the user.
Return value The return value flag of CVodeGetQuadDky is one of:
CV_SUCCESS CVodeGetQuadDky succeeded.
CV_MEM_NULL The pointer to cvode_mem was NULL.
CV_NO_QUAD Quadrature integration was not initialized.
CV_BAD_DKY The vector dkyQ is NULL.
CV_BAD_K $\quad \mathrm{k}$ is not in the range $0,1, \ldots, q_{u}$.
CV_BAD_T The time $t$ is not in the allowed range.
Notes In case of an error return, an error message is also sent to the error handler function.

### 5.7.4 Optional inputs for quadrature integration

CVODES provides the following optional input functions to control the integration of quadrature equations.

| CVodeSetQuadFdata |  |
| :---: | :---: |
| Call f | flag = CVodeSetQuadFdata(cvode_mem, fQ_data); |
| Description $\begin{aligned} & \text { T } \\ & \\ & \\ & \text { a }\end{aligned}$ | The function CVodeSetQuadFdata specifies the user-defined data block fQ_data and attaches it to the main CVODES memory block. |
| Arguments ${ }^{\text {c }}$ | cvode mem (void *) pointer to the CVODES memory block. fQ_data (void *) pointer to the user data. |
| Return value | The return value flag (of type int) is one of: CV_SUCCESS The optional value has been successfully set. CV_MEM_NULL The cvode_mem pointer is NULL. |
| Notes $\begin{array}{ll}\text { If } \\ & \text { h } \\ & \text { th }\end{array}$ | If $f Q \_$data is not specified, a NULL pointer is passed to all user-supplied functions that have it as an argument. Note that fQ _data can be the same as the pointer $f$ _data set through CVodeSetFdata. |
| CVodeSetQuadErrCon |  |
| Call f | flag = CVodeSetQuadErrCon(cvode_mem, errconQ, itolQ, reltolQ, abstolQ); |
| Description $\begin{array}{ll}\text { T } \\ & \text { sh } \\ & \text { to }\end{array}$ | The function CVodeSetQuadErrCon specifies whether or not the quadrature variables should be used in the step size control mechanism, and if so, specifies the integration tolerances for the quadrature variables. |
| Arguments | cvode mem (void *) pointer to the CVODES memory block. <br> errconQ (booleantype) specifies whether quadrature variables are included (TRUE) or not (FALSE) in the error control mechanism. If errconQ $=$ FALSE, the following three arguments are ignored. |
|  | itolQ (int) is either CV_SS or CV_SV, where itolQ = CV_SS indicates scalar relative error tolerance and scalar absolute error tolerance, while itolQ $=$ CV_SV indicates scalar relative error tolerance and vector absolute error tolerance. The latter choice is important when the absolute error tolerance needs to be different for each quadrature variable. <br> reltolQ (realtype $*$ ) is a pointer to the relative error tolerance. |

abstolQ (void *) is a pointer to the absolute error tolerance. If itolQ=CV_SS, abstolQ must be a pointer to a realtype variable. If itolQ = CV_SV, abstolQ must be an N_Vector variable.

Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_ILL_INPUT An input argument to CVodeSetQuadErrCon has an illegal value.
Notes By default, errconQ is set to FALSE.
It is illegal to call CVodeSetQuadErrCon before a call to CVodeQuadMalloc.

### 5.7.5 Optional outputs for quadrature integration

CVODES provides the following functions that can be used to obtain solver performance information related to quadrature integration.

## CVodeGetQuadNumRhsEvals

Call flag = CVodeGetQuadNumRhsEvals(cvode_mem, \&nfQevals);
Description The function CVodeGetQuadNumRhsEvals returns the number of calls made to the user's quadrature right-hand side function.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
nfQevals (long int) number of calls made to the user's $f Q$ function.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_QUAD Quadrature integration has not been initialized.

## CVodeGetQuadNumErrTestFails

Call flag = CVodeGetQuadNumErrTestFails(cvode_mem, \&nQetfails);
Description The function CVodeGetQuadNumErrTestFails returns the number of local error test failures due to quadrature variables.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
nQetfails (long int) number of error test failures due to quadrature variables.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_QUAD Quadrature integration has not been initialized.

CVodeGetQuadErrWeights
Call flag = CVodeGetQuadErrWeights (cvode_mem, eQweight);
Description The function CVodeGetQuadErrWeights returns the quadrature error weights at the current time.

Arguments cvode_mem (void *) pointer to the CVODES memory block.
eQweight (N_Vector) quadrature error weights at the current time.
Return value The return value flag (of type int) is one of:

CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_QUAD Quadrature integration has not been initialized.
Notes The user must allocate memory for eQweight.
If quadratures were not included in the error control mechanism (through a call to CVodeSetQuadErrCon with errconQ = TRUE), CVodeGetQuadErrWeights does not set the eQweight vector.

```
CVodeGetQuadStats
Call flag = CVodeGetQuadStats(cvode_mem, &nfQevals, &nQetfails);
```

Description The function CVodeGetQuadStats returns the CVODES integrator statistics as a group.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
nfQevals (long int) number of calls to the user's fR function.
nQetfails (long int) number of error test failures due to quadrature variables.
Return value The return value flag (of type int) is one of
CV_SUCCESS the optional output values have been successfully set.
CV_MEM_NULL the cvode_mem pointer is NULL.
CV_NO_QUAD Quadrature integration has not been initialized.

### 5.7.6 User-supplied function for quadrature integration

For integration of quadrature equations, the user must provide a function that defines the right-hand side of the quadrature equations. This function must be of type CVQuadRhsFn defined as follows:

## CVQuadRhsFn

Definition typedef int (*CVQuadRhsFn)(realtype $t$, N_Vector y, N_Vector yQdot, void *fQ_data);
Purpose This function computes the quadrature equation right-hand side for a given value of the independent variable $t$ and state vector $y$.

Arguments $t \quad$ is the current value of the independent variable.
$\mathrm{y} \quad$ is the current value of the dependent variable vector, $y(t)$.
yQdot is the output vector $f_{Q}(t, y)$.
$f Q \_$data is the $f Q \_$data pointer passed to CVodeSetQuadFdata.
Return value A CVQuadRhsFn should return 0 if successful, a positive value if a recoverable error occored (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverable (in which case the integration is halted and CV_QRHSFUNC_FAIL is returned).
Notes Allocation of memory for yQdot is automatically handled within CVODES.
Both y and yQdot are of type N_Vector, but they typically have different internal representations. It is the user's responsibility to access the vector data consistently (including the use of the correct accessor macros from each NVECTOR implementation). For the sake of computational efficiency, the vector functions in the two NVECTOR implementations provided with CVODES do not perform any consistency checks with respect to their N_Vector arguments (see $\S 8.1$ and $\S 8.2$ ).
There are two situations in which recovery is not possible even if CVQuadRhsFn function returns a recoverable error flag. This include the situation when this occurs at the very first call to the CVQuadRhsFn (in which case CVODES returns CV_FIRST_QRHSFUNC_ERR)
or if a recoverable error is reported when CVQuadRhsFn is called after an error test failure, while the linear multistep method order is equal to 1 (in which case CVODES returns CV_UNREC_QRHSFUNC_ERR).

### 5.8 Rootfinding

While solving the IVP, CVODES has the capability to find the roots of a set of user-defined functions. This section describes the user-callable functions used to initialize and define the rootfinding problem and to obtain solution information, and it also describes the required user-supplied function.

### 5.8.1 User-callable functions for rootfinding

```
CVodeRootInit
Call flag = CVodeRootInit(cvode_mem, nrtfn, g, g_data);
```

Description The function CVodeRootInit specifies that the roots of a set of functions $g_{i}(t, y)$ are to be found while the IVP is being solved.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block returned by CVodeCreate. nrtfn (int) is the number of root functions $g_{i}$.
$\mathrm{g} \quad$ (CVRootFn) is the C function which defines the nrtfn functions $g_{i}(t, y)$ whose roots are sought. See $\S 5.8 .2$ for details.
g_data (void $*)$ pointer to the user data for use by the user's root function $g$.
Return value The return value flag (of type int) is one of
CV_SUCCESS The call to CVodeRootInit was successful.
CV_MEM_NULL The cvode_mem argument was NULL.
CV_MEM_FAIL A memory allocation failed.
CV_ILL_INPUT The function g is NULL, but nrtfn>0.
Notes If a new IVP is to be solved with a call to CVodeReInit, where the new IVP has no rootfinding problem but the prior one did, then call CVodeRootInit with nrtfn $=0$.
There are two optional output functions associated with rootfinding.

## CVodeGetRootInfo

```
Call flag = CVodeGetRootInfo(cvode_mem, rootsfound);
```

Description The function CVodeGetRootInfo returns an array showing which functions were found to have a root.

Arguments cvode_mem (void *) pointer to the CVODES memory block.
rootsfound (int *) an int array of length nrtfn showing the indices of the user functions $g_{i}$ found to have a root. For $i=0, \ldots, n r t f n-1$, rootsfound $[i]=1$ if $g_{i}$ has a root, and 0 if not.

Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output values have been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes The user must allocate memory for the vector rootsfound.

```
CVodeGetNumGEvals
Call flag = CVodeGetNumGEvals(cvode_mem, &ngevals);
```

Description The function CVodeGetNumGEvals returns the cumulative number of calls made to the user-supplied root function $g$.
Arguments cvodemem (void $*$ ) pointer to the CVODES memory block.
ngevals (long int) number of calls made to the user's function $g$ thus far.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.

### 5.8.2 User-supplied function for rootfinding

If a rootfinding problem is to be solved during the integration of the ODE system, the user must supply a C function of type CVRootFn, defined as follows:

## CVRootFn

Definition typedef int (*CVRootFn)(realtype $t, N \_V e c t o r y, ~ r e a l t y p e ~ * g o u t, ~$ void *g_data);
Purpose This function implements a vector-valued function $g(t, y)$ such that the roots of the nrtfn components $g_{i}(t, y)$ are sought.
Arguments $t \quad$ is the current value of the independent variable. $\mathrm{y} \quad$ is the current value of the dependent variable vector, $y(t)$. gout is the output array, of length nrtfn, with components $g_{i}(t, y)$. g_data is the g_data pointer passed to CVodeRootInit.
Return value A CVRootFn should return 0 if successful or a non-zero value if an error occured (in which case the integration is halted and CVode returns CV_RTFUNC_FAIL).
Notes Allocation of memory for gout is automatically handled within CVODES.

### 5.9 Preconditioner modules

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, problemspecific preconditioner, CVODES provides a banded preconditioner in the module CVBANDPRE and a band-block-diagonal preconditioner module CVBBDPRE.

### 5.9.1 A serial banded preconditioner module

This preconditioner provides a band matrix preconditioner for use with any of the Krylov iterative linear solvers, in a serial setting. It uses difference quotients of the ODE right-hand side function $f$ to generate a band matrix of bandwidth $m_{l}+m_{u}+1$, where the number of super-diagonals ( $m_{u}$, the upper half-bandwidth) and sub-diagonals ( $m_{l}$, the lower half-bandwidth) are specified by the user, and uses this to form a preconditioner for use with the Krylov linear solver. Although this matrix is intended to approximate the Jacobian $\partial f / \partial y$, it may be a very crude approximation. The true Jacobian need not be banded, or its true bandwidth may be larger than $m_{l}+m_{u}+1$, as long as the banded approximation generated here is sufficiently accurate to speed convergence as a preconditioner.

In order to use the CVBANDPRE module, the user need not define any additional functions. Aside from the header files required for the integration of the ODE problem (see $\S 5.3$ ), to use the CVBANDPRE module, the main program must include the header file cvodes_bandpre.h which declares the needed function prototypes. The following is a summary of the usage of this module. Steps that are unchanged from the skeleton program presented in $\S 5.4$ are grayed out.

1. Set problem dimensions
2. Set vector of initial values

## 3. Create CVODES object

4. Allocate internal memory
5. Set optional inputs

## 6. Initialize the CVBANDPRE preconditioner module

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

```
bp_data = CVBandPrecAlloc(cvode_mem, N, mu, ml);
```

to allocate memory for and to initialize a data structure (pointed to by bp_data) to be passed to the appropriate CVSp* linear solver.

## 7. Attach the Krylov linear solver, one of:

```
flag = CVBPSpgmr(cvode_mem, pretype, maxl, bp_data);
```

flag = CVBPSpbcg(cvode_mem, pretype, maxl, bp_data);
flag = CVBPSptfqmr (cvode_mem, pretype, maxl, bp_data);

Each function CVBPSp* is a wrapper around the corresponding specification function CVSp* and performs the following actions:

- Attaches the CVSPILS linear solver to the main CVODES solver memory;
- Sets the preconditioner data structure for CVBANDPRE;
- Sets the preconditioner setup function for CVBANDPRE;
- Sets the preconditioner solve function for CVBANDPRE;

The arguments pretype and maxl are described below. The last argument of CVBPSp* is the pointer to the CVBANDPRE data returned by CVBandPrecAlloc.

## 8. Set linear solver optional inputs

Note that the user should not overwrite the preconditioner data, setup function, or solve function through calls to CVSp* optional input functions.
9. Advance solution in time
10. Deallocate memory for solution vector

## 11. Free the CVbandpre data structure

CVBandPrecFree(\&bp_data);
12. Free solver memory

The user-callable functions that initialize, attach, and deallocate the CVBANDPRE preconditioner module (steps 6, 7 and 11 above) are described in more detail below.

## CVBandPrecAlloc

Call bp_data $=$ CVBandPrecAlloc (cvode_mem, $N$, mu, ml);
Description The function CVBandPrecAlloc initializes and allocates memory for the CVBANDPRE preconditioner.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
$\mathrm{N} \quad$ (long int) problem dimension.
mu (long int) upper half-bandwidth of the Jacobian approximation.
ml (long int) lower half-bandwidth of the Jacobian approximation.
Return value If successful, CVBandPrecAlloc returns a pointer to the newly created CVBANDPRE memory block (of type void *). If an error occurred, CVBandPrecAlloc returns NULL.
Notes The banded approximate Jacobian will have nonzero elements only in locations ( $i, j$ ) with $-\mathrm{ml} \leq j-i \leq \mathrm{mu}$.

## CVBPSpgmr

Call
flag = CVBPSpgmr (cvode_mem, pretype, maxl, bp_data);
Description The function CVBPSpgmr links the CVBANDPRE data to the CVSPGMR linear solver and attaches the latter to the CVODES memory block.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
pretype (int) specifies the preconditioning type and must be either PREC LEFT or PREC_RIGHT.
$\operatorname{maxl}$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
bp_data (void $*$ ) pointer to the CVBANDPRE data structure.
Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The CVSPGMR initialization was successful.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
CVBANDPRE_PDATA_NULL The CVBANDPRE preconditioner has not been initialized.

CVBPSpbcg
Call flag = CVBPSpbcg(cvode_mem, pretype, maxl, bp_data);
Description The function CVBPSpbcg links the CVBANDPRE data to the CVSPBCG linear solver and attaches the latter to the CVODES memory block.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. pretype (int) preconditioning type. Must be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxl} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
bp_data (void $*$ ) pointer to the CVBANDPRE data structure.
Return value The return value flag (of type int) is one of CVSPILS_SUCCESS The CVSPBCG initialization was successful. CVSPILS_MEM_NULL The cvode_mem pointer is NULL. CVSPILS_ILL_INPUT The preconditioner type pretype is not valid. CVSPILS_MEM_FAIL A memory allocation request failed. CVBANDPRE_PDATA_NULL The CVBANDPRE preconditioner has not been initialized.

## CVBPSptfqmr

Call flag = CVBPSptfqmr (cvode_mem, pretype, maxl, bp_data);
Description The function CVBPSptfqmr links the CVBANDPRE data to the CVSPTFQMR linear solver and attaches the latter to the CVODES memory block.

| Arguments | cvode_mem (void $*)$ pointer to the CVODES memory block. |
| :--- | :--- | :--- |
| pretype | (int) preconditioning type. Must be one of PREC_LEFT or PREC_RIGHT. |
| maxl | (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use |
|  | the default value CVSPILS_MAXL $=5$. |

Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The CVSPTFQMR initialization was successful.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
CVBANDPRE_PDATA_NULL The CVBANDPRE preconditioner has not been initialized.

## CVBandPrecFree

Call CVBandPrecFree(\&bp_data);
Description The function CVBandPrecFree frees the pointer allocated by CVBandPrecAlloc.
Arguments The only argument passed to CVBandPrecFree is the pointer to the CVBANDPRE data structure (of type void *).
Return value The function CVBandPrecFree has no return value.
The following three optional output functions are available for use with the CVBANDPRE module:

## CVBandPrecGetWorkSpace

Call flag = CVBandPrecGetWorkSpace(bp_data, \&lenrwBP, \&leniwBP);
Description The function CVBandPrecGetWorkSpace returns the sizes of the CVBANDPRE real and integer workspaces.
Arguments bp_data (void *) pointer to the CVBANDPRE data structure.
lenrwBP (long int) the number of realtype values in the CVBANDPRE workspace.
leniwBP (long int) the number of integer values in the CVBANDPRE workspace.
Return value The return value flag (of type int) is one of:
CVBANDPRE_SUCCESS The optional output values have been successfully set.
CVBANDPRE_PDATA_NULL The CVBANDPRE preconditioner has not been initialized.
Notes In terms of problem size $N$ and $s m u=\min (N-1, m u+m l)$, the actual size of the real workspace is $(2 \mathrm{ml}+\mathrm{mu}+\mathrm{smu}+2) N$ realtype words, and the actual size of the integer workspace is $N$ integer words.
The workspaces referred to here exist in addition to those given by the corresponding CVSp***GetWorkSpace function.

## CVBandPrecGetNumRhsEvals

Call flag = CVBandPrecGetNumRhsEvals(bp_data, \&nfevalsBP);
Description The function CVBandPrecGetNumRhsEvals returns the number of calls made to the usersupplied right-hand side function for finite difference banded Jacobian approximation used within the preconditioner setup function.
Arguments bp_data (void $*$ ) pointer to the CVBANDPRE data structure.
nfevalsBP (long int) the number of calls to the user right-hand side function.
Return value The return value flag (of type int) is one of:
CVBANDPRE_SUCCESS The optional output value has been successfully set.

CVBANDPRE_PDATA_NULL The CVBANDPRE preconditioner has not been initialized.
Notes The counter nfevalsBP is distinct from the counter nfevalsLS returned by the corresponding CVSp***GetNumRhsEvals function, and also from nfevals, returned by CVodeGetNumRhsEvals. The total number of right-hand side function evaluations is the sum of all three of these counters.
CVBandPrecGetReturnFlagName
Call name $=$ CVBandPrecGetReturnFlagName (flag);

Description The function CVBandPrecGetReturnFlagName returns the name of the CVBANDPRE constant corresponding to flag.
Arguments The only argument, of type int is a return flag from a CVBANDPRE function.
Return value The return value is a string containing the name of the corresponding constant.

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

A principal reason for using a parallel ODE solver such as CVODES lies in the solution of partial differential equations (PDEs). Moreover, the use of a Krylov iterative method for the solution of many such problems is motivated by the nature of the underlying linear system of equations (3.5) that must be solved at each time step. The linear algebraic system is large, sparse and structured. However, if a Krylov iterative method is to be effective in this setting, then a nontrivial preconditioner needs to be used. Otherwise, the rate of convergence of the Krylov iterative method is usually unacceptably slow. Unfortunately, an effective preconditioner tends to be problem-specific.

However, we have developed one type of preconditioner that treats a rather broad class of PDEbased problems. It has been successfully used for several realistic, large-scale problems [20] and is included in a software module within the CVODES package. This module works with the parallel vector module NVECTOR_PARALLEL and is usable with any of the Krylov iterative linear solvers. It generates a preconditioner that is a block-diagonal matrix with each block being a band matrix. The blocks need not have the same number of super- and sub-diagonals and these numbers may vary from block to block. This Band-Block-Diagonal Preconditioner module is called CVbbdpre.

One way to envision these preconditioners is to think of the domain of the computational PDE problem as being subdivided into $M$ non-overlapping subdomains. Each of these subdomains is then assigned to one of the $M$ processes to be used to solve the ODE system. The basic idea is to isolate the preconditioning so that it is local to each process, and also to use a (possibly cheaper) approximate right-hand side function. This requires the definition of a new function $g(t, y)$ which approximates the function $f(t, y)$ in the definition of the ODE system (3.1). However, the user may set $g=f$. Corresponding to the domain decomposition, there is a decomposition of the solution vector $y$ into $M$ disjoint blocks $y_{m}$, and a decomposition of $g$ into blocks $g_{m}$. The block $g_{m}$ depends both on $y_{m}$ and on components of blocks $y_{m^{\prime}}$ associated with neighboring subdomains (so-called ghost-cell data). Let $\bar{y}_{m}$ denote $y_{m}$ augmented with those other components on which $g_{m}$ depends. Then we have

$$
\begin{equation*}
g(t, y)=\left[g_{1}\left(t, \bar{y}_{1}\right), g_{2}\left(t, \bar{y}_{2}\right), \ldots, g_{M}\left(t, \bar{y}_{M}\right)\right]^{T} \tag{5.1}
\end{equation*}
$$

and each of the blocks $g_{m}\left(t, \bar{y}_{m}\right)$ is uncoupled from the others.
The preconditioner associated with this decomposition has the form

$$
\begin{equation*}
P=\operatorname{diag}\left[P_{1}, P_{2}, \ldots, P_{M}\right] \tag{5.2}
\end{equation*}
$$

where

$$
\begin{equation*}
P_{m} \approx I-\gamma J_{m} \tag{5.3}
\end{equation*}
$$

and $J_{m}$ is a difference quotient approximation to $\partial g_{m} / \partial y_{m}$. This matrix is taken to be banded, with upper and lower half-bandwidths mudq and mldq defined as the number of non-zero diagonals above and below the main diagonal, respectively. The difference quotient approximation is computed using
mudq $+\mathrm{mldq}+2$ evaluations of $g_{m}$, but only a matrix of bandwidth mu $+\mathrm{ml}+1$ 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. The solution of the complete linear system

$$
\begin{equation*}
P x=b \tag{5.4}
\end{equation*}
$$

reduces to solving each of the equations

$$
\begin{equation*}
P_{m} x_{m}=b_{m} \tag{5.5}
\end{equation*}
$$

and this is done by banded LU factorization of $P_{m}$ followed by a banded backsolve.
Similar block-diagonal preconditioners could be considered with different treatments of the blocks $P_{m}$. For example, incomplete LU factorization or an iterative method could be used instead of banded LU factorization.

The CVBBDPRE module calls two user-provided functions to construct $P$ : a required function gloc (of type CVLocalFn) which approximates the right-hand side function $g(t, y) \approx f(t, y)$ and which is computed locally, and an optional function cfn (of type CVCommFn) which performs all interprocess communication necessary to evaluate the approximate right-hand side $g$. These are in addition to the user-supplied right-hand side function $f$. Both functions take as input the same pointer f_data that is passed by the user to CVodeSetFdata and that was passed to the user's function $f$, and neither function has a return value. The user is responsible for providing space (presumably within f_data) for components of $y$ that are communicated between processes by cfn, and that are then used by gloc, which is not expected to do any communication.

## CVLocalFn

Definition typedef int (*CVLocalFn) (long int Nlocal, realtype t, N_Vector y, N_Vector glocal, void *f_data);
Purpose This function computes $g(t, y)$. It loads the vector glocal as a function of t and y .
Arguments Nlocal is the local vector length.
$\mathrm{t} \quad$ is the value of the independent variable.
$\mathrm{y} \quad$ is the dependent variable.
glocal is the output vector.
f_data is the f_data pointer passed to CVodeSetFdata.
Return value A CVLocalFn should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverably (in which case the integration is halted and CVode returns CV LSETUP FAIL).
Notes This function assumes that all interprocess communication of data needed to calculate glocal has already been done, and that this data is accessible within f data.
The case where $g$ is mathematically identical to $f$ is allowed.

## CVCommFn

Definition typedef int (*CVCommFn) (long int Nlocal, realtype t, N_Vector y, void *f_data);

Purpose This function performs all interprocess communication necessary for the execution of the gloc function above, using the input vector y .
Arguments Nlocal is the local vector length.
$\mathrm{t} \quad$ is the value of the independent variable.
$\mathrm{y} \quad$ is the dependent variable.
f_data is the f_data pointer passed to CVodeSetFdata.
Return value $A$ CVCommFn should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverably (in which case the integration is halted and CVode returns CV_LSETUP_FAIL).

Notes The cfn function is expected to save communicated data in space defined within the data structure f_data.
Each call to the cfn function is preceded by a call to the right-hand side function $f$ with the same ( $t, y$ ) arguments. Thus, cfn can omit any communication done by $f$ if relevant to the evaluation of glocal. If all necessary comunication was done in $f$, then cfn $=$ NULL can be passed in the call to CVBBDPrecAlloc (see below).
Besides the header files required for the integration of the ODE problem (see §5.3), to use the CVBBDPRE module, the main program must include the header file cvodes_bbdpre.h which declares the needed function prototypes.

The following is a summary of the proper usage of this module. Steps that are unchanged from the skeleton program presented in $\S 5.4$ are grayed out.

1. Initialize MPI
2. Set problem dimensions
3. Set vector of initial values
4. Create CVODES object
5. Allocate internal memory
6. Set optional inputs

## 7. Initialize the CVBBDPRE preconditioner module

Specify the upper and lower half-bandwidths mudq and mldq, and mukeep and mlkeep, and call

```
bbd_data = CVBBDPrecAlloc(cvode_mem, local_N, mudq, mldq,
    mukeep, mlkeep, dqrely, gloc, cfn);
```

to allocate memory for and to initialize a data structure bbd_data (of type void *) to be passed to the Krylov linear solver selected (in the next step). The last two arguments passed to CVBBDPrecAlloc are the two user-supplied functions described above.

