# VNODE-LP

# A Validated Solver for Initial Value Problems in Ordinary Differential Equations

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# **Preface**

We present VNODE-LP, a C++ solver for computing bounds on the solution of an initial-value problem (IVP) for an ordinary differential equation (ODE). In contrast to traditional ODE solvers, which compute approximate solutions, this solver proves that a unique solution to a problem exists and then computes rigorous bounds that are guaranteed to contain it. Such bounds can be used to help prove a theoretical result, check if a solution satisfies a condition in a safety-critical calculation, or simply to verify the results produced by a traditional ODE solver.

This package is a successor of the VNODE [25], Validated Numerical ODE, package of N. Nedialkov. A distinctive feature of the present solver is that it is developed entirely using Literate Programming (LP) [17]. As a result, the correctness of VNODE-LP's implementation can be examined much easier than the correctness of VNODE—the theory, documentation, and source code of VNODE-LP are interwoven in this manuscript, which can be verified for correctness by a human expert, like in a peer-review process.

**Literate programming.** With LP, a program (or function) is normally subdivided into pieces of code or *chunks*, and each of them may be subdivided into smaller chunks. How they are divided and put together should be clear from the exposition.

The present document is produced by cweave [18] on LaTeX-like cweb files, which contain both LaTeX text and C++ code. The C++ code for VNODE-LP and all the examples are generated by running ctangle [18] on those files; see Figure 1.

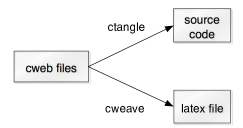


Figure 1. Producing C++ and LATEX files from cweb files

Structure. Part I describes the problem VNODE-LP solves, shows how it can

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be installed, and illustrates on several examples how VNODE-LP can be used. Parts II–V contain the implementation of this package.

If a reader is interested only in using VNODE-LP, then studying Part I should provide sufficient knowledge for using this package.

This document is *open*: errors found by a reader will be fixed and suggestions on improving it will be incorporated. Such suggestions can be on both exposition and code.

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N. Nedialkov July 26, 2006

# Part I Introduction, Installation, Use

## Chapter 1

# Introduction

# **1.1** The problem VNODE-LP solves

We consider the IVP

$$y'(t) = f(t, y), \quad y(t_0) = y_0, \qquad y \in \mathbb{R}^n, \ t \in \mathbb{R}.$$
 (1.1)

We denote the set of closed (finite) intervals on  $\mathbb{R}$  by

$$\mathbb{IR} = \{ \boldsymbol{a} = [\underline{\boldsymbol{a}}, \overline{\boldsymbol{a}}] \mid \underline{\boldsymbol{a}} \le x \le \overline{\boldsymbol{a}}, \ \underline{\boldsymbol{a}}, \ \overline{\boldsymbol{a}} \in \mathbb{R} \}.$$

An interval vector is a vector with interval components. We denote the set of n-dimensional interval vectors by  $\mathbb{IR}^n$ .

Given a point  $t_{end} \neq t_0$   $(t_{end} \in \mathbb{R})$  and  $\mathbf{y}_0 \in \mathbb{R}^n$ , the goal of VNODE-LP is to compute  $\mathbf{y}_{end} \in \mathbb{R}^n$  at  $t_{end}$  that contains the solution to (1.1) at  $t_{end}$  for all  $y_0 \in \mathbf{y}_0$ . If VNODE-LP cannot reach  $t_{end}$ , bounds on the solution at some  $t^*$  between  $t_0$  and  $t_{end}$  are returned.

This package is applicable to ODE problems for which derivatives of the solution y(t) exist to some order; that is, y(t) is sufficiently smooth. As a consequence, the code list of f should not contain functions such as branches, abs, or min.

In practice,  $t_0$  or  $t_{\rm end}$ , or both, may not be representable as floating-point numbers; for example the decimal 0.1 has an infinite binary representation. In this case, the user can set a machine-representable interval  $t_0$  [resp.  $t_{\rm end}$ ] containing  $t_0$  [resp.  $t_{\rm end}$ ].

# 1.2 On Literate Programming

The VNODE-LP package is a successor of VNODE [23, 25]. Both are written in C++. A major difference is that VNODE-LP is produced entirely, including this manuscript, using Literate Programming (LP) [17] and CWEB [18]. Why LP?

In general, interval methods produce results that can have the power of a mathematical proof. For example, when computing an enclosure of the solution of an IVP ODE, an interval method first proves that there exists a unique solution to the problem and then produces bounds that contain it. When solving a nonlinear equation, an interval method can prove that a region does not contain a solution or compute bounds that contain a unique solution to the problem.

However, if an interval method is not implemented correctly, it may not produce rigorous results. Furthermore, we cannot claim mathematical rigor if we miss to include even a single roundoff error in a computation. Therefore, it is of paramount importance to ensure that an interval algorithm is encoded correctly in a programming language.

In the author's opinion, interval software should be written such that it can be readily verified in a human peer-review process, like a mathematical proof is checked for correctness. The main goal of this work is to implement and document an interval solver for IVPs for ODEs such that its correctness can be verified by a reviewer.

To accomplish our goal, we have chosen the LP approach. The author has found LP particularly suitable for ensuring that an implementation of a numerical algorithm is a correct translation of its underlying theory into a programming language. Some of the benefits of employing LP follow.

- We can combine theory, source code, and documentation in a single document;
   we shall refer to it as an LP document.
- With LP, we can produce nearly "one-to-one" translation of the mathematical theory of a method into a computer program. In particular, we can split the theory into small pieces, translate each of them, and keep mathematical expressions and the corresponding code close together in a unified document. This facilitates verifying the correctness of smaller pieces and of a program as a whole.
- Since theory and implementation are in a single document, it is easier to keep them consistent, compared to having separate theory, source code, and documentation.

The user guide, theory, and source code of VNODE-LP are presented in the remainder of this document. The source code of VNODE-LP is extracted from source, *cweb* files using CWEB's [18] ctangle. This manuscript is produced by running cweave on these files and then calling LATEX.

If the correctness of this manuscript is confirmed by reviewers in a peer-review-like process, we may trust the correctness of the implementation of VNODE-LP, and accept the bounds it computes as rigorous. When claiming rigor, however, we presume that the operating system, compiler, and the packages VNODE-LP uses do not contain errors.

# 1.3 Applications

Applications of validated integration include, for example, the solution of Smale's 14th problem [30] and rigorous computation of asteroid orbits [7]. The (previous)

1.4. Limitations 5

VNODE package had been employed in applications such as rigorous multibody simulations [5], reliable surface intersection [22, 28], computing bounds on eigenvalues [8], parameter and state estimation [15], rigorous shadowing [11, 12], and theoretical computer science [3].

#### 1.4 Limitations

Generally, VNODE-LP is suitable for computing bounds on the solution of an IVP ODE with point initial conditions, or interval initial conditions with a sufficiently small width, over not very long time intervals. If the initial condition set is not small enough and/or long time integration is desired, the reader is referred to the Taylor models approach of Berz and Makino, and their COSY package. Alternatively, one can subdivide the initial interval vector (box)  $y_0$  into smaller boxes, perform integrations with them as initial conditions, and build an enclosure of the solution at the desired  $t_{\rm end}$ .

## 1.5 Prerequisites

A user of VNODE-LP does not need to know how the underlying methods work. It is sufficient to know that, if a and  $b \in \mathbb{IR}$  and  $\bullet \in \{+, -, \times, /\}$ , then VNODE-LP builds on the interval-arithmetic (IA) operations defined as

$$\boldsymbol{a} \bullet \boldsymbol{b} = \big\{ x \bullet y \mid x \in \boldsymbol{a}, \ y \in \boldsymbol{b} \big\},\tag{1.2}$$

where division is undefined if  $0 \in \mathbf{b}$ . This definition can be implemented, for example, as

$$a + b = [\underline{a} + \underline{b}, \overline{a} + \overline{b}],$$
  
 $a - b = [\underline{a} - \overline{b}, \overline{a} - \underline{b}],$   
 $a \times b = [\min{\{\underline{a}\underline{b}, \underline{a}\overline{b}, \overline{a}\underline{b}, \overline{a}\overline{b}\}, \max{\{\underline{a}\underline{b}, \underline{a}\overline{b}, \overline{a}\underline{b}, \overline{a}\overline{b}\}}], \text{ and }$   
 $a/b = [a, \overline{a}] \times [1/\overline{b}, 1/b], \quad 0 \notin b.$ 

On a computer, a and b are representable machine intervals, and the computed result of an IA operation must contain (1.2), provided no exceptions occur. For example, if intervals are represented by their endpoints, when computing a + b, the true  $\underline{a} + \underline{b}$  is rounded towards  $-\infty$ , and the true  $\overline{a} + \overline{b}$  is rounded towards  $+\infty$ .

From a language perspective, we have tried to avoid using advanced C++ techniques; however, basic knowledge of C++ is required.

The installation of VNODE-LP is explained in Chapter 2. Chapter 3 presents various examples of how VNODE-LP can be used. Chapter 4 lists and describes the functions available to a user of VNODE-LP. Chapter 5 contains descriptions of test cases. Various listings are given in Chapter 6.

# Chapter 2

# Installation

In this Chapter, we list the utilities and packages necessary for installing VNODE-LP, list successful installations, and then describe the installation process.

# 2.1 Prerequisites

The following utilities are needed:

- 1. gunzip (GNU unzip)
- 2. tar (tape archiver)
- 3. ar (for creating a library archive)
- 4. C++ compiler
- 5. GNU make
- 6. libg2c run-time library, if the GNU C++ compiler is used

Normally, 1-5 are present on a Unix-based system, while libg2c may need to be installed.

The following packages are used by VNODE-LP and must be installed before VNODE-LP is installed:

interval arithmetic: FILIB++ [19] or PROFIL/BIAS [16]

linear algebra: LAPACK [2] and BLAS [1]

## 2.2 Successful installations

To date VNODE-LP has been successfully compiled and installed as follows.

IA	Operating	Architecture	Compiler
package	system		
FILIB++	Linux	x86	gcc
	Solaris	Sparc	gcc
PROFIL	Linux	x86	gcc
	Solaris	Sparc	gcc
	Mac OSX	PowerPC	gcc
	Windows with Cygwin	x86	gcc

**Note.** At the time of writing this manuscript, the author has not been able to install FILIB++ correctly on Mac OS X. However, VNODE-LP compiles on it.

## 2.3 Installation process

The installation process consists of the following steps:

- 1. extracting the source code
- 2. preparing a configuration file
- 3. building the VNODE-LP library, examples, and tests
- 4. installing the library files

#### 2.3.1 Extracting the source code

VNODE-LP can be downloaded from www.cas.mcmaster.ca~/nedialk/vnodelp. The corresponding file is vnodelp.tar.gz. To extract the source files, type

tar -zxvf vnodelp.tar.gz

This will create the directory vnodelp and store the VNODE-LP files in it.

#### 2.3.2 Preparing a configuration file

The user has to prepare a *configuration file*, which contains information such as compiler, options, libraries, and various directory paths. There are four such files used by the author: MacOSXWithFilib, MacOSXWithProfil, LinuxWithFilib, and LinuxWithProfil, located in vnodelp/config. One can modify any of these files or create his own configuration file, where the variables described in Figure 2.1 should be set appropriately. The files MacOSXWithProfil and LinuxWithProfil are given in Figures 2.2 and 2.3.

variable	stores
CXX	name of C++ compiler
CXXFLAGS	C++ compiler flags
GPP_LIBS	GNU C++ standard library libstdc++ and
	the libg2c run-time library
LDFLAGS	linker flags
I_PACKAGE	FILIB_VNODE or PROFIL_VNODE
I_INCLUDE	name of the directory containing include files of the
	interval-arithmetic package
I_LIBDIR	name of the directory containing interval libraries
I_LIBS	names of interval libraries
MAX_ORDER	value for the maximum order VNODE-LP can use
L_LAPACK	name of the directory containing the LAPACK library
L_BLAS	name of the directory containing the BLAS library
LAPACK_LIB	name of the LAPACK library file
BLAS_LIB	name of the BLAS library file

Figure 2.1. Variables of a VNODE-LP configuration file

#### 2.3.3 Building the VNODE-LP library and examples

The makefile in vnodelp (see Figure 2.4) contains two variables that need to be set appropriately:

CONFIG\_FILE contains the name of the configuration file; and

INSTALL\_DIR contains the directory, where VNODE-LP should be installed.

After these variables are set appropriately, type

#### make

The library libvnode.a will be created in subdirectory vnodelp/lib, and the examples will be created in vnodelp/examples. Then, several test programs in subdirectory tests will be compiled and executed. If VNODE-LP compiles successfully and the tests pass, the following message should appear.

```
***********

*** VNODE-LP has compiled successfully

*** All tests have executed successfully

**********

If you have set the install directory, type
make install
```

```
CXX
         = g++
CXXFLAGS = -O2 -g -Wall -pedantic -Wno-deprecated
GPP\_LIBS = -lstdc++/sw/lib/libg2c.a
LDFLAGS += -bind_at_load -Wno
\# interval package
LPACKAGE = PROFIL_VNODE
I_{I}NCLUDE =
                 (HOME)/NUMLIB/Profil -2.1/src
                 (HOME)/NUMLIB/Profil -2.1/src/BIAS
                 (HOME)/NUMLIB/Profil -2.1/src/Base
                 (HOME)/NUMLIB/Profil -2.1/src/BIAS
I_LIBDIR =
                 (HOME)/NUMLIB/Profil -2.1/src/Base
                 (HOME)/NUMLIB/Profil -2.1/src/lr
I_LIBS
                 -lProfil -lBias -llr
MAX\_ORDER = 50
\# LAPACK and BLAS
           = $ (HOME) / NUMLIB/LAPACK
L_LAPACK
           = $ (HOME) / NUMLIB/LAPACK
L_BLAS
LAPACK\_LIB = -llapack\_MACOSX
BLAS_LIB
           = -lblas_MACOSX
# --- DO NOT CHANGE BELOW ----
INCLUDES = \$(addprefix -I, \$(I\_INCLUDE))
        -I$ (PWD) /FADBAD++
LIB\_DIRS = \$(addprefix -L, \$(I\_LIBDIR)
        (L_APACK) (L_BLAS)
CXXFLAGS += -D\{LPACKAGE\} \setminus
        -DMAXORDER=$ (MAX_ORDER) $ (INCLUDES)
LDFLAGS += (LIB\_DIRS)
LIBS = (I\_LIBS) (LAPACK\_LIB) (BLAS\_LIB)
        $(GPP_LIBS)
```

Figure 2.2. File config/MacOSXWithProfil

#### 2.3.4 Installing the library files

To install the library and the related include files, type

#### make install

This will create a subdirectory vnodelp of the directory stored in INSTALL\_DIR and subdirectories of vnodelp as follows:

```
CXX = gcc
CXXFLAGS = -O2 - Wall - Wno-deprecated - DNDEBUG
GPP\_LIBS = -lstdc++-lg2c
# interval package
I_PACKAGE = PROFIL_VNODE
I_{INCLUDE} =
        (HOME)/NUMLIB/Profil-2.0/include
        (HOME)/NUMLIB/Profil - 2.0/include/BIAS
        (HOME)/NUMLIB/Profil - 2.0/src/Base
I_LIBDIR = \frac{(HOME)}{NUMLIB}/Profil - 2.0/lib
I_LIBS
          = -lProfil -lBias -llr
MAX\_ORDER = 50
\# LAPACK and BLAS
L_LAPACK
L_BLAS
LAPACK\_LIB = -llapack
BLAS\_LIB = -lblas
# --- DO NOT CHANGE BELOW ----
INCLUDES = \$(addprefix -I, \$(I\_INCLUDE))
        -I$ (PWD) /FADBAD++
LIB_DIRS = \$(addprefix -L, \$(I_LIBDIR)
        $(L_LAPACK) $(L_BLAS))
CXXFLAGS \mathrel{+}{=} -D\$\{LPACKAGE\} \setminus
        -DMAXORDER=$(MAX_ORDER) $(INCLUDES)
LDFLAGS += (LIB\_DIRS)
LIBS = (I\_LIBS) *(LAPACK\_LIB) *(BLAS\_LIB)
        $(GPP_LIBS)
```

Figure 2.3. File config/LinuxWithProfil

directory	contains
lib	libvnode.a
include	libvnode.a's include files
config	configuration files
doc	documentation file vnode.pdf

Subsection 3.1.4 contains details about how to build user's programs.

```
# set CONFIG_FILE and INSTALL_DIR

CONFIG_FILE ?= MacOSXWithProfil
INSTALL_DIR ?= $ (HOME)

#---- DO NOT CHANGE BELOW ----
```

Figure 2.4. The first six lines of makefile in vnodelp

## Chapter 3

# **Examples**

We start with an example showing how a basic integration with VNODE-LP can be carried out, Section 3.1. In Section 3.2 we examine how VNODE-LP does on a simple scalar ODE. Section 3.3 contains an example of integrating a time-dependent system of ODEs and illustrates how this package can be used to check the correctness of the numerical results produced by a standard ODE method.

Section 3.4 outlines how to integrate with interval initial condition and output intermediate results. We describe how VNODE-LP outputs results at given time points in Section 3.5, and how parameters can be passed to an ODE problem in Section 3.6.

In Section 3.7, we show how an integration can be controlled, and in Section 3.8, we perform a simple study of the computational work versus the order of the method implemented in VNODE-LP. Section 3.9 contains a study of the computational work versus the size of the problem. Section 3.10 illustrates the stepsize behavior when integrating an orbit problem. Finally, Section 3.11 shows the stepsize behavior of VNODE-LP as the stiffness in an ODE increases.

# 3.1 Basic usage

In VNODE-LP, the user has to specify the right side of an ODE problem and provide a main program.

#### 3.1.1 Problem definition

An ODE must be specified by a template function for evaluating y' = f(t, y) of the form

```
18 \langle \text{template ODE function 18} \rangle \equiv \\ \text{template} \langle \text{typename var_type} \rangle \\ \text{void } ODEName(\text{int } n, \text{var_type } *yp, \text{const var_type } *y, \text{var_type } t, \\ \text{void } *param) \\ \{
```

```
/* body */
}
```

Here n is the size of the problem, t is the time variable, y is a pointer to input variables, yp is a pointer to output variables, and param is a pointer to additional parameters that can be passed to this function.