## 8. Attach the Krylov linear solver, one of:

```
flag = CVBBDSpgmr(cvode_mem, pretype, maxl, bbd_data);
flag = CVBBDSpbcg(cvode_mem, pretype, maxl, bbd_data);
flag = CVBBDSptfqmr(cvode_mem, pretype, maxl, bbd_data);
```

The function CVBPSp* is a wrapper around the corresponding specification function CVSp* and performs the following actions:

- Attaches the CVSPILS linear solver to the main CVODES solver memory;
-Sets the preconditioner data structure for CVBBDPRE;
- Sets the preconditioner setup function for CVBBDPRE;
- Sets the preconditioner solve function for CVBBDPRE;

The arguments pretype and maxl are described below. The last argument of CVBBDSp* is the pointer to the CVBBDPRE data returned by CVBBDPrecAlloc.
9. Set linear solver optional inputs

Note that the user should not overwrite the preconditioner data, setup function, or solve function through calls to CVSPILS optional input functions.
10. Advance solution in time
11. Deallocate memory for solution vector

## 12. Free the cVBBDPRE data structure

CVBBDPrecFree(\&bbd_data);
13. Free solver memory

## 14. Finalize MPI

The user-callable functions that initialize, attach, and deallocate the CVBBDPRE preconditioner module (steps 7, 8, and 12 above) are described next.


Description The function CVBBDPrecAlloc initializes and allocates memory for the CVBBDPRE preconditioner.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
local_N (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.
dqrely (realtype) the relative increment in components of y used in the difference quotient approximations. The default is dqrely $=\sqrt{\text { unit roundoff, which }}$ can be specified by passing dqrely $=0.0$.
gloc (CVLocalFn) the C function which computes the approximation $g(t, y) \approx$ $f(t, y)$.
cfn (CVCommFn) the optional C function which performs all interprocess communication required for the computation of $g(t, y)$.

Return value If successful, CVBBDPrecAlloc returns a pointer to the newly created CVBBDPRE memory block (of type void *). If an error occurred, CVBBDPrecAlloc returns NULL.
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 local_N-1, it is replaced with 0 or local_N-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 a greater efficiency.
Also, the half-bandwidths mukeep and mlkeep of the retained banded approximate Jacobian block may be even smaller, to reduce storage and computational costs further.

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

## CVBBDSpgmr

Call flag = CVBBDSpgmr (cvode_mem, pretype, maxl, bbd_data);
Description The function CVBBDSpgmr links the CVBBDPRE data to the CVSPGMR linear solver and attaches the latter to the CVODES memory block.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
pretype (int) preconditioning type. Must be either PREC_LEFT or PREC_RIGHT.
$\operatorname{maxl} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
bbd_data (void *) pointer to the CVBBDPRE data structure.
Return value The return value flag (of type int) is one of:

| CVSPILS_SUCCESS | The CVSPGMR initialization was successful. |
| :--- | :--- |
| CVSPILS_MEM_NULL | The cvode_mem pointer is NULL. |
| CVSPILS_ILL_INPUT | The preconditioner type pretype is not valid. |
| CVSPILS_MEM_FAIL | A memory allocation request failed. |
| CVBBDPRE_PDATA_NULL | The CVBBDPRE preconditioner has not been initialized. |

CVBBDSpbcg
Call
flag $=$ CVBBDSpbcg(cvode_mem, pretype, maxl, bbd_data);
Description The function CVBBDSpbcg links the CVBBDPRE data to the CVSPBCG linear solver and attaches the latter to the CVODES memory block.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
pretype (int) preconditioning type. Must be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxl} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
bbd_data (void $*$ ) pointer to the CVBBDPRE data structure.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The CVSPBCG initialization was successful.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
CVBBDPRE_PDATA_NULL The CVBBDPRE preconditioner has not been initialized.

## CVBBDSptfqmr

Call flag = CVBBDSptfqmr (cvode_mem, pretype, maxl, bbd_data);
Description The function CVBBDSptfqmr links the CVBBDPRE data to the CVSPTFQMR linear solver and attaches the latter to the CVODES memory block.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
pretype (int) preconditioning type. Must be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxl} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
bbd_data (void *) pointer to the CVBBDPRE data structure.
Return value The return value flag (of type int) is one of
CVSPILS_SUCCESS The CVSPTFQMR initialization was successful.
CVSPILS_MEM_NULL The cvode_mem pointer is NULL.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
CVBBDPRE_PDATA_NULL The CVBBDPRE preconditioner has not been initialized.

CVBBDPrecFree
Call CVBBDPrecFree(\&bbd_data);
Description The function CVBBDPrecFree frees the memory allocated by CVBBDPrecAlloc.
Arguments The only argument passed to CVBBDPrecFree is the address of the pointer to the CVBBDPRE data structure (of type void *).

Return value The function CVBBDPrecFree has no return value.
The cVbBDPRE module also provides a reinitialization function to allow solving a sequence of problems of the same size, with the same linear solver choice, provided there is no change in local $N$, mukeep, or mlkeep. After solving one problem, and after calling CVodeReInit to re-initialize CVODES for a subsequent problem, a call to CVBBDPrecReInit can be made to change any of the following: the half-bandwidths mudq and mldq used in the difference-quotient Jacobian approximations, the relative increment dqrely, or one of the user-supplied functions gloc and cfn. If there is a change in any of the linear solver inputs, an additional call to CVSpgmr, CVSpbcg, or CVSptfqmr, and/or one or more of the corresponding CVSp***Set*** functions, must also be made.

## CVBBDPrecReInit

Call flag = CVBBDPrecReInit(bbd_data, mudq, mldq, dqrely, gloc, cfn);
Description The function CVBBDPrecReInit re-initializes the CVBBDPRE preconditioner.
Arguments bbd_data (void *) pointer to the CVBBDPRE data structure.
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.
dqrely (realtype) the relative increment in components of y used in the difference quotient approximations. The default is dqrely $=\sqrt{\text { unit roundoff }}$, which can be specified by passing dqrely $=0.0$.
gloc (CVLocalFn) the C function which computes the approximation $g(t, y) \approx$ $f(t, y)$.
cfn (CVCommFn) the optional C function which performs all interprocess communication required for the computation of $g(t, y)$.
Return value The return value flag (of type int) is one of
CVBBDPRRE_SUCCESS The CVSPBCG re-initialization was successful.
CVBBDPRE_PDATA_NULL The CVBBDPRE preconditioner has not been initialized.
Notes If one of the half-bandwidths mudq or mldq is negative or exceeds the value local_N-1, it is replaced with 0 or local_N-1 accordingly.
The following two optional output functions are available for use with the CVBBDPRE module:

CVBBDPrecGetWorkSpace
Call flag = CVBBDPrecGetWorkSpace(bbd_data, \&lenrwBBDP, \&leniwBBDP);
Description The function CVBBDPrecGetWorkSpace returns the local CVBBDPRE real and integer workspace sizes.

Arguments bbd_data (void *) pointer to the CVBBDPRE data structure.
lenrwBBDP (long int) local number of realtype values in the CVBBDPRE workspace.
leniwBBDP (long int) local number of integer values in the CVBBDPRE workspace.
Return value The return value flag (of type int) is one of:
CVBBDPRE_SUCCESS The optional output values have been successfully set.
CVBBDPRE_PDATA_NULL The CVBBDPRE preconditioner has not been initialized.

Notes In terms of local_N and smu $=\min ($ local $N-1$, mukeep + mlkeep $)$, the actual size of the real workspace is ( $2 \mathrm{mlkeep}+$ mukeep $+\mathrm{smu}+2$ ) local_N realtype words, and the actual size of the integer workspace is local $N$ integer words. These values are local to each process.
The workspaces referred to here exist in addition to those given by the corresponding CVSp***GetWorkSpace function.

## CVBBDPrecGetNumGfnEvals

Call flag = CVBBDPrecGetNumGfnEvals(bbd_data, \&ngevalsBBDP);
Description The function CVBBDPrecGetNumGfnEvals returns the number of calls made to the usersupplied gloc function due to the finite difference approximation of the Jacobian blocks used within the preconditioner setup function.
Arguments bbd_data (void $*$ ) pointer to the CVBBDPRE data structure.
ngevalsBBDP (long int) the number of calls made to the user-supplied gloc function.
Return value The return value flag (of type int) is one of
CVBBDPRE_SUCCESS The optional output value has been successfully set.
CVBBDPRE_PDATA_NULL The CVBBDPRE preconditioner has not been initialized.

## CVBBDPrecGetReturnFlagName

Call name = CVBBDPrecGetReturnFlagName(flag);
Description The function CVBBDPrecGetReturnFlagName returns the name of the CVBBDPRE constant corresponding to flag.
Arguments The only argument, of type int is a return flag from a CVBBDPRE function.
Return value The return value is a string containing the name of the corresponding constant.
In addition to the ngevalsBBDP gloc evaluations, the costs associated with CVBBDPRE also include nlinsetups LU factorizations, nlinsetups calls to cfn , npsolves banded backsolve calls, and nfevalsLS right-hand side function evaluations, where nlinsetups is an optional CVODES output and npsolves and nfevalsLS are linear solver optional outputs (see §5.5.7).

## Chapter 6

## Using CVODES for Forward Sensitivity Analysis

This chapter describes the use of CVODES to compute solution sensitivities using forward sensitivity analysis. One of our main guiding principles was to design the CVODES user interface for forward sensitivity analysis as an extension of that for IVP integration. Assuming a user main program and user-defined support routines for IVP integration have already been defined, in order to perform forward sensitivity analysis the user only has to insert a few more calls into the main program and (optionally) define an additional routine which computes the right-hand side of the sensitivity systems (3.9). The only departure from this philosophy is due to the CVRhsFn type definition (§5.6). Without changing the definition of this type, the only way to pass values of the problem parameters to the ODE right-hand side function is to require the user data structure f_data to contain a pointer to the array of real parameters $p$.

CVODES uses various constants for both input and output. These are defined as needed in this chapter, but for convenience are also listed separately in Chapter 11.

We begin with a brief overview, in the form of a skeleton user program. Following that are detailed descriptions of the interface to the various user-callable routines and of the user-supplied routines that were not already described in $\S 5$.

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

The following is a skeleton of the user's main program (or calling program) as an application of CVODES. The user program is to have these steps in the order indicated, unless otherwise noted. For the sake of brevity, we defer many of the details to the later sections. As in $\S 5.4$, most steps are independent of the NVECTOR implementation used; where this is not the case, usage specifications are given for the two implementations provided with CVODES: steps marked with $[\mathbf{P}]$ correspond to NVECTOR_PARALLEL, while steps marked with [S] correspond to NVECTOR_SERIAL. Differences between the user main program in $\S 5.4$ and the one below start only at step (9).

First, note that no additional header files need be included for forward sensitivity analysis beyond those for IVP solution (§5.4).

1. [P] Initialize MPI
2. Set problem dimensions
3. Set initial values
4. Create CVODES object
5. Allocate internal memory

## 6. Set optional inputs

7. Attach linear solver module

## 8. Set linear solver optional inputs

## 9. Define the sensitivity problem

- Number of sensitivities (required)

Set Ns, the number of parameters with respect to which sensitivities are to be computed.

- Problem parameters (optional)

If CVODES will evaluate the right-hand sides of the sensitivity systems, set p , an array of Np real parameters upon which the IVP depends. Only parameters with respect to which sensitivities are (potentially) desired need to be included. Attach p to the user data structure f_data. For example, f_data->p = p;
If the user provides a function to evaluate the sensitivity right-hand side, $p$ need not be specified.

- Parameter list (optional)

If CVODES will evaluate the right-hand sides of the sensitivity systems, set plist, an array of Ns integer flags to specify the parameters p with respect to which solution sensitivities are to be computed. If sensitivities with respect to the $j$-th problem parameter are desired, set plist $_{i}=j$, for some $i=0, \ldots, N_{s}-1$.
If plist is not specified, CVODES will compute sensitivities with respect to the first Ns parameters; i.e., plist $_{i}=i, i=0, \ldots, N_{s}-1$.
If the user provides a function to evaluate the sensitivity right-hand side, plist need not be specified.

- Parameter scaling factors (optional)

If CVODES estimates tolerances for the sensitivity solution vectors (based on tolerances for the state solution vector) or if CVODES will evaluate the right-hand sides of the sensitivity systems using the internal difference-quotient function, the results will be more accurate if order of magnitude information is provided.
Set pbar, an array of Ns positive scaling factors. Typically, if $p_{i} \neq 0$, the value $\bar{p}_{\text {plist }_{i}}=\left|p_{i}\right|$ can be used.
If pbar is not specified, CVODES will use $\bar{p}_{i}=1.0$.
If the user provides a function to evaluate the sensitivity right-hand side and specifies tolerances for the sensitivity variables, pbar need not be specified.

Note that the names for p , pbar, plist, as well as the field $p$ of f data are arbitrary, but they must agree with the arguments passed to CVodeSetSensParams below.

## 10. Set sensitivity initial conditions

Set the Ns vectors ySO[i] of N initial values for sensitivities (for $i=0, \ldots, N s-1$ ).
First, create an array of Ns vectors by making the call
[S] ySO = N_VNewVectorArray_Serial(Ns, N);
[P] ySO = N_VNewVectorArray_Parallel(Ns, N);
and, for each $i=1, \cdots, \mathrm{Ns}$, load initial values for the $i$-th sensitivity vector into the structure defined by:
[S] NV_DATA_S (ySO [i])
[P] NV_DATA_P (ySO [i])

If the initial values for the sensitivity variables are already available in realtype arrays, create an array of Ns "empty" vectors by making the call
$[\mathbf{S}]$ ySO = N_VNewVectorArrayEmpty_Serial(Ns, N);
$[\mathbf{P}]$ ySO $=$ N_VNewVectorArrayEmpty_Parallel(Ns, N);
and then attach the realtype array ySO_i containing the initial values of the $i$-th sensitivity vector using
[S] N_VSetArrayPointer_Serial(ySO_i, ySO[i]);
[P] N_VSetArrayPointer_Parallel(ySO_i, ySO[i]);

## 11. Activate sensitivity calculations

Call flag $=$ CVodeSensMalloc(...); to activate forward sensitivity computations and allocate internal memory for CVODES related to sensitivity calculations (see §6.2.1).

## 12. Set sensitivity analysis optional inputs

Call CVodeSetSens* routines to change from their default values any optional inputs that control the behavior of CVODES in computing forward sensitivities.
13. Advance solution in time

## 14. Extract sensitivity solution

After each successful return from CVode, the solution of the original IVP is available in the y argument of CVode, while the sensitivity solution can be extracted into yS (which can be the same as ySO) by calling the routine flag = CVodeGetSens (cvode_mem, t, yS); (see §6.2.3).
15. Deallocate memory for solution vector

## 16. Deallocate memory for sensitivity vectors

Upon completion of the integration, deallocate memory for the vectors ySO:
[S] N_VDestroyVectorArray_Serial(yS0, Ns);
[P] N_VDestroyVectorArray_Parallel(ySO, Ns);
If yS was created from realtype arrays yS_i, it is the user's responsibility to also free the space for the arrays yS0_i.
17. Free user data structure
18. Free solver memory
19. Free vector specification memory

### 6.2 User-callable routines for forward sensitivity analysis

This section describes the CVODES functions, additional to those presented in $\S 5.5$, that are called by the user to setup and solve a forward sensitivity problem.

### 6.2.1 Forward sensitivity initialization and deallocation functions

Activation of forward sensitivity computation is done by calling CVodeSensMalloc. The form of the call to this routine is as follows:

```
CVodeSensMalloc
Call flag = CVodeSensMalloc(cvode_mem, Ns, ism, ySO);
```

Description The routine CVodeSensMalloc activates forward sensitivity computations and allocates internal memory related to sensitivity calculations.

Arguments cvode_mem (void *) pointer to the CVODES memory block returned by CVodeCreate. Ns (int) the number of sensitivities to be computed.
ism (int) a flag used to select the sensitivity solution method and can be CV_SIMULTANEOUS, CV_STAGGERED, or CV_STAGGERED1:

- In the CV_SIMULTANEOUS approach, the state and sensitivity variables are corrected at the same time. If CV_NEWTON was selected as the nonlinear system solution method, this amounts to performing a modified Newton iteration on the combined nonlinear system;
- In the CV_STAGGERED approach, the correction step for the sensitivity variables takes place at the same time for all sensitivity equations, but only after the correction of the state variables has converged and the state variables have passed the local error test;
- In the CV_STAGGERED1 approach, all corrections are done sequentially, first for the state variables and then for the sensitivity variables, one parameter at a time. If the sensitivity variables are not included in the error control, this approach is equivalent to CV_STAGGERED. Note that the CV_STAGGERED1 approach can be used only if the user-provided sensitivity right-hand side function is of type CVSensRhs1Fn (see $\S 6.3$ ).
ySO (N_Vector *) a pointer to an array of Ns vectors containing the initial values of the sensitivities.

Return value The return value flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeSensMalloc was successful.
CV_MEM_NULL The CVODES memory block was not initialized through a previous call to CVodeCreate.
CV_MEM_FAIL A memory allocation request has failed.
CV_ILL_INPUT An input argument to CVodeSensMalloc has an illegal value.
Notes If an error occured, CVodeSensMalloc also prints an error message to the file specified by the optional input errfp.
In terms of the problem size $N$, number of sensitivity vectors $N_{s}$, and maximum method order maxord, the size of the real workspace is increased by:

- Base value: lenrw $=$ lenrw $+(\operatorname{maxord}+5) N_{s} N$
- With itolS $=$ CV_SV (see CVodeSetSensTolerances): lenrw $=$ lenrw $+N_{s} N$
the size of the integer workspace is increased by:
- Base value: leniw $=$ leniw $+(\operatorname{maxord}+5) N_{s} N$
- With itolS $=$ CV_SV: leniw $=$ leniw $+N_{s} N$

The routine CVodeSensReInit, useful during the solution of a sequence of problems of same size, reinitializes the sensitivity-related internal memory and must follow a call to CVodeSensMalloc (and maybe a call to CVodeReInit). The number Ns of sensitivities is assumed to be unchanged since the call to CVodeSensMalloc. The call to the CVodeSensReInit function has the form:

## CVodeSensReInit

Call flag = CVodeSensReInit(cvode_mem, ism, ySO);
Description The routine CVodeSensReInit reinitializes forward sensitivity computations.
Arguments cvode_mem (void *) pointer to the CVODES memory block returned by CVodeCreate.
ism (int) a flag used to select the sensitivity solution method and can be CV_SIMULTANEOUS, CV_STAGGERED, or CV_STAGGERED1.
ySO (N_Vector *) a pointer to an array of Ns variables of type N_Vector containing the initial values of the sensitivities.

Return value The return value flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeReInit was successful.
CV_MEM_NULL The CVODES memory block was not initialized through a previous call to CVodeCreate.

CV_NO_SENS Memory space for sensitivity integration was not allocated through a previous call to CVodeSensMalloc.
CV_ILL_INPUT An input argument to CVodeSensReInit has an illegal value.
CV_MEM_FAIL A memory allocation request has failed.
Notes All arguments of CVodeSensReInit are the same as those of CVodeSensMalloc.
If an error occured, CVodeSensReInit also prints an error message to the file specified by the optional input errfp.
To deallocate all forward sensitivity-related memory (allocated in a prior call to CVodeSensMalloc), the user must call

CVodeSensFree
Call CVodeSensFree(cvode_mem);
Description The function CVodeSensFree frees the memory allocated for forward sensitivity computaions by a previous call to CVodeSensMalloc.

Arguments The argument is the pointer to the CVODES memory block (of type void *).
Return value The function CVodeSensFree has no return value.
Notes After a call to CVodeSensFree, forward sensitivity computations can be reactivated only by calling again CVodeSensMalloc.
To activate and deactivate forward sensitivity calculations for successive CVODES runs, without having to allocate and deallocate memory, the following function is provided:

## CVodeSensToggleOff

Call CVodeSensToggleOff (cvode_mem);
Description The function CVodeSensToggleOff deactivates forward sensitivity calculations. It does not deallocate sensitivity-related memory.

Arguments cvode_mem (void *) pointer to the memory previously allocated by CVodeMalloc.
Return value The return value flag of CVodeSensToggle is one of:
CV_SUCCESS CVodeSensToggleOff was successful.
CV_MEM_NULL cvode_mem was NULL.
Notes Since sensitivity-related memory is not deallocated, sensitivities can be reactivated at a later time (using CVodeSensReInit).

### 6.2.2 CVODE solver function

Even if forward sensitivity analysis was enabled, the call to the main solver function CVode is exactly the same as in $\S 5.5 .4$. However, in this case the return value flag can also be one of the following:
CV_SRHSFUNC_FAIL The sensitivity right-hand side function failed in an unrecoverable manner.
CV_FIRST_SRHSFUNC_FAIL The sensitivity right-hand side function failed at the first call.
CV_REPTD_SRHSFUNC_ERR Convergence tests occurred too many times due to repeated recoverable errors in the sensitivity right-hand side function. The CV_REPTD_RHSFUNC_ERR will also be returned if the sensitivity right-hand side function had repeated recoverable errors during the estimation of an initial step size.

CV_UNREC_SRHSFUNC_ERR The sensitivity right-hand function had a recoverable error, but no recovery was possible. This failure mode is rare, as it can occur only if the sensitivity right-hand side function fails recoverably after an error test failed while at order one.

### 6.2.3 Forward sensitivity extraction functions

If forward sensitivity computations have been initialized by a call to CVodeSensMalloc, or reinitialized by a call to CVSensReInit, then CVODES computes both a solution and sensitivities at time t. However, CVode will still return only the solution $y$ in $y$. Solution sensitivities can be obtained through one of the following functions:
CVodeGetSens
Call flag $=$ CVodeGetSens (cvode_mem, t, yS);

Description The function CVodeGetSens returns the sensitivity solution vectors after a successful return from CVode.
Arguments cvode_mem (void $*$ ) pointer to the memory previously allocated by CVodeMalloc.
t (realtype) specifies the time at which sensitivity information is requested. The time $t$ must fall within the interval defined by the last successful step taken by CVODES.
yS (N_Vector *) the computed forward sensitivity vectors.
Return value The return value flag of CVodeGetSens is one of:
CV_SUCCESS CVodeGetSens was successful.
CV_MEM_NULL cvode_mem was NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
CV_BAD_DKY yQ is NULL.
CV_BAD_T The time $t$ is not in the allowed range.
Notes In case of an error return, an error message is also printed.
The function CVodeGetSensDky computes the k-th derivatives of the interpolating polynomials for the sensitivity variables at time $t$. This function is called by CVodeGetSens with $k=0$, but may also be called directly by the user.
CVodeGetSensDky
Call flag $=$ CVodeGetSensDky (cvode_mem, t, k, dkyS);

Description The function CVodeGetSensDky returns derivatives of the sensitivity solution vectors after a successful return from CVode.

Arguments cvode_mem (void $*$ ) pointer to the memory previously allocated by CVodeMalloc.
t (realtype) specifies the time at which sensitivity information is requested. The time $t$ must fall within the interval defined by the last successful step taken by CVODES.
$\mathrm{k} \quad$ (int) order of derivatives.
dkyS
(N_Vector *) the vectors containing the derivatives. The space for dkys must be allocated by the user.
Return value The return value flag of CVodeGetSensDky is one of:
CV_SUCCESS CVodeGetSensDky succeeded.
CV_MEM_NULL The pointer to cvode_mem was NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
CV_BAD_DKY One of the vectors dkyS is NULL.
CV_BAD_K $\quad \mathrm{k}$ is not in the range $0,1, \ldots, q_{u}$.
CV_BAD_T The time $t$ is not in the allowed range.
Notes In case of an error return, an error message is also printed.
Forward sensitivity solution vectors can also be extracted separately for each parameter in turn through the functions CVodeGetSens1 and CVodeGetSensDky1, defined as follows:

```
CVodeGetSens1
Call flag = CVodeGetSens1(cvode_mem, t, is, yS);
```

Description The function CVodeGetSens1 returns the is-th sensitivity solution vector after a successful return from CVode.
Arguments cvode_mem (void $*$ ) pointer to the memory previously allocated by CVodeMalloc.
t (realtype) specifies the time at which sensitivity information is requested. The time $t$ must fall within the interval defined by the last successful step taken by CVODES.
is (int) specifies which sensitivity vector is to be returned ( $0 \leq$ is $<N_{s}$ ).
yS (N_Vector) the computed forward sensitivity vector.
Return value The return value flag of CVodeGetSens1 is one of:
CV_SUCCESS CVodeGetSens1 was successful.
CV_MEM_NULL cvode_mem was NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
CV_BAD_IS The index is is not in the allowed range.
CV_BAD_DKY yQ is NULL.
CV_BAD_T The time $t$ is not in the allowed range.
Notes In case of an error return, an error message is also printed.
CVodeGetSensDky1
Call flag $=$ CVodeGetSensDky1 (cvode_mem, t, k, is, dkyS);

Description The function CVodeGetSensDky1 returns the k-th derivative of the is-th sensitivity solution vector after a successful return from CVode.
Arguments cvode_mem (void $*$ ) pointer to the memory previously allocated by CVodeMalloc.
$\mathrm{t} \quad$ (realtype) specifies the time at which sensitivity information is requested. The time $t$ must fall within the interval defined by the last successful step taken by CVODES.
$\mathrm{k} \quad$ (int) order of derivative.
is
(int) specifies the sensitivity derivative vector to be returned ( $0 \leq$ is $<N_{s}$ ).

Table 6.1: Forward sensitivity optional inputs

| Optional input | Routine name | Default |
| :--- | :--- | :--- |
| Sensitivity right-hand side fct. and data | CVodeSetSensRhsFn | internal DQ |
| Sensitivity right-hand side fct. and data | CVodeSetSensRhs1Fn | internal DQ |
| Sensitivity scaling factors | CVodeSetSensPbar | NULL |
| DQ approximation method | CVodeSetSensDQMethod | 0.0 |
| Error control strategy | CVodeSetSensErrCon | FALSE |
| Sensitivity integration tolerances | CVodeSetSensTolerances | estimated |
| Maximum no. of nonlinear iterations | CVodeSetSensMaxNonlinIters | 3 |

dkyS (N_Vector) the vector containing the derivative. The space for dkyS must be allocated by the user.
Return value The return value flag of CVodeGetSensDky1 is one of:
CV_SUCCESS CVodeGetQuadDky1 succeeded.
CV_MEM_NULL The pointer to cvode_mem was NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
CV_BAD_DKY One of the vectors dkyS is NULL.
CV_BAD_IS The index is is not in the allowed range.
CV_BAD_K $\quad \mathrm{k}$ is not in the range $0,1, \ldots, q_{u}$.
CV_BAD_T The time $t$ is not in the allowed range.
Notes In case of an error return, an error message is also printed.