As an example, consider the Lorenz system

```
y'_1 = \sigma(y_2 - y_1)

y'_2 = y_1(\rho - y_3) - y_2

y'_3 = y_1 y_2 - \beta y_3,
```

where  $\sigma$ ,  $\rho$ , and  $\beta$  are constants. This system is encoded in the *Lorenz* function below. The constants have values  $\sigma = 10$ ,  $\beta = 8/3$ , and  $\rho = 28$ . We initialize beta with the interval containing 8/3: **interval**(8.0) creates an interval with endpoints 8.0, and **interval**(8.0)/3.0 is the interval containing 8/3. The last parameter, param, is not used here, but its role is discussed in Section 3.6.

#### 3.1.2 Main program

We give a simple main program and explain its parts.

```
20 \langle \text{simple main program } 20 \rangle \equiv \langle \text{Lorenz } 19 \rangle
int main()
{

\langle \text{set initial condition and endpoint } 21 \rangle
\langle \text{create AD object } 22 \rangle
\langle \text{create a solver } 23 \rangle
\langle \text{integrate (basic) } 24 \rangle
\langle \text{check if success } 25 \rangle
```

This code is used in chunks 20, 45, 48, 61, and 70.

<sup>&</sup>lt;sup>1</sup>The result of this division is the interval with endpoints 8/3 rounded toward  $-\infty$  and 8/3 rounded towards  $+\infty$ .

3.1. Basic usage 15

```
\ output results 26 \\
    return 0;
}
This code is used in chunk 27.
```

The initial condition and endpoint are represented as intervals in VNODE-LP. In this example, they are all point values stored as intervals. The components of **iVector** (interval vector) are accessed like a C/C++ array is accessed.

```
21 \langle set initial condition and endpoint 21\rangle \equiv const int n=3; interval t=0.0, tend=20.0; iVector y(n); y[0]=15.0; y[1]=15.0; y[2]=36.0;
```

This code is used in chunks 20, 42, 48, 58, and 61.

Then we create an AD object of type **FADBAD\_AD**. It is instantiated with data types for computing Taylor coefficients (TCs) of the ODE solution and TCs of the solution to the variational equation, respectively [23]. To compute these coefficients, we employ the FADBAD++ package [29]. The first parameter in the constructor of **FADBAD\_AD** is the size of the problem. The second and third parameters are the name of the template function, here *Lorenz*.

```
22 ⟨create AD object 22⟩ ≡
AD *ad = new FADBAD_AD(n, Lorenz, Lorenz);
This code is used in chunks 20, 45, 48, 61, and 68.
Now, we create a solver:
23 ⟨create a solver 23⟩ ≡
VNODE *Solver = new VNODE(ad);
```

This code is used in chunks 20, 35, 40, 45, 48, 58, 61, and 68.

This code is used in chunks 20, 35, 40, and 68.

The integration is carried out by the *integrate* function. It attempts to compute bounds on the solution at tend. When integrate returns, either t = tend or  $t \neq tend$ . In both cases, y contains the ODE solution at t.

```
24 \langle integrate (basic) 24 \rangle \equiv Solver→integrate(t, y, tend);
This code is used in chunks 20, 35, 40, and 68.

We check if an integration is successful by calling Solver→successful():
25 \langle check if success 25 \rangle \equiv if (¬Solver→successful())
cout \langle "VNODE-LP_\(\text{could}\)_\(\text{not}\)_\(\text{reach}\)_\(\text{tend} \langle endl;
```

Finally, we output the computed enclosure of the solution at t by 26  $\langle$  output results  $26 \rangle \equiv cout \ll "Solution\_enclosure\_at\_t\_=\_" \ll t \ll endl; printVector(y); This code is used in chunks 20, 35, 40, and 47.$ 

#### 3.1.3 Files

The code of VNODE-LP is in the namespace **vnodelp**. The interface to VNODE-LP is stored in the file **vnode.h**, which must be included in any file using VNODE-LP. We store our program in the file **basic.cc**.

```
27 ⟨basic.cc 27⟩ ≡
#include <ostream>
#include "vnode.h"
using namespace std;
using namespace vnodelp;
⟨simple main program 20⟩
```

#### 3.1.4 Building an executable

We describe how basic.cc is compiled and linked with the VNODE-LP library. The subdirectory user\_program of vnodelp contains the files basic.cc and makefile, which is given in Figure 3.1. We consider this file here.

Figure 3.1. makefile in vnodelp/user\_program

The directory where vnodelp resides is set in INSTALL\_DIR, and the configuration file is set in CONFIG\_FILE. The variables CXXFLAGS and LDFLAGS need not be changed. Finally, the rule for building basic is given. To create the executable file basic, type make in vnodelp/user\_program.

3.1. Basic usage

#### 3.1.5 Output

The output of basic is

```
Solution enclosure at t = [20,20]
14.30[38159449608937,44694855332662]
9.5[785941360078012,801274302834650]
39.038[2373597549516,4111043348412]
```

These results are interpreted as

$$y(20) \in \begin{pmatrix} [14.3038159449608937, 14.3044694855332662] \\ 9.5785941360078012, 9.5801274302834650] \\ [39.0382373597549516, 39.0384111043348412] \end{pmatrix}. \tag{3.1}$$

For comparison, if we integrate this problem with MAPLE using dsolve with options method=taylorseries and abserr=Float(1,-18), and with Digits := 20, we obtain

$$y(20) \approx \begin{pmatrix} 14.304146251277895001 \\ 9.5793690774871976695 \\ 39.038325167739731729 \end{pmatrix},$$

which is contained in the bounds (3.1).

#### Remarks

- 1. All numerical results in this manuscript are produced with PROFIL on 1.25 GHz PowerPC G4 with MacOS X, 512 MB RAM, and 512KB L2 cache.
- 2. The output format is due to the PROFIL/BIAS [16] interval-arithmetic package.
- 3. On different architectures, or with different IA packages on the same architecture, the computed results are likely to differ, but they must contain the true results.

#### 3.1.6 Standard coding

All source-code files in the VNODE-LP distribution, except the test programs in subdirectory vnodelp/tests, are generated with ctangle from CWEB. Since a user may not use LP, we also give the "standard" C++ code of basic.cc in Figure 3.2.

For the remaining examples, we do not explain how they are compiled. For details, see the makefile in Figure 6.1. This file is in the directory vnodelp/examples. Also, we do not provide "standard" code of the corresponding C++ files; if needed, it can be extracted from the ctangle generated files \*.cc in vnodelp/examples.

```
#include <ostream>
#include "vnode.h"
using namespace std;
using namespace vnodelp;
template<typename var_type>
void Lorenz(int n, var_type*yp, const var_type*y, var_type t,
             void*param)
  interval sigma (10.0), rho (28.0);
  interval beta = interval (8.0)/3.0;
  yp[0] = sigma*(y[1]-y[0]);
  yp[1] = y[0]*(rho-y[2]) - y[1];
  yp[2] = y[0] * y[1] - beta * y[2];
int main()
  const int n = 3;
  interval t = 0.0, tend = 20.0;
  iVector y(n);
  y[0] = 15.0;
  y[1] = 15.0;
  y[2] = 36.0;
  AD *ad= new FADBAD_AD(n, Lorenz, Lorenz);
  VNODE *Solver= new VNODE(ad);
  Solver->integrate(t,y,tend);
  if (! Solver -> successful())
    cout << "VNODE-LP_could_not_reach_t_=_"<< tend << endl;
  cout << "Solution_enclosure_at_t_=_"<< t << endl;
  printVector(y);
  return 0;
```

Figure 3.2. The "standard" C++ code of basic.cc

#### 3.2 One-dimensional ODE

VNODE-LP is designed to be a general-purpose ODE interval solver. Nevertheless, it should handle scalar ODEs. This example illustrates how VNODE-LP deals with the simple problem

```
y' = -y, y(0) = 1, t_{\text{end}} = 20.
    We write
32 \langle scalar ODE example 32\rangle \equiv
      template\langle typename\ var\_type\rangle
      void Scalar Example (int n, var_type *yp, const var_type *y, var_type t,
                void *param)
        yp[0] = -y[0];
   This code is used in chunk 35.
      The initial condition and endpoint are set in
33 \langle set scalar ODE initial condition and endpoint 33 \rangle \equiv
      const int n = 1;
      interval t = 0.0, tend = 20.0:
                       /* number of state variables is 1 */
      iVector y(n);
      y[0] = 1.0;
   This code is used in chunk 35.
      To create the necessary AD object, we call
34 \langle create scalar AD object 34\rangle \equiv
      AD *ad = new FADBAD\_AD(n, Scalar Example, Scalar Example);
   This code is used in chunk 35.
      The main program is
35 \langle \text{scalar.cc} 35 \rangle \equiv
    #include <ostream>
    #include "vnode.h"
      using namespace std;
      using namespace vnodelp;
      \langle \text{ scalar ODE example } 32 \rangle
      int main()
         (set scalar ODE initial condition and endpoint 33)
         (create scalar AD object 34)
         (create a solver 23)
```

```
⟨integrate (basic) 24⟩
⟨check if success 25⟩
⟨output results 26⟩
return 0;
}
```

The output of this program is 0.000000002061153[6,7], which must enclose  $e^{-20}$ . Using MAPLE with a 30-digit computation, we obtain for  $e^{-20}$ 

```
\underline{0.2061153}62243855782796594038016\cdot 10^{-8}
```

(the digits that coincide are underlined), which is contained in the interval computed by VNODE-LP.

# 3.3 Time-dependent ODE

We show an example of integrating a time-dependent ODE. We choose the E1 problem from the DETEST test set [14]. The problem is

```
y_1' = y_2
y_2' = -\left(\frac{y_2}{t+1} + \left(1 - \frac{0.25}{(t+1)^2}\right)y_1\right)
y_1(0) = 0.6713967071418030,
y_2(0) = 0.09540051444747446,
t_{\text{end}} = 20.
37 \langle \text{DETEST E1 } 37 \rangle \equiv
\text{template} \langle \text{typename var_type} \rangle
\text{void DETEST_E1(int } n, \text{var_type } *yp, \text{const var_type } *y, \text{var_type } t, \text{void } *param)
\{ \text{var_type } t1 = t + 1.0;
yp[0] = y[1];
yp[0] = -(y[1]/t1 + (1.0 - 0.25/(t1 * t1)) *y[0]);
\}
This code is used in chunk 40.
```

We store the initial condition as intervals containing the corresponding decimal values. The function <code>string\_to\_interval</code> converts a decimal string to a machine interval that contains the decimal value stored in the string.

```
\langle set E1 initial condition and endpoint 38 \rangle \equiv const int n = 2; interval t = 0.0, tend = 20.0; iVector y(n);
```

```
y[0] = string\_to\_interval("0.6713967071418030");
      y[1] = string\_to\_interval("0.09540051444747446");
   This code is used in chunk 40.
      To create an AD object, we call
39 \langle \text{ create E1 39} \rangle \equiv
      AD *ad = new FADBAD\_AD(n, DETEST\_E1, DETEST\_E1);
   This code is used in chunk 40.
      The main program is
40 \langle E1.cc 40 \rangle \equiv
    #include <ostream>
    #include "vnode.h"
      using namespace std;
      using namespace vnodelp;
      〈DETEST E1 37〉
      int main()
         (set E1 initial condition and endpoint 38)
         (create E1 39)
         (create a solver 23)
         (integrate (basic) 24)
         \langle \text{ check if success } 25 \rangle
         (output results 26)
        return 0;
      The output of this program is
   Solution enclosure at t = [20,20]
   0.14567236007282[02,87]
    -0.0988350019557[410,507]
```

We have also computed a numerical solution using MATLAB's ode45. The corresponding programs are in Figure 6.3, and the output is

We have underlined the digits that coincide in the VNODE-LP and MATLAB output.

Since VNODE-LP includes all possible errors in its computation, including conversion errors in the input from decimal to binary, we have bounds on the true solution at t=20. Using these bounds, we can check the accuracy of the numerical solution found by MATLAB's ode45.

#### 3.4 Interval initial conditions

Suppose we want to compute bounds on the solution of the Lorenz problem for all

$$y(0) \in \begin{pmatrix} 15 + [-10^{-4}, 10^{-4}] \\ 15 + [-10^{-4}, 10^{-4}] \\ 36 + [-10^{-4}, 10^{-4}] \end{pmatrix}.$$

We set an interval initial condition by

42  $\langle$  set interval initial condition and endpoint 42 $\rangle$   $\equiv$   $\langle$  set initial condition and endpoint 21 $\rangle$ 

interval  $eps = interval(-1, 1)/1 \cdot 10^4$ ;

$$y[0] += eps;$$
  
 $y[1] += eps;$   
 $y[2] += eps;$ 

This code is used in chunk 45.

Before presenting the rest of the code, we introduce some notation. We denote the radius of a by

$$r(\boldsymbol{a}) = (\overline{\boldsymbol{a}} - \underline{\boldsymbol{a}})/2.$$

The midpoint of a is denoted by

$$m(\mathbf{a}) = (\overline{\mathbf{a}} + \underline{\mathbf{a}})/2.$$

(In machine arithmetic, the true  $\overline{a} - \underline{a}$  is rounded upward, and the true  $(\overline{a} + \underline{a})/2$  is rounded to the nearest.) Radius and midpoint are defined componentwise for interval vectors.

Informally, the *global excess* at a point  $t^*$  is the overestimation in the computed bounds over the true solution set at  $t^*$  [23]. VNODE-LP computes an estimate, which is a nonnegative number, of the global excess in the computed bounds on each solution component, and also the max norm of these estimates.

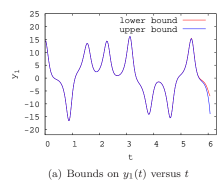
When stepping in time from  $t_0$  to  $t_{\rm end}$ , VNODE-LP computes bounds on the solution at points that are machine numbers, except possibly at  $t_{\rm end}$ , which may be a machine interval with a nonzero radius. In this example, we output intermediate results during the integration. We tell the integrator to return after each step is completed by calling setOneStep(on). This is convenient when we want to access intermediate results, for example, for plotting solutions, stepsize, etc. In the code below, we record such results in a file, which is later used by gnuplot to generate the plots in Figure 3.3.

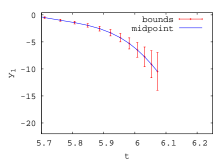
43  $\langle \text{ indicate single step } 43 \rangle \equiv Solver \rightarrow setOneStep(on);$ This code is used in chunks 44 and 61. Now we integrate and record in the file lorenzi.out the midpoint and the radius of the computed bounds on  $y_1(t)$  for each t selected by the solver. We also output an estimate on the *global excess*. The function getGlobalExcess returns the max norm of a vector with estimates on the global excess for each solution component. We exit the **while** loop below if this estimate exceeds 15 (This number is chosen so we can visualize the divergence of the computed bounds; see Figure 3.3(b).)

```
44 (integrate with interval initial condition 44) \equiv
       (indicate single step 43)
      ofstream outFile("lorenzi.out", ios::out);
      while (t \neq tend) {
         Solver \rightarrow integrate(t, y, tend);
         if (Solver \rightarrow successful() \land Solver \rightarrow getGlobalExcess() \le 15.0) {
           outFile \ll midpoint(t) \ll "\t"
                \ll midpoint(y[0]) \ll "\t"
                \ll rad(y[0]) \ll "\t"
                \ll Solver \rightarrow getGlobalExcess() \ll endl;
         else break;
      outFile.close();
   This code is used in chunk 45.
      The main program is
45 \langle integi.cc 45 \rangle \equiv
    #include <fstream>
    #include "vnode.h"
      using namespace std;
      using namespace vnodelp;
      (Lorenz 19)
      int main()
         (set interval initial condition and endpoint 42)
         (create AD object 22)
         (create a solver 23)
         (integrate with interval initial condition 44)
         return 0;
```

In Figure 3.3(a) and (b), we plot the lower and upper bounds on  $y_1$ . Their divergence is clearly seen in (b). In (c) we plot the logarithm of the estimate on the global excess versus t. On this problem, the bounds become too wide as t goes beyond  $\approx 6.0.7$ .

The gnuplot file employed to generate this plot is given in Figure 6.2.





(b) Bounds on  $y_1(t)$  versus t; "midpoint" lines connect the midpoints of the computed intervals

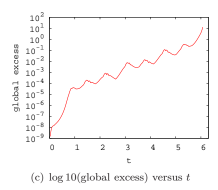


Figure 3.3. Plots generated using integi.cc

# 3.5 Producing intermediate results

We show how we can output enclosures on the solution at given points, for example, at t = 0.1, 0.2, 0.3, and t = 10. The decimal value 0.1 cannot be stored exactly as an IEEE floating-point number—we store 0.1 as an interval.

```
47 \langle integrate with intermediate output 47 \rangle \equiv interval step = string\_to\_interval("0.1");
tend = 0.0;
for (int i = 1; i \leq 3; i++) {
tend += step;
Solver \rightarrow integrate(t, y, tend);
\langle output results 26 \rangle
}
tend = 10;
Solver \rightarrow integrate(t, y, tend);
\langle output results 26 \rangle
```

This code is used in chunk 48.

```
The main program is
48 \langle intermediate.cc 48 \rangle \equiv
  #include <iostream>
  #include "vnode.h"
    using namespace std;
   using namespace vnodelp;
    ⟨Lorenz 19⟩
   int main()
     (set initial condition and endpoint 21)
     (create AD object 22)
     (create a solver 23)
     (integrate with intermediate output 47)
     return 0;
   The output is
  9.5199890775031[033,833]
  1.172296185059[1790,3086]
  36.286934318697[3267,4618]
  2.870955583196[2392,4120]
  -1.446088378503[6309,8119]
  27.476193552015[1645,3422]
  0.339508665582[1531,3150]
  -0.925287772052[5888,8462]
  20.901865632568[5157,7041]
  Solution enclosure at t = [10,10]
  -5.909806 [3819893544,7254843408]
  -11.34140 [28468979033,34573302108]
  9.08017 [76297270730,80129492258]
      left point
```

and right point

## 3.6 ODE control

## 3.6.1 Passing data to an ODE

Suppose that we want to pass the constants to the Lorenz system. We can encapsulate them in the structure

```
51 \langle \text{constants Lorenz } 51 \rangle \equiv
struct LorenzConsts {
   interval beta;
   double rho, sigma;
  };
This code is used in chunk 58.

We can set \sigma, \beta, and \rho in the main programs as

52 \langle \text{set ODE parameters } 52 \rangle \equiv
LorenzConsts p;
  p.sigma = 10.0;
  p.beta = \text{interval}(8.0)/3.0;
  p.rho = 28.0;
This code is used in chunk 58.
```

We can access parameters for the ODE through the **void** pointer *param*. The user has to ensure that such parameters are properly stored and later extracted through *param*.

```
53 \langle \text{passing parameters to Lorenz 53} \rangle \equiv \text{template} \langle \text{typename var_type} \rangle
void Lorenz2 (\text{int } n, \text{var_type } *yp, \text{const var_type } *y, \text{var_type } t, \text{void } *param)
 \left\{ \text{LorenzConsts } *p = (\text{LorenzConsts } *) \; param; \text{interval } beta = p \text{-} beta; \text{double } sigma = p \text{-} sigma; \text{double } rho = p \text{-} rho; \text{yp}[0] = sigma * (y[1] - y[0]); \text{yp}[1] = y[0] * (rho - y[2]) - y[1]; \text{}
```

3.6. ODE control 27

```
yp[2] = y[0] * y[1] - beta * y[2]; }
```

This code is used in chunk 58.