### 6.2.4 Optional inputs for forward sensitivity analysis

Optional input variables that control the computation of sensitivities can be changed from their default values through calls to CVodeSetSens* functions. Table 6.1 lists all forward sensitivity optional input functions in CVODES which are described in detail in the remainder of this section.

```
CVodeSetSensRhsFn
Call flag = CVodeSetSensRhsFn(cvode_mem, fS, data);
```

Description The function CVodeSetSensRhsFn specifies the user-supplied C function used to evaluate the sensitivity right-hand sides (for all parameters at once) and he user data block for use by the user-supplied sensitivity right-hand side function.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
fS (CVSensRhsFn) user-defined sensitivity right-hand side function.
fS_data (void *) pointer to the user data.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes This type of function is not compatible with the CV_STAGGERED1 sensitivity solution method (argument ism to CVodeSensMalloc). The compatibility test is performed at the first step in CVode.
Passing $\mathrm{fS}=$ NULL indicates using the default internal difference quotient sensitivity righthand side routine.

The pointer fS_data can be the same as the pointer f_data, specified in a prior call to CVodeSetFdata (see §5.5.5.1) and passed to the user's right-hand side function $f$.

```
CVodeSetSensRhs1Fn
Call flag = CVodeSetSensRhs1Fn(cvode_mem, fS, data);
```

Description The function CVodeSetSensRhs1Fn specifies the user-supplied C function used to evaluate the sensitivity right-hand sides (one parameter at a time) and he user data block for use by the user-supplied sensitivity right-hand side function.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
fS (CVSensRhs1Fn) user-defined sensitivity right-hand side function.
fS_data (void *) pointer to the user data.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes This type of sensitivity right-hand side function must be used when the CV STAGGERED1 sensitivity solution method is selected through CVodeSensMalloc, but can also be used with the other two choices (CV_SIMULTANEOUS and CV_STAGGERED).

Passing $\mathrm{fS}=$ NULL indicates using the default internal difference quotient sensitivity righthand side routine.
The pointer fS_data can be the same as the pointer f_data, specified in a prior call to CVodeSetFdata (see §5.5.5.1) and passed to the user's right-hand side function $f$.

## CVodeSetSensParams

Call flag = CVodeSetSensParams(cvode_mem, p, pbar, plist);
Description The function CVodeSetSensParams specifies problem parameter information for sensitivity calculations.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
$\mathrm{p} \quad($ realtype $*)$ a pointer to the array of real problem parameters used to evaluate $f(t, y, p)$. If non-NULL, p must point to a field in the user's data structure $f \_d a t a$ passed to the righ-hand side function. (See §6.1).
pbar (realtype *) an array of Ns positive scaling factors. If non-NULL, pbar must have all its components $>0.0$. (See $\S 6.1$ ).
plist (int $*)$ an array of Ns non-negative flags to specify which parameters to use in estimating the sensitivity equations. If non-NULL, plist must have all components $\geq 0$. (See §6.1).
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
CV_ILL_INPUT An argument has an illegal value.
Notes This function must be preceeded by a call to CVodeSensMalloc.

CVodeSetSensDQMethod
Call flag = CVodeSetSensDQMethod(cvode_mem, DQtype, DQrhomax);
Description The function CVodeSetSensDQMethod specifies the difference quotient strategy in the case in which the right-hand side of the sensitivity equations are to be computed by CVODES.
Arguments cvode_mem (void *) pointer to the CVODES memory block.

DQtype (int) specifies the difference quotient type and can be one of CV_CENTERED or CV_FORWARD.
DQrhomax (realtype) positive value of the selection parameter used in deciding switching between a simultaneous or separate approximation of the two terms in the sensitivity right-hand side.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_ILL_INPUT An argument has an illegal value.
Notes If DQrhomax $=0.0$, then no switching is performed. The approximation is done simultaneously using either centered or forward finite differences, depending on the value of DQtype. For values of DQrhomax $\geq 1.0$ the simultaneous approximation is used whenever the estimated finite difference perturbations for states and parameters are within a factor of DQrhomax and the separate approximation is used otherwise. Note that a value $\operatorname{DQrhomax}<1.0$ will effectively disable switching. See $\S 3.2$ for more details.
The default value are DQtype=CV_CENTERED and DQrhomax $=0.0$.
CVodeSetSensErrCon
Call flag $=$ CVodeSetSensErrCon(cvode_mem, errconS);

Description The function CVodeSetSensErrCon specifies the error control strategy for sensitivity variables.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
errcons (booleantype) specifies whether sensitivity variables are included (TRUE) or not (FALSE) in the error control mechanism.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes By default, errconS is set to FALSE. If errconS=TRUE then both state variables and sensitivity variables are included in the error tests. If errconS $=$ FALSE then the sensitivity variables are excluded from the error tests. Note that, in any event, all variables are considered in the convergence tests.

## CVodeSetSensTolerances

Call flag = CVodeSetSensTolerances (cvode_mem, itolS, reltolS, abstolS);
Description The function CVodeSetSensTolerances specifies the integration tolerances for sensitivity variables.
Arguments cvodemem (void *) pointer to the CVODES memory block.
itolS (int) is one of CV_SS, CV_SV, or CV_EE, where itolS $=$ CV_SS indicates scalar relative error tolerance and scalar absolute error tolerance, while itolS $=$ CV_SV indicates scalar relative error tolerance and vector absolute error tolerance. If itolS $=$ CV_EE, the arguments reltolS and abstolS are ignored and CVODES will estimate tolerances for the sensitivity variables based on the state tolerances and the scaling factors $\bar{p}$.
reltolS (realtype) is the relative error tolerance.
abstolS (void $*)$ is a pointer to the absolute error tolerance. If itolS $=C V \_S S$, then abstolS must be a pointer to an array of realtype variables. If itolS $=$ CV_SV, then abstolS must be an array of Ns variables of type N_Vector. In the latter case, abstolS should be created and set in the same manner as the vectors of initial values for the sensitivity variables (see $\S 6.1$ ).

Table 6.2: Forward sensitivity optional outputs

| Optional output | Routine name |
| :--- | :--- |
| No. of calls to sensitivity r.h.s. function | CVodeGetNumSensRhsEvals |
| No. of calls to r.h.s. function for sensitivity | CVodeGetNumRhsEvalsSens |
| No. of sensitivity local error test failures | CVodeGetNumSensErrTestFails |
| No. of calls to lin. solv. setup routine for sens. | CVodeGetNumSensLinSolvSetups |
| Error weight vector for sensitivity variables | CVodeGetSensErrWeights |
| No. of sens. nonlinear solver iterations | CVodeGetNumSensNonlinSolvIters |
| No. of sens. convergence failures | CVodeGetNumSensNonlinSolvConvFails |
| No. of staggered nonlinear solver iterations | CVodeGetNumStgrSensNonlinSolvIters |
| No. of staggered convergence failures | CVodeGetNumStgrSensNonlinSolvConvFails |

Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional values have been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
CV_ILL_INPUT An input argument to CVodeSetSensTolerances has an illegal value.
Notes The default behavior is for CVODES to estimate appropriate integration tolerances for the sensitivity variables based on the state tolerances and the scaling factors $\bar{p}$. See $\S 3.2$ for details.
It is the user's responsibility to provide compatible itolS and abstolS arguments.
This function must be preceeded by a call to CVodeSensMalloc.
CVodeSetSensMaxNonlinIters
Call flag = CVodeSetSensMaxNonlinIters(cvode_mem, maxcors);

Description The function CVodeSetSensMaxNonlinIters specifies the maximum number of nonlinear solver iterations for sensitivity variables per step.

Arguments cvode_mem (void *) pointer to the CVODES memory block.
maxcors (int) maximum number of nonlinear solver iterations allowed per step.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
Notes The default value is 3 .

### 6.2.5 Optional outputs for forward sensitivity analysis

Optional output functions that return statistics and solver performance information related to forward sensitivity computations are listed in Table 6.2 and described in detail in the remainder of this section.

## CVodeGetNumSensRhsEvals

Call flag = CVodeGetNumSensRhsEvals (cvode_mem, \&nfSevals);
Description The function CVodeGetNumSensRhsEvals returns the number of calls to the sensitivity right-hand side function.

Arguments cvode_mem (void *) pointer to the CVODES memory block.
nfSevals (long int) number of calls to the sensitivity right-hand side function.

Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
Notes In order to accommodate any of the three possible sensitivity solution methods, the default internal finite difference quotient functions evaluate the sensitivity right-hand sides one at a time. Therefore, nfSevals will always be a multiple of the number of sensitivity parameters (the same as the case in which the user supplies a routine of type CVSensRhs1Fn).

```
CVodeGetNumRhsEvalsSens
Call flag = CVodeGetNumRhsEvalsSens(cvode_mem, &nfevalsS);
```

Description The function CVodeGetNumRhsEvalsSEns returns the number of calls to the user's righthand side function due to the internal finite difference approximation of the sensitivity right-hand sides.
Arguments cvode_mem (void *) pointer to the CVODES memory block.
nfevalsS (long int) number of calls to the user right-hand side function.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
Notes This counter is incremented only if the internal finite difference approximation routines are used for the evaluation of the sensitivity right-hand sides.
CVodeGetNumSensErrTestFails

Call flag = CVodeGetNumSensErrTestFails(cvode_mem, \&nSetfails);
Description The function CVodeGetNumSensErrTestFails returns the number of local error test failures for the sensitivity variables that have occured.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. nSetfails (long int) number of error test failures.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
Notes This counter is incremented only if the sensitivity variables have been included in the error test (see CVodeSetSensErrCon in §6.2.4). Even in that case, this counter is not incremented if the ism=CV_SIMULTANEOUS sensitivity solution method has been used.

CVodeGetNumSensLinSolvSetups
Call flag = CVodeGetNumSensLinSolvSetups (cvode mem, \&nlinsetupsS);
Description The function CVodeGetNumSensLinSolvSetups returns the number of calls to the linear solver setup function due to forward sensitivity calculations.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. nlinsetupsS (long int) number of calls to the linear solver setup function.
Return value The return value flag (of type int) is one of:

CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
Notes This counter is incremented only if Newton iteration has been used and if either the ism=CV_STAGGERED or the ism=CV_STAGGERED1 sensitivity solution method has been specified in the call to CVodeSensMalloc (see §6.2.1).

## CVodeGetSensStats

Call flag = CVodeGetSensStats(cvode_mem, \&nfSevals, \&nfevalsS, \&nSetfails, \&nlinsetupsS);
Description The function CVodeGetSensStats returns all of the above sensitivity-related solver statistics as a group.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
nfSevals (long int) number of calls to the sensitivity right-hand side function.
nfevalsS (long int) number of calls to the user-supplied right-hand side function.
nSetfails (long int) number of error test failures.
nlinsetupss (long int) number of calls to the linear solver setup function.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output values have been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.

## CVodeGetSensErrWeights

Call flag = CVodeGetSensErrWeights (cvode_mem, eSweight);
Description The function CVodeGetSensErrWeights returns the sensitivity error weights at the current time. These are the reciprocals of the $W_{i}$ of (3.7) for the sensitivity variables.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
eSweight (N_Vector_S) pointer to the array of error weight vectors.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
Notes The user must allocate memory for eweightS.

## CVodeGetNumSensNonlinSolvIters

Call flag = CVodeGetNumSensNonlinSolvIters(cvodemem, \&nSniters);
Description The function CVodeGetNumSensNonlinSolvIters returns the number of nonlinear iterations performed for sensitivity calculations.

Arguments cvode mem (void *) pointer to the CVODES memory block. nSniters (long int) number of nonlinear iterations performed.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.

Notes This counter is incremented only if the ism was CV_STAGGERED or CV_STAGGERED1 in the call to CVodeSensMalloc (see §6.2.1).
In the CV_STAGGERED1 case, the value of nSniters is the sum of the number of nonlinear iterations performed for each sensitivity equation. These individual counters can be obtained through a call to CVodeGetNumStgrSensNonlinSolvIters (see below).

## CVodeGetNumSensNonlinSolvConvFails

Call flag = CVodeGetNumSensNonlinSolvConvFails(cvode_mem, \&nSncfails);
Description The function CVodeGetNumSensNonlinSolvConvFails returns the number of nonlinear convergence failures that have occurred for sensitivity calculations.

Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block. nSncfails (long int) number of nonlinear convergence failures.

Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
Notes This counter is incremented only if the ism was CV_STAGGERED or CV_STAGGERED1 in the call to CVodeSensMalloc (see §6.2.1).
In the CV_STAGGERED1 case, the value of nSncfails is the sum of the number of nonlinear convergence failures that occured for each sensitivity equation. These individual counters can be obtained through a call to CVodeGetNumStgrSensNonlinConvFails (see below).

## CVodeGetSensNonlinSolvStats

Call flag = CVodeGetSensNonlinSolvStats (cvode_mem, \&nSniters, \&nSncfails);
Description The function CVodeGetSensNonlinSolvStats returns the sensitivity-related nonlinear solver statistics as a group.
Arguments cvode_mem (void $*$ ) pointer to the CVODES memory block.
nSniters (long int) number of nonlinear iterations performed.
nSncfails (long int) number of nonlinear convergence failures.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output values have been successfully set.
CV_MEM_NULL The cvode mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.

## CVodeGetNumStgrSensNonlinSolvIters

Call flag = CVodeGetNumStgrSensNonlinSolvIters (cvode_mem, nSTGR1niters);
Description The function CVodeGetNumStgrSensNonlinSolvIters returns the number of nonlinear (functional or Newton) iterations performed for each sensitivity equation separately, in the CV_STAGGERED1 case.
Arguments cvode mem (void *) pointer to the CVODES memory block.
nSTGR1niters (long int *) an array (of dimension Ns) which will be set with the number of nonlinear iterations performed for each sensitivity system individually.
Return value The return value flag (of type int) is one of:

CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
Notes The user must allocate space for nSTGR1niters.

## CVodeGetNumStgrSensNonlinSolvConvFails

Call flag = CVodeGetNumStgrSensNonlinSolvConvFails(cvode_mem, nSTGR1ncfails);
Description The function CVodeGetNumStgrSensNonlinSolvConvFails returns the number of nonlinear convergence failures that have occurred for each sensitivity equation separately, in the CV_STAGGERED1 case.
Arguments cvode mem (void $*$ ) pointer to the CVODES memory block.
nSTGR1ncfails (long int $*$ ) an array (of dimension Ns) which will be set with the number of nonlinear convergence failures for each sensitivity system individually.
Return value The return value flag (of type int) is one of:
CV_SUCCESS The optional output value has been successfully set.
CV_MEM_NULL The cvode_mem pointer is NULL.
CV_NO_SENS Forward sensitivity analysis was not initialized.
Notes The user must allocate space for nSTGR1ncfails.

### 6.3 User-supplied routines for forward sensitivity analysis

In addition to the required and optional user-supplied routines described in §5.6, when using CVODES for forward sensitivity analysis, the user has the option of providing a routine that calculates the right-hand side of the sensitivity equations (3.9).

By default, CVODES uses difference quotient approximation routines for the right-hand sides of the sensitivity equations. However, CVODES allows the option for user-defined sensitivity right-hand side routines (which also provides a mechanism for interfacing CVODES to routines generated by automatic differentiation).

### 6.3.1 Sensitivity equations right-hand side (all at once)

If the CV_SIMULTANEOUS or CV_STAGGERED approach was selected in the call to CVodeSensMalloc, the user may provide the right-hand sides of the sensitivity equations (3.9), for all sensitivity parameters at once, through a function of type CVSensRhsFn defined by:

## CVSensRhsFn

Definition typedef int (*CVSensRhsFn)(int Ns, realtype t, N_Vector y, N_Vector ydot, N_Vector *yS, N_Vector *ySdot, void *fS_data, N_Vector tmp1, N_Vector tmp2);
Purpose This function computes the sensitivity right-hand side for all sensitivity equations at once. It must compute the vectors $(\partial f / \partial y) s_{i}(t)+\left(\partial f / \partial p_{i}\right)$ and store them in ySdot [i].
Arguments $t \quad$ is the current value of the independent variable.
$\mathrm{y} \quad$ is the current value of the state vector, $y(t)$.
ydot is the current value of the right-hand side of the state equations.
yS contains the current values of the sensitivity vectors.

```
ySdot is the output of CVSensRhsFn. On exit it must contain the sensitivity right- hand side vectors.
f_data is a pointer to user data - the same as the fS_data parameter passed to
    CVodeSetSensRhsFn.
tmp1
tmp2 are N_Vectors which can be used as temporary storage.
```

Return value A CVSensRhsFn should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverably (in which case the integration is halted and CV_SRHSFUNC_FAIL is returned).

Notes A sensitivity right-hand side function of type CVSensRhsFn is not compatible with the CV_STAGGERED1 approach.
Allocation of memory for ySdot is handled within CVODES.
For efficiency considerations, the right-hand side function is not evaluated at the converged solution of the nonlinear solver. Therefore, a recoverable error in CVSensRhsFn at that point cannot be corrected (as it will occur when the right-hand side function is called the first time during the following integration step and a successful step cannot be undone).
There are two situations in which recovery is not possible even if CVSensRhsFn function returns a recoverable error flag. This include the situation when this occurrs at the very first call to the CVSensRhsFn (in which case CVODES returns CV_FIRST_SRHSFUNC_ERR) or if a recoverable error is reported when CVSensRhsFn is called after an error test failure, while the linear multistep method order is equal to 1 (in which case CVODES returns CV_UNREC_SRHSFUNC_ERR).

### 6.3.2 Sensitivity equations right-hand side (one at a time)

Alternatively, the user may provide the sensitivity right-hand sides, one sensitivity parameter at a time, through a function of type CVSensRhs1Fn. Note that a sensitivity right-hand side function of type CVSensRhs1Fn is compatible with any valid value of the CVodeSensMalloc argument ism, and is required if ism=CV_STAGGERED1. The type CVSensRhs1Fn is defined by

| CVSensRhs |  |
| :---: | :---: |
| Definition | ```typedef int (*CVSensRhs1Fn)(int Ns, realtype t, N_Vector y, N_Vector ydot, int iS, N_Vector yS, N_Vector ySdot, void *fS_data, N_Vector tmp1, N_Vector tmp2);``` |
| Purpose | This function computes the sensitivity right-hand side for one sensitivity equation at a time. It must compute the vector $(\partial f / \partial y) s_{i}(t)+\left(\partial f / \partial p_{i}\right)$ for $i=\mathrm{iS}$ and store it in ySdot. |
| Arguments | t is the current value of the independent variable. <br> y is the current value of the state vector, $y(t)$. <br> ydot is the current value of the right-hand side of the state equations. <br> iS is the index of the parameter for which the sensitivity right-hand side must be <br> computed.  |
|  | ```yS contains the current value of the iS-th sensitivity vector. ySdot is the output of CVSensRhs1Fn. On exit it must contain the iS-th sensitivity right-hand side vector.``` |

```
f_data is a pointer to user data - the same as the fS_data parameter passed to
    CVodeSetSensRhs1Fn.
tmp1
tmp2 are N_Vectors which can be used as temporary storage.
```

Return value A CVSensRhsFn should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverably (in which case the integration is halted and CV_SRHSFUNC_FAIL is returned).
Notes Allocation of memory for ySdot is handled within CVODES.
For efficiency considerations, the right-hand side function is not evaluated at the converged solution of the nonlinear solver. Therefore, a recoverable error in CVSensRhs1Fn at that point cannot be corrected (as it will occur when the right-hand side function is called the first time during the following integration step and a successful step cannot be undone).
There are two situations in which recovery is not possible even if CVSensRhs1Fn function returns a recoverable error flag. This include the situation when this occurrs at the very first call to the CVSensRhs1Fn (in which case CVODES returns CV FIRST _SRHSFUNC ERR) or if a recoverable error is reported when CVSensRhs 1 Fn is called after an error test failure, while the linear multistep method order is equal to 1 (in which case CVODES returns CV_UNREC_SRHSFUNC_ERR).

### 6.4 Note on using partial error control

For some problems, when sensitivities are excluded from the error control test, the behavior of CVODES may appear at first glance to be erroneous. One would expect that, in such cases, the sensitivity variables would not influence in any way the step size selection. A comparison of the solver diagnostics reported for cvsdenx and the second run of the cvsfwddenx example in [18] indicates that this may not always be the case.

The short explanation of this behavior is that the step size selection implemented by the error control mechanism in CVODES is based on the magnitude of the correction calculated by the nonlinear solver. As mentioned in $\S 6.2 .1$, even with partial error control selected in the call to CVodeSensMalloc, the sensitivity variables are included in the convergence tests of the nonlinear solver.

When using the simultaneous corrector method (§3.2), the nonlinear system that is solved at each step involves both the state and sensitivity equations. In this case, it is easy to see how the sensitivity variables may affect the convergence rate of the nonlinear solver and therefore the step size selection. The case of the staggered corrector approach is more subtle. After all, in this case (ism=CV_STAGGERED or ism=CV_STAGGERED1 in the call to CVodeSensMalloc), the sensitivity variables at a given step are computed only once the solver for the nonlinear state equations has converged. However, if the nonlinear system corresponding to the sensitivity equations has convergence problems, CVODES will attempt to improve the initial guess by reducing the step size in order to provide a better prediction of the sensitivity variables. Moreover, even if there are no convergence failures in the solution of the sensitivity system, CVODES may trigger a call to the linear solver's setup routine which typically involves reevaluation of Jacobian information (Jacobian approximation in the case of CVDENSE and CVBAND, or preconditioner data in the case of CVSPGMR). The new Jacobian information will be used by subsequent calls to the nonlinear solver for the state equations and, in this way, potentially affect the step size selection.

When using the simultaneous corrector method it is not possible to decide whether nonlinear solver convergence failures or calls to the linear solver setup routine have been triggered by convergence problems due to the state or the sensitivity equations. When using one of the staggered corrector methods however, these situations can be identified by carefully monitoring the diagnostic information provided through optional outputs. If there are no convergence failures in the sensitivity nonlinear
solver, and none of the calls to the linear solver setup routine were made by the sensitivity nonlinear solver, then the step size selection is not affected by the sensitivity variables.

Finally, the user must be warned that the effect of appending sensitivity equations to a given system of ODEs on the step size selection (through the mechanisms described above) is problemdependent and can therefore lead to either an increase or decrease of the total number of steps that CVODES takes to complete the simulation. At first glance, one would expect that the impact of the sensitivity variables, if any, would be in the direction of increasing the step size and therefore reducing the total number of steps. The argument for this is that the presence of the sensitivity variables in the convergence test of the nonlinear solver can only lead to additional iterations (and therefore a smaller final correction), or to additional calls to the linear solver setup routine (and therefore more up-to-date Jacobian information), both of which will lead to larger steps being taken by CVODES. However, this is true only locally. Overall, a larger integration step taken at a given time may lead to step size reductions at later times (due to either nonlinear solver convergence failures or error test failures).

## Chapter 7

## Using CVODES for Adjoint Sensitivity Analysis

This chapter describes the use of CVODES to compute sensitivities of derived functions using adjoint sensitivity analysis. As mentioned before, the adjoint sensitivity module of CVODES provides the infrastructure for integrating backward in time any system of ODEs that depends on the solution of the original IVP, by providing various interfaces to the main CVODES integrator, as well as several supporting user-callable functions. For this reason, in the following sections we refer to the backward problem and not to the adjoint problem when discussing details relevant to the ODEs that are integrated backward in time. The backward problem can be the adjoint problem (3.17) or (3.20), and can be augmented with some quadrature differential equations.

CVODES uses various constants for both input and output. These are defined as needed in this chapter, but for convenience are also listed separately in Chapter 11.

We begin with a brief overview, in the form of a skeleton user program. Following that are detailed descriptions of the interface to the various user-callable functions and of the user-supplied functions that were not already described in $\S 5$.

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

The following is a skeleton of the user's main program as an application of CVODES. The user program is to have these steps in the order indicated, unless otherwise noted. For the sake of brevity, we defer many of the details to the later sections. As in $\S 5.4$, most steps are independent of the NVECTOR implementation used; where this is not the case, usage specifications are given for the two implementations provided with CVODES: steps marked with $[\mathbf{P}]$ correspond to NVECTOR_PARALLEL, while steps marked with $[\mathbf{S}]$ correspond to NVECTOR_SERIAL.

## 1. Include necessary header files

The cvodes.h header file also defines additional types, constants, and function prototypes for the adjoint sensitivity module user-callable functions. In addition, the main program should include an NVECTOR implementation header file (nvector_serial.h or nvector_parallel.h for the two implementations provided with CVODES) and, if Newton iteration was selected, the main header file of the desired linear solver module.
2. [P] Initialize MPI

## Forward problem

3. Set problem dimensions for the forward problem

## 4. Set initial conditions for the forward problem

5. Create CVODES object for the forward problem
6. Allocate internal memory for the forward problem
7. Set optional inputs for the forward problem
8. Attach linear solver module for the forward problem
9. Set linear solver optional inputs for the forward problem

## 10. Allocate space for the adjoint computation

Call cvadjmem = CVadjMalloc() to allocate memory for the combined forward-backward problem (see $\S 7.2 .1$ for more details). This call requires Nd, the number of steps between two consecutive checkpoints. CVadjMalloc also specifies the type of interpolation used (see §3.3.1).

## 11. Integrate forward problem

Call CVodeF, a wrapper for the CVODES main integration function CVode, either in CV NORMAL mode to the time tout or in CV_ONE_STEP mode inside a loop (if intermediate solutions of the forward problem are desired (see $\S 7.2 .2)$ ). The final value of tret, denoted tfinal, is then the maximum allowable value for the endpoint $t_{1}$.

## Backward problem

## 12. Set problem dimensions for the backward problem

[ $\mathbf{S}]$ set NB, the number of variables in the backward problem
$[\mathbf{P}]$ set NB and NBlocal

## 13. Set final values for the backward problem

Set the vector yBO of final values for the backward problem.

## 14. Create CVODES object for the backward problem

Call CVodeCreateB, a wrapper for CVodeCreate, to create the CVODES memory block and specify the solution method (linear multistep method and nonlinear solver iteration type) for the backward problem. Unlike CVodeCreate, the function CVodeCreateB does not return a pointer to the newly created memory block. Instead, this pointer is attached to the adjoint memory block (returned by CVadjMalloc and passed as the first argument to CVodeCreateB).

## 15. Allocate memory for the backward problem

Call CVodeMallocB, a wrapper for CVodeMalloc, to allocate internal memory and initialize CVODES at tBO for the backward problem (see §7.2.3).

## 16. Set optional inputs for the backward problem

Call CVodeSet*B functions to change from their default values any optional inputs that control the behavior of CVODES. Unlike their counterparts for the forward problem, these functions take as their first argument the adjoint memory block returned by CVadjMalloc.

## 17. Attach linear solver module for the backward problem

If Newton iteration is chosen, initialize the linear solver module for the backward problem by calling the appropriate wrapper function: CVDenseB, CVBandB, CVDiagB, CVSpgmrB, CVSpbcgB, or CVSptfqmr (see §7.2.4). Note that it is not required to use the same linear solver module for both the forward and the backward problems; for example, the forward problem could be solved with the CVDENSE linear solver and the backward problem with CVSPGMR.

## 18. Initialize quadrature calculation

If additional quadrature equations must be evaluated, call CVodeQuadMallocB, a wrapper around CVodeQuadMalloc, to initialize and allocate memory for quadrature integration. Optionally, call CVodeSetQuad*B functions to change from their default values optional inputs that control the integration of quadratures during the backward phase.

## 19. Integrate backward problem

Call CVodeB, a second wrapper around the CVODES main integration function CVode, to integrate the backward problem from tB0 (see §7.2.5). This function can be called either in CV_NORMAL or CV_ONE_STEP mode. Typically, CVodeB will be called in CV_NORMAL mode with an end time equal to the initial time of the forward problem.

## 20. Extract quadrature variables

If applicable, call CVodeGetQuadB, a wrapper around CVodeGetQuad, to extract the values of the quadrature variables at the time returned by the last call to CVodeB.

## 21. Deallocate memory

Upon completion of the backward integration, call all necessary deallocation functions. These include appropriate destructors for the vectors y and yB, a call to CVodeFree to free the CVODES memory block for the forward problem, and a call to CVadjFree (see $\S 7.2 .1$ ) to free the memory allocated for the combined problem. Note that CVadjFree also deallocates the CVODES memory for the backward problem.

## 22. Finalize MPI

[P] If MPI was initialized by the user main program, call MPI Finalize();
The above user interface to the adjoint sensitivity module in CVODES was motivated by the desire to keep it as close as possible in look and feel to the one for ODE IVP integration. Note that if steps (12)-(20) are not present, a program with the above structure will have the same functionality as one described in $\S 5.4$ for integration of ODEs, albeit with some overhead due to the checkpointing scheme.

### 7.2 User-callable functions for adjoint sensitivity analysis

### 7.2.1 Adjoint sensitivity allocation and deallocation functions

After the setup phase for the forward problem, but before the call to CVodeF, memory for the combined forward-backward problem must be allocated by a call to the function CVadjMalloc. The form of the call to this function is

```
CVadjMalloc
```

Call cvadj_mem = CVadjMalloc(cvode_mem, Nd, interpType);
Description The function CVadjMalloc allocates internal memory for the combined forward and backward integration, other than the CVODES memory block. Space is allocated for the $N_{d}$ interpolation data points and a linked list of checkpoints is initialized.

Arguments cvode_mem (void *) is the CVODES memory block for the forward problem returned by a previous call to CVodeCreate.
Nd (long int) is the number of integration steps between two consecutive checkpoints.
interpType (int) specifies the type of interpolation used and can be CV_POLYNOMIAL or CV_HERMITE, indicating variable-degree polynomial and cubic Hermite interpolation, respectively (see §3.3.1).

Return value If successful, CVadjMalloc returns a pointer of type void *. The user does not need to access this memory block but must pass it to other adjoint module user-callable functions. In case of failure (cvode_mem is NULL, an input has an illegal value, or a memory request fails), CVadjMalloc prints an error message to the standard output stream stderr and returns NULL.

Notes The user must set Nd so that all data needed for interpolation of the forward problem solution between two checkpoints fits in memory. CVadjMalloc attempts to allocate space for $(2 \mathrm{Nd}+3)$ variables of type N_Vector.

CVadjFree
Call CVadjFree(\&cvadj_mem);
Description
Argument
CVadjFree(\&cvadj_mem);

The function CVadjFree frees the memory allocated by a previous call to CVadjMalloc.
The only argument is the address of the pointer to the adjoint memory block (of type void *).
Return value The function CVadjFree has no return value.
Notes This function frees all memory allocated by CVadjMalloc. This includes workspace memory, the linked list of checkpoints, memory for the interpolation data, as well as the CVODES memory for the backward integration phase.
If the user wishes to change the interpolation method after the call to CVadjMalloc, this can be done through a call to the function CVadsSetInterpType. The form of the call to this function is

```
CVadjSetInterpType
Call flag = CVadjSetInterpType(cvadjmem, interpType);
```

Description The function CVadjSetInterpType resets the interpolation method to interpType.
Arguments cvadj mem (void $*$ ) pointer to the adjoint memory block.
interpType (int) specifies the type of interpolation used and can be CV POLYNOMIAL or CV_HERMITE, indicating variable-degree polynomial and cubic Hermite interpolation, respectively (see §3.3.1).
Return value The return flag (of type int) is one of:
CV_SUCCESS The call to CVadjSetInterpType was successful.
CV_ADJMEM_NULL The cvadj_mem argument was NULL.
CV_ILL_INPUT The input argument interpType has an illegal value.
CV_MEM_FAIL An error occured while trying to allocate memory for the interpolation module.

Notes The function CVadjSetInterpType performs memory deallocation and allocation.

### 7.2.2 Forward integration function

The function CVodeF is very similar to the CVODES function CVode (see $\S 5.5 .4$ ) in that it integrates the solution of the forward problem and returns the solution in y. At the same time, however, CVodeF stores checkpoint data every Nd integration steps. CVodeF can be called repeatedly by the user. The call to this function has the form

CVodeF
Call flag = CVodeF (cvadj mem, tout, yout, tret, itask, ncheck);
Description The function CVodeF integrates the forward problem over an interval in $t$ and saves checkpointing data.

Arguments cvadjmem (void *) pointer to the adjoint memory block.
tout (realtype) the next time at which a computed solution is desired.
yout (N_Vector) the computed solution vector.
tret (realtype *) the time reached by the solver.
itask (int) a flag indicating the job of the solver for the next step. The CV_NORMAL task is to have the solver take internal steps until it has reached or just passed the user-specified tout parameter. The solver then interpolates in order to return an approximate value of $y$ (tout). The CV_ONE_STEP option tells the solver to just take one internal step and return the solution at the point reached by that step. The CV_NORMAL_TSTOP and CV_ONE_STEP_TSTOP modes are similar to CV_NORMAL and CV_ONE_STEP, respectively, except that the integration never proceeds past the value tstop (specified through the function CVodeSetStopTime).
Return value On return, CVodeF returns a vector yout and a corresponding independent variable value $t=*$ tret, such that yout is the computed value of $y(t)$. Additionally, it returns in ncheck the number of checkpoints saved. The return value flag (of type int) will be one of the following. For more details see §5.5.4.
CV_SUCCESS CVodeF succeeded.
CV_TSTOP_RETURN CVodeF succeeded by reaching the optional stopping point.
CV_NO_MALLOC The function CVodeMalloc has not been previously called.
CV_ILL_IPUT One of the inputs to CVodeF is illegal.
CV_TOO_MUCH_WORK The solver took mxstep internal steps but could not reach tout.
CV_TOO_MUCH_ACC The solver could not satisfy the accuracy demanded by the user for some internal step.
CV_ERR_FAILURE Error test failures occurred too many times during one internal time step or occurred with $|h|=h_{\text {min }}$.
CV_CONV_FAILURE Convergence test failures occurred too many times during one internal time step or occurred with $|h|=h_{\text {min }}$.
CV_LSETUP_FAIL The linear solver's setup function failed in an unrecoverable manner.
CV_LSOLVE_FAIL The linear solver's solve function failed in an unrecoverable manner.
CV_ADJMEM_NULL The cvadj mem argument was NULL.
CV_MEM_FAIL A memory allocation request has failed (in an attempt to allocate space for a new checkpoint).
Notes All failure return values are negative and therefore a test flag<0 will trap all CVodeF failures.

At this time, CVodeF stores checkpoint information in memory only. Future versions will provide for a safeguard option of dumping checkpoint data into a temporary file as needed. The data stored at each checkpoint is basically a snapshot of the CVODES internal memory block and contains enough information to restart the integration from that time and to proceed with the same step size and method order sequence as during the forward integration.
In addition, CVodeF also stores interpolation data between consecutive checkpoints so that, at the end of this first forward integration phase, interpolation information is already available from the last checkpoint forward. In particular, if no check points were necessary, there is no need for the second forward integration phase.
Although it is legal to define a value for tstop and then call CVodeF in CV_NORMAL_TSTOP or CV_ONE_STEP_TSTOP modes, after a return with flag = TSTOP_RETURN, the integration should not be continued (no tstop information is stored at checkpoints).

It is illegal to change the integration tolerances between consecutive calls to CVodeF, as this information is not captured in the checkpoints data.


### 7.2.3 Backward problem initialization functions

The functions CVodeCreateB and CVodeMallocB must be called in the order listed. They instantiate a CVODES solver object, provide problem and solution specifications, and allocate internal memory for the backward problem.

## CVodeCreateB

Call flag = CVodeCreateB(cvadj mem, lmm, iter);
Description The function CVodeCreateB instantiates a CVODES solver object and specifies the solution method for the backward problem.
Arguments cvadjmem (void *) pointer to the adjoint memory block returned by CVadjMalloc.
lmm (int) specifies the linear multistep method and must be one of two possible values: CV_ADAMS or CV_BDF.
iter (int) specifies the type of nonlinear solver iteration and must be either CV_NEWTON or CV_FUNCTIONAL.

Return value If successful, CVodeCreateB stores a pointer to the newly created CVODES memory block (of type void *) for the backward problem. The return flag (of type int) is one of:

CV_SUCCESS The call to CVodeCreateB was successful.
CV_ADJMEM_NULL The cvadj mem argument was NULL.
CV_MEM_FAIL An error occured while trying to create the CVODES memory block for the backward problem.
The function CVodeMallocB is essentially a call to CVodeMalloc with some particularization for backward integration as described below.

## CVodeMallocB

Call flag = CVodeMallocB (cvadj mem, fB, tBO, yBO, itolB, reltolB, abstolB);
Description The function CVodeMallocB provides required problem and solution specifications, allocates internal memory, and initializes CVODES for the backward problem.
Arguments cvadj_mem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc. $\mathrm{fB} \quad(\mathrm{CVRhsFnB})$ is the C function which computes $f B$, the right-hand side of the backward ODE problem. This function has the form $f B(t, y, y B, y B d o t$, f_dataB) (for full details see $\S 7.3$ ).
tB0 (realtype) specifies the endpoint where final conditions are provided for the backward problem.
yB0 (N_Vector) is the final value of the backward problem.
itolB (int) is one of CV_SS or CV_SV where itol=CV_SS indicates scalar relative error tolerance and scalar absolute error tolerance, while itol=CV_SV indicates scalar relative error tolerance and vector absolute error tolerance.
reltolB (realtype) is the relative error tolerance.
abstolB (void $*$ ) is a pointer to the absolute error tolerance. If itolB $=\mathrm{CV}$ _SS, abstolB must be a pointer to a realtype variable. If itolB $=\mathrm{CV} \_\mathrm{SV}$, abstolB must be an N_Vector variable.
Return value The return flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeMallocB was successful.
CV_ADJMEM_NULL The cvadj_mem argument was NULL.
CV_BAD_TBO The final time tBO is outside the interval over which the forward problem was solved.

CV_MEM_NULL The CVODES memory block for the backward problem was not initialized through a previous call to CVodeCreateB.
CV_MEM_FAIL A memory allocation request has failed.
CV_ILL_INPUT An input argument to CVodeMallocB has an illegal value.
Notes It is the user's responsibility to provide compatible itolB and abstolB arguments.
The memory allocated by CVodeMallocB is deallocated by the function CVadjFree.
For the case when it is needed to solve several different backward problems corresponding to the same original problem, cVODES provides a mechanism to reuse the existing checkpoints. The function CVodeReInitB reinitializes the CVODES memory block for the backward problem, where a prior call to CVodeMallocB has been made with the same problem size NB. CVodeReInitB performs the same input checking and initializations that CVodeMallocB does, but does no memory allocation, assuming that the existing internal memory is sufficient for the new problem. Note that CVReInitB is essentially a wrapper for CVodeReInit and so all details given for CVodeReInit in $\S 5.5 .8$ apply. The call to the CVodeReInitB function has the form

```
CVodeReInitB
Call flag = CVodeReInitB(cvadj_mem, fB, tBO, yBO, itolB, reltolB, abstolB);
Description The function CVodeReInitB provides required problem specifications and reinitializes
    CVODES for the backward problem.
Arguments cvadjmem (void *) pointer to the adjoint memory block returned by CVadjMalloc.
    fB (CVRhsFnB) is the C function which computes }fB\mathrm{ , the right-hand side of the
    backward ODE problem. This function has the form fB(t,y, yB, yBdot,
    f_dataB) (for full details see §7.3).
    tB0 (realtype) specifies the endpoint where final conditions are provided for
    the backward problem.
    yB0 (N_Vector) is the final value of the backward problem.
    itolB (int) is either CV_SS or CV_SV, where itol=CV_SS indicates scalar relative
        error tolerance and scalar absolute error tolerance, while itol=CV_SV indi-
        cates scalar relative error tolerance and vector absolute error tolerance.
    reltolB (realtype) is the relative error tolerance.
    abstolB (void *) is a pointer to the absolute error tolerance. If itolB = CV_SS,
        abstolB must be a pointer to a realtype variable. If itolB = CV_SV,
        abstolB must be an N_Vector variable.
```

Return value The return value flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeReInitB was successful.
CV_ADJMEM_NULL The cvadj_mem argument was NULL.
CV_BAD_TBO The final time tBO is outside the interval over which the forward problem was solved.

CV_MEM_NULL The CVODES memory block for the backward problem was not initialized through a previous call to CVodeCreateB.
CV_NO_MALLOC Memory space for the CVODES memory block for the backward problem was not allocated through a previous call to CVodeMallocB.
CV_ILL_INPUT An input argument to CVodeReInitB has an illegal value.
Notes It is the user's responsibility to provide compatible itolB and abstolB arguments.

### 7.2.4 Linear solver initialization functions for backward problem

All linear solver modules in CVODES provide additional specification functions for the case in which Newton iteration is selected for the solution of the backward problem. The initialization functions
described in $\S 5.5 .3$ cannot be directly used since the optional user-defined Jacobian-related functions have different prototypes for the backward problem than for the forward problem (see §7.3).

The following six wrapper functions can be used to initialize one of the linear solver modules for the backward problem. Their arguments are identical to those of the functions in $\S 5.5 .3$ with the exception of their first argument which must be the pointer to the adjoint memory block returned by CVadjMalloc.

```
flag = CVDenseB(cvadj_mem, nB);
flag = CVDiagB(cvadj_mem);
flag = CVBandB(cvadj_mem, nB, mupperB, mlowerB);
flag = CVSpgmrB(cvadj_mem, pretypeB, maxlB);
flag = CVSpbcgB(cvadj_mem, pretypeB, maxlB);
flag = CVSptfqmrB(cvadj_mem, pretypeB, maxlB);
flag = CVSpbcgB(cvadj_mem, pretypeB, maxlB);
flag = CVSptfqmrB(cvadj_mem, pretypeB, maxlB);
```

Their return value flag (of type int) can have any of the return values of their counterparts. If the cvadj mem argument was NULL, flag will be CVDENSE_ADJMEM_NULL, CVDIAG_ADJMEM_NULL, CVBAND_ADJMEM_NULL, or CVSPILS_ADJMEM_NULL.

### 7.2.5 Backward integration function

The function CVodeB performs the integration of the backward problem. It is essentially a wrapper for the CVODES main integration function CVode and, in the case in which checkpoints were needed, it evolves the solution of the backward problem through a sequence of forward-backward integrations between consecutive checkpoints. The first run integrates the original IVP forward in time and stores interpolation data; the second run integrates the backward problem backward in time and performs the required interpolation to provide the solution of the IVP to the backward problem.

The call to this function has the form

## CVodeB

Call flag $=$ CVodeB(cvadj mem, tBout, yBout, tBret, itaskB);
Description The function CVodeB integrates the backward ODE problem over an interval in $t$.
Arguments cvadj_mem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc.
tBout (realtype) the next time at which a computed solution is desired.
yBout (N_Vector) the computed solution vector of the backward problem.
tBret (realtype *) the time reached by the solver.
itaskB (int) a flag indicating the job of the solver for the next step. The CV_NORMAL task is to have the solver take internal steps until it has reached or just passed the user specified tBout parameter. The solver then interpolates in order to return an approximate value of $y B$ (tBout). The CV_ONE_STEP option tells the solver to just take one internal step and return the solution at the point reached by that step.
Return value On return, CVodeB returns a vector yBout and a corresponding independent variable value $t=*$ tBret, such that yBout is the computed value of the solution of the backward problem.

The return value flag (of type int) will be one of the following. For more details see §5.5.4.
CV_SUCCESS CVodeB succeeded.
CV_NO_MALLOC The CVODES memory for the backward problem was NULL.
CV_ILL_INPUT One of the inputs to CVode is illegal.
CV_BAD_ITASK The itaskB argument has an illegal value.

|  | CV_TOO_MUCH_WORK CV_TOO_MUCH_ACC | The solver took mxstep internal steps but could not reach tBout. The solver could not satisfy the accuracy demanded by the user for some internal step. |
| :---: | :---: | :---: |
|  | CV_ERR_FAILURE | Error test failures occurred too many times during one internal time step. |
|  | CV_CONV_FAILURE | Convergence test failures occurred too many times during one internal time step. |
|  | CV_LSETUP_FAIL | The linear solver's setup function failed in an unrecoverable manner. |
|  | CV_SOLVE_FAIL | The linear solver's solve function failed in an unrecoverable manner. |
|  | CV_ADJMEM_NULL | The cvadj mem argument was NULL. |
|  | CV_BCKMEM_NULL | The cvodes memory for the backward problem was not created through a call to CVodeCreateB. |
|  | CV_BAD_TBOUT | The desired output time tBout is outside the interval over which the forward problem was solved. |
|  | CV_REIFWD_FAIL | Reinitialization of the forward problem failed at the first checkpoint (corresponding to the initial time of the forward problem). |
|  | CV_FWD_FAIL | An error occured during the integration of the forward problem. |
| Notes | All failure return v failures. | values are negative and therefore a test $\mathrm{flag}<0$ will trap all CVodeB |

### 7.2.6 Optional input functions for the backward problem

### 7.2.6.1 Main solver optional input functions

The adjoint module in CVODES provides wrappers for most of the optional input functions defined in $\S 5.5 .5 .1$. The only difference is that the first argument of the optional input functions for the backward problem is the pointer to the adjoint memory block, cvadj mem, of type void $*$, returned by CVadjMalloc. The optional input functions defined for the backward problem are:

```
flag = CVodeSetErrHandlerB(cvadj_mem, ehfunB, eh\_dataB);
flag = CVodeSetErrFileB(cvadj_mem, errfpB);
flag = CVodeSetIterTypeB(cvadj_mem, iterB);
flag = CVodeSetFdataB(cvadj_mem, f_dataB);
flag = CVodeSetMaxOrdB(cvadj_mem, maxordB);
flag = CVodeSetMaxNumStepsB(cvadj_mem, mxstepsB);
flag = CVodeSetStabLimDetB(cvadj_mem, stldetB);
flag = CVodeSetInitStepB(cvadj_mem, hinB);
flag = CVodeSetMinStepB(cvadj_mem, hminB);
flag = CVodeSetMaxStepB(cvadj_mem, hmaxB);
```

Their return value flag (of type int) can have any of the return values of their counterparts, but it can also be CV_ADJMEM_NULL if the cvadj mem argument was NULL.

### 7.2.6.2 Dense linear solver

Optional inputs for the CVDENSE linear solver module can be set for the backward problem through the following function:

```
CVDenseSetJacFnB
Call flag = CVDenseSetJacFnB(cvadj_mem, djacB, jac_dataB);
```

Description The function CVDenseSetJacFnB specifies the dense Jacobian approximation function to be used for the backward problem and the pointer to user data.
Arguments cvadj_mem (void *) pointer to the adjoint memory block.
djacB (CVDenseJacFnB) user-defined dense Jacobian approximation function. jac_dataB (void $*$ ) pointer to the user-defined data structure.
Return value The return value flag (of type int) is one of:
CVDENSE_SUCCESS The optional value has been successfuly set.
CVDENSE_MEM_NULL The CVODES solver memory block was not created through a call to CVodeCreateB.
CVDENSE_LMEM_NULL The CVDENSE linear solver has not been initialized through a call to CVDenseB.
CVDENSE_ADJMEM_NULL The cvadjmem argument was NULL.
Notes The function type CVDenseJacFnB is described in §7.3.

### 7.2.6.3 Band linear solver

Optional inputs for the CVBAND linear solver module can be set for the backward problem through the following function:

```
CVBandSetJacFnB
Call flag = CVBandSetJacFnB(cvadj mem, bjacB, jac_dataB);
```

Description The function CVBandSetJacFnB specifies the banded Jacobian approximation function to be used for the backward problem and the pointer to user data.

Arguments cvadjmem (void *) pointer to the adjoint memory block.
bjacB (CVBandJacFnB) user-defined banded Jacobian approximation function.
jac_dataB (void $*$ ) pointer to the user-defined data structure.
Return value The return value flag (of type int) is one of:
CVBAND_SUCCESS The optional value has been successfuly set.
CVBAND_MEM_NULL The CVODES solver memory block was not created through a call to CVodeCreateB.
CVBAND_LMEM_NULL The CVBAND linear solver has not been initialized through a call to CVBandB.
CVBAND_ADJMEM_NULL The cvadj_mem argument was NULL.
Notes The function type CVBandJacFnB is described in $\S 7.3$.

### 7.2.6.4 SPILS linear solvers

Optional inputs for the CVSPILS linear solver module can be set for the backward problem through the following functions:

```
CVSpilsSetPreconditionerB
Call flag = CVSpilsSetPreconditionerB(cvadjmem, psolveB, psetupB, p_dataB);
```

Description The function CVSpilsSetPrecSolveFnB specifies the preconditioner setup and solve functions and the pointer to user data for the backward integration.
Arguments cvadjmem (void *) pointer to the adjoint memory block.
psolveB (CVSpilsPrecSolveFnB) user-defined preconditioner solve function.
psetupB (CVSpilsPrecSetupFnB) user-defined preconditioner setup function.
p_dataB (void $*$ ) pointer to the user-defined data structure.
Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The optional value has been successfuly set.

CVSPILS_MEM_NULL The CVODES solver memory block was not created through a call to CVodeCreateB.
CVSPILS_LMEM_NULL The CVSPGMR linear solver has not been initialized.
CVSPILS_ADJMEM_NULL The cvadj_mem argument was NULL.
Notes The function types CVSpilsPrecSolveFnB and CVSpilsPrecSetupFnB are described in §7.3.
CVSpilsSetJacTimesVecFnB
Call flag = CVSpilsSetJacTimesVecFnB(cvadj_mem, jtimesB, jac_data);

Description The function CVSpilsSetJacTimesFnB specifies the Jacobian-vector product function to be used and te pointer to user data.
Arguments cvadjmem (void *) pointer to the adjoint memory block.
jtimesB (CVSpilsJacTimesVecFnB) user-defined Jacobian-vector product function. jac_dataB (void $*$ ) pointer to the user-defined data structure.
Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The optional value has been successfuly set.
CVSPILS_MEM_NULL The CVODES solver memory block was not created through a call to CVodeCreateB.
CVSPILS_LMEM_NULL The CVSPGMR linear solver has not been initialized.
CVSPILS_ADJMEM_NULL The cvadj_mem argument was NULL.
Notes The function type CVSpilsJacTimesVecFnB is described in §7.3.
CVSpilsSetGSTypeB
Call flag $=$ CVSpilsSetGSType(cvadj mem, gstypeB);

Description The function CVSpilsSetGSTypeB specifies the type of Gram-Schmidt orthogonalizadion to be used with CVSPGMR. This must be one of the enumeration constants MODIFIED_GS or CLASSICAL_GS. These correspond to using modified Gram-Schmidt and classical Gram-Schmidt, respectively.
Arguments cvadjmem (void *) pointer to the adjoint memory block. gstypeB (int) type of Gram-Schmidt orthogonalization.
Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The optional value has been successfuly set.
CVSPILS_MEM_NULL The CVODES solver memory block was not created through a call to CVodeCreateB.
CVSPILS_LMEM_NULL The CVSPGMR linear solver has not been initialized.
CVSPILS_ILL_INPUT The Gram-Schmidt orthogonalization type gstypeB is not valid. CVSPILS_ADJMEM_NULL The cvadj_mem argument was NULL.
Notes The default value is MODIFIED_GS.
This option is available only with CVSPGMR.