To pass parameters to the ODE, we create an AD object with fourth parameter the address of p:

54  $\langle$  create problem object with parameters 54 $\rangle \equiv$ 

```
AD *ad = new FADBAD\_AD(n, Lorenz2, Lorenz2, \&p);
```

This code is used in chunk 58.

## 3.6.2 Integration with parameter change

We illustrate how to integrate with changing  $\beta$ . First, we integrate from t0 to tend and output  $m(y_j)$  into a file;  $y_j$  is the computed enclosure at  $t_j$ .

```
\langle \text{simple integration } 55 \rangle \equiv \\ Solver \rightarrow setOneStep(on); \\ \textbf{ofstream } outFile1 ("odeparam1.out", ios::out); \\ \textbf{while } (t \neq tend) \ \{ \\ Solver \rightarrow integrate(t, y, tend); \\ outFile1 \ll midpoint(y[0]) \ll " \t" \\ \ll midpoint(y[1]) \ll " \t" \\ \ll midpoint(y[2]) \ll endl; \\ \} \\ outFile1.close(); \\ \text{This code is used in chunk } 58.
```

Now, we

- 1. integrate from  $t\theta$  to tend/2,
- 2. change  $\beta$ , and
- 3. integrate to tend.

Before calling *integrate* again, we call setFirstEntry. This call ensures that certain internal data structures are initialized. If setFirstEntry is not called, integrate would use data corresponding to the last computed solution from the most recent call to integrate.

```
56 \langle integrate from t to tend/2 56 \rangle \equiv t = 0.0; y[0] = 15; y[1] = 15; y[2] = 36; interval tend2 = tend/2.0; Solver \rightarrow setFirstEntry(); while (t \neq tend2) Solver \rightarrow integrate(t, y, tend2); This code is used in chunk 57.
```

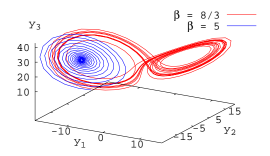
When changing  $\beta$ , the integration continues with the last computed solution, which is computed with the previous value for  $\beta$ . After a parameter is changed, ad-eval(&p) must be called. This ensures that some internal data structures are updated, to reflect the change in the ODE.

```
57 \langle integrate with resetting constants 57 \rangle \equiv
      \langle \text{ integrate from } t \text{ to } tend/2 \text{ 56} \rangle
                          /* change \beta */
      p.beta = 5.0;
                          /* must be called to update internal data structures */
      ad \rightarrow eval(\&p);
      ofstream outFile2("odeparam2.out", ios:: out);
      while (t \neq tend) {
         Solver \rightarrow integrate(t, y, tend);
         outFile2 \ll midpoint(y[0]) \ll "\t"
              \ll midpoint(y[1]) \ll "\t"
              \ll midpoint(y[2]) \ll endl;
      outFile2.close();
    This code is used in chunk 58.
      The main program is
58 \langle \text{odeparam.cc} 58 \rangle \equiv
    #include <fstream>
    #include "vnode.h"
      using namespace std;
      using namespace vnodelp;
       (constants Lorenz 51)
      (passing parameters to Lorenz 53)
      int main()
         (set initial condition and endpoint 21)
         (set ODE parameters 52)
          create problem object with parameters 54
          create a solver 23
         \langle \text{ simple integration 55} \rangle
         (integrate with resetting constants 57)
         return 0;
```

In Figure 3.4, we plot  $m(\boldsymbol{y}_j)$  in  $(y_1, y_2, y_3)$  coordinates corresponding to  $\beta = 8/3$  and  $\beta = 5$ . The gnuplot file for producing this plot is in Figure 6.4.

## 3.7 Integration control

We start by introducing various facts related to the integration process of VNODE-LP. Then we show ways of controlling it.



**Figure 3.4.** Midpoints of the computed bounds with  $\beta = 8/3$  from 0 to 20; and with  $\beta = 8/3$  changed to 5 at t = 10

The method implemented in VNODE-LP is a one-step method based on Taylor series and the Hermite-Obreschkoff scheme [23]. These are high-order methods, where typical values for the order, denoted here by p, can be in the range of 20 to 30; see Section 3.8.

VNODE-LP steps in time from  $t_0$  to  $t_{\rm end}$ , where the jth integration step,  $j \geq 1$ , is from  $t_{j-1}$  to  $t_j$ . The associated stepsize is  $h_j = t_j - t_{j-1}$ . Each of these  $t_j$  is a representable machine number, except  $t_0$  and  $t_{\rm end}$ , which can be machine intervals  $t_0$  and  $t_{\rm end}$  containing the true  $t_0$  and  $t_{\rm end}$ , respectively. For simplicity of the exposition, we assume point values  $t_0$  and  $t_{\rm end}$ .

In addition to computing bounds  $\boldsymbol{y}_j$  on the solution at each  $t_j$ , VNODE-LP also computes bounds for all  $t \in \boldsymbol{T}_j = [t_{j-1}, t_j]$ , or  $\boldsymbol{T}_j = [t_j, t_{j-1}]$  if the integration is in negative direction. We denote such bounds by  $\widetilde{\boldsymbol{y}}_j$ , and refer to them as a priori bounds [23]; we shall also refer to  $\boldsymbol{y}_j$  as tight bounds.

On the first integration step, VNODE-LP determines initial stepsize and the magnitude of the smallest stepsize that is allowed,  $h_{\min}$ . Then, on each step, VNODE-LP automatically selects a stepsize subject to absolute and relative error tolerances, atol and rtol. If the selected stepsize  $h_j$  is such that  $|h_j| < h_{\min}$ , the integration cannot continue, and VNODE-LP returns.

The following parameters can be changed by the user.

parameter	default value
$\overline{p}$	20
atol	$10^{-12}$
rtol	$10^{-12}$
$h_{\min}$	computed by VNODE-LP

The functions for changing them are given in Section 4.4. The order p does not

<sup>&</sup>lt;sup>2</sup>VNODE-LP selects  $h_j$  and then finds  $t_j$ , where in computer arithmetic the true  $t_{j-1} + h_j$  is rounded toward zero when computing  $t_j$ .

vary during an integration. By default p=20, but its value can be changed at the beginning of an integration. As pointed out earlier, p must be between 3 and the value set to MAX\_ORDER in the makefile for building the library; cf. Figures 2.2 and 2.3. If the user sets a value for  $h_{\min}$ , then this value is used by *integrate*.

```
If we wish to set, for example,
            \text{rtol} = \text{atol} = 10^{-14},
              p = 40, and
           h_{\min} = 10^{-5},
    we proceed with (cweave typesets 1e-14 as 1 \cdot 10^{-14})
60 \langle set control data for the solver 60 \rangle \equiv
       Solver \rightarrow set Tols (1 \cdot 10^{-14}, 1 \cdot 10^{-14});
       Solver \rightarrow setOrder(40);
       Solver \rightarrow setHmin(1 \cdot 10^{-5});
    This code is used in chunk 61.
       We write the main program
61 \langle integctrl.cc 61 \rangle \equiv
    #include <fstream>
    #include "vnode.h"
       using namespace std;
       using namespace vnodelp;
       (Lorenz 19)
       int main()
          \langle set initial condition and endpoint 21 \rangle
          (create AD object 22)
           create a solver 23
          (set control data for the solver 60)
          open files 62
          (indicate single step 43)
          (output initial condition 64)
          while (t \neq tend) {
            Solver \rightarrow integrate(t, y, tend);
            (output solution 65)
          (close files 63)
         return 0;
```

## Writing into files

We output data into three files, lorenz.tight, lorenz.apriori and lorenz.step. In lorenz.tight, each line contains (rounded in decimal to output precision)

$$m(\boldsymbol{t}_j)$$
  $\underline{\boldsymbol{y}}_{1,j}$   $\overline{\boldsymbol{y}}_{1,j}$   $w(\boldsymbol{y}_{1,j})$ 

where the subscripts 1, j refers to the jth computed solution for component  $y_1$ ;  $\mathbf{w}(\boldsymbol{y}_{1,j}) = 2\mathbf{r}(\boldsymbol{y}_{1,j})$  is the width, or diameter of  $\boldsymbol{y}_{1,j}$ . This width can be viewed as the *global excess* in the computed  $\boldsymbol{y}_{1,j}$ .

In lorenz.apriori, we output the a apriori enclosures and the corresponding time intervals in a form suitable for gnuplot to produce boxes denoting these a priori bounds; see Figure 3.5(b). The function getAprioriEncl returns  $\tilde{\boldsymbol{y}}_{j}$ , and getT returns  $T_{j}$ .

```
In lorenz.step, each line is
```

```
m(t_i) h_i
```

The function getStepsize returns  $h_j$ .

```
62 ⟨open files 62⟩ ≡
    ofstream outFile1("lorenz.tight", ios::out);
    ofstream outFile2("lorenz.step", ios::out);
    ofstream outFile3("lorenz.apriori", ios::out);
```

This code is used in chunk 61.

```
63 \langle \text{close files 63} \rangle \equiv outFile1.close(); outFile2.close(); outFile3.close();
```

This code is used in chunk 61.

In the code below, *inf* returns the left point of an interval, and *sup* returns the right point of an interval. (The output goes through C++'s stream output, so the endpoints are not rounded outward.)

```
64 \langle \text{ output initial condition } 64 \rangle \equiv \\ outFile1 \ll midpoint(t) \ll "\t" \\ \ll inf(y[0]) \ll "\t" \ll sup(y[0]) \ll "\t" \ll width(y[0]) \ll endl; This code is used in chunk 61.
```

```
65 \langle \text{ output solution } 65 \rangle \equiv \\ outFile1 \ll midpoint(t) \ll "\t" \\ \ll inf(y[0]) \ll "\t" \ll sup(y[0]) \ll "\t" \ll width(y[0]) \ll endl; \\ outFile2 \ll midpoint(t) \ll "\t" \ll Solver \rightarrow getStepsize() \ll endl; \\ \textbf{iVector } Y = Solver \rightarrow getAprioriEncl(); \\ \textbf{interval } Tj = Solver \rightarrow getT(); \\ outFile3 \ll inf(Tj) \ll "\t" \ll inf(Y[0]) \ll endl; \\ outFile3 \ll inf(Tj) \ll "\t" \ll sup(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll sup(Y[0]) \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll sup(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll endl; \\ outFile3 \ll sup(Tj) \ll "\t" \ll inf(Y[0]) \ll endl \ll
```

```
\begin{array}{l} outFile3 \ll \inf{(Tj)} \ll \text{"} \texttt{\t^{"}} \ll \sup{(Y[0])} \ll endl; \\ outFile3 \ll \sup{(Tj)} \ll \text{"} \texttt{\t^{"}} \ll \sup{(Y[0])} \ll endl \ll endl; \end{array}
```

This code is used in chunk 61.

## Plots

To visualize the results in these files, we produce the plots in Figure 3.5. In Figure 3.5(a) and (b), the upper and lower tight bounds cannot be distinguished when plotted. In (b), the a priori bounds are shown as boxes. The gnuplot file for generating this figure is displayed in Figure 6.5

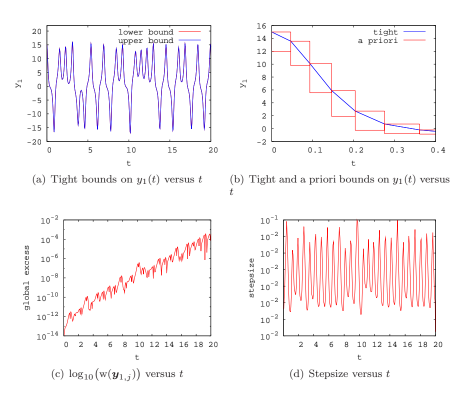


Figure 3.5. Plots generated using integctrl.cc

## 3.8 Work versus order

We show in Figure 3.6 how the computing time depends on the order for various tolerance when integrating the Lorenz system. We consider orders p = 5, 6, ..., 40 and tolerances atol =  $\text{rtol} = 10^{-7}, 10^{-8}, ..., 10^{-13}$ .

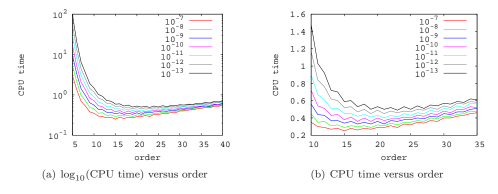


Figure 3.6. Plots generated using orderstudy.cc

The gnuplot file employed to generate these plots is given in Figure 6.6.

```
The main program is
68 \langle main program for order study 68 \rangle \equiv
                                   \textbf{static double} \ \ tol[\,] = \{1 \cdot 10^{-7}, 1 \cdot 10^{-8}, 1 \cdot 10^{-9}, 1 \cdot 10^{-10}, 1 \cdot 10^{-11}, 1 \cdot 10^{-12}, 
                                   int main()
                                                  const int n=3;
                                                    (create AD object 22)
                                                     (create a solver 23)
                                                  iVector y(n);
                                                   for (int i = 0; i < 7; i ++)
                                                                  Solver \rightarrow set Tols(tol[i]);
                                                                 (create file name 69)
                                                                 ofstream outFile(file_name.c_str(), ios::out);
                                                                 cout \ll " \_ \mathsf{tol} \_ = \_ " \ll tol[i] \ll
                                                                                             ":\squarewriting\squareinto\square" \ll file_name \ll "\square..." \ll endl;
                                                                 for (int p = 5; p \le 40; p++)
                                                                              Solver \rightarrow setOrder(p);
                                                                             interval t = 0.0, tend = 10.0;
```

```
y[0] = 15.0;
             y[1] = 15.0;
             y[2] = 36.0;
             Solver \rightarrow setFirstEntry();
             double time = getTime();
             \langle \text{ integrate (basic) } 24 \rangle
             (check if success 25)
             time = getTotalTime(time, getTime());
             outFile \ll p \ll "\t" \ll time \ll endl;
           outFile.close();
        return 0;
   This code is used in chunk 70.
      We create a file name for each tolerance value by
69 \langle create file name 69 \rangle \equiv
      string prefix("order");
      std::stringstream num(std::stringstream::out);
      num \ll tol[i];
      string file\_name = prefix + num.str() + ".out";
   This code is used in chunk 68.
      We store all this into
70 \langle \text{orderstudy.cc} 70 \rangle \equiv
   #include <fstream>
   #include <sstream>
   #include <string>
   #include <cstdlib>
   #include "vnode.h"
      using namespace std;
      using namespace vnodelp;
      (Lorenz 19)
      (main program for order study 68)
```

## 3.9 Work versus problem size

We investigate how the computing time depends on the size of the problem. We consider the DETEST problem C3 [14]

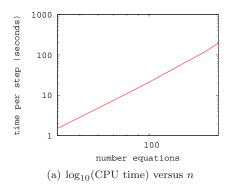
$$y' = \begin{pmatrix} -2 & 1 & 0 & 0 & \cdots & 0 \\ 1 & -2 & 1 & 0 & \cdots & 0 \\ 0 & 1 & -2 & 1 & \cdots & 0 \\ & & \vdots & & & \\ 0 & \cdots & & 1 & -2 & 1 \\ 0 & \cdots & & 0 & 1 & -2 \end{pmatrix} y$$

$$(3.2)$$

```
with y(0) = (1, 0, \dots, 0)^T. We integrate with problem sizes n = 40, 60, \dots, 200 for
   t = 0 \text{ to } t = 5.
         The C++ program is
71 \langle \text{detest\_c3.cc} 71 \rangle \equiv
   #include <ostream>
   #include "vnode.h"
     using namespace std;
     using namespace vnodelp;
     template \(\text{typename var_type}\)
     void DETEST_C3(int n, var_type *yp, const var_type *y, var_type t,
               void *param)
        yp[0] = -2.0 * y[0] + y[1];
        for (int i = 1; i < n - 1; i ++) {
          yp[i] = y[i-1] - 2.0 * y[i] + y[i+1];
        yp[n-1] = y[n-2] - 2.0 * y[n-1];
     int main()
        for (int n = 40; n \le 200; n += 20) {
          cout \ll n;
          interval t = 0.0, tend = 5;
          iVector y(n);
          for (int i = 0; i < n; i ++) y[i] = 0;
          y[0] = 1;
          AD *ad = new FADBAD\_AD(n, DETEST\_C3, DETEST\_C3);
          VNODE *Solver = new VNODE(ad);
          double time\_start = getTime();
          Solver \rightarrow integrate(t, y, tend);
          double time\_end = getTime();
```

```
cout \ll "_{\sqcup \sqcup}" \ll getTotalTime(time\_start, \\ time\_end) \ll "_{\sqcup \sqcup \sqcup}" \ll Solver \neg steps \ll endl; \\ \mathbf{delete} \ Solver; \\ \mathbf{delete} \ ad; \\ \} \\ \mathbf{return} \ 0; \\ \}
```

In Figure 3.7(a) and (b), we display the CPU time versus n. The gnuplot file for generating this figure is in Figure 6.7. It is not difficult to see that the computing



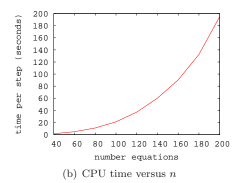


Figure 3.7. CPU time versus n for Problem 3.2. VNODE-LP takes 8 steps for each n.

time grows like  $n^3$ .

## 3.10 Stepsize behavior

We consider the orbit problem (see for example [4])

$$y_1'' = y_1 + 2y_2' - \widehat{\mu} \frac{y_1 + \mu}{D_1} - \mu \frac{y_1 - \widehat{\mu}}{D_2},$$
  

$$y_2'' = y_2 - 2u_1' - \widehat{\mu} \frac{y_2}{D_1} - \mu \frac{y_2}{D_2},$$
(3.3)

where

$$\mu = 0.012277471, \quad \hat{\mu} = 1 - \mu,$$
(3.4)

$$D_1 = ((y_1 + \mu)^2 + y_2^2)^{3/2}$$
, and (3.5)

$$D_2 = \left( (y_1 - \widehat{\mu})^2 + y_2^2 \right)^{3/2}. \tag{3.6}$$

We integrate this problem with

```
y_1(0) = 0.994,

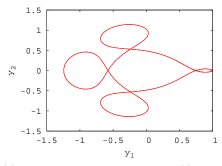
y_2(0) = 0,

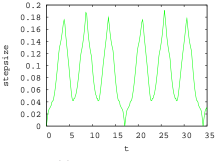
y'_1(0) = 0,

y'_2(0) = -2.00158510637908252240537862224,
(3.7)
```

and  $t_{\rm end} = 35$ , which corresponds to slightly more than two periods.

In Figure 3.8, we plot  $y_2$  versus  $y_1$  and the stepsize versus t. (The gnuplot file is in Figure 6.8.)





(a) Midpoints of the bounds on  $y_2(t)$  versus  $y_1(t)$ 

(b) Stepsize versus t

Figure 3.8. Plots generated using orbit.cc

The C++ program follows.

```
73 \langle \text{orbit.cc} 73 \rangle \equiv
   #include <fstream>
   #include <sstream>
   #include <string>
   #include <cstdlib>
   #include "vnode.h"
     using namespace std;
     using namespace vnodelp;
     template (typename var_type)
     void Orbit(int n, var\_type *yp, const var\_type *y, var\_type t,
               void *param)
        interval mu = string\_to\_interval("0.012277471");
        interval mu_h = 1.0 - mu;
        \mathbf{var\_type} \ \mathtt{D1} = pow(sqr(y[0] + mu) + sqr(y[1]), \mathbf{interval}(1.5));
        var_type D2 = pow(sqr(y[0] - mu_h) + sqr(y[1]), interval(1.5));
        yp[0] = y[2];
        yp[1] = y[3];
```

```
yp[2] = y[0] + 2.0 * y[3] - mu\_h * (y[0] + mu)/D1 - mu * (y[0] - mu\_h)/D2;
  yp[3] = y[1] - 2.0 * y[2] - mu\_h * y[1]/D1 - mu * y[1]/D2;
int main()
  const int n=4;
  iVector y(n);
  y[0] = string\_to\_interval("0.994");
  y[1] = 0;
  y[2] = 0;
  y[3] = string\_to\_interval("-2.00158510637908252240537862224");
  interval t = 0.0, tend = 35;
  AD *ad = new FADBAD\_AD(n, Orbit, Orbit);
  VNODE *Solver = new VNODE(ad);
  ofstream outFileSol("orbit_sol.out", ios::out);
  ofstream outFileStep("orbit_step.out", ios::out);
  outFileSol \ll midpoint(y[0]) \ll "\t" \ll midpoint(y[1]) \ll endl;
  Solver \rightarrow setOneStep(on);
  while (t \neq tend) {
    Solver \rightarrow integrate(t, y, tend);
    outFileSol \ll midpoint(y[0]) \ll "\t" \ll midpoint(y[1]) \ll endl;
    outFileStep \ll midpoint(t) \ll "\t"
         \ll Solver \rightarrow getStepsize() \ll endl;
  }
  outFileSol.close();
  outFileStep.close();
  return 0;
```

## 3.11 Stiff problems

We illustrate how VNODE-LP behaves when integrating a stiff problem. We integrate Van der Pol's equation (written as a first-order system)

$$y'_{1} = y_{2} y'_{2} = \mu(1 - y_{1}^{2})y_{2} - y_{1}$$
(3.8)

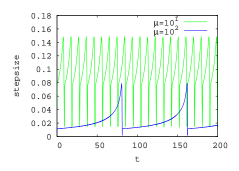
with

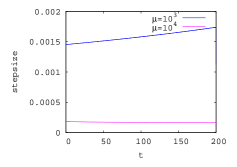
$$y(0) = (2,0)^T, \quad t_{\text{end}} = 200.$$
 (3.9)

We perform integrations with  $\mu = 10, 10^2, 10^3$ , and  $10^4$ . In Table 3.1, we show the number of steps and CPU time used by VNODE-LP. In Figure 3.9, we plot the stepsizes versus t. As can be seen from this table and figure, VNODE-LP is not efficient when this problem becomes stiff. In general, VNODE-LP works well on

$\mu$	steps	time (secs)
$10^{1}$	2377	2.4
$10^{2}$	11697	11.6
$10^{3}$	126459	124.4
$10^{4}$	1180844	1182.7

**Table 3.1.** Number of steps and CPU time used by VNODE-LP on (3.8–3.9)





**Figure 3.9.** Stepsize versus t on (3.8–3.9) for  $\mu = 10, 10^2, 10^3, 10^4$ 

non-stiff and mildly stiff problems. A more detailed study regarding stiff problems can be found in [23].

The program used to produce the numerical results for this problem follows. The gnuplot file for generating the plots is in Figure 6.9.

```
74 \langle vanderpol.cc 74 \rangle \equiv
#include <fstream>
#include <iomanip>
#include <sstream>
#include <string>
#include "vnode.h"

using namespace std;
using namespace vnodelp;

template \langle typename \ var\_type \rangle
void VDP(int \ n, var\_type \ *yp, const \ var\_type \ *y, var\_type \ t, void \ *param)
{
    double *MU = (double *) param;
    yp[0] = y[1];
    yp[1] = (*MU) * (1 - sqr(y[0])) * y[1] - y[0];
}

int main()
{
```

```
const int n=2;
double MU = 10.0;
AD *ad = new FADBAD\_AD(n, VDP, VDP, \&MU);
VNODE *Solver = new VNODE(ad);
   /* file for storing Table data */
ofstream outSteps("vdp_nosteps.out", ios::out);
outSteps \ll fixed \ll showpoint \ll setprecision(1);
for (int i = 1; i \le 4; i ++) {
            /* file for storing stepsizes */
  string prefix("vdp_step");
  std::stringstream num(std::stringstream::out);
  num \ll i;
  string file\_name = prefix + num.str() + ".out";
  ofstream outStepSizes(file\_name.c\_str(), ios :: out);
  cout \ll " \sqcup \texttt{MU} \sqcup = \sqcup " \ll \texttt{MU} \ll
       "\_writing\_into\_" \ll file\_name \ll "..." \ll endl;
  Solver \rightarrow setFirstEntry();
  Solver \rightarrow setOneStep(on);
  interval t = 0, tend = 200;
  iVector y(n);
  y[0] = 2.0, y[1] = 0;
  interval t\_prev = t;
  double time = getTime();
  while (t \neq tend) {
    if (midpoint(t - t\_prev) > 0.01 \lor t \equiv tend) {
       outStepSizes \ll midpoint(t) \ll "\t" \ll Solver\neg getStepsize() \ll endl;
       t\_prev = t;
    }
  outStepSizes.close();
  time = getTotalTime(time, getTime());
  outSteps \ll "$10^{"} \ll i \ll "}$" \ll "\t_{\omega}"
       \ll Solver \rightarrow getNoSteps() \ll " \t_{\sqcup} \&_{\sqcup}"
       \ll time \ll "\t" \ll "\\" \ll endl;
  MU *= 10.0;
  ad \rightarrow eval(\&MU);
outSteps.close();
return 0;
```

## Chapter 4

## Interface

First, we list the data types used by VNODE-LP that are of interest to the user. Then we list and describe briefly the public functions of the VNODE-LP solver.

## 4.1 Interval data type

VNODE-LP does not call the functions of these packages directly. Instead, it implements wrapper functions; see below. To build VNODE-LP on a new IA package, only these functions need to be implemented using this new package.

## 4.2 Wrapper functions

In the descriptions that follow, a, b, and c denote corresponding intervals in mathematical notation. We assume that the endpoints of the input intervals are representable machine numbers and no exceptions occur when a result is computed.

```
double inf (const interval &a) returns \underline{a}.
double sup (const interval &a) returns \overline{a}.
```

```
double midpoint(\mathbf{const\ interval\ }\&a)
      returns (\underline{a} + \overline{a})/2 rounded to the nearest.
 double width (const interval &a)
      returns \overline{a} - \underline{a} rounded to +\infty.
 double mag(const interval \& a)
      returns \max\{|\underline{a}|, |\overline{a}|\}.
 bool subseteq(\mathbf{const} \ \mathbf{interval} \ \&a, \mathbf{const} \ \mathbf{interval} \ \&b)
      returns true if a \subseteq b; false otherwise.
 bool interior(\mathbf{const} \ \mathbf{interval} \ \&a, \mathbf{const} \ \mathbf{interval} \ \&b)
      returns true if a is in the interior of b; false otherwise.
 bool disjoint(\mathbf{const}\ \mathbf{interval}\ \&a, \mathbf{const}\ \mathbf{interval}\ \&b)
      returns true if a \cap b = \emptyset; false otherwise.
 bool intersect (interval &c, const interval &a, const interval &b)
      returns true if a \cap b \neq \emptyset and sets c = a \cap b; false if a \cap b = \emptyset and leaves the
      input c unchanged.
 interval pi()
      returns an interval containing \pi.
 interval pow (const interval &a, const interval &b)
      returns an interval containing \{x^y \mid x \in \mathbf{a}, y \in \mathbf{b}\}.
 interval pow (const interval &a, int k)
      returns an interval containing \{x^k \mid x \in a\}.
Each of the functions e that follows returns \{e(x) \mid x \in a\}.
 interval exp(const interval \& a)
 interval log(const interval \& a)
 interval sqr(\mathbf{const} \ \mathbf{interval} \ \&a)
 interval sqrt (const interval &a)
 interval sin(const interval &a)
 interval cos(const interval \& a)
 interval tan(const interval \& a)
 interval asin(const interval \& a)
 interval acos (const interval &a)
 interval atan(const interval &a)
```

4.3. Interval vector 43

Finally,

```
interval string\_to\_interval(const char *s)
```

returns an interval that contains the decimal number that is stored in the character string input.

## 4.3 Interval vector

```
VNODE-LP uses interval vector, iVector, defined as
```

```
80 \langle interval vector 80 \rangle \equiv 
#include <vector>
#include "vnodeinterval.h"
    using namespace std;
    using namespace v_bias;
    typedef vector < interval > iVector;
```

This code is used in chunk 122.

## 4.4 Solver's public functions

The present solver is implemented by the class **VNODE**. We explain briefly its constructor and public member functions.

## 4.4.1 Constructor

```
VNODE(AD *ad)
```

constructs a **VNODE** object. Here *ad* is a pointer to an object of a class derived from the **AD** class; see Subsection 4.5. Currently, there is only one such class, **FADBAD\_AD**, and *ad* is a pointer to an object of this class.

## 4.4.2 Integrator

```
void integrate (interval &t, iVector &y, interval tend)
```

By default, integrate(t, y, tend) tries to compute an enclosure of the solution to an ODE problem at tend. If successful, y contains such an enclosure at t = tend. If an integration is not successful, y is an enclosure of the computed solution at  $t \neq tend$ .

The initial and end points,  $t_0$  and  $t_{\rm end}$ , can be stored as intervals containing their true values. Normally, the corresponding interval for  $t_0$  [resp.  $t_{\rm end}$ ] should be of the width of (at most) a few machine numbers. If we denote these intervals by  $t_0$  and  $t_{\rm end}$ , integrate requires that  $t_0 \cap t_{\rm end} = \emptyset$ . If  $t_0 \cap t_{\rm end} \neq \emptyset$ , integrate returns without performing an integration.

#### 4.4.3 Set functions

## void $setTols(\mathbf{double}\ a, \mathbf{double}\ r = 0)$

sets atol to a and rtol to r. The latter parameter has a default value of 0. For example,  $setTols(1 \cdot 10^{-10})$  sets atol =  $10^{-10}$  and rtol = 0.

The code roughly controls the "drift" away from the true solution at each integration step to be of size atol +  $\|y\|_{\infty}$  · rtol, where  $\|y\|$  is a measure of the size of current solution. This drift is accounted for by the enclosure, thus the size of the enclosure at the end point being roughly proportional to atol +  $\|y\|_{\infty}$  · rtol.

Default values are atol =  $10^{-12}$  and rtol =  $10^{-12}$ .

## void setOrder(int p)

sets the order to p. It must be between 3 and the value set to MAX\_ORDER in the makefile for building the library; cf. Figures 2.2 and 2.3. If p is not in this range, *integrate* returns without performing an integration.

Experience suggests that order in the range of about 20 to 30 results in efficient integration.

Default value is 20.

## void setHmin(double h)

sets the value of the magnitude of minimum stepsize allowed to h. If minimum stepsize is not set, or setHmin(0) is called, integrate computes such.

## void setFirstEntry()

indicates to *integrate* that this is a first entry into it. If this function is called before *integrate*, the latter will perform various initializations before time-stepping. When *integrate* is called for the first time, setFirstEntry() need not be called before integrate.

## void setOneStep(stepAction action)

tells the integrator whether to stop,  $action \equiv on$  or continue,  $action \equiv off$ , after each step it takes. If setOneStep(on) is called before integrate, the latter will return after each step it takes. To turn off this feature, call setOneStep(off). In this case, if integrate is re-entered, it will not stop after each step it takes (except the last one).

## 4.4.4 Get functions

#### bool successful() const

returns true if an integration is successful and false otherwise. If integrate has not been called, successful() returns true by default.

int getMaxOrder() returns the maximum order allowed in an integration. This is the value set in MAX\_ORDER in the configuration file.

#### double qetStepsize() const

returns the value of the most recent stepsize.

```
 \begin{array}{c} \textbf{double} \ \textit{getNoSteps}() \ \textbf{const} \\ \text{returns the number of successful steps taken by VNODE-LP.} \\ \textbf{const iVector} \ \& \textit{getAprioriEncl}() \ \textbf{const} \\ \text{returns the last computed} \ \widetilde{\boldsymbol{y}}_j. \\ \textbf{const interval} \ \& \textit{getT}() \ \textbf{const} \\ \text{returns the last computed} \ \boldsymbol{T}_j. \\ \textbf{double} \ \textit{getGlobalExcess}() \ \textbf{const} \\ \text{returns an estimate of the global excess in the most recent computed enclosure.} \\ \textbf{double} \ \textit{getGlobalExcess}(\textbf{int} \ i) \ \textbf{const} \\ \text{returns an estimate of the global excess in the} \ \textit{ith} \ \textbf{component of the most recent computed enclosure.} \\ \end{array}
```

## 4.5 Constructing an AD object

VNODE-LP computes Taylor coefficients for the ODE and its variational equation. The user has to construct an object for computing such coefficients by

```
AD *ad = new FADBAD\_AD(n, function\_name, function\_name)
or
AD *ad = new FADBAD\_AD(n, function\_name, function\_name, param)
```

Here n is the size of the problem, function\_name is the name of the template function, as described earlier, and param is a pointer to parameters that need to be passed to the ODE problem. After a parameter is changed, void eval(void \*p) must be called to update internal structures in the AD object.

## 4.6 Some helpful functions

```
template (class T) void printVector(const T &v, const char *s = 0)
    prints the components of a vector v on the standard output. If the second
    parameter is given, printVector prints it before the content of v.

double getTime()
    returns the current time measured as user time.

double getTotalTime(double start_time, double end_time)
    returns end_time - start_time rounded to the nearest.
```

## Chapter 5

# **Testing**

The code for the test cases is located in subdirectory tests. We give a brief description of each test.

## 5.1 General tests

File test0.cc

With the tests in this file, we check if the functions described in Subsection 4.2 compile and execute.

File test01.cc

We check if the elementary functions FADBAD++ uses compile and execute.

## 5.2 Linear problems

## 5.2.1 Constant coefficient problems

We consider

$$y_1' = y_2 y_2' = -y_1$$
 (5.1)

with

$$y(0) = (1,1)^T. (5.2)$$

The true solution is

$$y(t) = \begin{pmatrix} \cos(t) & \sin(t) \\ -\sin(t) & \cos(t) \end{pmatrix} y(0). \tag{5.3}$$

#### File test1.cc

We integrate (5.1, 5.2) from t = 0 to  $t_{\text{end}} = 10000$ . At each integration point  $t_j$ , selected by the solver, we evaluate (5.3) in interval arithmetic and check if the resulting enclosure intersects with the computed bounds. This test succeeds if they intersect for all  $t_j$ .

## File test2.cc

We integrate (5.1, 5.2) and check that at  $t = 2k\pi$ , for k = 2, 4, ..., 1000, y(0) is contained in the computed bounds. If so, this test is successful.

#### File test3.cc

We integrate (5.1, 5.2), but with  $t_{\text{end}} = -10000$ . This test is successful if the true solution (5.3), evaluated in interval arithmetic, and the computed bounds intersect at each point selected by the solver.

#### File test4.cc

We integrate (5.1, 5.2) and check that at  $-2k\pi$ , for  $k=2,4,\ldots,1000,\ y(0)$  is contained in the computed bounds. If so, this test is successful.

#### File test5.cc

We consider

$$y_1' = y_1 - 2y_2 y_2' = 3y_1 - 4y_2$$
 (5.4)

with

$$y(0) = (1, -1)^T. (5.5)$$

The true solution is

$$y(t) = \begin{cases} 5e^{-t} - 4e^{-2t} \\ 5e^{-t} - 6e^{-2t}. \end{cases}$$
 (5.6)

We integrate (5.4, 5.5) and check that, at each point selected by the solver, the true (evaluated in interval arithmetic (5.6)) and computed solutions intersect. If they do, we accept this test as successful.

## 5.2.2 Time-dependent problems

#### File test6.cc

The problem is

$$y'_{1} = \sin(t+10)y_{1} - 2y_{2} - y_{3} - \cos(t)$$

$$y'_{2} = 3y_{1} - 4\cos(t^{2})y_{2} - \cos(t)$$

$$y'_{3} = e^{-t^{2}}y_{1} - e^{-t^{2}}y_{2} - \sin(t)$$

$$y(0) \in ([0, 5], [-2, 6], [5, 12])^{T}, \quad t_{\text{end}} = 20.$$

We compute an enclosure at  $t_{\text{end}} = 20$  with the above initial condition set. Then we compute bounds at  $t_{\text{end}}$  for each corner of the initial box. These bounds must intersect with the enclosure resulting from  $([0, 5], [-2, 6], [5, 12])^T$ .

We have also computed (accurate) approximate solutions using MAPLE for each corner of the initial box. At  $t_{\rm end}$ , each approximate solution must be inside the bounds computed with the same corner point.