CVSpilsSetDeltB
Call flag = CVSpilsSetDeltB (cvadjmem, deltB);
Description The function CVSpilsSetDeltB specifies the factor by which the Krylov linear solver's convergence test constant is reduced from the Newton iteration test constant.
Arguments cvadj_mem (void *) pointer to the adjoint memory block.
deltB (realtype) the value of the convergence test constant reduction factor.
Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The optional value has been successfuly set.
CVSPILS_MEM_NULL The CVODES solver memory block was not created through a call to CVodeCreateB.
CVSPILS_LMEM_NULL The CVSPGMR linear solver has not been initialized.
CVSPILS_ILL_INPUT The factor deltB is negative.
CVSPILS_ADJMEM_NULL The cvadj_mem argument was NULL.
Notes The default value is 0.05 . Passing a value deltB $=0.0$ also indicates using the default value.

```
CVSpilsSetMaxlB
Call flag = CVSpilsSetMaxlB(cvadj mem, maxlB);
```

Description The function CVSpilsSetMaxlB resets maximum Krylov subspace dimension for the Bi-CGStab or TFQMR methods.

Arguments cvadj_mem (void *) pointer to the adjoint memory block.
$\operatorname{maxlB} \quad$ (realtype) maximum dimension of the Krylov subspace.
Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The optional value has been successfuly set.
CVSPILS_MEM_NULL The CVODES solver memory block was not created through a call to CVodeCreateB.
CVSPILS_LMEM_NULL The CVSPGMR linear solver has not been initialized.
CVSPILS_ILL_INPUT The factor deltB is negative.
CVSPILS_ADJMEM_NULL The cvadj_mem argument was NULL.
Notes The maximum subspace dimension is initially specified in the call to CVSpbcgB or CVSptfqmrB. The call to CVSpilsSetMaxlB is needed only if maxl is being changed from its previous value.

This option is available only for the CVSPBCG and CVSPTFQMR linear solvers.

## CVSpilsSetPrecTypeB

Call flag = CVSpilsSetPrecTypeB (cvadj mem, pretypeB);
Description The function CVSpilsSetPrecTypeB resets the type of preconditioning to be used.
Arguments cvadjmem (void $*$ ) pointer to the adjoint memory block.
pretypeB (int) specifies the type of prconditioning and must be one of: PREC_NONE, PREC_LEFT, PREC_RIGHT, or PREC_BOTH.
Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The optional value has been successfuly set.
CVSPILS_MEM_NULL The CVODES solver memory block was not created through a call to CVodeCreateB.
CVSPILS_LMEM_NULL The CVSPGMR linear solver has not been initialized.
CVSPILS_ILL_INPUT The preconditioner type pretype is not valid.
CVSPILS_ADJMEM_NULL The cvadj_mem argument was NULL.
Notes The preconditioning type is initially specified in the call to the linear solver specification function (see $\S 7.2 .4$ ). The call to CVSpilsSetPrecTypeB is needed only if pretypeB is being changed from its previous value.

### 7.2.7 Optional output functions for the backward problem

The user of the adjoint module in CVODES has access to any of the optional output functions described in $\S 5.5 .7$, both for the main solver and for the linear solver modules. The first argument of these CVodeGet* and CV*Get* functions is the CVODES memory block for the backward problem. In order to call any of these functions, the user must first call the following function to obtain a pointer to this memory block:

## CVadjGetCVodeBmem

Call cvode_memB = CVadjGetCVodeBmem (cvadj_mem);
Description The function CvadjGetCVodeBmem returns a pointer to the CVODES memory block for the backward problem.
Arguments The argument cvadjmem (of type void $*$ ) is a pointer to the adjoint memory block returned by CVadjMalloc.
Return value The return value, cvode_memB (of type void *), is a pointer to the CVODES memory for the backward problem.
Notes The user should not modify in any way cvode_memB.

### 7.2.8 Backward integration of pure quadrature equations

### 7.2.8.1 Backward quadrature initialization functions

The function CVodeQuadMallocB initializes and allocates memory for the backward integration of quadrature equations. It has the following form:
CVodeQuadMallocB
Call flag $=$ CVodeQuadMallocB(cvadj mem, fQB, yQBO);

Description The function CVodeQuadMallocB provides required problem specifications, allocates internal memory, and initializes backward quadrature integration.
Arguments cvadjmem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc.
f QB (CVQuadRhsFnB) is the C function which computes $f Q B$, the right-hand side of the backward quadrature equations. This function has the form $\mathrm{fQB}(\mathrm{t}$, $\left.y, y B, q B d o t, f Q \_d a t a B\right)$ (for full details see below).
yQBO (N_Vector) is the value of the quadrature variables at tBS.
Return value The return value flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeQuadMallocB was successful.
CV_MEM_NULL The CVODES solver memory block was not created through a previous call to CVodeCreateB.
CV_MEM_FAIL A memory allocation request has failed.
CV_ADJMEM_NULL The cvadj mem argument was NULL.
The integration of quadrature equations during the backward phase can be re-initialized by calling
CVodeQuadReInitB
Call flag $=$ CVodeQuadReInitB (cvadj mem, fQB, yQBO);

Description The function CVodeQuadReInitB re-initializes the backward quadrature integration.
Arguments cvadj mem (void $*$ ) pointer to the adjoint memory block.
f QB (CVQuadRhsFnB) is the C function which computes $f Q B$, the right-hand side of the backward quadrature equations.
yQBO (N_Vector) is the value of the quadrature variables at iBO.

Return value The return value flag (of type int) will be one of the following:
CV_SUCCESS The call to CVodeReInitB was successful.
CV_MEM_NULL The CVODES solver memory block was not created through a previous call to CVodeCreateB.

CV_NO_QUAD Quadrature integration was not activated through a previous call to CVodeQuadMallocB.
CV_ADJMEM_NULL The cvadj mem argument was NULL.

### 7.2.8.2 Backward quadrature extraction function

To extract the values of the quadrature variables at the last return time of CVodeB, CVODES provides a wrapper for the function CVodeGetQuad (see §5.7.3). The call to this function has the form

```
CVodeGetQuadB
Call flag = CVodeGetQuadB(cvadjmem, yQB);
```

Description The function CVodeGetQuadB returns the quadrature solution vector after a successful return from CVodeB.

Arguments cvadj_mem (void *) pointer to the adjoint memory returned by CVadjMalloc.
yQB (N_Vector) the computed backward quadrature vector.
Return value The return value flag of CVodeGetQuadB is one of:
CV_SUCCESS CVodeGetQuadB was successful.
CV_MEM_NULL The CVODES solver memory block was not created through a previous call to CVodeCreateB.
CV_NO_QUAD Quadrature integration was not initialized through a previous call to CVodeQuadMallocB..
Cv_BAD_DKY yQB is NULL.
CV_ADJMEM_NULL The cvadj mem argument was NULL.

### 7.2.8.3 Optional input/output functions for backward quadrature integration

Optional values controlling the backward integration of quadrature equations can be changed from their default values through calls to one of the following functions which are wrappers for the corresponding optional input functions defined in §5.7.4:

```
flag = CVodeSetQuadFdataB(cvadj_mem, fQ_dataB);
flag = CVodeSetQuadErrConB(cvadj_mem, errconQB, itolQB, reltolQB, abstolQB);
```

Their return value flag (of type int) can have any of the return values of their counterparts, but it can also be CV_ADJMEM_NULL if the cvadj mem argument was NULL.

Access to optional outputs related to backward quadrature integration can be obtained by calling the corresponding CVodeGetQuad* functions (see §5.7.5). A pointer to the CVODES memory block for the backward problem, required as the first argument of these functions, can be obtained through a call to the functions CVadjGetCVodeBmem (see §7.2.7).

### 7.2.9 Optional output from the adjoint module

### 7.2.9.1 Checkpoint information function

For debugging purposes, CVODES provides a function CVadjGetCheckPointsInfo which returns partial information from the linked list of checkpoints generated by CVodeF. The call to this function has the form:

```
CVadjGetCheckPointsInfo
Call flag = CVadjCheckPointsList(cvadj_mem, ckpnt);
```

Description The function CVadjGetCheckPointsList returns a structure array with checkpoint information.
Arguments cvadj_mem (void $*$ ) pointer to the adjoint memory returned by CVadjMalloc.
ckpnt (CVadjCheckPointRec *) an array of ncheck +1 structures with checkpoint information, where ncheck is the numebr of checkpoints returned by CVodeF.
Return value The return value flag of CVadjGetCheckPointsInfo is one of:
CV_SUCCESS CVadjGetCheckPointsInfo was successful.
CV_ADJMEM_NULL The cvadj mem argument was NULL.
Notes For an example of using CVadjGetCheckPointsInfo, see the cvadjdenx example.
The type CVadjCheckPointRec is defined in the header file cvodes.h:

```
typedef struct {
    void *my_addr;
    void *next_addr;
    realtype t0;
    realtype t1;
    long int nstep;
    int order;
    realtype step;
} CVadjCheckPointRec;
```

The fields in this structure have the following meanings:
my_addr Address of current checkpoint.
next_addr Address of next checkpoint.
t0
t1 Time interval between current and next checkpoint.
nstep Step number at which the current checkpoint was saved.
order Linear multistep method order at the current checkpoint.
step Integration stepsize at current checkpoint.

### 7.2.9.2 Interpolation data

Fo debugging purposes, CVODEA provides two extraction functions which return the data stored for interpolation purposes.


Return value The return value flag is one of:

CV_SUCCESS CVadjGetDataPointHermite was successful.
CV_ILL_INPUT The interpolation type was not cubic Hermite.
CV_ADJMEM_NULL The cvadj mem argument was NULL.
Notes It is the user's responsibility to allocate space for y and yd.

```
CVadjGetDataPointPolynomial
Call int = CVadjGetDataPointPolynomial(cvadjmem, which, &t, order, y);
Description The function CVadjGetDataPointPolynomial returns the time and two vectors associated with the which interpolation data point.
Arguments cvadjmem (void *) pointer to the adjoint memory returned by CVadjMalloc.
which (long int) index of the intepolation data point.
t (realtype *)
order (int)
yd (N_Vector) time, method order, and solution of the forward problem stored for interpolation purposes at the which data point.
```

Return value The return value flag is one of:
CV_SUCCESS CVadjGetDataPointPolynomial was successful.
CV_ILL_INPUT The interpolation type was not variable-order polynomial.
CV_ADJMEM_NULL The cvadj mem argument was NULL.
Notes It is the user's responsibility to allocate space for y .

### 7.2.9.3 Return flag name

The names of constants associated with CVODEA-specific return values can be obtained by calling the following function:


Description The function CVadjGetReturnFlagName returns the name of the CVODEA constant corresponding to flag.

Arguments The only argument, of type int is a return flag from a CVODEA function.
Return value The return value is a string containing the name of the corresponding constant.

### 7.3 User-supplied functions for adjoint sensitivity analysis

In addition to the required ODE right-hand side function and any optional functions for the forward problem, when using the adjoint sensitivity module in CVODES, the user must supply one function defining the backward problem ODE and, optionally, functions to supply Jacobian-related information (if Newton iteration is chosen) and one or two functions that define the preconditioner (if one of the CVSPILS solvers is selected) for the backward problem. Type definitions for all these user-supplied functions are given below.

### 7.3.1 ODE right-hand side for the backward problem

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

## CVRhsFnB

Definition typedef int (*CVRhsFnB) (realtype t, N_Vector y, N_Vector yB, N_Vector yBdot, void *f_dataB);

Purpose This function computes the right-hand side of the backward problem ODE system. This could be (3.17) or (3.20).
Arguments $t \quad$ is the current value of the independent variable.
$y \quad$ is the current value of the forward solution vector.
$y B \quad$ is the current value of the dependent variable vector.
yBdot is the output vector containing the right-hand side of the backward ODE problem.
f_dataB is a pointer to user data - the same as the f_dataB parameter passed to CVodeSetFdataB.
Return value $A$ CVRhsFnB should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecovdrably (in which case the integration is halted and CVodeB returns CV_RHSFUNC.FAIL).

Notes Allocation of memory for yBdot is handled within CVODES.
The $y$, $y B$, and yBdot arguments are all of type N_Vector, but yB and yBdot typically have different internal representations from y. It is the user's responsibility to access the vector data consistently (including the use of the correct accessor macros from each NVECTOR implementation). For the sake of computational efficiency, the vector functions in the two NVECTOR implementations provided with CVODES do not perform any consistency checks with respect to their N_Vector arguments (see $\S 8.1$ and $\S 8.2$ ).
The f_dataB pointer is passed to the user's dB function every time it is called and can be the same as the f_data pointer used for the forward problem.

Before calling the user's CVRhsFnB, CVODEA needs to evaluate (through interpolation) the values of the states from the forward integration. If an error occurs in the interporation, CVODEA triggers an unrecoverable failure in the righ-hand side function which will halt the integration and CVodeB will return CV_RHSFUNC_FAIL.

### 7.3.2 Quadrature right-hand side for the backward problem

The user must provide a function of type CVQuadRhsFnB defined by

CVQuadRhsFnB
Definition typedef int (*CVQuadRhsFnB) (realtype t, N_Vector y, N_Vector yB, N_Vector qBdot, void $* f$ Q_dataB) ;
Purpose This function computes the quadrature equation right-hand side for the backward problem.

Arguments $t$ is the current value of the independent variable.
$y \quad$ is the current value of the forward solution vector.
$\mathrm{yB} \quad$ is the current value of the dependent variable vector.
qBdot is the output vector containing the right-hand side of the backward quadratore equations.
$f Q_{d}$ data is a pointer to user data - the same as the $f Q_{\text {_data }}$ parameter passed to CVodeSetQuadFdataB.

Return value $A$ CVQuadRhsFnB should return 0 if successful, a positive value if a recoverable erfor occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverably (in which case the integration is halted and CVodeB returns CV_QRHSFUNC_FAIL).

Notes Allocation of memory for qBdot is handled within CVODES.
The $\mathrm{y}, \mathrm{yB}$, and yQBdot arguments are all of type N_Vector, but they typically all have different internal representations. It is the user's responsibility to access the vector data consistently (including the use of the correct accessor macros from each NVECTOR implementation). For the sake of computational efficiency, the vector functions in the two NVECTOR implementations provided with CVODES do not perform any consistency checks with repsect to their N_Vector arguments (see $\S 8.1$ and $\S 8.2$ ).
The $f Q \_$dataB pointer is passed to the user's $f Q B$ function every time it is called and can be the same as the f_data pointer used for the forward problem.

Before calling the user's CVQuadRhsFnB, CVODEA needs to evaluate (through interpolation) the values of the states from the forward integration. If an error occurrs in the interpolation, CVODEA triggers an unrecoverable failure in the quadrature righ-hand side function which will halt the integration and CVodeB will return CV_QRHSFUNCFAIL.

### 7.3.3 Jacobian information for the backward problem (direct method with dense Jacobian)

If the direct linear solver with dense treatment of the Jacobian is selected for the backward problem (i.e. CVDenseB is called in step 17 of $\S 7.1$ ), the user may provide, through a call to CVDenseSetJacFnB (see $\S 7.2 .6$ ), a function of the following type:

```
CVDenseJacFnB
Definition typedef int (*CVDenseJacFnB)(long int nB, DenseMat JB, realtype t,
                                    N_Vector y, N_Vector yB, N_Vector fyB,
                                    void *jac_dataB, N_Vector tmp1B,
    N_Vector tmp2B, N_Vector tmp3B);
```

Purpose This function computes the dense Jacobian of the backward problem (or an approximation to it). If the backward problem is the adjoint of the original IVP, then this Jacobian is just the transpose of $J=\partial f / \partial y$ with a change in sign.

| Arguments | nB | is the backward problem size. |
| :--- | :--- | :--- |
| J | is the output Jacobian matrix. |  |
| t | is the current value of the independent variable. |  |
| y | is the current value of the forward solution vector. |  |
| yB | is the current value of the dependent variable vector. |  |
| fyB | is the current value of the right-hand side of the backward problem. |  |
| jac_dataB | is a pointer to user data - the same as the jac_dataB parameter passed to |  |
| CVDenseSetJacDataB. |  |  |

Return value $A$ CVDenseJacFnB should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct, while CVDENSE sets last_flag on CVDENSE_JACFUNC_RECVR), or a negative value if it failed unrecoverably (in which case the integration is halted, CVodeB returns CV_LSETUP FAIL and CVDENSE sets last_flag on CVDENSE_JACFUNC_UNRECVR).
Notes A user-supplied dense Jacobian function must load the nB by nB dense matrix JB with an approximation to the Jacobian matrix at the point $(\mathrm{t}, \mathrm{y}, \mathrm{yB})$, where y is the solution of the original IVP at time $t$ and $y B$ is the solution of the backward problem at the same time. Only nonzero elements need to be loaded into JB as this matrix is set to
zero before the call to the Jacobian function. The type of JB is DenseMat. The user is referred to §5.6.4 for details regarding accessing a DenseMat object.
Before calling the user's CVDenseJacFnB, CVODEA needs to evaluate (through interpolation) the values of the states from the forward integration. If an error occurrs in the interpolation, CVODEA triggers an unrecoverable failure in the Jacobian function which will halt the integration (CVodeB returns CV LSETUP FAIL and CVDENSE sets last_flag on CVDENSE_JACFUNC_UNRECVR).

### 7.3.4 Jacobian information for the backward problem (direct method with banded Jacobian)

If the direct linear solver with banded treatment of the Jacobian is selected for the backward problem (i.e. CVBandB is called in step 17 of $\S 7.1$ ), the user may provide, through a call to CVBandSetJacFnB (see §7.2.6), a function the following type:

```
CVBandJacFnB
Definition typedef int (*CVBandJacFnB)(long int nB,
                                    long int mupperB, long int mlowerB,
                                    BandMat JB, realtype t, N_Vector y,
                                    N_Vector yB, N_Vector fyB, void *jac_dataB,
                                    N_Vector tmp1B, N_Vector tmp2B,
                                    N_Vector tmp3B);
```

Purpose This function computes the banded Jacobian of the backward problem (or a banded approximation to it).
Arguments $\mathrm{nB} \quad$ is the backward problem size.
mlowerB
mupperB are the lower and upper half-bandwidth of the Jacobian.
JB is the output Jacobian matrix.
$\mathrm{t} \quad$ is the current value of the independent variable.
$\mathrm{y} \quad$ is the current value of the forward solution vector.
$y B \quad$ is the current value of the dependent variable vector.
$\mathrm{fyB} \quad$ is the current value of the right-hand side of the backward problem.
jac_dataB is a pointer to user data - the same as the jac_dataB parameter passed to CVBandSetJacDataB.
tmp1B
tmp2B
tmp3B are pointers to memory allocated for variables of type N_Vector which can be used by CVBandJacFnB as temporary storage or work space.
Return value $A$ CVBandJacFnB should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct, while CVBAND sets last_flag on CVBAND_JACFUNC_RECVR), or a negative value if it failed unrecoverably (in which case the integration is halted, CVodeB returns CV_LSETUP_FAIL and CVDENSE sets last_flag on CVBAND_JACFUNC_UNRECVR).
Notes A user-supplied band Jacobian function must load the band matrix JB (of type BandMat) with the elements of the Jacobian at the point $(\mathrm{t}, \mathrm{y}, \mathrm{yB})$, where y is the solution of the original IVP at time $t$ and $y B$ is the solution of the backward problem at the same time. Only nonzero elements need to be loaded into JB because JB is preset to zero before the call to the Jacobian function. More details on the accessor macros provided for a BandMat object and on the rest of the arguments passed to a function of type CVBandJacFnB are given in §5.6.5.

Before calling the user's CVBandJacFnB, CVODEA needs to evaluate (through interpolatimon) the values of the states from the forward integration. If an error occurs in the interpolation, CVODEA triggers an unrecoverable failure in the Jacobian function which will halt the integration (CVodeB returns CV_LSETUP_FAIL and CVBAND sets last_flag on CVBAND_JACFUNC_UNRECVR).

### 7.3.5 Jacobian information for the backward problem (matrix-vector produt)

If one of the Krylov iterative linear solvers SPGMR, SPBCG, or SPTFQMR is selected (CVSp*B is called in step 17 of $\S 7.1$ ), the user may provide a function of type CVSpilsJacTimesVecFnB in the following form:

## CVSpilsJacTimesVecFnB

Definition typedef int (*CVSpilsJacTimesVecFnB) (N_Vector vB, N_Vector JvB, realtype t, N_Vector y, N_Vector yB, N_Vector yB, void *jac_dataB, N_Vector tmpB);

Purpose This function computes the action of the Jacobian on a given vector vB for the backward problem (or an approximation to it).

Arguments vB is the vector by which the Jacobian must be multiplied to the right.
$\mathrm{JvB} \quad$ is the output vector computed.
$\mathrm{t} \quad$ is the current value of the independent variable.
$\mathrm{y} \quad$ is the current value of the forward solution vector.
$\mathrm{yB} \quad$ is the current value of the dependent variable vector.
$\mathrm{fyB} \quad$ is the current value of the right-hand side of the backward problem.
jac_dataB is a pointer to user data - the same as the jac_dataB parameter passed to CVSp*SetJacTimesVecFnB.
$\operatorname{tmpB} \quad$ is a pointer to memory allocated for a vector which can be used for work space.

Return value The return value of a function of type CVSpilsJtimesFnB should be 0 if successful or nonzero if an error was encountered, in which case the integration is halted.

Notes A user-supplied Jacobian-vector product function must load the vector JvB with the result of the product between the Jacobian of the backward problem at the point ( $\mathrm{t}, \mathrm{y}$, $y B)$ and the vector $v B$. Here, $y$ is the solution of the original IVP at time $t$ and $y B$ is the solution of the backward problem at the same time. The rest of the arguments are equivalent to those passed to a function of type CVSpilsJacTimesVecFn (see §5.6.6). If the backward problem is the adjoint of $\dot{y}=f(t, y)$, then this function is to compute $-(\partial f / \partial y)^{T} v_{B}$.

### 7.3.6 Preconditioning for the backward problem (linear system solution)

If preconditioning is used during integration of the backward problem, then the user must provide a C function to solve the linear system $P z=r$, where $P$ may be either a left or a right preconditioner matrix. This function must be of type CVSpilsPrecSolveFnB defined by

Definition typedef int (*CVSpilsPrecSolveFnB) (realtype t, N_Vector y, N_Vector yB, N_Vector fyB, N_Vector rB, N_Vector zB, realtype gammaB, realtype deltaB, int lrB, void *P_dataB, N_Vector tmpB);

Purpose This function solves the preconditioning system $P z=r$ for the backward problem.
Arguments $t \quad$ is the current value of the independent variable.
$\mathrm{y} \quad$ is the current value of the forward solution vector.
$y B \quad$ is the current value of the dependent variable vector.
$\mathrm{fyB} \quad$ is the current value of the right-hand side of the backward problem.
$r B \quad$ is the right-hand side vector of the linear system.
$\mathrm{zB} \quad$ is the output vector computed.
gammaB is the scalar $\gamma$ appearing in the Newton matrix.
deltaB is an input tolerance to be used if an iterative method is employed in the solution.
$\operatorname{lrB} \quad$ is an input flag indicating whether the preconditioner solve function is to use the left preconditioner ( $\operatorname{lr}=1$ ) or the right preconditioner ( $\operatorname{lr}=2$ );
P_dataB is a pointer to user data - the same as the P_dataB parameter passed to the function CVSp*SetPreconditionerB.
$\operatorname{tmpB} \quad$ is a pointer to memory allocated for a vector which can be used for work space.
Return value The return value of a preconditioner solve function for the backward problem should be 0 if successful, positive for a recoverable error (in which case the step will be retried), or negative for an unrecoverable error (in which case the integration is halted).

### 7.3.7 Preconditioning for the backward problem (Jacobian data)

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

```
CVSpilsPrecSetupFnB
Definition typedef int (*CVSpilsPrecSetupFnB)(realtype t, N_Vector y, N_Vector yB,
                                    N_Vector fyB, booleantype jokB,
                                    booleantype *jcurPtrB,
                                    realtype gammaB, void *P_dataB,
                                    N_Vector tmp1B, N_Vector tmp2B,
                                    N_Vector tmp3B);
Purpose This function preprocesses and/or evaluates Jacobian-related data needed by the preconditioner for the backward problem.
Arguments The arguments of a CVSpilsPrecSetupFnB are as follows:
\(t \quad\) is the current value of the independent variable.
\(y \quad\) is the current value of the forward solution vector.
\(\mathrm{yB} \quad\) is the current value of the dependent variable vector.
fyB is the current value of the right-hand side of the backward problem.
jokB is an input flag indicating whether Jacobian-related data needs to be recomputed.
jcurPtrB is a pointer to an output integer flag which is to be set to TRUE if Jacobian data was recomputed, or to FALSE if Jacobian data was not recomputed but saved data was still reused.
gammaB is the scalar \(\gamma\) appearing in the Newton matrix.
```

P_dataB is a pointer to user data - the same as the P_dataB parameter passed to CVSpilsSetPreconditionerB.
tmp1B
tmp2B
tmp3B are pointers to memory allocated for vectors which can be used as temporary storage or work space.
Return value The return value of a preconditioner setup function for the backward problem should be 0 if successful, positive for a recoverable error (in which case the step will be retried), or negative for an unrecoverable error (in which case the integration is halted).

### 7.4 Using CVODES preconditioner modules for the backward problem

As on the forward integration phase, the efficiency of Krylov iterative methods for the solution of linear systems can be greatly enhanced through preconditioning. Both preconditioner modules provided with SUNDIALS, the banded preconditioner CVBANDPRE and the band-block-diagonal preconditioner module CVBBDPRE, provide interface functions through which they can be used on the backward integration phase.

### 7.4.1 Using the banded preconditioner CVBANDPRE

The adjoint module in CVODES offers an interface to the banded preconditioner module CVBANDPRE described in section §5.9.1. This preconditioner provides a band matrix preconditioner based on difference quotients of the backward problem right-hand side function $f B$. It generates a banded approximation to the Jacobian with $m_{l B}$ sub-diagonals and $m_{u B}$ super-diagonals to be used with one of the Krylov linear solvers.