## 5.3 Nonlinear problems

#### File test\_n1.cc

We integrate the Lorenz system with  $y(0) = (15, 15, 36)^T$  from 0 to 1. Denote the enclosure at t = 1 by  $y_{(1)}$ . We have also computed an approximate solution with MAPLE. Denote it by  $\hat{y}_{(1)}$ . We check first if

$$\widehat{y}_{(1)} \in y_{(1)}.$$
 (5.7)

Then we integrate this system with  $y(1) \in \mathbf{y}_{(1)}$  and  $t_{\text{end}} = 0$ . Denote the computed enclosure at t = 0 by  $\mathbf{y}_{(0)}$ . We check if

$$y(0) \in \boldsymbol{y}_{(0)}. \tag{5.8}$$

Finally, we integrate with  $y(0) \in \boldsymbol{y}_{(0)}$  and  $t_{\text{end}} = 1$ . Denote the computed enclosure by  $\boldsymbol{y}_{(1)}^*$ . We check if

$$\hat{y}_{(1)} \in y_{(1)}^* \quad \text{and} \quad y_{(1)}^* \cap y_{(1)} \neq 0.$$
 (5.9)

This test is successful if of (5.7), (5.8), and (5.9) hold.

#### File test\_n2.cc

We integrate a three-body problem from 0 to 1 and then from 1 to 0. The initial condition at t = 0 must be contained in the computed bounds.

## File test\_n3.cc

The same three-body problem is integrated from 0 to 1 with orders from 10 to the value set in MAX\_ORDER. If all the computed enclosures with these orders intersect, we accept this test as successful.

#### File test\_n4.cc

We integrate [20]

$$y'' + cy' + \sin(y) = b\cos(t),$$

written as a first-order system

$$y'_1 = y_2$$
  
 $y'_2 = b\cos(t) - cy_2 - \sin(y_1)$ 

with

$$y(0) \in ([0,0], [1.9999, 2.0001])^T, \quad t_{\text{end}} = 8,$$

and c = 0 and b = 0.

We compare the computed enclosure by VNODE-LP and AWA [20]. If these enclosure intersect, we accept this test as successful.

#### File test\_n5.cc

This is the restricted three-body test problem from AWA:

$$x'' = x + 2y' - l \frac{x+m}{((x+m)^2 + y^2)^{3/2}} - m \frac{(x-1)}{((x-1)^2 + y^2)^{3/2}}$$
$$x'' = y - 2x' - l \frac{y}{((x+m)^2 + y^2)^{3/2}} - m \frac{y}{((x-1)^2 + y^2)^{3/2}},$$

where m = 1/82.45 and l = 1 - m. The initial values are

$$x(0) = 1.2,$$
  
 $x'(0) = 0,$   
 $y(0) = 0,$  and  
 $y'(0) = -1.04935750983.$ 

We integrate from 0 to 6.192169331396. Again, this test is successful if the computed enclosures by VNODE-LP and AWA intersect.

## File test\_n6.cc

We integrate the Pleiades problem from the Test Set for IVP Solvers [21]. At the end point, we subtract the reference solution, given in [21], from the computed bounds. If the max norm of the resulting interval vector is  $\leq 10^{-2}$ , we accept this test as successful.

# Chapter 6

# Listings

```
# CONFIG_FILE is set in vnodelp/makefile and exported in this
# file. vnodelp/makefile calls this makefile.
include ... / config / $ (CONFIG_FILE)
CXXFLAGS += -I.../include
                                 # compiler flags
LDFLAGS += -L../lib
                                 \# library flags
         = -lvnode \$(I\_LIBS) \$(LAPACK\_LIB) \setminus
LIBS
        $(BLAS_LIB) $(GPP_LIBS) # libraries
EXAMPLES = orbit vanderpol basic E1 scalar basic
        intermediate integetrl odeparam integi
        order detest_c3
examples: $(EXAMPLES)
E1: E1.o
        $(CXX) $(LDFLAGS) -o $@ E1.o $(LIBS)
scalar: scalar.o
        $(CXX) $(LDFLAGS) -o $@ scalar.o $(LIBS)
basic:
        basic.o
        $(CXX) $(LDFLAGS) -o $@ basic.o $(LIBS)
intermediate: intermediate.o
        $(CXX) $(LDFLAGS) -o $@ intermediate.o $(LIBS)
integetrl: integetrl.o
        $(CXX) $(LDFLAGS) -o $@ integctrl.o $(LIBS)
odeparam: odeparam.o
        $(CXX) $(LDFLAGS) -o $@ odeparam.o $(LIBS)
integi: integi.o
        $(CXX) $(LDFLAGS) -o $@ integi.o $(LIBS)
order: orderstudy.o
        $(CXX) $(LDFLAGS) -o $@ orderstudy.o $(LIBS)
detest_c3: detest_c3.o
        $(CXX) $(LDFLAGS) -o $@ detest_c3.o $(LIBS)
vanderpol: vanderpol.o
        $(CXX) $(LDFLAGS) -o $@ vanderpol.o $(LIBS)
orbit: orbit.o
        $(CXX) $(LDFLAGS) -o $@ orbit.o $(LIBS)
clean:
        @-\$(RM) *.o *.out core.* \$(EXAMPLES)
cleanall:
        @-\$(RM) *.o *.cc *.out core.* \$(EXAMPLES)
```

Figure 6.1. The makefile in the examples directory

```
\# file basici.gp
set terminal postscript eps enh color solid "Courier" 28
set xlabel "t"
set ylabel "y_1"
set output 'lorenzil.eps'
plot [0:6.3][-22:25] \setminus
'lorenzi.out' u 1:($2+$3) \
        title 'lower_bound' w l lt 1 lw 2,\
'lorenzi.out' u 1:(\$2-\$3)\
        title 'upper_bound' w 1 1t 3 lw 2
set output 'lorenzi2.eps'
set format y "%g"
set xtics 5.7,0.1,6.3
set ylabel "y_1" 0
plot [5.7:6.25][-22:0.4]
'lorenzi.out' u 1:2:3 \setminus
        title 'bounds'
                         w errorbars lw 2,\
'lorenzi.out' u 1:2
        title 'midpoint' w lines lt 3 lw 2
set output 'lorenzi_excess.eps'
set ylabel "global_excess" 2
set logscale y
set xtics 1
set format y "10^{%L}"
plot [0:6.3] 'lorenzi.out' u 1:4 \
        notitle w l lt 1 lw 2
```

Figure 6.2. The gnuplot file for generating the plot in Figure 3.3

```
function dy = E1(t,y)
dy = zeros(2,1);
t1 = t+1;
dy(1) = y(2);
dy(2) = -(y(2)/t1 + (1.0- 0.25/(t1*t1))*y(1));

clear;
options = odeset('RelTol',1e-10,'AbsTol',1e-10);

y(1) = 0.6713967071418030;
y(2) = 0.09540051444747446;

[T,Y] = ode45(@E1,[0 20],y,options);
format long
Y(end,1:2)'
```

Figure 6.3. The MATLAB code for the DETEST E1 problem

Figure 6.4. The gnuplot file for generating the plots in Figure 3.4

```
\# file integetrl.gp
set terminal postscript eps enh color solid "Courier" 28
set ylabel "y_1"
set xlabel "t"
set output 'lorenz.eps'
plot [][-20:22] \setminus
 'lorenz.tight' u 1:2 title 'lower_bound' w l lw 2,\
 'lorenz.tight' u 1:3 title 'upper_bound' w l lt 3 lw 2
set output 'lorenz2.eps'
set xtics 0, 0.1, 0.4
plot [0:0.4]\
 'lorenz.tight'
                u 1:2 title 'tight'
                                          w 1 lt 3 lw 2,\
 'lorenz.tight' u 1:3 notitle
                                          w 1 1t 3 lw 2,\
 'lorenz.apriori' u 1:2 title
                               'a_priori' w l lt 1 lw 3
set output 'lorenz_err.eps'
set xtics 0,2,20
set ylabel "global_excess" 2
set logscale y
set format y "10^{\frac{3}{2}}"
plot 'lorenz.tight' u 1:4 notitle w l lw 2
set output 'lorenz_step.eps'
set nologscale
set ylabel "stepsize"
plot 'lorenz.step' u 1:2 notitle w l lw 2
```

Figure 6.5. The gnuplot file for generating the plots in Figure 3.5

```
# file orderstudy.gp
set terminal postscript eps enh color solid "Courier" 28
set ylabel "CPU_time"
set xlabel "order"
set output 'order.eps'
plot
     [10:35]
       'order1e-07.out' u 1:2 title '10^{-7}'
w 1 1t 1 lw 2,\
      'order1e -08.out' u 1:2 title '10^{-8}'
w 1 1t 2 lw 2, \setminus
      'order1e -09.out 'u 1:2 title '10^{-}\{-9\}'
w 1 1t 3 lw 2,\
      'order1e-10.out' u 1:2 title '10^{-10}' w l lt 4 lw 2,\
      'order1e-11.out' u 1:2 title '10^{-11}' w l lt 5 lw 2,\
      'order1e-12.out' u 1:2 title '10^{-12}' w 1 lt 9 lw 2,\
      'order
1e -13.\mathrm{out}' u 1:2 \mathbf{title} '10^{-13}' w l lt 7 lw 2
set logscale y
set format y "10^{3}{%L}"
set output 'timeorder.eps'
plot 'order1e -07.out' u 1:2 title '10^{-7}'
                                               w l lt 1 lw 2, \
     'order1e -08.out' u 1:2 title '10^{-8}'
                                               w l lt 2 lw 2, \
     'order1e-09.out' u 1:2 title '10^{-9}'
                                               w 1 1t 3 lw 2,\
     'order1e-10.out' u 1:2 title '10^{-10}' w l lt 4 lw 2,\
     "order1e-11.out" u 1:2 title "10^{-11}" w 1 1t 5 lw 2,
     "order1e-12.out" u 1:2 title
                                    '10^{-12}' w l lt 9 lw 2,\
                                    10^{-13}, w l lt 7 lw 2
     "order1e-13.out" u 1:2 title
```

Figure 6.6. The gnuplot file for generating the plots in Figure 3.6

```
# file work.gp
set terminal postscript eps enh color solid "Courier" 28

# model of the work
f(x) = a + b*x

fit f(x) "work.out" using (log($1)):(log($2)) via a,b

set xlabel "number_equations"
set ylabel "time_per_step_(seconds)"
set xrange [40:200]

set output 'work.eps'
plot 'work.out' using 1:2 notitle with lines

set logscale
set output 'worklog.eps'
plot 'work.out' using 1:2 notitle with lines
```

Figure 6.7. The gnuplot file for generating the plots in Figure 3.7

```
# file orbit.gp
set terminal postscript eps enh color solid "Courier" 28

set xlabel "y_1"
set ylabel "y_2"

set output 'orbit_sol.eps'
plot 'orbit_sol.out' u 1:2 notitle w l lt 1 lw 2

set xlabel "t"
set ylabel "stepsize"
set output 'orbit_step.eps'
plot 'orbit_step.out' u 1:2 notitle w l lt 2 lw 2
```

Figure 6.8. The gnuplot file for generating the plots in Figure 3.8

Figure 6.9. The gnuplot file for generating the plots in Figure 3.9

# Part II Third-party Components

# **Packages**

The VNODE-LP package builds on

- LAPACK [2] and BLAS [1],
- interval-arithmetic (IA) package FILIB++ [19] or PROFIL/BIAS [16], and
- automatic differentiation (AD) package FADBAD++ [29].

The interfaces to the IA package are kept as small as possible, which allows a new package to be introduced without substantial programming effort.

Chapter 8 presents the implementation of interfaces to FILIB++ and PRO-FIL/BIAS. Chapter 9 discusses functions for changing the rounding mode. The AD in VNODE-LP is implemented through abstract classes, which are described in Chapter 17. Implementation of these classes using FADBAD++ is given in Chapter 22.

# IA package

The basic data type in VNODE-LP is **interval**. This package can be built on top of FILIB++ [19] or PROFIL [16], or potentially other packages. The user can select which of these packages to use; for more details see the installation instructions in Section 2.3.

Each of these IA packages can be replaced, provided that the new package supplies an interval data type with overloaded arithmetic operations and elementary functions working with interval arguments. To incorporate a new IA package, the body of the functions described in Section 4, and implemented below, need to be implemented using the new package.

## **8.1** Functions calling FILIB++

```
inline double inf (const interval &a) {
    return a.inf();
}
inline double sup(const interval &a) {
    return a.sup();
}
inline double midpoint(const interval &a) {
    return a.mid();
}
inline double width(const interval &a) {
    return a.diam();
}
inline double mag(const interval &a) {
    return a.diam();
}
```

```
inline bool subseteq(\mathbf{const} \ \mathbf{interval} \ \&a, \mathbf{const} \ \mathbf{interval} \ \&b) {
  return filib::subset(a, b);
inline bool interior(\mathbf{const} \ \mathbf{interval} \ \&a, \mathbf{const} \ \mathbf{interval} \ \&b) {
  return filib :: interior(a, b);
inline bool disjoint (const interval &a, const interval &b) {
  return filib :: disjoint(a, b);
inline bool intersect(interval \&c, const interval \&a, const interval \&b)
  if (filib:: disjoint(a, b)) return false;
  c = filib :: intersect(a, b);
  return true;
inline interval pi() {
  return filib::interval(double)::PI();
inline interval pow (const interval &a, const interval &b) {
  return filib :: pow(a, b);
inline interval pow (const interval &a, int b) {
  return filib:: power(a, b);
inline interval exp(const interval \& a) {
  return filib :: exp(a);
inline interval log(const interval \& a) {
  return filib::log(a);
inline interval sqr(const interval \& a) {
  return filib::sqr(a);
inline interval sqrt (const interval &a) {
  return filib::sqrt(a);
inline interval sin(const interval \& a) {
  return filib::sin(a);
inline interval cos(const interval \& a) {
  return filib :: cos(a);
```

```
inline interval tan(const interval \& a) {
                   return filib::tan(a);
         inline interval asin(const interval \& a) {
                   return filib:: asin(a);
         inline interval acos (const interval &a) {
                   return filib:: acos(a);
         inline interval atan(const interval \& a) {
                   return filib:: atan(a);
          inline interval string_to_interval(const char *s)
                   std :: cerr \ll "\n\n_! ***_! WARNING_! ***_! \n";
                   \mathrm{std} :: \mathit{cerr} \ll " \_ \_ \_ Conversion \_ from \_ a \_ string \_ to \_ an \_ int \setminus String \_ to \_ int \_ int \setminus String \_ to \_ an \_ int \_ in
                                      erval_containing_it_\n";
                   std::cerr \ll "_{ \sqcup \sqcup \sqcup \sqcup \sqcup \sqcup} has_{\sqcup} not_{\sqcup} been_{\sqcup} implemented_{\sqcup} in_{\sqcup} filib++. \n";
                   std::cerr \ll "_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}For_{\sqcup}this_{\sqcup}feature,_{\sqcup}consider_{\sqcup}using_{\sqcup}PROFIL/BIAS.\n";
                   "_{\sqcup}is_{\sqcup}converted_{\sqcup}to_{\sqcup}double.\n";
                   OT_BE_CORRECT_***_\n\n";
                   double a;
                   std::istringstream\,iss(s);
                   iss \gg a;
                   return interval(a);
This code is used in chunk 115.
```

# **8.2 Functions calling PROFIL**

```
114 \( \) functions calling PROFIL \( 114 \) \( \) inline double \( inf \) (const interval \( &a \) \( \) return \( Inf (a); \) \( \) inline double \( sup \) (const interval \( &a \) \( \) \( \) return \( Sup (a); \) \( \) inline double \( midpoint \) (const interval \( &a \) \( \) return \( Mid (a); \) \( \) inline double \( width \) (const interval \( &b \) \( \) \( \) inline double \( width \) (const interval \( &b \) \( \) \( \)
```

```
return Diam(b);
inline double mag(const interval \& a) {
  return Abs(a);
inline bool subseteq(\mathbf{const} \ \mathbf{interval} \ \&a, \mathbf{const} \ \mathbf{interval} \ \&b) {
  return a \leq b;
inline bool interior (const interval &a, const interval &b) {
  return (Inf(a)) > (Inf(b)) \wedge ((Sup(a)) < (Sup(b)));
inline bool disjoint (const interval &a, const interval &b) {
  interval c;
  return \neg Intersection(c, a, b);
inline bool intersect (interval &c, const interval &a, const interval &b)
  return Intersection(c, a, b);
inline interval pi() {
  return ArcCos(-1.0);
inline interval pow(const interval \& a, int b) {
  return Power(a, b);
inline interval pow(const interval \& a, const interval \& b) {
  return Power(a, b);
inline interval exp(const interval \& a) {
  return Exp(a);
inline interval log(const interval \& a) {
  return Log(a);
inline interval sqr(const interval \& a) {
  return Sqr(a);
inline interval sqrt (const interval &a) {
  return Sqrt(a);
inline interval sin(const interval \& a) {
  return Sin(a);
```

```
inline interval cos(const interval \& a) {
        return Cos(a);
      inline interval tan(const interval \& a) {
        return Tan(a);
      inline interval asin(const interval \& a) {
        return ArcSin(a);
      inline interval acos (const interval &a) {
        return ArcCos(a);
      inline interval atan (const interval &a) {
        return Arc Tan(a);
      inline interval string_to_interval(const char *s)
        return Enclosure(s);
    This code is used in chunk 115.
    Files
    The interface to the IA package is stored in
115 \langle \text{vnodeinterval.h} \ 115 \rangle \equiv
    #ifndef VNODEINTERVAL_H
    #define VNODEINTERVAL_H
    #ifdef PROFIL_VNODE
    #include <Interval.h>
    #include <Functions.h>
    #include <LongReal.h>
    #include <LongInterval.h>
      namespace v_bias {
         (interval data type (PROFIL) 76)
         (functions calling PROFIL 114)
    #endif
    #ifdef FILIB_VNODE
    #include <interval/interval.hpp>
    #include <iostream>
    #include <sstream>
    #include <string>
      namespace v_bias {
         (interval data type (FILIB++) 77)
         ⟨functions calling FILIB++ 113⟩
```

# Changing the rounding mode

For changing the rounding mode, VNODE-LP calls the functions below.

```
void round_nearest()
sets the rounding mode to the nearest.
void round_down()
sets the rounding mode to -∞.
void round_up()
sets the rounding mode to ∞.
```

Depending on the selected IA package, they call corresponding functions either from PROFIL/BIAS or FILIB++. A particular implementation is selected by the value of the variable I\_PACKAGE in the configuration file; see Subsection 2.3.2.

# 9.1 Changing the rounding mode using FILIB++

Changing the rounding mode using FILIB++ is done through **round\_control** defined as

```
117 \( \text{rounding control type (FILIB++)} \) 117 \\ \) \( \text{typedef filib::rounding_control} < \text{double} \), \( true > \text{round_control}; \)
This code is used in chunk 120.
```

The necessary functions are implemented as follows.

```
118 ⟨changing the rounding mode (FILIB++) 118⟩ ≡
    inline void round_nearest() {
        round_control::tonearest();
    }
    inline void round_down() {
        round_control::downward();
    }
}
```

#endif

```
inline void round_up() {
   round_control::upward();
}
This code is used in chunk 120.
```

## 9.2 Changing the rounding mode using BIAS

Similarly, we implement functions for changing the rounding mode using BIAS functions.

```
119 \langle changing the rounding mode (BIAS) 119 \rangle \equiv
      inline void round_nearest() {
         BiasRoundNear();
      inline void round_down() {
         BiasRoundDown();
      inline void round_up() {
         BiasRoundUp();
    This code is used in chunk 120.
    Files
    The above functions are stored in
120 \langle \text{vnoderound.h} 120 \rangle \equiv
    #ifndef VNODEROUND_H
    \#define VNODEROUND_H
    #ifdef FILIB_VNODE
    #include <rounding_control/rounding_control_double.hpp>
      namespace v_bias {
         ⟨rounding control type (FILIB++) 117⟩
         ⟨ changing the rounding mode (FILIB++) 118⟩
    #endif
    #ifdef PROFIL_VNODE
    #include "Bias0.h"
      namespace v_bias {
         (changing the rounding mode (BIAS) 119)
    #endif
```

# Part III

# Linear Algebra and Related Functions

# **Vectors and Matrices**

```
The VNODE-LP package works with point and interval vectors and matrices,
     namely pVector, iVector, pMatrix, and iMatrix. Here "p" is for point and
     "i" is for interval. The iVector type was defined in Subsection 4.3; pVector,
    pMatrix, and iMatrix are defined as
122 \langle vector and matrix types 122\rangle \equiv
       (interval vector 80)
       typedef vector (double) pVector;
       typedef vector (double) pMatrix;
       typedef vector (vector (interval)) iMatrix;
    This code is used in chunk 127.
     Size and memory allocation
      template \langle class\ Matrix \rangle void sizeM(Matrix\ \&A, unsigned\ int\ n)
          allocates space for an n \times n matrix.
      template \langle class\ Matrix \rangle sizeM (const Matrix &A)
          returns the size of a square matrix A.
      template \langle class\ Vector \rangle unsigned int size\ V(const\ Vector\ \&a)
          allocate space for an n vector.
      template \langle class\ Vector \rangle unsigned int size\ V(const\ Vector\ \&a)
          returns the size of a vector.
124 \langle \text{size/allocation } 124 \rangle \equiv
       template \langle class\ Matrix \rangle inline void sizeM(Matrix\ \&A, unsigned\ int\ n)
         A.resize(n);
         for (unsigned int i = 0; i < A.size(); i ++ A[i].resize(n);
```

```
template \langle class\ Matrix \rangle inline unsigned int sizeM(const\ Matrix\ \&A)
         return A.size();
       template \langle class\ Vector \rangle inline void sizeV(Vector\ \&a, unsigned\ int\ n)
         a.resize(n);
       template \langle class\ Vector \rangle inline unsigned int size\ V(const\ Vector\ \&a)
         return a.size();
    This code is used in chunk 127.
    Files
127 \langle \text{vector\_matrix.h} \quad 127 \rangle \equiv
    \#ifndef\ VECTOR\_MATRIX
    #define VECTOR_MATRIX
    #include <vector>
    #include "vnodeinterval.h"
       using namespace std;
       using namespace v_bias;
       namespace v_blas {
          (vector and matrix types 122)
          ⟨size/allocation 124⟩
    #endif
```

# **Basic functions**

We provide "generic" functions for operations involving matrices and vectors. Below, A, B, and C are  $n \times n$  matrices, x, y, and z are n vectors, and a is a scalar.

## 11.1 Vector operations

```
template (class Vector, class scalar)
    void setV(\mathbf{Vector} \ \&z, \mathbf{scalar} \ a)
    sets each component of z to a.
template (class Vector, class Vector)
    void assign V (Vector &z, const Vector &x)
    copies x to z
template (class Vector, class scalar)
    void scale V (Vector &z, scalar a)
    multiplies each element of z by a.
void addViVi(iVector \&z, const iVector \&x)
    adds z and x and stores the result in z.
void addViVp (iVector &z, const pVector &x)
    adds z and x and stores the result in z.
void sub ViVp(iVector \&z, const pVector \&x)
    subtracts x from z and stores the result in z.
void sub ViVi(iVector \&z, const iVector \&x)
    subtracts x from z and stores the result in z.
void addViVi(iVector \&z, const iVector \&x, const iVector \&y)
    adds x and y and stores the result in z.
```

```
void addViVp (iVector &z, const iVector &x, const pVector &y)
          adds x and y and stores the result in z.
     void subViVp(iVector \&z, const iVector \&x, const pVector \&y)
          subtracts y from x and stores the result in z.
     double inf\_normV(\mathbf{const\ iVector\ }\&z)
          returns \max |z_i|, where |a_i| = \max\{|\underline{a}_i|, |\overline{a}_i|\}.
     double inf\_normV(\mathbf{const\ pVector\ }\&z)
          returns ||z||.
     template (class scalar, class Vector1, class Vector2)
          inline void dot_product(scalar &r, const Vector1 &a, const Vector2 &b)
          computes the dot product of a and b and stores it in r. For input point vectors,
          this dot product is computed in the current rounding mode.
     double norm2 (const pVector &v)
          returns the two norm of a point vector. For a point vector, this norm is
          computed in the current rounding mode.
129 \langle vector operations 129 \rangle \equiv
       template (class Vector, class scalar)
                 inline void set V(\textbf{Vector } \&z, \textbf{scalar } a)
         fill(z.begin(), z.end(), a);
       template (class Vector1, class Vector2)
                 inline void assign V (Vector1 &z, const Vector2 &x)
         for (unsigned int i = 0; i < sizeV(z); i++) z[i] = x[i];
       template (class Vector, class scalar)
                 inline void scale V (Vector &z, scalar a)
         for (unsigned int i = 0; i < size V(z); i++) z[i] *= a;
       inline void addViVi(iVector \&z, const iVector \&x)
         transform(z.begin(), z.end(), x.begin(), z.begin(), plus(v_bias::interval)());
       inline void addViVp(iVector \&z, const pVector \&x)
         transform(z.beqin(), z.end(), x.beqin(), z.beqin(), plus(v_bias::interval)());
```

```
inline void sub ViVp (iVector &z, const pVector &x)
  transform(z.begin(), z.end(), x.begin(), z.begin(),
       minus\langle v\_bias :: interval \rangle ());
inline void sub ViVi(iVector \&z, const iVector \&x)
  transform(z.begin(), z.end(), x.begin(), z.begin(),
      minus(v_bias::interval)());
inline void addViVi (iVector &z, const iVector &x, const iVector &y)
  transform(x.begin(), x.end(), y.begin(), z.begin(), plus(v_bias::interval)());
inline void addViVp (iVector &z, const iVector &x, const pVector &y)
  transform(x.begin(), x.end(), y.begin(), z.begin(), plus(v_bias::interval)());
inline void sub ViVp (iVector &z, const iVector &x, const pVector &y)
  transform(x.begin(), x.end(), y.begin(), z.begin(),
       minus\langle v_bias :: interval \rangle ());
inline double inf\_normV (const iVector &z)
  double s = 0;
  for (unsigned int i = 0; i < sizeV(z); i++)
    if (\mathbf{v\_bias} :: mag(z[i]) > s) s = \mathbf{v\_bias} :: mag(z[i]);
  return s;
inline double inf\_normV(\mathbf{const}\ \mathbf{pVector}\ \&z)
  double s = 0;
  for (unsigned int i = 0; i < size V(z); i++)
    if (fabs(z[i]) > s) s = fabs(z[i]);
  return s;
template (class scalar, class Vector1, class Vector2)
          inline void dot_product(scalar &r, const Vector1 &a, const
         Vector2 \&b)
  r = 0.0;
  for (unsigned int i = 0; i < a.size(); i++) r += a[i] * b[i];
```

```
\begin{array}{l} \textbf{inline double} \ norm2 (\textbf{const pVector} \ \&v) \ \{\\ \textbf{double} \ s; \\ dot\_product(s,v,v); \\ \textbf{return} \ sqrt(s); \\ \} \\ \\ \textbf{This code is used in chunk } 137. \end{array}
```

# 11.2 Matrix/vector operations

#### This code is used in chunk 137.

# 11.3 Matrix operations

```
template \( class Matrix, class scalar \)
    void setM (Matrix &C, scalar a)
    sets each component of C to a.

template \( class Matrix \) void setId (Matrix &C)
    sets C to the identity matrix.

template \( class Matrix1, class Matrix2 \)
    void assignM (Matrix1 &C, const Matrix2 &A)
    copies A to C.
```

```
template \langle class Matrix, class scalar \rangle void scaleM(Matrix \&C, scalar a)
          multiplies each component of C by a.
      template \langle class\ Matrix \rangle void transpose(Matrix\ \&C, const\ Matrix\ \&A)
          stores the transpose of A in C.
      template \langle class Matrix \rangle void addId(Matrix \&C)
          adds the identity matrix to C.
      template \langle class \ Matrix \rangle \ void \ subFromId(Matrix \& C)
          subtracts C from the identity matrix and stores the result in C.
      void addMiMi(iMatrix \&C, const iMatrix \&A)
          adds C and A and stores the result in C.
      void subMiMp (iMatrix &C, const pMatrix &A)
          subtracts A from C and stores the result in C.
      void multMiMi(iMatrix \&C, const\ iMatrix \&A, const\ iMatrix \&B)
          multiplies A and B and stored the result in C.
      void multMiMp (iMatrix &C, const iMatrix &A, const pMatrix &B)
          multiplies A and B and stored the result in C.
      double inf\_normM (const iMatrix &C)
          computes ||C||_{\infty}. The computation is in the current rounding mode. For
          example, if an upper bound on ||C||_{\infty} is desired, the rounding mode must be
          set to +\infty before calling inf\_normM.
132 \langle \text{ matrix operations } 132 \rangle \equiv
       template (class Matrix, class scalar)
                 inline void setM(Matrix \&C, scalar a)
         for (unsigned int i = 0; i < C.size(); i ++ ) set V(C[i], a);
       template (class Matrix)
                inline void setId (Matrix &C)
         setM(C, 0.0);
         for (unsigned int i = 0; i < sizeM(C); i++) C[i][i] = 1.0;
       template (class Matrix1, class Matrix2)
                 inline void assignM(Matrix1 &C, const Matrix2 &A)
         for (unsigned int i = 0; i < C.size(); i ++ ) assign V(C[i], A[i]);
```

```
template (class Matrix, class scalar)
         inline void scaleM(Matrix \&C, scalar a)
  for (unsigned int i = 0; i < C.size(); i ++ ) scale V(C[i], a);
template (class Matrix)
         inline void transpose(Matrix \&C, const Matrix \&A)
  for (unsigned int i = 0; i < C.size(); i++) setColumn(C, A[i], i);
template (class Matrix)
         inline void addId(Matrix \&C)
  for (unsigned int i = 0; i < sizeM(C); i++) C[i][i] += 1.0;
template (class Matrix)
         inline void subFromId(Matrix \&C)
  unsigned int n = sizeM(C);
  for (unsigned int i = 0; i < n; i \leftrightarrow)
    for (unsigned int j = 0; j < n; j +++) C[i][j] = -C[i][j];
  for (unsigned int i = 0; i < n; i++) C[i][i] += 1.0;
inline void addMiMi (iMatrix &C, const iMatrix &A)
  for (unsigned int i = 0; i < C.size(); i \leftrightarrow ddViVi(C[i], A[i]);
inline void subMiMp (iMatrix &C, const pMatrix &A)
  for (unsigned int i = 0; i < C.size(); i \leftrightarrow sub ViVp(C[i], A[i]);
inline void multMiMi(iMatrix \&C, const iMatrix \&A, const iMatrix \&B)
  unsigned int n = sizeM(A);
  for (unsigned int i = 0; i < n; i++)
    for (unsigned int j = 0; j < n; j \leftrightarrow j) {
      C[i][j] = 0.0;
      for (unsigned int k = 0; k < n; k++) C[i][j] += A[i][k] * B[k][j];
}
inline void multMiMp (iMatrix &C, const iMatrix &A, const pMatrix &B)
  unsigned int n = sizeM(A);
```

```
for (unsigned int i = 0; i < n; i +++)
           for (unsigned int j = 0; j < n; j \leftrightarrow) {
              C[i][j] = 0.0;
             for (unsigned int k = 0; k < n; k++) C[i][j] += A[i][k] * B[k][j];
      template \langle class \ Matrix \rangle in line double inf\_normM (const Matrix &C)
         unsigned int n = sizeM(C);
         double m=0;
         for (unsigned int i = 0; i < n; i \leftrightarrow) {
           double s = 0:
           for (unsigned int j = 0; j < n; j \leftrightarrow v-bias:: mag(C[i][j]);
           if (s > m) m = s;
         return m;
    This code is used in chunk 137.
     11.4
               Get/set column
      template (class Vector, class Matrix)
          void getColumn(Vector \&z, const Matrix \&C, unsigned int j)
          stores the jth column of C in z.
      template (class Matrix, class Vector)
          void setColumn(Matrix \&C, const Vector \&z, unsigned int j)
          sets the jth column of C to z.
134 \langle \text{get/set column } 134 \rangle \equiv
      template (class Vector, class Matrix)
       void getColumn(Vector \&z, const Matrix \&C, unsigned int j)
       {
         for (unsigned int i = 0; i < sizeM(C); i++) z[i] = C[i][j];
      template (class Matrix, class Vector)
       void setColumn(Matrix \&C, const Vector \&z, unsigned int j)
         for (unsigned int i = 0; i < sizeM(C); i++) C[i][j] = z[i];
    This code is used in chunk 137.
```

#### 11.5 Conversions

The following two functions are convenient when calling LAPACK.

```
template (class scalar, class Matrix)
         void matrix2pointer(scalar *M, const Matrix &C)
         copies C into an array pointed to by M. The matrix at M is in a column-
         major form. That is, the (i, j) element of the matrix in the array at M is at
         jn+i, where n is the size of the matrix.
     template (class Matrix, class scalar)
         void pointer2matrix(Matrix \&C, const scalar *M)
         copies a matrix stored in an array pointed to by M into a matrix C. The
         matrix at M is in a column-major form. That is, the (i,j) element of the
         matrix in the array at M is at jn + i, where n is the size of the matrix.
135 \langle \text{matrix2pointer } 135 \rangle \equiv
      template (class scalar, class Matrix)
                inline void matrix2pointer(scalar *M, const Matrix &C)
        unsigned int n = sizeM(C);
        for (unsigned int j = 0; j < n; j \leftrightarrow j
           This code is used in chunk 137.
136 \langle pointer2matrix 136 \rangle \equiv
      template (class Matrix, class scalar)
                inline void pointer2matrix(Matrix \&C, const scalar *M)
        unsigned int n = sizeM(C);
        for (unsigned int j = 0; j < n; j ++)
           for (unsigned int i = 0; i < n; i++) C[i][j] = M[j * n + i];
    This code is used in chunk 137.
    Files
    We store the above functions in
137 \langle basiclinalg.h 137 \rangle \equiv
    #ifndef BASICLINALG_H
    #define BASICLINALG_H
    #include <algorithm>
    #include "vector_matrix.h"
      namespace v_blas {
         (vector operations 129)
         (matrix times vector 131)
         matrix operations 132
         ⟨get/set column 134⟩
```

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```
\begin{array}{c} \langle\, matrix2pointer\,\, 135\,\rangle \\ \langle\, pointer2matrix\,\, 136\,\rangle \\ \langle\, print\,\, vector\,\, 366\,\rangle \\ \big\} \\ \#endif \end{array}
```

# Interval functions

## 12.1 Inclusion

This subseteq function returns true if, for two interval vectors a and b,  $a \subseteq b$ , and returns false otherwise.

```
139 \langle check vector inclusion 139 \rangle \equiv
inline bool subseteq(\mathbf{const\ iVector}\ \&a, \mathbf{const\ iVector}\ \&b)
{
    return equal(a.begin(), a.end(), b.begin(), \mathbf{v\_bias} :: subseteq);
}
See also chunk 141.
This code is used in chunk 151.
```

#### 12.2 Interior

The *interior* function returns true if, for two interval vectors a and b, a is componentwise in the interior of b, and returns false otherwise.

```
nentwise in the interior of b, and returns false otherwise.

(check if in the interior 140) ≡
    inline bool interior(const iVector &a, const iVector &b)

{
        return equal(a.begin(), a.end(), b.begin(), v_bias::interior);
    }

This code is used in chunk 151.

The disjoint function returns true if, for two interval vectors a and b, a ∩ b = ∅, and returns false otherwise.

141 ⟨check vector inclusion 139⟩ +≡
    inline bool disjoint(const iVector &a, const iVector &b)
    {
        return equal(a.begin(), a.end(), b.begin(), v_bias::disjoint);
    }
```

## 12.3 Radius

This code is used in chunk 151.

```
The radius of an interval is
142 \langle \operatorname{rad} (\operatorname{interval}) 142 \rangle \equiv
        inline double rad (const interval &a)
           return 0.5 * \mathbf{v\_bias} :: width(a);
     This code is used in chunk 151.
        We compute the radius of an interval vector by
143 \langle \text{ rad (vector) } 143 \rangle \equiv
        inline void rad(\mathbf{pVector} \& r, \mathbf{const} \ \mathbf{iVector} \& v)
           transform(v.begin(), v.end(), r.begin(), \mathbf{v\_bias} :: rad);
     This code is used in chunk 151.
                  Width
     12.4
     We compute the width of an interval vector by
144 \langle \text{ width } 144 \rangle \equiv
        inline void width(\mathbf{pVector} \& r, \mathbf{const} i\mathbf{Vector} \& a)
           transform(a.begin(), a.end(), r.begin(), \mathbf{v\_bias} :: width);
     This code is used in chunk 151.
     12.5
                  Midpoints
145 \langle midpoint of an interval vector 145 \rangle \equiv
        inline void midpoint (pVector &r, const iVector &a)
           transform(a.begin(), a.end(), r.begin(), \mathbf{v\_bias} :: midpoint);
     This code is used in chunk 151.
146 \langle midpoint of an interval matrix 146 \rangle \equiv
        inline void midpoint(pMatrix \& R, const iMatrix \& A)
           for (unsigned int i = 0; i < A.size(); i++) midpoint(R[i], A[i]);
```

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## 12.6 Intersection

If interval vectors x and y intersect, we store their intersection in z and return true. Otherwise, we return false.