In order to use the CVBANDPRE module in the solution of the backward problem, the user need not define any additional functions. First, the user must initialize the CVBANDPRE module by calling


Return value If successful, CVBandPrecAlloc stores a pointer to the newly created CVBANDPRE memory block. The return value flag (of type int) is one of:

CVBANDPRE_SUCCESS The call to CVBandPrecAllocB was successful.
CVBANDPRE_MEM_FAIL An error occured while trying to create the CVBANDPRE memory block.
CVBANDPRE_ADJMEM NULL The cvadj mem argument was NULL.
To specify the use of the CVSPGMR linear solver module with the CVBANDPRE preconditioner module, make the following call:

## CVBPSpgmrB

Call flag = CVBPSpgmrB(cvadj_mem, pretypeB, maxlB);
Description The function CVBPSpgmrB links the CVBANDPRE data to the CVSPGMR linear solver and attaches the latter to the CVODES memory block for the backward problem.
Arguments cvadjmem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc. pretypeB (int) preconditioning type. Can be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxlB} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
Return value The return value flag (of type int) is one of:

| CVSPILS_SUCCESS | The CVSPGMR initialization was successful. |
| :--- | :--- |
| CVSPILS_MEM_NULL | The CVODES memory block for the backward problem was not <br> initialized through a previous call to CVodeCreateB. |
| CVSPILS_ILL_INPUT | The preconditioner type pretypeB is not valid. |
| CVSPILS_MEM_FAIL | A memory allocation request failed. |
| CVBANDPRE_PDATA_NULL | The CVBANDPRE preconditioner has not been initialized. |
| CVBANDPRE_ADJMEM_NULL | The cvadj mem argument was NULL. |

To specify the use of the CVSPBCG linear solver module with the CVBANDPRE preconditioner module, make the following call:

## CVBPSpbcgB

Call flag = CVBPSpbcgB(cvadj mem, pretypeB, maxlB);
Description The function CVBPSpbcgB links the CVBANDPRE data to the CVSPBCG linear solver and attaches the latter to the CVODES memory block for the backward problem.
Arguments cvadjmem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc. pretypeB (int) preconditioning type. Can be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxlB} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
Return value The return value flag (of type int) is one of:

| CVSPILS_SUCCESS | The CVSPBCG initialization was successful. |
| :--- | :--- |
| CVSPILS_MEM_NULL | The CVODES memory block for the backward problem was not <br> initialized through a previous call to CVodeCreateB. |
| CVSPILS_ILL_INPUT | The preconditioner type pretypeB is not valid. |
| CVSPILS_MEM_FAIL | A memory allocation request failed. |
| CVBANDPRE_PDATA_NULL | The CVBANDPRE preconditioner has not been initialized. |
| CVBANDPRE_ADJMEM_NULL | The cvadj mem argument was NULL. |

To specify the use of the CVSPTFQMR linear solver module with the CVBANDPRE preconditioner module, make the following call:

## CVBPSptfqmrB

Call flag = CVBPSptfqmrB (cvadj mem, pretypeB, maxlB);
Description The function CVBPSptfqmrB links the CVBANDPRE data to the CVSPTFQMR linear solver and attaches the latter to the CVODES memory block for the backward problem.
Arguments cvadj.mem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc. pretypeB (int) preconditioning type. Can be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxlB} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.

Return value The return value flag (of type int) is one of: CVBANDPRE_ADJMEM_NULL The CVSPTFQMR initialization was successful.
CVSPILS_MEM_NULL The CVODES memory block for the backward problem was not initialized through a previous call to CVodeCreateB.
CVSPILS_ILL_INPUT The preconditioner type pretypeB is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
CVBANDPRE_PDATA_NULL The CVBANDPRE preconditioner has not been initialized.
CVBANDPRE_ADJMEM_NULL The cvadj mem argument was NULL.
To deallocate the CVBANDPRE preconditioner module memory used for the backward integration, make the following call:

## CVBandPrecFreeB

Call CVBandPrecFreeB(cvadj mem);
Description The function CVBandPrecFreeB frees the memory allocated by CVBandPrecAllocB.
Arguments The only argument passed to CVBandPrecFreeB is the pointer to the CVODEA memory block (of type void *).

Return value The function CVBandPrecFreeB has no return value.
Notes The call to CVBandPrecFreeB must preceed that to CVadjFree.
For more details on CVBANDPRE see §5.9.1.

### 7.4.2 Using the band-block-diagonal preconditioner CVBBDPRE

The adjoint module in CVODES offers an interface to the band-block-diagonal preconditioner module CVBBDPRE described in section §5.9.2. This generates a preconditioner that is a block-diagonal matrix with each block being a band matrix and can be used with one of the Krylov linear solvers and with the parallel vector module NVECTOR_PARALLEL.

In order to use the CVBBDPRE module in the solution of the backward problem, the user must define one or two additional functions, described at the end of this section.

### 7.4.2.1 Usage of CVBBDPRE for the backward problem

The cVBbDPRE module is initialized by calling

```
CVBBDPrecAllocB
Call flag = CVBBDPrecAllocB(cvadj mem, NlocalB, mudqB, mldqB,
    mukeepB, mlkeepB, dqrelyB, glocB, cfnB);
```

Description The function CVBBDPrecAllocB initializes and allocates memory for the CVBBDPRE preconditioner for the backward problem.
Arguments cvadjmem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc.
NlocalB (long int) local vector dimension for the backward problem.
mudqB (long int) upper half-bandwidth to be used in the difference-quotient Jacobian approximation.
mldqB (long int) lower half-bandwidth to be used in the difference-quotient Jacobian approximation.
mukeepB (long int) upper half-bandwidth of the retained banded approximate Jacobian block.
mlkeepB (long int) lower half-bandwidth of the retained banded approximate Jacobian block.
dqrelyB (realtype) the relative increment in components of yB used in the difference quotient approximations. The default is dqrelyB $=\sqrt{\text { unit roundoff, which }}$ can be specified by passing dqrely $=0.0$.
glocB (CVLocalFnB) the C function which computes the approximation $g_{B}(t, y)$ to the right-hand side of the backward problem.
cfnB (CVCommFnB) the optional C function which performs all interprocess communication required for the computation of $g_{B}(t, y)$.
Return value If successful, CVBBDPrecAlloc stores a pointer to the newly created CVBBDPRE memory block. The return value flag (of type int) is one of:
CVBBDPRE_SUCCESS The call to CVBBDPrecAllocB was successful.
CVBBDPRE_PDATA_NULL An error occured while trying to create the CVBBDPRE memory block.
CVBBDPRE_ADJMEM_NULL The cvadj_mem argument was NULL.
To specify the use of the CVSPGMR linear solver module with the CVBBDPRE preconditioner module, make the following call:

## CVBBDSpgmrB

Call flag $=$ CVBBDSpgmrB (cvadj mem, pretypeB, maxlB);
Description The function CVBBDSpgmrB links the CVBBDPRE data to the CVSPGMR linear solver and attaches the latter to the CVODES memory block for the backward problem.
Arguments cvadjmem (void *) pointer to the adjoint memory block returned by CVadjMalloc. pretypeB (int) preconditioning type. Can be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxlB} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
Return value The return value flag (of type int) is one of:

| CVSPILS_SUCCESS | The CVSPGMR initialization was successful. |
| :--- | :--- |
| CVSPILS_MEM_NULL | The CVODES memory block for the backward problem was not <br> initialized through a previous call to CVodeCreateB. |
| CVSPILS_ILL_INPUT | The preconditioner type pretypeB is not valid. |
| CVSPILS_MEM_FAIL | A memory allocation request failed. |
| CVBBDPRE_PDATA_NULL | The CVBBDPRE preconditioner has not been initialized. |
| CVBBDPRE_ADJMEM_NULL | The cvadj_mem argument was NULL. |

To specify the use of the CVSPBCG linear solver module with the CVBBDPRE preconditioner module, make the following call:

CVBBDSpbcgB
Call flag $=$ CVBBDSpbcgB (cvadjmem, pretypeB, maxlB);
Description The function CVBBDSpbcgB links the CVBBDPRE data to the CVSPBCG linear solver and attaches the latter to the CVODES memory block for the backward problem.
Arguments cvadjmem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc.
pretypeB (int) preconditioning type. Can be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxlB} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.
Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The CVSPBCG initialization was successful.
CVSPILS_MEM_NULL The CVODES memory block for the backward problem was not initialized through a previous call to CVodeCreateB.

CVSPILS_ILL_INPUT The preconditioner type pretypeB is not valid.
CVSPILS_MEMFAIL A memory allocation request failed.
CVBBDPRE_PDATA_NULL The CVBBDPRE preconditioner has not been initialized.
CVBBDPRE_ADJMEM_NULL The cvadj_mem argument was NULL.
To specify the use of the CVSPTFQMR linear solver module with the CVBBDPRE preconditioner module, make the following call:

## CVBBDSptfqmrB

Call flag = CVBBDSptfqmrB(cvadj mem, pretypeB, maxlB);
Description The function CVBBDSptfqmrB links the CVBBDPRE data to the CVSPTFQMR linear solver and attaches the latter to the CVODES memory block for the backward problem.

Arguments cvadjmem (void *) pointer to the adjoint memory block returned by CVadjMalloc. pretypeB (int) preconditioning type. Can be one of PREC_LEFT or PREC_RIGHT.
$\operatorname{maxlB} \quad$ (int) maximum dimension of the Krylov subspace to be used. Pass 0 to use the default value CVSPILS_MAXL $=5$.

Return value The return value flag (of type int) is one of:
CVSPILS_SUCCESS The CVSPTFQMR initialization was successful.
CVSPILS_MEM NULL The CVODES memory block for the backward problem was not initialized through a previous call to CVodeCreateB.
CVSPILS_ILL_INPUT The preconditioner type pretypeB is not valid.
CVSPILS_MEM_FAIL A memory allocation request failed.
CVBBDPRE_PDATA_NULL The CVBBDPRE preconditioner has not been initialized.
CVBBDPRE_ADJMEM_NULL The cvadj_mem argument was NULL.
To reinitialize the CVBBDPRE preconditioner module for the backward problem call the following function:

## CVBBDPrecReInitB

```
Call flag = CVBBDPrecReInitB(cvadj mem, mudqB, mldqB, dqrelyB, glocB, cfnB);
```

Description The function CVBBDPrecReInitB reinitializes the CVBBDPRE preconditioner for the backward problem.
Arguments cvadjmem (void $*$ ) pointer to the adjoint memory block returned by CVadjMalloc.
mudqB (long int) upper half-bandwidth to be used in the difference-quotient Jacobian approximation.
mldqB (long int) lower half-bandwidth to be used in the difference-quotient Jacobian approximation.
dqrelyB (realtype) the relative increment in components of $y B$ used in the difference quotient approximations.
glocB (CVLocalFnB) the C function which computes the approximation $g_{B}(t, y)$ to the right-hand side of the backward problem.
cfnB (CVCommFnB) the optional C function which performs all interprocess communication required for the computation of $g_{B}(t, y)$.
Return value The return value flag (of type int) is one of:
CVBBDPRRE_SUCCESS The CVBBDPRE initialization was successful.
CVBBDPRE_PDATA_NULL The CVBBDPRE preconditioner has not been initialized.
CVBBDPRE_ADJMEM_NULL The cvadj_mem argument was NULL.
To deallocate the CVBBDPRE preconditioner module memory used for the backward integration, make the following call:

## CVBBDPrecFreeB

Call CVBBDPrecFreeB(cvadj mem);
Description The function CVBBDPrecFreeB frees the memory allocated by CVBBDPrecAllocB.
Arguments The only argument passed to CVBBDPrecFreeB is the pointer to the CVODEA memory block (of type void $*$ ).

Return value The function CVBBDPrecFreeB has no return value.
Notes The call to CVBBDPrecFreeB must preceed that to CVadjFree.
For more details on CVBBDPRE see §5.9.2.

### 7.4.2.2 User-supplied functions for CVBBDPRE

To use the CVBBDPRE module, the user must supply one or two functions which the module calls to construct the preconditioner: a required function glocB (of type CVLocalFnB) which approximates the right-hand side of the backward problem and which is computed locally, and an optional function cfnB (of type CVCommFnB) which performs all interprocess communication necessary to evaluate this approximate right-hand side (see §5.9.2). The prototypes for these two functions are described below.

## CVLocalFnB

Definition typedef int (*CVLocalFnB) (long int NlocalB, realtype t, N_Vector $y$, N_Vector $y B$, N_Vector $g B$, void *f_dataB);
Purpose This function loads the vector $g B$ as a function of $t, y$, and $y B$.
Arguments NlocalB is the local vector length for the backward problem.
$t \quad$ is the value of the independent variable.
$\mathrm{y} \quad$ is the current value of the forward solution vector.
$\mathrm{yB} \quad$ is the current value of the dependent variable vector.
$g B \quad$ is the output vector.
f_dataB is a pointer to user data - the same as the $f$ dataB parameter passed to CVodeSetFdataB.

Return value $A$ CVLocalFnB should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverably (in which case the integration is halted and CVodeB returns CV LSETUP FAIL).
Notes This routine assumes that all interprocess communication of data needed to calculate gB has already been done, and this data is accessible within $f$ _dataB.
Before calling the user's CVLocalFnB, CVODEA needs to evaluate (through interpolation) the values of the states from the forward integration. If an error occurrs in the interpolation, CVODEA triggers an unrecoverable failure in the preconditioner setup function which will halt the integration (CVodeB returns CV LSETUP FAIL).

## CVCommFnB

Definition typedef int ( $*$ CVCommFnB) (long int NlocalB, realtype $t$, N_Vector $\left.y, N+V e c t o r ~ y B, ~ v o i d ~ * f \_d a t a B\right) ; ~$
Purpose This function performs all interprocess communications necessary for the execution of the glocB function above, using the input vectors $y$ and $y B$.
Arguments NlocalB is the local vector length.
$t \quad$ is the value of the independent variable.
$\mathrm{y} \quad$ is the current value of the forward solution vector.
$y B \quad$ is the current value of the dependent variable vector.
f_dataB is a pointer to user data - the same as the f_dataB parameter passed to CVodeSetFdataB.
Return value $A$ CVCommFn should return 0 if successful, a positive value if a recoverable error occurred (in which case CVODES will attempt to correct), or a negative value if it failed unrecoverably (in which case the integration is halted and CVodeB returns CV LSETUP_FAIL).
Notes The cfnB function is expected to save communicated data in space defined within the structure f_dataB.

Each call to the cfnB function is preceded by a call to the function that evaluates the right-hand side of the backward problem with the same $t, y$, and $y B$ arguments. If there is no additional communication needed, then pass $\mathrm{cfnB}=$ NULL to CVBBDPrecAllocB.

## Chapter 8

## Description of the NVECTOR module

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

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

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

The _generic_N_Vector_Ops structure is essentially a list of pointers to the various actual vector operations, and is defined as

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

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

The generic NVECTOR module 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 8.1 contains a complete list of all vector operations defined by the generic NVECTOR module.
Finally, note that the generic NVECTOR module defines the functions N_VCloneVectorArray and N_VCloneEmptyVectorArray. Both functions create (by cloning) an array of count variables of type N_Vector, each of the same type as an existing N_Vector. Their prototypes are

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

and their definitions are based on the implementation-specific N_VClone and N_VCloneEmpty operations, respectively.

An array of variables of type N_Vector can be destroyed by calling N_VDestroyVectorArray, whose prototype is

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

and whose definition is based on the implementation-specific N_VDestroy operation.
A particular implementation of the NVECTOR module must:

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

Table 8.1: Description of the NVECTOR operations

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


| continued from last page |  |
| :---: | :---: |
| Name | Usage and Description |
| N_VScale | N_VScale(c, x, z); <br> Scales the N_Vector x by the scalar c and returns the result in $\mathrm{z}: z_{i}=$ $c x_{i}, i=0, \ldots, n-1$. |
| N_VAbs | N_VAbs (x, z) ; <br> Sets the components of the N_Vector $z$ to be the absolute values of the components of the N_Vector $\mathrm{x}: y_{i}=\left\|x_{i}\right\|, i=0, \ldots, n-1$. |
| N_VInv | N_VInv(x, z) ; <br> Sets the components of the N_Vector $\mathbf{z}$ to be the inverses of the components of the N_Vector $\mathrm{x}: z_{i}=1.0 / x_{i}, i=0, \ldots, n-1$. 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); <br> Adds the scalar b to all components of x and returns the result in the N_Vector $\mathrm{z}: z_{i}=x_{i}+b, i=0, \ldots, n-1$. |
| N_VDotProd | $\mathrm{d}=\mathrm{N} \_\operatorname{VDotProd}(\mathrm{x}, \mathrm{y}) \text {; }$ <br> Returns the value of the ordinary dot product of x and $\mathrm{y}: d=\sum_{i=0}^{n-1} x_{i} y_{i}$. |
| N_VMaxNorm | $\mathrm{m}=\mathrm{N}$ _VMaxNorm(x); <br> Returns the maximum norm of the N_Vector $\mathrm{x}: m=\max _{i}\left\|x_{i}\right\|$. |
| N_VWrmsNorm | $\mathrm{m}=\mathrm{N} \_$VWrmsNorm $(\mathrm{x}, \mathrm{w})$ <br> Returns the weighted root-mean-square norm of the N_Vector x with weight vector w: $m=\sqrt{\left(\sum_{i=0}^{n-1}\left(x_{i} w_{i}\right)^{2}\right) / n}$. |
| N_VWrmsNormMask | $\mathrm{m}=\mathrm{N}$ _VWrmsNormMask(x, w, id); <br> Returns the weighted root mean square norm of the N_Vector x with weight vector w built using only the elements of x corresponding to nonzero elements of the N_Vector id: $m=\sqrt{\left(\sum_{i=0}^{n-1}\left(x_{i} w_{i} \operatorname{sign}\left(i d_{i}\right)\right)^{2}\right) / n}$ |
| N_VMin | $\mathrm{m}=\mathrm{N} \_\operatorname{VMin}(\mathrm{x}) ;$ <br> Returns the smallest element of the N_Vector $\mathrm{x}: m=\min _{i} x_{i}$. |
| N_VWL2Norm | $\mathrm{m}=\mathrm{N}$ _VWL2Norm ( $\mathrm{x}, \mathrm{w}$ ) ; <br> Returns the weighted Euclidean $\ell_{2}$ norm of the N_Vector x with weight vector w: $m=\sqrt{\sum_{i=0}^{n-1}\left(x_{i} w_{i}\right)^{2}}$. |
| N_VL1Norm | $m=\text { N_VL1Norm }(x) \text {; }$ <br> Returns the $\ell_{1}$ norm of the N_Vector $\mathrm{x}: m=\sum_{i=0}^{n-1}\left\|x_{i}\right\|$. |
| N_VCompare | N_VCompare (c, x, z); <br> Compares the components of the N_Vector x to the scalar c and returns an N_Vector $z$ such that: $z_{i}=1.0$ if $\left\|x_{i}\right\| \geq c$ and $z_{i}=0.0$ otherwise. |
|  | continued on next page |


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

### 8.1 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 serial version.

- NV_CONTENT_S

This routine gives access to the contents of the serial vector N_Vector.
The assignment v_cont $=$ NV_CONTENT_S(v) sets v_cont to be a pointer to the serial N_Vector content structure.
Implementation:
\#define NV_CONTENT_S(v) ( (N_VectorContent_Serial) (v->content) )

- NV_OWN_DATA_S, NV_DATA_S, NV_LENGTH_S

These macros give individual access to the parts of the content of a serial N_Vector.
The assignment v_data $=$ NV DATA_S(v) sets v_data to be a pointer to the first component of the data for the N_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 $r=N V \_I t h \_S(v, i)$ sets $r$ to be the value of the i-th component of $v$. The assignment NV_Ith_S $(v, i)=r$ sets the value of the $i-t h$ 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 8.1. Their names are obtained from those in Table 8.1 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_VCloneVectorArrayEmpty_Serial

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

N_Vector $*$ N_VCloneVectorArrayEmpty_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_VCloneVectorArrayEmpty_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_VCloneVectorArrayEmpty_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.


### 8.2 The NVECTOR_PARALLEL implementation

The parallel implementation of the NVECTOR module provided with SUNDIALS, NVECTOR_PARALLEL, defines the content field of N_Vector to be a structure containing the global and local lengths of the vector, a pointer to the beginning of a contiguous local data array, an MPI communicator, an a boolean flag own_data indicating ownership of the data array data.

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

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

- NV_CONTENT_P

This macro gives access to the contents of the parallel vector N_Vector.
The assignment v_cont = NV_CONTENT_P(v) sets v_cont to be a pointer to the N_Vector content structure of type struct _N_VectorParallelContent.
Implementation:
\#define NV_CONTENT_P(v) ( (N_VectorContent_Parallel) (v->content) )

- NV_OWN_DATA_P, NV_DATA_P, NV_LOCLENGTH_P, NV_GLOBLENGTH_P

These macros give individual access to the parts of the content of a parallel N_Vector.
The assignment v_data $=$ NV_DATA_P (v) sets v_data to be a pointer to the first component of the local data for the N_Vector v . The assignment NV_DATA_P(v) = v_data sets the component array of $v$ to be v_data by storing the pointer v_data.
The assignment v_llen = NV_LOCLENGTH_P (v) sets v_llen to be the length of the local part of v . The call NV_LENGTH $P(\mathrm{v})=$ llen_v sets the local length of v to be llen_v.
The assignment v_glen $=$ NV_GLOBLENGTH_P $(v)$ sets $v \_g l e n ~ t o ~ b e ~ t h e ~ g l o b a l ~ l e n g t h ~ o f ~ t h e ~ v e c t o r ~$ v . The call NV_GLOBLENGTH_P $(\mathrm{v})=$ glen_v sets the global length of v to be glen_v.
Implementation:

```
#define NV_OWN_DATA_P(v) ( NV_CONTENT_P(v)->own_data )
#define NV_DATA_P(v) ( NV_CONTENT_P(v)->data )
```

```
#define NV_LOCLENGTH_P(v) ( NV_CONTENT_P(v)->local_length )
```

\#define NV_GLOBLENGTH_P(v) ( NV_CONTENT_P(v)->global_length )

- NV_COMM_P

This macro provides access to the MPI communicator used by the NVECTOR_PARALLEL vectors. Implementation:

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

- NV_Ith_P

This macro gives access to the individual components of the local data array of an N_Vector.
The assignment $r=N V \_I t h P(v, i)$ sets $r$ to be the value of the i-th component of the local part of $v$. The assignment NV_Ith_P $(v, i)=r$ sets the value of the $i$-th component of the local part of $v$ to be $r$.
Here $i$ ranges from 0 to $n-1$, where $n$ is the local length.
Implementation:

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

The NVECTOR_PARALLEL module defines parallel implementations of all vector operations listed in Table 8.1 Their names are obtained from those in Table 8.1 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_Vector N_VNew_Parallel(MPI_Comm comm,
    long int local_length,
    long int global_length);
```


## - N_VNewEmpty_Parallel

This function creates a new parallel N_Vector with an empty (NULL) data array.
N_Vector N_VNewEmpty_Parallel(MPI_Comm comm, long int local_length, long int global_length);

## - N_VMake_Parallel

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

```
N_Vector N_VMake_Parallel(MPI_Comm comm,
    long int local_length,
    long int global_length,
    realtype *v_data);
```

- 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_VCloneVectorArrayEmpty_Parallel

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

N_Vector *N_VCloneVectorArrayEmpty_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_VCloneVectorArrayEmpty_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_VCloneVectorArrayEmpty 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.


### 8.3 NVECTOR functions used by CVODES

In Table 8.2 below, we list the vector functions in the NVECTOR module within the CVODES package. The table also shows, for each function, which of the code modules uses the function. The cVODES column shows function usage within the main integrator module, while the remaining seven columns show function usage within each of the six CVODES linear solvers (CVSPILS stands for any of CVSPGMR, CVSPBCG, or CVSPTFQMR), the CVBANDPRE and CVBBDPRE preconditioner modules, and the CVODEA adjoint sensitivity module.

There is one subtlety in the cVSPILS column hidden by the table, explained here for the case of the CVSPGMR module). The dot product function N_VDotProd is called both within the implementation file cvodes_spgmr.c for the CVSPGMR solver and within the implementation files sundials_spgmr.c and sundials_iterative.c for the generic SPGMR solver upon which the CVSPGMR solver is implemented. Also, although N_VDiv and N_VProd are not called within the implementation file cvodes_spgmr.c, they are called within the implementation file sundials_spgmr.c and so are required by the CVSPGMR solver module. This issue does not arise for the other three CVODES linear solvers because the generic DENSE and BAND solvers (used in the implementation of CVDENSE and CVBAND) do not make calls to any vector functions and CVDIAG is not implemented using a generic diagonal solver.

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

The vector functions listed in Table 8.1 that are not used by CVODES are: N_VWL2Norm, N_VL1Norm, N_VWrmsNormMask, N_VConstrMask, N_VCloneEmpty, and N_VMinQuotient. Therefore a user-supplied NVECTOR module for CVODES could omit these six kernels.

Table 8．2：List of vector functions usage by CVODES code modules

|  | $\begin{aligned} & 0 \\ & \text { n } \\ & 0 \\ & 0 \\ & 0 \end{aligned}$ | $\begin{aligned} & \text { 思 } \\ & \stackrel{i}{⿴ 囗 十} \\ & 0 \end{aligned}$ | 号 $\substack{\infty \\ 0 \\ 0}$ | $\begin{aligned} & 0 \\ & 4 \\ & 0 \\ & 0 \\ & \hline \end{aligned}$ | $\begin{aligned} & \text { n } \\ & \text { An } \\ & 0 \\ & 0 \end{aligned}$ |  | $\begin{aligned} & \text { N } \\ & \\ & \\ & 0 \\ & 0 \\ & 0 \\ & \hline 0 \end{aligned}$ | ¢ 1 0 0 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| N＿VClone | $\checkmark$ |  |  | $\checkmark$ | $\checkmark$ |  |  | $\checkmark$ |
| N＿VDestroy | $\checkmark$ |  |  | $\checkmark$ | $\checkmark$ |  |  | $\checkmark$ |
| N＿VSpace | $\checkmark$ |  |  |  |  |  |  |  |
| N＿VGetArrayPointer |  | $\checkmark$ | $\checkmark$ |  |  | $\checkmark$ | $\checkmark$ |  |
| N＿VSetArrayPointer |  | $\checkmark$ |  |  |  |  |  |  |
| N＿VLinearSum | $\checkmark$ | $\checkmark$ |  | $\checkmark$ | $\checkmark$ |  |  | $\checkmark$ |
| N＿VConst | $\checkmark$ |  |  |  | $\checkmark$ |  |  |  |
| N＿VProd | $\checkmark$ |  |  | $\checkmark$ | $\checkmark$ |  |  |  |
| N＿VDiv | $\checkmark$ |  |  | $\checkmark$ | $\checkmark$ |  |  |  |
| N＿VScale | $\checkmark$ | $\checkmark$ | $\checkmark$ | $\checkmark$ | $\checkmark$ | $\checkmark$ | $\checkmark$ | $\checkmark$ |
| N＿VAbs | $\checkmark$ |  |  |  |  |  |  |  |
| N＿VInv | $\checkmark$ |  |  | $\checkmark$ |  |  |  |  |
| N＿VAddConst | $\checkmark$ |  |  | $\checkmark$ |  |  |  |  |
| N＿VDotProd |  |  |  |  | $\checkmark$ |  |  |  |
| N＿VMaxNorm | $\checkmark$ |  |  |  |  |  |  |  |
| N＿VWrmsNorm | $\checkmark$ | $\checkmark$ | $\checkmark$ |  | $\checkmark$ | $\checkmark$ | $\checkmark$ |  |
| N＿VMin | $\checkmark$ |  |  |  |  |  |  |  |
| N＿VCompare |  |  |  | $\checkmark$ |  |  |  |  |
| N＿VInvTest |  |  |  | $\checkmark$ |  |  |  |  |

## Chapter 9

## Providing Alternate Linear Solver Modules

The central CVODES module interfaces with the linear solver module to be used 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 and allocate memory specific to the linear solver;
- lsetup: preprocess and evaluate 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 those described in $\S 5.5 .3)$ which will attach the above four functions to the main CVODES memory block. The cVodes memory block is a structure defined in the header file cvodes_impl.h. A pointer to such a structure is defined as the type CVodeMem. The four fields in a CvodeMem structure that must point to the linear solver's functions are cv_linit, cv_lsetup, cv_lsolve, and cv_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 consistency with the existing CVODES linear solver modules, we recommend that the return value of the specification function be 0 for a successful return or a negative value if an error occurs (the pointer to the main CVODES memory block is NULL, an input is illegal, the NVECTOR implementation is not compatible, a memory allocation fails, etc.)

To facilitate data exchange between the four interface functions, the field cv_lmem in the CVODES memory block can be used to attach a linear solver-specific memory block.

To be used during the backward integration with the CVODEA module, a linear solver module must also provide an additional user-callable specification function (like those described in §7.2.4) which will attach the four functions to the CVODES memory block for the backward integration. Note that this block (of type struct CVodeMemRec) is not directly accessible to the user, but rather is itself a field (cvb_mem) in the CVODEA memory block. The CVODEA memory block is a structure defined in the header file cvodea_impl.h. A pointer to such a structure is defined as the type CVadjMem. The specification function for backward integration should also return a negative value if the adjoint CVODEA memory block is NULL.

An additional field (ca_lmemB) in the CVODEA memory block provides a hook-up for optionally attaching a linear solver-specific memory block.

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

### 9.1 Initialization function

The type definition of linit is

## linit

Definition int (*linit) (CVodeMem cv_mem);
Purpose The purpose of linit is to complete linear solver-specific initializations, such as counters and statistics.

Arguments cv_mem is the CVODES memory pointer of type CVodeMem.
Return value An linit function should return 0 if it has successfully initialized the CVODES linear solver and -1 otherwise.

### 9.2 Setup function

The type definition of 1setup is

## lsetup

Definition