```
147 (intersection of interval vectors 147) \equiv
inline bool intersect(iVector \&z, const iVector \&x, const iVector \&y) {
    interval c;
    for (unsigned int i=0;\ i < sizeV(y);\ i++) {
        bool b=\mathbf{v_-bias}::intersect(c,x[i],y[i]);
        if (\neg b) return false;
        else z[i]=c;
    }
    return true;
```

This code is used in chunk 151.

# 12.7 Computing h such that $[0, h]a \subseteq b$

#### 12.7.1 The interval case

Given intervals  $\boldsymbol{a}$  and  $\boldsymbol{b}$ , where  $0 \in \boldsymbol{b}$ , we wish to compute the largest machine-representable  $h \geq 0$  such that

```
[0,h] \mathbf{a} \subseteq \mathbf{b}.
```

We consider the following cases.

- 1.  $\underline{a} = \overline{a}$ .
  - (a) If  $\underline{a} = \overline{a} = 0$ , then we set h to be the largest representable machine number.
  - (b) If  $\underline{a} = \overline{a} > 0$ , then  $[0, h]a = [0, h\overline{a}] \subseteq b$  iff  $h \leq \overline{b}/\overline{a}$ .
  - (c) If  $\underline{a} = \overline{a} < 0$ , then  $[0, h]a = [\underline{a}h, 0] \subseteq b$  iff  $h \leq \underline{b}/\underline{a}$ .
- 2.  $\underline{a} \neq \overline{a}$ .
  - (a)  $\underline{a} \geq 0$ . Then  $\overline{a} > 0$  and  $[0, h]a = [0, h\overline{a}] \subseteq b$  iff  $h \leq \overline{b}/\overline{a}$ .
  - (b)  $\overline{a} \leq 0$ . Then  $\underline{a} < 0$  and  $[0, h]a = [h\underline{a}, 0] \subseteq b$  iff  $h \leq \underline{b}/\underline{a}$ .

We summarize the above cases in the following procedure:

- 1. If  $\underline{a} = \overline{a} = 0$  then set h to be the largest representable machine number;
- 2. else
  - (a) if  $\underline{a} \geq 0$  then  $h = \nabla(\overline{b}/\overline{a})$
  - (b) else  $h = \nabla(\underline{b}/\underline{a})$ .

```
149 \langle h \text{ such that } [0,h]a \subseteq b \text{ (intervals) } 149 \rangle \equiv \\ \# \text{include} < \text{climits} > \\ \text{using namespace std;} \\ \text{using namespace v_bias;} \\ \text{inline double } compH(\text{const v_bias}::\text{interval } \&a, \text{const v_bias}::\text{interval } \&b) \\ \{ \\ \text{if } (inf(a) \equiv 0 \land sup(a) \equiv 0) \text{ return numeric_limits} \langle \text{double} \rangle :: max(); \\ round\_down(); \\ \text{if } (inf(a) \geq 0) \text{ return } sup(b)/sup(a); \\ \text{return } inf(b)/inf(a); \\ \} \\ \text{This code is used in chunk 150.}
```

#### 12.7.2 The interval vector case

Given two interval vectors a and b, where b contains the vector with zero component, we compute the largest machine-representable  $h \ge 0$  such that

```
[0,h]\boldsymbol{a} \subset \boldsymbol{b}.
150 \langle h \text{ such that } [0, h] \boldsymbol{a} \subseteq \boldsymbol{b} \text{ (interval vectors) } 150 \rangle \equiv
         \langle h \text{ such that } [0, h] \boldsymbol{a} \subseteq \boldsymbol{b} \text{ (intervals) } 149 \rangle
         double compH(\mathbf{const\ iVector\ }\&a,\mathbf{const\ iVector\ }\&b)
            double hmin = compH(a[0], b[0]);
            for (unsigned int i = 1; i < sizeV(a); i \leftrightarrow j {
               double h = compH(a[i], b[i]);
               if (h < hmin) hmin = h;
            return hmin;
      This code is used in chunk 152.
      Files
151 \langle intvfuncs.h 151 \rangle \equiv
      #ifndef INTVFUNC_H
      #define INTVFUNC_H
      #include "vector_matrix.h"
         namespace v_bias {
            (rad (interval) 142)
         } namespace v_blas {
            using namespace v_bias;
```

```
\langle check if in the interior 140\rangle
           ⟨ check vector inclusion 139 ⟩
           (width 144)
           (midpoint of an interval vector 145)
           (midpoint of an interval matrix 146)
           \langle intersection of interval vectors 147\rangle
           ⟨rad (vector) 143⟩
           double compH(\mathbf{const\ iVector\ }\&a,\mathbf{const\ iVector\ }\&b);
     \#endif
152 \langle intvfuncs.cc 152 \rangle \equiv
     #include <limits>
      #include <algorithm>
     \#include "vnodeinterval.h"
      #include "vnoderound.h"
     #include "vector_matrix.h"
        namespace v_blas {
           \langle h \text{ such that } [0,h] \boldsymbol{a} \subseteq \boldsymbol{b} \text{ (interval vectors) } 150 \rangle
```

# **QR** factorization

In our implementation of VNODE-LP, we need to compute the QR factorization of an  $n \times n$  matrix. We employ the routines DGEQRF and DORGQR from LAPACK. DGEQRF computes a QR factorization of a real  $m \times n$  matrix A. DORGQR generates an  $m \times n$  real matrix Q with (approximately) orthonormal columns, which is defined from the elementary reflectors returned by DGEQRF. If computeQR is successful, true is returned; otherwise false is returned.

```
153 \langle compute QR factorization 153\rangle \equiv
       extern "C"
         void dgeqrf_{-}(int *m, int *n, double *A, int *lda, double *tau, double
              *work, int *lwork, int *info);
         void dorgqr_{-}(int *m, int *n, int *k, double *A, int *lda, double
              *tau, double *work, int *lwork, int *info);
       }
       bool compute QR(\mathbf{pMatrix} \& Q, \mathbf{const} \ \mathbf{pMatrix} \ \& A)
         int n = sizeM(A);
         int m=n;
         int lda = n;
         int k = n;
         int info;
                                  /* lwork has to be n *(optimal block size). The value
         int lwork = 10 * n;
              10 is somewhat random. */
         double *tau = new double[n];
         double *work = new double[lwork];
         double *M =  new double [n * n];
         v_bias::round_nearest();
         matrix2pointer(M, A);
         dgeqrf_{\bullet}(\&m,\&n,M,\&lda,tau,work,\&lwork,\&info);
```

```
if (info \equiv 0) {
            dorgqr\_(\&m,\&n,\&k,M,\&lda,tau,work,\&lwork,\&info);
            if (info \equiv 0) pointer2matrix(Q, M);
          \mathbf{delete}[]M;
          delete[] work;
          \mathbf{delete}[] \ tau;
          if (info \equiv 0) return true;
          return false;
     This code is used in chunk 154.
     Files
154 \langle qr.cc 154 \rangle \equiv
     \#include "basiclinalg.h"
     #include "vnoderound.h"
       namespace vnodelp {
          using namespace v_blas;
          \langle compute QR factorization 153\rangle
```

# Matrix inverse

The **MatrixInverse** class provides functions for computing an approximate inverse of a point matrix, enclosing the inverse of a point matrix and enclosing the inverse of a floating-point approximation to an orthogonal matrix.

## 14.1 Matrix inverse class

```
156 \langle class MatrixInverse 156 \rangle \equiv
      class MatrixInverse {
      public:
        MatrixInverse(int n);
        bool invertMatrix (pMatrix & Ainv, const pMatrix & A);
        bool encloseMatrixInverse (iMatrix & Ainv, const pMatrix & A);
        bool orthogonalInverse (iMatrix & Ainv, const pMatrix & A);
        \simMatrixInverse();
        int iterations;
      private:
        bool encloseLS (iVector &x, const pMatrix &A, const iVector &b0,
             const iMatrix &B, double beta);
        pVector radx;
        iVector b\theta, x1, x;
        iMatrix B, Ci;
        pMatrix C;
        double *M;
        int *ipiv;
        double *work;
        int lwork;
      };
    This code is used in chunk 172.
```

invertMatrix tries to compute a floating-point approximation to the inverse of A (if it exists). If successful, invertMatrix stores the result in Ainv and returns true; otherwise, it returns false.

encloseMatrixInverse tries to enclose the inverse of A (if it exists). If successful, encloseMatrixInverse stores the result in Ainv and returns true; otherwise, it returns false.

orthogonalInverse tries to enclose the inverse of a floating-point approximation to an orthogonal matrix. If successful, orthogonalInverse stores the result in Ainv and returns true; otherwise it returns false.

# 14.2 Computing $A^{-1}$

• info success if info  $\equiv 0$ 

**v\_bias**::round\_nearest();

if  $(info \neq 0)$  { #ifdef VNODE\_DEBUG

 $dgetrf_{-}(\&n,\&n,M,\&lda,ipiv,\&info);$ 

First, we compute the LU factorization of A using LAPACK's dgetrf. Then, using this LU factorization, we try to compute the inverse of A using LAPACK's dgetri.

```
printMessage("Could_not_invert_a_matrix");
#endif
      return false;
         /*
   • lwork size of work. We set lwork = 2 * n in the constructor
   • work array of size lwork
   • info success if info \equiv 0
    dgetri\_(\&n, M, \&lda, ipiv, work, \&lwork, \&info);
    if (info \neq 0) {
#ifdef VNODE_DEBUG
      printMessage("Could_not_invert_a_matrix");
#endif
      return false;
    pointer2matrix(Ainv, M);
    return true;
This code is used in chunk 173.
```

## 14.3 Enclosing the solution of a linear system

We try to enclose the solution to the linear system Ax = b,  $A \in \mathbb{R}^{n \times n}$ ,  $b \in \mathbb{R}^n$  in encloseLS. It implements Krawczyk's method; see for example [10]. By  $C \in \mathbb{R}^{n \times n}$  we denote a preconditioner. This method works if  $||I - CA||_{\infty} = \beta < 1$ . Let

$$\alpha = \frac{\|Cb\|_{\infty}}{1 - \beta}.$$

Then

$$x \in ([-\alpha, \alpha], \dots, [-\alpha, \alpha])^T$$
.

This shows how to compute an initial box that is guaranteed to contain x. Then Krawczyk's iteration is

$$\boldsymbol{x}^{(i+1)} = \left(Cb + (I - CA)\boldsymbol{x}^{(i)}\right) \cap \boldsymbol{x}^{(i)}.$$

The *encloseLS* function follows. In the comment inside it, before the horizontal line, we list the input variables and what they contain. The output is described after this line.

When describing code in this manuscript, we shall mix variable names and math symbols, where appropriate. Such a mixture may be viewed as an abuse of notation, but the author has found it helpful. For example  $b\theta \ni Cb$  below means that the interval vector  $b\theta$  contains the true Cb. Similarly,  $beta \ge \|I - CA\|_{\infty}$  denotes that the value of the input variable beta must be greater or equal the true value of  $\|I - CA\|_{\infty}$ .

```
160 \langle enclose the solution to Ax = b 160 \rangle \equiv
       bool MatrixInverse :: encloseLS (iVector &x,
                const pMatrix &A, const iVector &b\theta,
                const iMatrix &B, double beta)
             /*
                    A is the matrix from Ax = b
                   b\theta \ni Cb
                    B \ni I - CA
                  beta \ge ||I - CA||_{\infty}, beta must be < 1
                     x contains the solution to Ax = b if true is returned
         (compute initial box 161)
         ⟨ do Krawczyk's iteration 165⟩
         return true;
    This code is used in chunk 173.
    14.3.1 Initial box
    The input b0 contains Cb, and the input B contains I - CA. Also, beta \ge ||I - CA||
```

 $CA\|_{\infty}$ . We compute  $a \geq \|Cb\|_{\infty}/(1-\beta)$  and set the initial box containing the solution.

```
161 \langle compute initial box 161 \rangle \equiv
         \mathbf{v\_bias} :: round\_down();
         double a = 1 - beta; /* a \le 1 - \beta */
         double a1 = inf\_normV(b0); /* a1 \ge ||Cb||_{\infty} */
         \mathbf{v}_{\mathbf{bias}} :: round_{\mathbf{u}p}();
         a = a1/a;
                          /* a \ge ||Cb||_{\infty}/(1-\beta) */
         set V(x, \mathbf{v\_bias} :: \mathbf{interval}(-a, a));
      This code is used in chunk 160.
```

#### 14.3.2 Krawczyk's iteration

We implement  $Cb+(I-CA)x^{(i)}$ . That is, we program the expression x1=b0+B\*x.

```
162 \langle Krawczyk's iteration 162 \rangle \equiv
        multMiVi(x1, B, x);
        addViVi(x1,b0);
     See also chunk 163.
```

This code is used in chunk 165.

Now we can intersect x1 and x. The result is stored in x, if  $b \equiv true$ . Otherwise, we return false.

```
163 \langle Krawczyk's iteration 162\rangle +\equiv
       bool b = intersect(x, x, x1);
       if (b \equiv false) {
     #ifdef VNODE_DEBUG
          printMessage("x_{\sqcup}and_{\sqcup}x1_{\sqcup}do_{\sqcup}not_{\sqcup}intersect");
     #endif
          return false;
       For the while loop that follows, we need to compute the sum of the radii of the
     components of x.
164 \langle \text{sum radii } 164 \rangle \equiv
       rad(radx, x);
       v_bias::round_nearest();
        sum\_radii = accumulate(radx.begin(), radx.end(), 0.0);
     This code is used in chunk 165.
       Now, we compose the whole iteration. Initially, we set sum_old_radii to the
     largest machine number. The while loop below iterates as far as sum_radii <
     mult * sum\_old\_radii. The factor mult = (1 + beta)/2 is the same as in [10].
165 \langle do Krawczyk's iteration 165 \rangle \equiv
       double sum\_old\_radii = \mathbf{numeric\_limits} \langle \mathbf{double} \rangle :: max();
       double sum_radii;
        (sum radii 164)
        round\_up();
       int max\_iterations = 20;
       int counter = 0;
       double mult = (1 + beta)/2;
       while (sum\_radii < mult * sum\_old\_radii \land counter < max\_iterations) {
          ⟨Krawczyk's iteration 162⟩
          sum\_old\_radii = sum\_radii;
          (sum radii 164)
          counter ++:
        iterations = counter;
     This code is used in chunk 160.
```

#### 14.4 Enclosing the inverse of a general point matrix

To enclose the inverse of a point matrix A, we enclose the solution to

```
Ax_i = e_i for i = 1, \ldots, n,
```

where  $e_i$  is the *i*th unit vector. Then, the *i*th column of the inverse is  $x_i$ .

First we compute a floating-point inverse of A. If invertMatrix fails, encloseMatrixInverse returns false. Then we compute beta that is needed for encloseLS and call this function for each  $e_i$ .

```
166 \langle enclose the inverse of a matrix 166 \rangle \equiv
        bool MatrixInverse:: encloseMatrixInverse(iMatrix & Ainv,
                  const pMatrix \&A)
          bool b = invertMatrix(C, A);
          if (b \equiv false) return false;
          \langle \text{ find beta 167} \rangle
          (enclose each column 169)
     #ifdef VNODE_DEBUG
          iMatrix B = Ainv;
          multMiMp(B, Ainv, A);
          int n = sizeM(A);
          for (int i = 0; i < n; i ++)
             for (int j = 0; j < n; j ++) {
               interval b = B[i][j];
               if (i \equiv j) assert (v_bias::subseteq(interval(1.0), b));
               else assert(\mathbf{v\_bias} :: subseteq(\mathbf{interval}(0.0), b));
     #endif
          return true;
     This code is used in chunk 173.
        We compute B such that it contains I - CA.
167 \langle \text{ find beta 167} \rangle \equiv
        assignM(Ci, C);
        multMiMp(B, Ci, A);
        subFromId(B);
     See also chunk 168.
     This code is used in chunks 166 and 170.
        Now we find beta \geq \beta. If beta \geq 1, we return false, as Krawczyk's iteration
     cannot proceed.
168 \langle \text{ find beta } 167 \rangle + \equiv
        \mathbf{v\_bias} :: round\_up();
        double beta = inf\_normM(B);
        if (beta \ge 1) return false;
```

Finally, we can call encloseLS for each right side  $e_i$ . getColumn extracts the ith column of C in  $b\theta$ . setColumn sets the interval vector containing the corresponding solution to  $Ax = e_i$ .

#### 14.5 Enclosing the inverse of an orthogonal matrix

We have a floating-point approximation for an orthogonal matrix. Normally, its transpose is not the same as the inverse of this matrix. Hence, we need to enclose it.