```
int (*lsetup)(CVodeMem cv_mem, int convfail, N_Vector ypred,
    N_Vector fpred, booleantype *jcurPtr,
    N_Vector vtemp1, N_Vector vtemp2, N_Vector vtemp3);
```

Purpose The job of lsetup is to prepare the linear solver for subsequent calls to lsolve. It may recompute Jacobian-related data if it is deemed necessary.
Arguments cv_mem is the CVODES memory pointer of type CVodeMem.
convfail is an input flag used to indicate any problem that occurred during the solution of the nonlinear equation on the current time step for which the linear solver is being used. This flag can be used to help decide whether the Jacobian data kept by a CVODES linear solver needs to be updated or not. Its possible values are:

- NO_FAILURES: this value is passed to lsetup if either this is the first call for this step, or the local error test failed on the previous attempt at this step (but the Newton iteration converged).
- FAIL_BAD_J: this value is passed to lsetup if (a) the previous Newton corrector iteration did not converge and the linear solver's setup function indicated that its Jacobian-related data is not current, or (b) during the previous Newton corrector iteration, the linear solver's solve function failed in a recoverable manner and the linear solver's setup function indicated that its Jacobian-related data is not current.
- FAIL_OTHER: this value is passed to lsetup if during the current internal step try, the previous Newton iteration failed to converge even though the linear solver was using current Jacobian-related data.
ypred is the predicted y vector for the current CVODES internal step.
fpred is the value of the right-hand side at ypred, i.e. $f\left(t_{n}, y_{\text {pred }}\right)$.
jcurPtr is a pointer to a boolean to be filled in by lsetup. The function should set *jcurPtr = TRUE if its Jacobian data is current after the call, and should set $*$ jcurPtr $=$ FALSE if its Jacobian data is not current. If lsetup calls for
reevaluation of Jacobian data (based on convfail and CVODES state data), it should return $*$ jcurPtr $=$ TRUE unconditionally; otherwise an infinite loop can result.
vtemp1
vtemp2
vtemp3 are temporary variables of type N_Vector provided for use by lsetup.
Return value An lsetup function should return 0 if successful, a positive value for a recoverable error, and a negative value for an unrecoverable error.


### 9.3 Solve function

The type definition of lsolve is

## lsolve

Definition int (*lsolve)(CVodeMem cv_mem, N_Vector b, N_Vector weight, N_Vector ycur, N_Vector fcur);
Purpose The function lsolve must solve the linear equation $M x=b$, where $M$ is some approximation to $I-\gamma J, J=(\partial f / \partial y)\left(t_{n}, y_{c u r}\right)$ (see Eq.(3.6)), and the right-hand side vector $b$ is input.

Arguments cv_mem is the CVODES memory pointer of type CVodeMem.
$\mathrm{b} \quad$ is the right-hand side vector $b$. The solution is to be returned in the vector b . weight is a vector that contains the error weights. These are the $W_{i}$ of Eq.(3.7).
ycur is a vector that contains the solver's current approximation to $y\left(t_{n}\right)$.
fcur is a vector that contains $f\left(t_{n}, y_{c u r}\right)$.
Return value An lsolve function should return a positive value for a recoverable error and a negative value for an unrecoverable error. Success is indicated by a 0 return value.

### 9.4 Memory deallocation function

The type definition of lfree is

## lfree

Definition void (*lfree) (CVodeMem cv_mem);
Purpose The function lfree should free up any memory allocated by the linear solver.
Arguments The argument cv_mem is the CVODES memory pointer of type CVodeMem.
Return value An 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 10

## Generic Linear Solvers in SUNDIALS

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

- The DENSE matrix package, which includes the matrix type DenseMat, macros and functions for DenseMat matrices, and functions for small dense matrices treated as simple array types.
- The BAND matrix package, which includes the matrix type BandMat, macros and functions for BandMat matrices.
- The SPGMR package, which includes a solver for the scaled preconditioned GMRES method.
- The spbcg package, which includes a solver for the scaled preconditioned Bi-CGStab method.
- The SPTFQMR package, which includes a solver for the scaled preconditioned TFQMR method.

For reasons related to installation, the names of the files involved in these generic solvers begin with the prefix sundials_. But despite this, each of the solvers is in fact generic, in that it is usable completely independently of SUNDIALS.

For the sake of space, the functions for DenseMat and BandMat matrices and the functions in SPGMR, SPBCG and SPTFQMR are only summarized briefly, since they are less likely to be of direct use in connection with CVODES. The functions for small dense matrices are fully described, because we expect that they will be useful in the implementation of preconditioners used with the combination of CVODES and the CVSPGMR, CVSPBCG or CVSPTFQMR solver.

### 10.1 The DENSE module

Relative to the SUNDIALS srcdir, the files comprising the DENSE generic linear solver are as follows:

- header files (located in srcdir/include/sundials) sundials_dense.h sundials_smalldense.h sundials_types.h sundials_math.h sundials_config.h
- source files (located in srcdir/src/sundials)
sundials_dense.c sundials_smalldense.c sundials math.c
Only two of the preprocessing directives in the header file sundials_config.h are relevant to the DENSE package by itself (see $\S 2.4$ 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 ABS macro and RAbs function.

The eight files listed above can be extracted from the sundials srcdir and compiled by themselves into a DENSE library or into a larger user code.

### 10.1.1 Type DenseMat

The type DenseMat is defined to be a pointer to a structure with the number of rows, number of columns, and a data field:

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

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

### 10.1.2 Accessor Macros

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

- DENSE_ELEM

Usage : DENSE_ELEM (A,i,j) = a_ij; or a_ij = DENSE_ELEM(A,i,j);
DENSE_ELEM references the ( $\mathrm{i}, \mathrm{j}$ )-th element of the $M \times N$ DenseMat A, $0 \leq \mathrm{i}<M, 0 \leq \mathrm{j}<N$.

- DENSE_COL

Usage : col_j = DENSE_COL(A,j);
DENSE_COL references the $j$-th column of the $M \times N$ DenseMat A, $0 \leq j<N$. 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 ( $\mathbf{i}, \mathrm{j})$-th element of A is referenced by col_j[i].

### 10.1.3 Functions

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

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


### 10.1.4 Small Dense Matrix Functions

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

- denalloc
denalloc ( $\mathrm{m}, \mathrm{n}$ ) 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 denalloc returns NULL. The underlying type of the dense matrix returned is realtype**. If we allocate a dense matrix realtype** a by $a=$ denalloc $(m, n)$, then $a[j][i]$ references the ( $i, j$ )-th element of the matrix $\mathrm{a}, 0 \leq \mathrm{i}<\mathrm{m}, 0 \leq \mathrm{j}<n$, and $\mathrm{a}[\mathrm{j}]$ is a pointer to the first element in the j -th column of a. The location a[0] contains a pointer to $m \times n$ contiguous locations which contain the elements of a.


## - denallocpiv

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

- denGETRF
denGETRF ( $a, m, n, p$ ) 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. $\mathrm{p}[\mathrm{k}]$ contains the row number of the pivot element chosen at the beginning of elimination step $\mathrm{k}, \mathrm{k}=0,1, \ldots, \mathrm{n}-1$.
2. If the unique LU factorization of a is given by $P a=L U$, 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.
denGETRF 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.

## - denGETRS

denGETRS ( $\mathrm{a}, \mathrm{n}, \mathrm{p}, \mathrm{b}$ ) solves the n by n linear system $a x=b$. It assumes that a (of size $\mathrm{n} \times \mathrm{n}$ ) has been LU-factored and the pivot array $p$ has been set by a successful call to denGETRF ( $a, n, n, p$ ). The solution $x$ is written into the b array.

- denzero
denzero ( $\mathrm{a}, \mathrm{m}, \mathrm{n}$ ) sets all the elements of the m by n dense matrix a to be 0.0 ;
- dencopy
dencopy ( $\mathrm{a}, \mathrm{b}, \mathrm{m}, \mathrm{n}$ ) copies the m by n dense matrix a into the m by n dense matrix b ;
- denscale
denscale ( $\mathrm{c}, \mathrm{a}, \mathrm{m}, \mathrm{n}$ ) scales every element in the m by n dense matrix a by c ;
- denaddI
denaddI ( $\mathrm{a}, \mathrm{n}$ ) increments the n by n dense matrix a by the identity matrix;
- denfreepiv
denfreepiv(p) frees the pivot array p allocated by denallocpiv;
- denfree
denfree (a) frees the dense matrix a allocated by denalloc;
- denprint
denprint ( $a, m, n$ ) prints the $m$ by $n$ dense matrix a to standard output as it would normally appear on paper. It is intended as a debugging tool with small values of $n$. The elements are printed using the $\% \mathrm{~g}$ option. A blank line is printed before and after the matrix.


### 10.2 The BAND module

Relative to the SUNDIALS srcdir, the files comprising the BAND generic linear solver are as follows:

- header files (located in srcdir/include/sundials)
sundials_band.h
sundials_types.h sundials_math.h sundials_config.h
- source files (located in srcdir/src/sundials)
sundials_band.c sundials_math.c
Only two of the preprocessing directives in the header file sundials_config.h are required to use the BAND package by itself (see $\S 2.4$ 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 of the SUNDIALS realtype and booleantype types and the macro RCONST, while the sundials_math. h header file is needed for the MIN, MAX, and ABS macros and RAbs function.

The six files listed above can be extracted from the SUNDIALS srcdir and compiled by themselves into a BAND library or into a larger user code.

### 10.2.1 Type BandMat

The type BandMat is the type of a large band matrix A (possibly distributed). It is defined to be a pointer to a structure defined by:

```
typedef struct {
    long int size;
    long int mu, ml, smu;
    realtype **data;
} *BandMat;
```

The fields in the above structure are:

- size is the number of columns (which is the same as the number of rows);
- $m u$ is the upper half-bandwidth, $0 \leq m u \leq$ size -1 ;
- $m l$ is the lower half-bandwidth, $0 \leq m l \leq$ size -1 ;
- $s m u$ is the storage upper half-bandwidth, $m u \leq s m u \leq s i z e-1$. The BandGBTRF routine writes the LU factors into the storage for A . The upper triangular factor U , however, may have an upper half-bandwidth as big as $\min ($ size $-1, m u+m l)$ because of partial pivoting. The $s m u$ field holds the upper half-bandwidth allocated for A.
- data is a two dimensional array used for component storage. The elements of a band matrix of type BandMat 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.
If we number rows and columns in the band matrix starting from 0 , then
- data[0] is a pointer to $(s m u+m l+1) *$ size contiguous locations which hold the elements within the band of A
- data[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 $s m u-m u$ (to access the uppermost element within the band in the j -th column) to $s m u+m l$ (to access the lowest element within the band in the j-th column). Indices from 0 to $s m u-m u-1$ give access to extra storage elements required by BandGBTRF.
- data[j][i-j+smu] is the $(i, j)$-th element, $j-m u \leq i \leq j+m l$.

The macros below allow a user to 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 into the j-th column of elements can be obtained via the BAND_COL macro. Users should use these macros whenever possible.

See Figure 10.1 for a diagram of the BandMat type.

### 10.2.2 Accessor Macros

The following three macros are defined by the BAND module to provide access to data in the BandMat 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 $0 \leq \mathrm{i}, \mathrm{j} \leq N-1$. The location $(i, j)$ should further satisfy $j-(A->m u) \leq i \leq j+(A->m l)$.


Figure 10.1: Diagram of the storage for a band matrix of type BandMat. Here A is an $N \times N$ band matrix of type BandMat 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 $\mathrm{A}(\mathrm{i}, \mathrm{j})$. The greyed out areas of the underlying component storage are used by the BandGBTRF and BandGBTRS routines.

- 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, $0 \leq$ $\mathrm{j} \leq N-1$. The type of the expression BAND_COL $(\mathrm{A}, \mathrm{j})$ is realtype $*$. The pointer returned by the call BAND_COL $(A, j)$ can be treated as an array which is indexed from $-(A->m u)$ to ( $A->m l)$.

- BAND_COL_ELEM

Usage : BAND_COL_ELEM (col_j,i,j) = a_ij; or $a_{-} i j=$ BAND_COL_ELEM (col_j,i,j);
This macro references the ( $\mathbf{i}, \mathrm{j}$ )-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 $j-(A->m u)$ $\leq i \leq j+(A->m l)$.

### 10.2.3 Functions

The following functions for BandMat matrices are available in the BAND package. For full details, see the header file sundials_band.h.

- BandAllocMat: allocation of a BandMat matrix;
- BandAllocPiv: allocation of a pivot array for use with BandGBTRF/BandGBTRS;
- BandGBTRF: LU factorization with partial pivoting;
- BandGBTRS: solution of $A x=b$ using LU factorization;
- BandZero: load a matrix with zeros;
- BandCopy: copy one matrix to another;
- BandScale: scale a matrix by a scalar;
- BandAddI: increment a matrix by the identity matrix;
- BandFreeMat: free memory for a BandMat matrix;
- BandFreePiv: free memory for a pivot array;
- BandPrint: print a BandMat matrix to standard output.


### 10.3 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 (SPBCG and SPTFQMR). For full details, including usage instructions, see the header files sundials_spgmr.h and sundials_iterative.h.

Relative to the SUNDIALS srcdir, the files comprising the SPGMR generic linear solver 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 $\S 2.4$ 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 MAX and ABS macros and RAbs and RSqrt functions.

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 SPGMR package can only be used in conjunction with an actual NVECTOR implementation library, such as the NVECTOR_SERIAL or NVECTOR_PARALLEL provided with SUNDIALS.

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.

### 10.3.1 Functions

The following functions are available in the SPGMR package:

- SpgmrMalloc: allocation of memory for SpgmrSolve;
- SpgmrSolve: solution of $A x=b$ 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.


### 10.4 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 SPBCG package can only be used in conjunction with an actual NVECTOR implementation library, such as the NVECTOR_SERIAL or NVECTOR_PARALLEL provided with SUNDIALS.

The files needed to use the SPBCG module by itself are the same as for the SPGMR module, with sundials_spbcgs. (h,c) replacing sundials_spgmr. (h, c).

### 10.4.1 Functions

The following functions are available in the SPBCG package:

- SpbcgMalloc: allocation of memory for SpbcgSolve;
- SpbcgSolve: solution of $A x=b$ by the SPBCG method;
- SpbcgFree: free memory allocated by SpbcgMalloc.


### 10.5 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 SPTFQMR package can only be used in conjunction with an actual NVECTOR implementation library, such as the NVECTOR_SERIAL or NVECTOR_PARALLEL provided with SUNDIALS.

The files needed to use the SPTFQMR module by itself are the same as for the SPGMR module, with sundials_sptfqmr. (h, c) replacing sundials_spgmr. (h, c).

### 10.5.1 Functions

The following functions are available in the SPTFQMR package:

- SptfqmrMalloc: allocation of memory for SptfqmrSolve;
- SptfqmrSolve: solution of $A x=b$ by the SPTFQMR method;
- SptfqmrFree: free memory allocated by SptfqmrMalloc.


## Chapter 11

## CVODES 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.

### 11.1 CVODES input constants

|  | CVODES main solver module |  |
| :--- | :--- | :--- |
| CV_ADAMS | 1 | Adams-Moulton linear multistep method. |
| CV_BDF | 2 | BDF linear multistep method. |
| CV_FUNCTIONAL | 1 | Nonlinear system solution through functional iterations. |
| CV_NEWTON | 2 | Nonlinear system solution through Newton iterations. |
| CV_SS | 1 | Scalar relative tolerance, scalar absolute tolerance. |
| CV_SV | 2 | Scalar relative tolerance, vector absolute tolerance. |
| CV_EE | 2 | Estimated relative tolerance and absolute tolerance for sen- |
|  |  | sitivity variables. |
| CV_NORMAL | 1 | Solver returns at specified output time. |
| CV_ONE_STEP | 2 | Solver returns after each successful step. <br> CV_NORMAL_TSTOP |
|  | 3 | Solver returns at specified output time, but does not proceed |
| CV_ONE_STEP_TSTOP | 4 | past the specified stopping time. <br>  <br> Solver returns after each successful step, but does not pro- <br> ceed past the specified stopping time. |
| CV_SIMULTANEOUS | 1 | Simultaneous corrector forward sensitivity method. <br> CV_STAGGERED |
| CV_STAGGERED1 | 2 | Staggered corrector forward sensitivity method. <br> CV_CENTERED |
|  | 1 | Staggered (variant) corrector forward sensitivity method. |
| Central difference quotient approximation $\left(2^{\text {nd }}\right.$ order) of the |  |  |

CVODEA adjoint solver module
CV_HERMITE 1 Use Hermite interpolation.

CV_POLYNOMIAL 2 Use variable-degree polynomial interpolation.

## Iterative linear solver module

| PREC_NONE | 0 | No preconditioning |
| :--- | :--- | :--- |
| PREC_LEFT | 1 | Preconditioning on the left only. |
| PREC_RIGHT | 2 | Preconditioning on the right only. |
| PREC_BOTH | 3 | Preconditioning on both the left and the right. |
| MODIFIED_GS | 1 | Use modified Gram-Schmidt procedure. |
| CLASSICAL_GS | 2 | Use classical Gram-Schmidt procedure. |

### 11.2 CVODES output constants

| CVODES main solver module |  |  |
| :---: | :---: | :---: |
| CV_SUCCESS | 0 | Successful function return. |
| CV_TSTOP_RETURN | 1 | CVode succeeded by reaching the specified stopping point. |
| CV_ROOT_RETURN | 2 | CVode succeeded and found one or more roots. |
| CV_TOO_MUCH_WORK | -1 | The solver took mxstep internal steps but could not reach tout. |
| CV_TOO_MUCH_ACC | -2 | The solver could not satisfy the accuracy demanded by the user for some internal step. |
| CV_ERR_FAILURE | -3 | Error test failures occurred too many times during one internal time step or minimum step size was reached. |
| CV_CONV_FAILURE | -4 | Convergence test failures occurred too many times during one internal time step or minimum step size was reached. |
| CV_LINIT_FAIL | -5 | The linear solver's initialization function failed. |
| CV_LSETUP_FAIL | -6 | The linear solver's setup function failed in an unrecoverable manner. |
| CV_LSOLVE_FAIL | -7 | The linear solver's solve function failed in an unrecoverable manner. |
| CV_RHSFUNC_FAIL | -8 | The right-hand side function failed in an unrecoverable manner. |
| CV_FIRST_RHSFUNC_ERR | -9 | The right-hand side function failed at the first call. |
| CV_REPTD_RHSFUNC_ERR | -10 | The right-hand side function had repetead recoverable errors. |
| CV_UNREC_RHSFUNC_ERR | -11 | The right-hand side function had a recoverable error, but no recovery is possible. |
| CV_RTFUNC_FAIL | -12 | The rootfinding function failed in an unrecoverable manner. |
| CV_MEM_FAIL | -20 | A memory allocation failed. |
| CV_MEM_NULL | -21 | The cvode_mem argument was NULL. |
| CV_ILL_INPUT | -22 | One of the function inputs is illegal. |
| CV_NO_MALLOC | -23 | The cVode memory block was not allocated by a call to CVodeMalloc. |
| CV_BAD_K | -24 | The derivative order $k$ is larger than the order used. |
| CV_BAD_T | -25 | The time $t$ s outside the last step taken. |
| CV_BAD_DKY | -26 | The output derivative vector is NULL. |
| CV_TOO_CLOSE | -27 | The output and initial times are too close to each other. |
| CV_NO_QUAD | -30 | Quadrature integration was not activated. |
| CV_QRHSFUNC_FAIL | -31 | The quadrature right-hand side function failed in an unrecoverable manner. |


| CV_FIRST_QRHSFUNC_ERR | -32 | The quadrature right-hand side function failed at the first <br> call. |
| :--- | :--- | :--- |
| CV_REPTD_QRHSFUNC_ERR | -33 | The quadrature ight-hand side function had repetead recov- <br> erable errors. |
| CV_UNREC_QRHSFUNC_ERR | -34 | The quadrature right-hand side function had a recoverable <br> error, but no recovery is possible. <br> The sensitivity index is larger than the number of sensitivi- <br> ties computed. |
| CV_BAD_IS | -40 | -41 |
| CV_NO_SENS | -42 | Forward sensitivity integration was not activated. <br> The sensitivity right-hand side function failed in an unre- <br> coverable manner. <br> The sensitivity right-hand side function failed at the first <br> call. |
| CV_SRHSFUNC_FAIL | -43 | The sensitivity ight-hand side function had repetead recov- <br> erable errors. <br> The sensitivity right-hand side function had a recoverable <br> error, but no recovery is possible. |
| CV_REPTD_SRHSFUNC_ERR | -44 |  |


|  | CVODEA adjoint solver module |  |
| :--- | :--- | :--- |
| CV_ADJMEM_NULL | -101 | The cvadj_mem argument was NULL. |
| CV_BAD_TBO | -103 | The final time for the adjoint problem is outside the interval <br> over which the forward problem was solved. <br> The cvodes memory for the backward problem was not cre- <br> ated. |
| CV_BCKMEM_NULL | -104 | -105 |
| Reinitialization of the forward problem failed at the first |  |  |
| checkpoint. |  |  |
| CV_REIFWD_FAIL | -106 | An error occured during the integration of the forward prob- <br> lem. |
| CV_FWD_FAIL | -107 | Wrong task for backward integration. <br> The desired output time is outside the interval over which <br> CV_BAD_ITASK |
| CV_BAD_TBOUT | -108 | the forward problem was solved. <br> CV_GETY_BADT |

```
CVBAND_SUCCESS 0 Successful function return.
CVBAND_MEM_NULL -1 The cvode_mem argument was NULL.
CVBAND_LMEM_NULL
    -2 The CVBAND linear solver has not been initialized.
CVBAND_ILL_INPUT
CVBAND_MEM_FAIL
CVBAND_JACFUNC_UNRECVR
CVBAND_JACFUNC_RECVR
CVBAND_ADJMEM_NULL
CVBAND_LMEMB_NULL
```

0 Successful function return.
-1 The cvode_mem argument was NULL.
-2 The cVBAND linear solver has not been initialized.
-3 The cVBand solver is not compatible with the current NVECTOR module, or an input value was illegal.
-4 A memory allocation request failed.
-5 The Jacobian function failed in an unrecoverable manner.
-6 The Jacobian function had a recoverable error.
-101 The cvadj mem argument was NULL.
-102 The CVBAND linear solver has not been initialized for the backward integration.

|  | CVDIAG linear solver module |  |
| :--- | ---: | :--- |
| CVDIAG_SUCCESS | 0 | Successful function return. |
| CVDIAG_MEM_NULL | -1 | The cvode_mem argument was NULL. |
| CVDIAG_LMEM_NULL | -2 | The CVDIAG linear solver has not been initialized. |
| CVDIAG_ILL_INPUT | -3 | The CVDIAG solver is not compatible with the current NVEC- |
|  |  | TOR module. |
| CVDIAG_MEM_FAIL | -4 | A memory allocation request failed. |
| CVDIAG_ADJMEM_NULL | -101 | The cvadj mem argument was NULL. |

CVSPILS linear solver modules

|  | CVSPILS linear solver modules |  |
| :--- | ---: | :--- |
| CVSPILS_SUCCESS | 0 | Successful function return. |
| CVSPILS_MEM_NULL | -1 | The cvode_mem argument was NULL. |
| CVSPILS_LMEM_NULL | -2 | The linear solver has not been initialized. |
| CVSPILS_ILL_INPUT | -3 | The solver is not compatible with the current NVECTOR |
|  |  | module, or an input value was illegal. |
| CVSPILS_MEM_FAIL | -4 | A memory allocation request failed. |
| CVSPILS_ADJMEM_NULL | -101 | The cvadj_mem argument was NULL. |
| CVSPILS_LMEMB_NULL | -102 | The linear solver has not been initialized for the backward |
|  |  | integration. |

## 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 function 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 function failed unrecoverably.
```


## 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 function 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. |
| SPBCG_PSET_FAIL_UNREC | -4 | The preconditioner setup function 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 function 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 function failed unrecoverably. |

CVBANDPRE preconditioner module

| CVBANDPRE_SUCCESS | 0 | Successful function return. |
| :--- | ---: | :--- |
| CVBANDPRE_PDATA_NULL | -11 | The preconditioner module has not been initialized. |
| CVBANDPRE_RHSFUNC_UNRECVR | -12 | The right-hand side function failed unrecoverably. |
| CVBANDPRE_ADJMEM_NULL | -111 | The cvadj_mem argument was NULL. |
| CVBANDPRE_MEM_FAIL | -112 | A memory allocation failed. |

## CVBBDPRE preconditioner module

| CVBBDPRE_SUCCESS | 0 | Successful function return. |
| :--- | ---: | :--- |
| CVBBDPRE_PDATA_NULL | -11 | The preconditioner module has not been initialized. |
| CVBBDPRE_FUNC_UNRECVR | -12 | A user supplied function failed unrecoverably. |
| CVBBDPRE_ADJMEM_NULL | -111 | The cvadj_mem argument was NULL. |
| CVBBDPRE_PDATAB_NULL | -112 | The CVBBDPRE preconditionr module has not been initial- <br> ized for the backward integration. |
| CVBBDPRE_MEM_FAIL | -113 | A memory allocation failed. |

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## Index

Adams method, 15
adjoint sensitivity analysis
checkpointing, 23
implementation in CVODES, 24, 29
mathematical background, 21-24
quadrature evaluation, 127
right-hand side evaluation, 126
BAND generic linear solver
functions, 159
macros, 157-159
type BandMat, 157
BAND_COL, 71, 159
BAND_COL_ELEM, 71, 159
BAND_ELEM, 71, 157
BandMat, 33, 71, 129, 157
BDF method, 15
Bi-CGStab method, 39, 51, 122, 160
BIG_REAL, 32, 143
CLASSICAL_GS, 51, 121
CV_ADAMS, $35,66,116$
CV_ADJMEM_NULL, 114-117, 119, 123-126
CV_BAD_DKY, 52, 77, 98-100
Cv_BAD_DKY, 124
CV_BAD_IS, 99, 100
CV_BAD_ITASK, 118
CV_BAD_K, 52, 77, 99, 100
CV_BAD_T, 52, 77, 98-100
CV_BAD_TBO, 116, 117
CV_BAD_TBOUT, 119
CV_BCKMEM_NULL, 119
CV_BDF, 35, 66, 116
CV_CENTERED, 102
CV_CONV_FAILURE, 41, 115, 119
CV_EE, 102
CV_ERR_FAILURE, 41, 115, 119
CV_FIRST_QRHSFUNC_ERR, 79
CV_FIRST_QRHSFUNC_FAIL, 76
CV_FIRST_RHSFUNC_ERR, 68
CV_FIRST_RHSFUNC_FAIL, 41
CV_FIRST_SRHSFUNC_ERR, 108, 109
CV_FIRST_SRHSFUNC_FAIL, 98
CV_FORWARD, 102
CV_FUNCTIONAL, 35, 47, 116

CV_FWD_FAIL, 119
CV_HERMITE, 113, 114
CV_ILL_INPUT, 36, 40, 44, 45, 47, 67, 78, 96, 97, $101-103,114,117,118,126$
CV_ILL_IPUT, 115
CV_LINIT_FAIL, 41
CV_LSETUP_FAIL, $41,69,70,86,115,119,128,129$, 137, 138
CV_LSOLVE_FAIL, 41, 115
CV_MEM_FAIL, 36, 75, 96, 97, 114-117, 123
CV_MEM_NULL, $36,40,43-48,52,54-59,67,75,77-$ 81, 96-107, 117, 123, 124
CV_NEWTON, 35, 47, 116
CV_NO_MALLOC, 40, 67, 115, 117, 118
CV_NO_QUAD, 75, 77-79, 124
CV_NO_SENS, 97-101, 103-107
CV_NORMAL, 40, 112, 115, 118
CV_NORMAL_TSTOP, 40, 115
CV_ONE_STEP, 40, 112, 115, 118
CV_ONE_STEP_TSTOP, 40, 115
CV_POLYNOMIAL, 113, 114
CV_QRHSFUNC_FAIL, 76, 79, 127
CV_REIFWD_FAIL, 119
CV_REPTD_QRHSFUNC_ERR, 76
CV_REPTD_RHSFUNC_ERR, 41
CV_REPTD_SRHSFUNC_ERR, 98
CV_RHSFUNC_FAIL, 41, 68, 127
CV_ROOT_RETURN, 40
CV_RTFUNC_FAIL, 41, 81
CV_SIMULTANEOUS, 27, 96, 107
CV_SOLVE_FAIL, 119
CV_SRHSFUNC_FAIL, 98, 108, 109
CV_SS, 35, 47, 67, 77, 102, 116, 117
CV_STAGGERED, 27, 96, 107
CV_STAGGERED1, 29, 96, 108
CV_SUCCESS, 36, 40, 43-48, 52, 54-59, 67, 75-81, 96-107, 114-118, 123-126
CV_SV, $35,47,67,77,102,116,117$
CV_TOO_CLOSE, 41
CV_TOO_MUCH_ACC, 41, 115, 119
CV_TOO_MUCH_WORK, 41, 115, 119
CV_TSTOP_RETURN, 40, 115
CV_UNREC_QRHSFUNC_ERR, 80
CV_UNREC_RHSFUNC_ERR, 41, 68, 76
CV_UNREC_SRHSFUNC_ERR, 98, 108, 109

CV_WARNING, 68
CV_WF, 35, 67
CVadjCheckPointRec, 125
CVadjFree, 114
CVadjGetCheckPointsInfo, 124
CVadjGetCheckPointsList, 125
CvadjGetCVodeBmem, 123
CVadjGetDataPointHermite, 125
CVadjGetDataPointPolynomial, 126
CVadjGetReturnFlagName, 126
CVadjMalloc, 112, 113
CVadjSetInterpType, 114
CVBAND linear solver
Jacobian approximation used by, 49
memory requirements, 61
NVECTOR compatibility, 38
optional input, 49, 120
optional output, 61-62
selection of, 38
CVBand, 34, 37, 38, 70
CVBAND_ADJMEM_NULL, 120
CVBAND_ILL_INPUT, 38
CVBAND_JACFUNC_RECVR, 70, 129
CVBAND_JACFUNC_UNRECVR, 70, 129, 130
CVBAND_LMEM_NULL, 49, 61, 62, 120
CVBAND_MEM_FAIL, 38
CVBAND_MEM_NULL, $38,49,61,62,120$
CVBAND_SUCCESS, $38,49,61,62,120$
CVBandB, 129
CVBandDQJac, 49
CVBandGetLastFlag, 62
CVBandGetNumJacEvals, 61
CVBandGetNumRhsEvals, 61
CVBandGetReturnFlagName, 62
CVBandGetWorkSpace, 61
CVBandJacFn, 70
CVBANDPRE preconditioner
description, 81
optional output, 84-85
usage, 81-82
usage with adjoint module, 132-134
user-callable functions, 82-84, 132-134
CVBANDPRE_ADJMEM_NULL, 132-134
CVBANDPRE_MEM_FAIL, 132
CVBANDPRE_PDATA_NULL, 83-85, 133, 134
CVBANDPRE_SUCCESS, 84, 132
CVBandPrecAlloc, 82
CVBandPrecAllocB, 132
CVBandPrecFree, 84
CVBandPrecFreeB, 134
CVBandPrecGetNumRhsEvals, 84
CVBandPrecGetReturnFlagName, 85
CVBandPrecGetWorkSpace, 84
CVBandSetJacFn, 49

CVBandSetJacFnB, 120
CVBBDPRE preconditioner
description, 85-86
optional output, 90-91
usage, 87-88
usage with adjoint module, 134-138
user-callable functions, 88-90, 134-137
user-supplied functions, 86-87, 137-138
CVBBDPRE_ADJMEM_NULL, 135, 136
CVBBDPRE_PDATA_NULL, 89-91, 135, 136
CVBBDPRE_SUCCESS, 90, 91, 135, 136
CVBBDPrecAlloc, 88
CVBBDPrecAllocB, 134
CVBBDPrecFree, 90
CVBBDPrecFreeB, 137
CVBBDPrecGetNumGfnEvals, 91
CVBBDPrecGetReturnFlagName, 91
CVBBDPrecGetWorkSpace, 90
CVBBDPrecReInit, 90
CVBBDPrecReInitB, 136
CVBBDSpbcg, 89
CVBBDSpbcgB, 135
CVBBDSpgmr, 88
CVBBDSpgmrB, 135
CVBBDSptfqmr, 89
CVBBDSptfqmrB, 136
CVBPSpbcg, 83
CVBPSpbcgB, 133
CVBPSpgmr, 83
CVBPSpgmrB, 133
CVBPSptfqmr, 83
CVBPSptfqmrB, 133
CVDENSE linear solver
Jacobian approximation used by, 48
memory requirements, 59
NVECTOR compatibility, 37
optional input, 48, 119-120
optional output, 59-61
selection of, 37
CVDense, 34, 37, 69
CVDENSE_ADJMEM_NULL, 120
CVDENSE_ILL_INPUT, 38
CVDENSE_JACFUNC_RECVR, 69, 128
CVDENSE_JACFUNC_UNRECVR, 69, 128, 129
CVDENSE_LMEM_NULL, 48, 59, 60, 120
CVDENSE_MEMFAIL, 38
CVDENSE_MEM_NULL, $38,48,59,60,120$
CVDENSE_SUCCESS, $38,48,59,60,120$
CVDenseB, 128
CVDenseDQJac, 48
CVDenseGetLastFlag, 60
CVDenseGetNumJacEvals, 60
CVDenseGetNumRhsEvals, 60
CVDenseGetReturnFlagName, 61

CVDenseGetWorkSpace, 59
CVDenseJacFn, 69
CVDenseSetJacFn, 48
CVDenseSetJacFnB, 119
CVDIAG linear solver
Jacobian approximation used by, 38
memory requirements, 62
optional output, 62-63
selection of, 38
CVDiag, 34, 37, 38
CVDIAG_ILL_INPUT, 38
CVDIAG_LMEM_NULL, 63
CVDIAG_MEM_FAIL, 38
CVDIAG_MEM_NULL, 38, 62, 63
CVDIAG_SUCCESS, 38, 62, 63
CVDiagGetLastFlag, 63
CVDiagGetNumRhsEvals, 63
CVDiagGetReturnFlagName, 63
CVDiagGetWorkSpace, 62
CVErrHandlerFn, 68
CVEwtFn, 69
CVODE, 1
CVode, 34, 40
CVodeB, 113, 118
CVodeCreate, 35
CVodeCreateB, 112, 116
CVodeF, 112, 114
CVodeFree, 34, 36
CVodeGetActualInitStep, 56
CVodeGetCurrentOrder, 56
CVodeGetCurrentStep, 56
CVodeGetCurrentTime, 57
CVodeGetDky, 52
CVodeGetErrWeights, 57
CVodeGetEstLocalErrors, 58
CVodeGetIntegratorStats, 58
CVodeGetLastOrder, 55
CVodeGetLastStep, 56
CVodeGetNonlinSolvStats, 59
CVodeGetNumErrTestFails, 55
CVodeGetNumGEvals, 81
CVodeGetNumLinSolvSetups, 55
CVodeGetNumNonlinSolvConvFails, 59
CVodeGetNumNonlinSolvIters, 58
CVodeGetNumRhsEvals, 55
CVodeGetNumRhsEvalsSEns, 104
CVodeGetNumSensErrTestFails, 104
CVodeGetNumSensLinSolvSetups, 104
CVodeGetNumSensNonlinSolvConvFails, 106
CVodeGetNumSensNonlinSolvIters, 105
CVodeGetNumSensRhsEvals, 103
CVodeGetNumStabLimOrderReds, 57
CVodeGetNumSteps, 54
CVodeGetNumStgrSensNonlinSolvConvFails, 107

CVodeGetNumStgrSensNonlinSolvIters, 106
CVodeGetQuad, 124
CVodeGetQuadB, 113
CVodeGetQuadDky, 76
CVodeGetQuadErrWeights, 78
CVodeGetQuadNumErrTestFails, 78
CVodeGetQuadNumRhsEvals, 78
CVodeGetQuadStats, 79
CVodeGetReturnFlagName, 59
CVodeGetRootInfo, 80
CVodeGetSens, 95
CVodeGetSensDky, 98
CVodeGetSensErrWeights, 105
CVodeGetSensNonlinSolvStats, 106
CVodeGetSensStats, 105
CVodeGetTolScaleFactor, 57
CVodeGetWorkSpace, 54
CVodeMalloc, 35, 66
CVodeMallocB, 112, 116, 117
CVodeQuadMalloc, 75
CVodeQuadMallocB, 123
CVodeReInit, 66
CVodeReInitB, 117
CVodeRootInit, 80
CVODES
brief description of, 1
motivation for writing in C, 2
package structure, 27
relationship to CVODE, PVODE, 1-2
relationship to VODE, VODPK, 1
CVODES linear solvers
built on generic solvers, 37
CVBAND, 38
CVDENSE, 37
CVDIAG, 38
CVSPBCG, 39
CVSPGMR, 39
CVSPTFQMR, 39
header files, 32
implementation details, 30
list of, 29
NVECTOR compatibility, 31
selecting one, 37
usage with adjoint module, 117-118
cvodes.h, 32
cvodes_band.h, 33
cvodes_dense.h, 32
cvodes_diag.h, 33
cvodes_spbcgs.h, 33
cvodes_spgmr.h, 33
cvodes_sptfqmr.h, 33
CVodeSensFree, 97
CVodeSensMalloc, 95, 96
CVodeSensReInit, 96, 97

CVodeSensToggleOff, 97
CVodeSetErrFile, 43
CVodeSetErrHandlerFn, 43
CVodeSetEwtFn, 48
CVodeSetFdata, 43
CVodeSetInitStep, 45
CVodeSetIterType, 47
CVodeSetMaxConvFails, 46
CVodeSetMaxErrTestFails, 46
CVodeSetMaxHnilWarns, 44
CVodeSetMaxNonlinIters, 46
CVodeSetMaxNumSteps, 44
CVodeSetMaxOrder, 44
CVodeSetMaxStep, 45
CVodeSetMinStep, 45
CVodeSetNonlinConvCoef, 47
CVodeSetQuadErrCon, 77
CVodeSetQuadFdata, 77
CVodeSetSensDQMethod, 101
CVodeSetSensErrCon, 102
CVodeSetSensMaxNonlinIters, 103
CVodeSetSensParams, 101
CVodeSetSensRhs1Fn, 101
CVodeSetSensRhsFn, 100
CVodeSetSensTolerances, 102
CVodeSetStabLimDet, 44
CVodeSetStopTime, 46
CVodeSetTolerances, 47
CVQuadRhsFn, 75, 79
CVQuadRhsFnB, 123, 127
CVRhsFn, 35, 67
CVRhsFnB, 116, 117, 126
CVRootFn, 81
CVSensRhs1Fn, 108
CVSensRhsFn, 107
CVSPBCG linear solver
Jacobian approximation used by, 49
memory requirements, 63
optional input, 49-51, 120-122
optional output, 63-66
preconditioner setup function, 49, 72, 131
preconditioner solve function, 49, 72, 130
selection of, 39
CVSpbcg, 34, 37, 39
CVSPGMR linear solver
Jacobian approximation used by, 49
memory requirements, 63
optional input, 49-51, 120-122
optional output, 63-66
preconditioner setup function, 49, 72, 131
preconditioner solve function, $49,72,130$
selection of, 39
CVSpgmr, 34, 37, 39
CVSPILS_ADJMEM_NULL, 121, 122

CVSPILS_ILL_INPUT, 39, 40, 50, 51, 83, 84, 89, 121, 122, 133-136
CVSPILS_LMEM_NULL, 50, 51, 64-66, 121, 122
CVSPILS_MEM_FAIL, 39, 40, 83, 84, 89, 133-136
CVSPILS_MEM_NULL, $39,40,50,51,64-66,83,84$, 89, 121, 122, 133-136
CVSPILS_SUCCESS, 39, 40, 50, 51, 64-66, 83, 84, 89, 120-122, 133, 135, 136
CVSpilsDQJtimes, 49
CVSpilsGetLastFlag, 66
CVSpilsGetNumConvFails, 64
CVSpilsGetNumJtimesEvals, 65
CVSpilsGetNumLinIters, 64
CVSpilsGetNumPrecEvals, 64
CVSpilsGetNumPrecSolves, 65
CVSpilsGetNumRhsEvals, 65
CVSpilsGetReturnFlagName, 66
CVSpilsGetWorkSpace, 64
CVSpilsJacTimesVecFn, 71
CVSpilsJacTimesVecFnB, 130
CVSpilsPrecSetupFn, 72
CVSpilsPrecSetupFnB, 131
CVSpilsPrecSolveFn, 72
CVSpilsPrecSolveFnB, 130
CVSpilsSetDelt, 51
CVSpilsSetDeltB, 121
CVSpilsSetGSType, 51
CVSpilsSetGSTypeB, 121
CVSpilsSetJacTimesFn, 50
CVSpilsSetJacTimesFnB, 121
CVSpilsSetMaxl, 51
CVSpilsSetMaxlB, 122
CVSpilsSetPreconditioner, 50
CVSpilsSetPrecSolveFnB, 120
CVSpilsSetPrecType, 50
CVSpilsSetPrecTypeB, 122
CVSPTFQMR linear solver
Jacobian approximation used by, 49
memory requirements, 63
optional input, 49-51, 120-122
optional output, 63
preconditioner setup function, 49, 72, 131
preconditioner solve function, 49, 72, 130
selection of, 39
CVSptfqmr, 34, 37, 39
denaddI, 156
denalloc, 155
denallocpiv, 155
dencopy, 156
denfree, 156
denfreepiv, 156
denGETRF, 155
denGETRS, 156
denprint, 156
denscale, 156
DENSE generic linear solver
functions
large matrix, 154-155
small matrix, 155-156
macros, 154
type DenseMat, 154
DENSE_COL, 70, 154
DENSE_ELEM, 70, 154
DenseMat, 32, 69, 129, 154
denzero, 156
e_data, 69
eh_data, 68
error control
order selection, 18
sensitivity variables, 20
step size selection, 17-18
error messages, 41, 43
user-defined handler, 43, 68
f_data, $43,68,86$
f_dataB, $127,137,138$
forward sensitivity analysis
absolute tolerance selection, 20-21
correction strategies, 19-20, 27, 96, 97
mathematical background, 18-21
right hand side evaluation, 21
right-hand side evaluation, 21, 107-109
fQ_data, 77, 79
fQ_dataB, 127
fS_data, 108, 109
g_data, 81
generic linear solvers
BAND, 156
DENSE, 153
SPBCG, 160
SPGMR, 159
SPTFQMR, 161
use in Cvodes, 30
GMRES method, 39, 159
Gram-Schmidt procedure, 51, 121
half-bandwidths, $38,70-71,82,88$
header files, $32,81,87$
itask, 34, 40, 115
iter, 35, 47
itol, $35,116,117$
itolQ, 77
itols, 102
Jacobian approximation function
band
difference quotient, 49
user-supplied, 49, 70-71
user-supplied (backward), 120, 129
dense
difference quotient, 48
user-supplied, 48, 69-70
user-supplied (backward), 119, 128
diagonal
difference quotient, 38
Jacobian times vector
difference quotient, 49
user-supplied, 50
Jacobian-vector product
user-supplied, 71-72
user-supplied (backward), 121, 130
linit, 150
1mm, 35, 66
LSODE, 1
$\max 1,39,40,83,84,89$
maxord, 44, 66
memory requirements
CVBAND linear solver, 61
CVBANDPRE preconditioner, 84
CVBBDPRE preconditioner, 91
CVDENSE linear solver, 59
CVDIAG linear solver, 62
CVODES solver, 75, 96
CVODES solver, 54
CVSPGMR linear solver, 63
MODIFIED_GS, 51, 121
MPI, 4
N_VCloneEmptyVectorArray, 140
N_VCloneVectorArray, 140
N_VCloneVectorArray_Parallel, 146
N_VCloneVectorArray_Serial, 144
N_VCloneVectorArrayEmpty_Parallel, 146
N_VCloneVectorArrayEmpty_Serial, 144
N_VDestroyVectorArray, 140
N_VDestroyVectorArray_Parallel, 147
N_VDestroyVectorArray_Serial, 144
N_Vector, 32, 139
N_VMake_Parallel, 146
N_VMake_Serial, 144
N_VNew_Parallel, 146
N_VNew_Serial, 144
N_VNewEmpty_Parallel, 146
N_VNewEmpty_Serial, 144
N_VPrint_Parallel, 147
N_VPrint_Serial, 144
nonlinear system
definition, 15-16

Newton convergence test, 17
Newton iteration, 16-17
NV_COMM_P, 146
NV_CONTENT_P, 145
NV_CONTENT_S, 143
NV_DATA_P, 145
NV_DATA_S, 143
NV_GLOBLENGTH_P, 145
NV_Ith_P, 146
NV_Ith_S, 144
NV_LENGTH_S, 143
NV_LOCLENGTH_P, 145
NV_OWN_DATA_P, 145
NV_OWN_DATA_S, 143
NVECTOR module, 139
nvector_parallel.h, 32
nvector_serial.h, 32
optional input
backward solver, 119
band linear solver, 49, 120
dense linear solver, 48, 119-120
forward sensitivity, 100-103
iterative linear solver, 49-51, 120-122
quadrature integration, 77-78, 124
solver, 43-48
optional output
backward solver, 123
band linear solver, 61-62
band-block-diagonal preconditioner, 90-91
banded preconditioner, 84-85
checkpoint information, 124
dense linear solver, 59-61
diagonal linear solver, 62-63
forward sensitivity, 103-107
interpolated quadratures, 76
interpolated sensitivities, 98
interpolated solution, 52
interpolation data, 125
iterative linear solver, 63-66
quadrature integration, 78-79, 124
solver, 52-59
output mode, 18, 40, 115, 118
partial error control
explanation of CVODES behavior, 109
portability, 32
PREC_BOTH, 39, 40, 50, 122
PREC_LEFT, $39,40,50,83,84,89,122,133,135$, 136
PREC_NONE, 39, 40, 50, 122
PREC_RIGHT, 39, 40, 50, 83, 84, 89, 122, 133, 135, 136
preconditioning
advice on, 24, 29
band-block diagonal, 85
banded, 81
setup and solve phases, 29
user-supplied, 49-50, 72, 120-121, 130, 131
pretype, $39,40,50,83,84,89$
pretypeB, 122, 133, 135, 136
PVODE, 1
RCONST, 32
realtype, 32
reinitialization, 66, 117
right-hand side function, 67
backward problem, 126
forward sensitivity, 107-109
quadrature backward problem, 127
quadrature equations, 79
Rootfinding, 25, 34, 80
SMALL_REAL, 32
SPBCG generic linear solver
description of, 160
functions, 160
SPGMR generic linear solver
description of, 159
functions, 160
support functions, 160
SPTFQMR generic linear solver
description of, 161
functions, 161
Stability limit detection, 24
step size bounds, 45
sundials_nvector.h, 32
sundials_types.h, 32
TFQMR method, 40, 51, 122, 161
tolerances, 16, 35, 36, 47, 69, 77, 102, 116, 117
UNIT_ROUNDOFF, 32
User main program
Adjoint sensitivity analysis, 111
CVBANDPRE usage, 81
CVBBDPRE usage, 87
forward sensitivity analysis, 93
integration of quadratures, 74
IVP solution, 33
vode, 1
VODPK, 1
weighted root-mean-square norm, 16


[^0]:    ${ }^{1}$ Files for both the serial and parallel versions of CVODES are included in the distribution. For users in a serial computing environment, the files specific to parallel environments (which may be deleted) are as follows: all files in src/nvec_par/; cvodes_bbdpre.c, cvodes_bbdpre_impl.h (in src/cvodes/); cvodes_bbdpre.h (in include/cvodes/); all files in examples/cvodes/parallel/. (By "serial version" of CVODES we mean the CVODES solver with the serial NVECTOR module attached, and similarly for "parallel version".)

[^1]:    ${ }^{1}$ The degree of the interpolation polynomial is always that of the current BDF order for the forward interpolation at the first point to the right of the time at which the interpolated value is sought (unless too close to the $i$-th checkpoint, in which case it uses the BDF order at the right-most relevant point). However, because of the FLC BDF implementation (see $\S 3.1$ ), the resulting interpolation polynomial is only an approximation to the underlying BDF interpolant.

    The Hermite cubic interpolation option is present because it was implemented chronologically first and it is also used by other adjoint solvers (e.g. DASPKADJOINT). The variable-degree polynomial is more memory-efficient (it requires only half of the memory storage of the cubic Hermite interpolation) and is more accurate. The accuracy differences are minor when using BDF (since the maximum method order cannot exceed 5), but can be significant for the Adams method for which the order can reach 12 .