```
170 \langle enclose the inverse of an orthogonal matrix 170 \rangle \equiv bool MatrixInverse :: orthogonalInverse (iMatrix &Ainv, const pMatrix &A)  \{ \\ transpose(C,A); \\ \langle \text{ find beta 167} \rangle \\ \langle \text{ enclose each column 169} \rangle \\ \text{return } true; \\ \}
```

This code is used in chunk 173.

#### 14.6 Constructor and destructor

```
171 \langle MatrixInverse constructor/destructor 171\rangle \equiv MatrixInverse :: MatrixInverse (int n) \{
M = \mathbf{new \ double}[n*n];
ipiv = \mathbf{new \ int}[n];
lwork = 2*n;
work = \mathbf{new \ double}[lwork];
size V (radx, n);
size V (b0, n);
size V (x1, n);
size V (x, n);
size M (B, n);
size M (Ci, n);
```

```
sizeM(C, n);
       MatrixInverse: \sim MatrixInverse()
         delete[] work;
         delete[] ipiv;
         \mathbf{delete}[]M;
    This code is used in chunk 173.
    Files
172 \langle matrixinverse.h 172 \rangle \equiv
    \#ifndef MATRIXINVERSE_H
    #define MATRIXINVERSE_H
    #include <numeric>
    #include "vnodeinterval.h"
    #include "vnoderound.h"
    #include "vector_matrix.h"
      namespace v_blas {
         using namespace v_bias;
         ⟨ class MatrixInverse 156 ⟩
    #endif
173 \langle matrixinverse.cc 173 \rangle \equiv
    #include <climits>
    #include "matrixinverse.h"
    #include "basiclinalg.h"
    #include "intvfuncs.h"
    #include "debug.h"
      using namespace std;
             namespace v_blas {
         ⟨ MatrixInverse constructor/destructor 171 ⟩
          \langle \text{ compute } A^{-1} | 158 \rangle
          (enclose the solution to Ax = b 160)
          (enclose the inverse of a matrix 166)
         (enclose the inverse of an orthogonal matrix 170)
```

# Part IV Solver Implementation

### **Structure**

We list the classes in VNODE-LP along with brief descriptions. These classes are depicted in Figure 15.1.

**Solution** contains a representation of the solution at each time point.

**Apriori** contains a representation of an enclosure of the solution over an integration step.

Control stores various data for controlling an integration.

**AD\_ODE** provides functions for generating Taylor coefficients for the solution to an ODE.

**AD\_VAR** provides functions for generating Taylor coefficients for the solution to the variational equation.

AD aggregates objects of AD\_ODE and AD\_VAR.

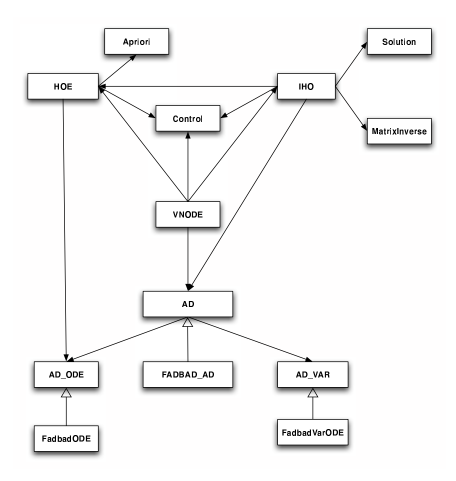
FadbadODE, FadbadVarODE, and FADBAD\_AD are implementations of AD\_ODE, AD\_VAR, and AD, respectively.

**HOE** implements the High-Order Enclosure method [27] for enclosing the solution to an ODE and computing a priori bounds.

**IHO** implements the Interval Hermite-Obreschkoff method [23] for computing tight bounds on the solution.

**VNODE** implements the overall integrator.

MatrixInverse provides functions for computing the inverse of a matrix.



**Figure 15.1.** Classes in VNODE-LP. The triangle arrows denote inheritance relations; the normal arrows denote uses relations.

# Solution enclosure representation

The present solver implements a one-step method. The initial point  $t_0$  and endpoint  $t_{\text{end}}$  are stored as intervals. For example, if  $t_0$  is representable, the interval is  $t_0 = [t_0, t_0]$ ; otherwise,  $t_0$  is an interval containing  $t_0$ . The interval containing  $t_{\text{end}}$  is denoted by  $t_{\text{end}}$ .

Internally, VNODE-LP select points that are machine-representable numbers. For simplicity in the considerations that follow, we denote such a point by the interval  $t_j$ , which contains  $t_j$ . At  $t_0$ , the user provides  $y_0$ . On each integration step, we maintain two types of representations of an enclosure of an ODE solution: tight enclosure at  $t_j$  and an a priori enclosure over  $[\underline{t}_j, \overline{t}_{j+1}]$  (or  $[\underline{t}_{j+1}, \overline{t}_j]$  if the integration is in negative direction).

#### 16.1 Tight enclosure

We maintain three representations as enclosures on the solution at  $t_i$ :

```
\begin{aligned} &\{u_j + S_j \alpha + A_j r \mid \alpha \in \boldsymbol{\alpha}, \ r \in r_j\} \\ &\{u_j + S_j \alpha + Q_j r \mid \alpha \in \boldsymbol{\alpha}, \ r \in r_{\text{QR},j}\}, \quad \text{and} \\ &\boldsymbol{y}_j. \end{aligned}
```

Here,  $u_j, \alpha, r \in \mathbb{R}^n$ ;  $S_j, A_j, Q_j \in \mathbb{R}^{n \times n}$ ; and  $\boldsymbol{r}_j, \boldsymbol{r}_{QR,j}, \boldsymbol{\alpha}, \boldsymbol{y}_j \in \mathbb{R}^n$ . The first set corresponds to the P (parallelepiped) method, and the second set corresponds to the QR method [24]. The use of these sets is discussed in detail in Chapter 20.

```
178 \langle \text{tight enclosure representation } 178 \rangle \equiv \\ \text{class Solution } \{\\ \text{public:} \\ \text{Solution(int } n); \\ \text{void } init(\text{const v\_bias::interval } \&t\theta, \text{const iVector } \&y\theta); \\ \text{v\_bias::interval } t; \\ \text{pVector } u;
```

```
iVector y;
iVector alpha, r, rQR;
pMatrix S, A, Q;
};
This code is used in chunk 184.
```

Solution::Solution(int n)

179  $\langle \text{ create Solution } 179 \rangle \equiv$ 

The constructor of the **Solution** class allocates memory for the vector and matrix members of this class.

```
size V(u, n);
      size V(alpha, n);
      size V(r, n);
      sizeV(rQR, n);
      size V(y, n);
      sizeM(S, n);
      sizeM(A, n);
      sizeM(Q, n);
This code is used in chunk 185.
   Initially, when j = 0, we set
           u_0 = \mathbf{m}(\boldsymbol{y}_0),
           \boldsymbol{\alpha} = \boldsymbol{y}_0 - u_0,
           \mathbf{r}_0 = 0,
       r_{\text{QR},0} = 0,
           S_0 = I,
          A_0 = I, and
          Q_0 = I.
```

When computing the midpoint of  $y\theta = y_0$ , we find a point vector u that is the rounded to the nearest true midpoint of  $y\theta$ .

```
180 \( \text{initialize Solution } 180 \) \( \sim \text{void Solution} :: init(\const \ \mathbf{v_bias} :: interval \ &t\theta, \const \ iVector \ &y\theta) \) \\ \{ \quad t = t\theta; \quad y = y\theta; \quad midpoint(u, y\theta); \quad /* \quad u \pi \midpoint(u, y\theta); \quad /* \quad u \pi \midpoint(u, y\theta); \quad /* \quad alpha \geq \mathbf{y}_0 - u \quad */ \quad set V(r, 0.0); \quad set V(rQR, 0.0); \quad set Id(S); \end{alpha} \)
```

```
 \begin{array}{c} setId(Q);\\ setId(A);\\ \end{array} \} This code is used in chunk 185.
```

#### 16.2 A priori enclosure

On each integration step, we validate existence and uniqueness of a solution and compute a priori bounds  $\tilde{y}_i$  such that

```
y(t;t_j,y_j) \in \widetilde{\boldsymbol{y}}_i
      for all t \in \widetilde{\boldsymbol{t}}_j := \boldsymbol{t}_j + [0,1]h_j and all y_j \in \boldsymbol{y}_j. Here, h_j \in \mathbb{R} is a stepsize selected by
      VNODE-LP.
             The interval \widetilde{t}_j and the interval vector \widetilde{\boldsymbol{y}}_j are stored in
181 \langle a priori enclosure representation 181 \rangle \equiv
         class Apriori {
         public:
            Apriori(int n);
            void init(const \ v\_bias::interval \ t\theta, const \ iVector \ y\theta);
            v_bias::interval t;
            iVector y;
      This code is used in chunk 184.
         The constructor allocates a vector y
182 \langle \text{ create Apriori } 182 \rangle \equiv
         Apriori::Apriori(int n) {
            size V(y, n);
      This code is used in chunk 185.
         When initializing an Apriori object, we set t and y.
183 (initialize Apriori 183) \equiv
         void Apriori::init(const \ v\_bias::interval \ t\theta, const \ iVector \ y\theta) {
            t = t\theta;
            y = y\theta;
      This code is used in chunk 185.
      Files
184 \langle solution.h 184\rangle \equiv
      #ifndef SOLUTION_H
```

```
\#define SOLUTION_H
       name space\ vnodelp\ \{
         using namespace v_bias;
         using namespace v_blas;
         \langle tight enclosure representation 178\rangle
         \langle a priori enclosure representation 181\rangle
    \#endif
185 \langle solution.cc 185\rangle
    \#include "vnodeinterval.h"
    #include "basiclinalg.h"
    #include "solution.h"
    #include "intvfuncs.h"
       name space\ vnodelp\ \{
         \langle \text{ create Solution } 179 \rangle
         (initialize Solution 180)
         ⟨ create Apriori 182⟩
         (initialize Apriori 183)
```

# Taylor coefficient computation

We provide an abstract class, **AD\_ODE**, for generating Taylor coefficients (TCs) for the solution to an ODE and an abstract class, **AD\_VAR**, for generating TCs for the solution to the ODE's variational equation. Concrete implementations of these classes using FADBAD++ [6] are discussed in Chapter 22.

#### 17.1 Taylor coefficients for an ODE solution

```
187 〈AD_ODE 187〉≡
    class AD_ODE {
    public:
        virtual void set(const v_bias::interval &t0, const iVector &y0,
            const v_bias::interval &h, int k) = 0;
        virtual void compTerms() = 0;
        virtual void sumTerms(iVector &sum, int m) = 0;
        virtual void getTerm(iVector &term, int i) const = 0;
        virtual v_bias::interval getStepsize() const = 0;
        virtual void eval(void *param) = 0;
        virtual ~AD_ODE() {}
    };
    This code is used in chunk 194.
```

Brief descriptions of the above functions follow. The kth TC of the solution to (1.1) at  $(t^*, y^*)$  is denoted by  $f^{[i]}(t^*, y^*)$ ; cf. [23].

set initializes a TC computation by setting a point of expansion  $t_0$ ,  $y_0$ , enclosure on the stepsize h, and order k.

compTerms encloses

$$hf^{[1]}(t_0, y_0), h^2f^{[2]}(t_0, y_0), \dots, h^kf^{[k]}(t_0, y_0).$$

sumTerms encloses  $\sum_{i=0}^{m} \mathbf{h}^{i} f^{[i]}(\mathbf{t}_{0}, \mathbf{y}_{0})$ , where  $m \leq k$ . The result is stored in the parameter sum.

getTerm obtains the ith term,  $\boldsymbol{h}^i f^{[i]}(t_0, \boldsymbol{y}_0)$ , where  $i \leq k$ .

getStepsize returns h.

eval evaluates f(t, y) and rebuilds the computational graph. param is a pointer to parameters that can be passed to f.

### 17.2 Taylor coefficients for the solution of the variational equation

Brief descriptions of the above functions follow.

set initializes a TC computation by setting a point of expansion  $t_0$ ,  $y_0$ , stepsize h, and order k.

compTerms encloses

This code is used in chunk 195.

$$h \frac{\partial f^{[1]}}{\partial y}(t_0, y_0), h^2 \frac{\partial f^{[2]}}{\partial y}(t_0, y_0), \dots, h^k \frac{\partial f^{[k]}}{\partial y}(t_0, y_0).$$

sumTerms encloses

$$\sum_{i=0}^{m} \boldsymbol{h}^{i} \frac{\partial f^{[i]}}{\partial y} (\boldsymbol{t}_{0}, \boldsymbol{y}_{0}),$$

where  $m \leq k$ . The result is stored in the parameter sum.

getTerm obtains the ith term,  $h^i \frac{\partial f^{[i]}}{\partial y}(t_0, y_0)$ , where  $i \leq k$ .

eval evaluates f(t, y) and rebuilds the computational graph. param is a pointer to parameters that can be passed to f.

17.3. AD class 111

#### 17.3 AD class

```
It is convenient to encapsulate the above classes into
191 \langle \text{encapsulated AD 191} \rangle \equiv
       class AD {
       public:
         AD(int n, AD\_ODE *a, AD\_VAR *av);
         void eval(\mathbf{void} *p);
         virtual int getMaxOrder() const = 0;
             /* maximum order that is allowed */
       public:
         int size;
         \mathbf{AD\_ODE} * tayl\_coeff\_ode;
         AD_VAR * tayl\_coeff\_var;
       };
    This code is used in chunk 196.
       The constructor of AD sets the size of the problem and pointers to objects of
     AD_ODE and AD_VAR.
192 \langle implementation of encapsulated AD 192 \rangle \equiv
       inline AD::AD(int n, AD_ODE *a, AD_VAR *av)
       : size(n), tayl\_coeff\_ode(a), tayl\_coeff\_var(av) { }
     See also chunk 193.
    This code is used in chunk 196.
       The eval function calls the corresponding eval functions of AD_ODE and AD_VAR
193 (implementation of encapsulated AD 192) +\equiv
       inline void AD :: eval(void *p)
         tayl\_coeff\_ode \rightarrow eval(p);
         tayl\_coeff\_var \rightarrow eval(p);
    Files
    The above classes are stored in ad_ode.h, ad_var.h, and allad.h, respectively.
194 \langle ad\_ode.h 194 \rangle \equiv
     #ifndef AD_ODE_H
     #define AD_ODE_H
     #include "vnodeinterval.h"
     #include "vector_matrix.h"
       namespace vnodelp {
         using namespace v_bias;
         using namespace v_blas;
```

```
\langle AD_ODE 187\rangle
     \#endif
195 \langle ad_var.h 195 \rangle \equiv
     \#ifndef AD_VAR_H
     \#\mathbf{define}\ \mathtt{AD\_VAR\_H}
     #include "vnodeinterval.h"
     #include "vector_matrix.h"
       namespace vnodelp {
          using namespace v_bias;
          using namespace v_blas;
          \langle AD_VAR 189\rangle
     \#endif
196 \langle allad.h 196\rangle \equiv
     \#\mathbf{ifndef}\ \mathtt{ALLAD}_\mathtt{H}
     \#define ALLAD_H
     \#include "ad_ode.h"
     #include "ad_var.h"
       name space\ vnodelp\ \{
          (encapsulated AD 191)
          \langle implementation of encapsulated AD 192\rangle
     \# endif
```

### **Control data**

We store various data needed to control an integration in a **Control** class. An integration in VNODE-LP is carried out by the *integrate* function of **VNODE**; see Chapter 21.

#### 18.1 Indicator type

First, we introduce an enumerated  $\mathbf{Ind}$  data type, where a variable of this type can take the following values:

value	description		
first_entry	indicates a first entry into integrate		
success	$integrate$ has reached $t_{\mathrm{end}}$ successfully		
failure	an error has occurred in $integrate$		

```
198 ⟨indicator type 198⟩ ≡
          typedef enum {
          first_entry, success, failure
        } Ind;
```

This code is used in chunk 201.

#### 18.2 Interrupt type

Similarly, we have an  ${\bf Interrupt}$  type, where a variable can take values as described below.

value	description		
no	$integrate$ tries to reach $t_{ m end}$		
$before\_accept$	integrate takes a step and returns before accepting this step		

```
199 ⟨interrupt type 199⟩ ≡

typedef enum {

no, before_accept
} Interrupt;

This code is used in chunk 201.
```

#### 18.3 Control data

In the **Control** class, we store the following data:

name	default value	description
ind	$\mathit{first\_entry}$	indicator variable
interrupt	no	indicates if interrupts are requested
order	20	order of the method
atol	$10^{-12}$	absolute error tolerance
rtol	$10^{-12}$	relative error tolerance
hmin	0	magnitude of the minimum stepsize allowed. If a positive value is set by the user, this value for <i>hmin</i> will be used in an integration. Otherwise, the solver computes a minimum stepsize as discussed in Subsection 21.2.3.

```
200 \langle \operatorname{control class}\ 200 \rangle \equiv  class Control {
    public:
        Ind ind;
        Interrupt interrupt;
        unsigned int order;
        double atol,\ rtol;
        double hmin;
        Control():
            ind\ (first\_entry),
            interrupt\ (no),
            order\ (20),
            atol\ (1\cdot 10^{-12}),\ rtol\ (1\cdot 10^{-12}),
        hmin\ (0)\ \{\}
};
```

This code is used in chunk 201.

#### Files

We store  $\mathbf{Ind}$ ,  $\mathbf{Interrupt}$ , and  $\mathbf{Control}$  in

18.3. Control data

```
201 \( \) control.h \( 201 \) \( \) \\ #ifndef CONTROL_H \\
#define CONTROL_H \\
namespace vnodelp \{ \( \) \( \) indicator type 198 \\ \( \) \( \) \( \) control class 200 \( \) \\ \\ \\ \\\
#endif
```

## Computing a priori bounds

We have implemented the HOE method [27] to compute a priori bounds on the solution of an ODE problem. In Section 19.1 we summarize the relevant theory. The **HOE** class is given in Section 19.2. The implementation of the HOE method is in Section 19.3. The rest of the functions of this class is in Section 19.4.

#### 19.1 Theory background

The HOE method is based on the following two results; cf. [9, 27].

1. If  $h_j$  and  $\widetilde{\boldsymbol{y}}_j$  are such that  $\boldsymbol{y}_j \subseteq \operatorname{int}(\widetilde{\boldsymbol{y}}_j)$  and

$$\sum_{i=0}^{k-1} (t-t_j)^i f^{[i]}(t_j, y_j) + (t-t_j)^k f^{[k]}(t_j + [0, 1]h_j, \widetilde{\boldsymbol{y}}_j) \subseteq \widetilde{\boldsymbol{y}}_j$$

for all  $t \in t_j + [0,1]h_j$  and all  $y_j \in \boldsymbol{y}_j$ , then

$$y' = f(t, y), \quad y(t_j) = y_j \in \mathbf{y}_j \tag{19.1}$$

has a unique solution

$$y(t;t_j,y_j) \in \widetilde{\boldsymbol{y}}_i$$

for all  $t \in t_j + [0,1]h_j$  and all  $y_j \in \mathbf{y}_j$ .

2. Let  $h_{j,0} \neq 0$  and let  $\boldsymbol{p}_j$  be an interval vector enclosing the set

$$\mathcal{P}_{j} = \left\{ \sum_{i=0}^{k-1} (t - t_{j})^{i} f^{[i]}(t_{j}, y_{j}) \mid t \in t_{j} + [0, 1] h_{j,0}, \ y_{j} \in \mathbf{y}_{j} \right\}.$$
 (19.2)

That is,  $\mathcal{P}_j \subseteq \boldsymbol{p}_i$ .

Let  $u_j$  be such that

$$\widetilde{\pmb{y}}_j = \pmb{p}_j + \pmb{u}_j,$$
 and 
$$\pmb{y}_j \subseteq \text{int}(\widetilde{\pmb{y}}_j).$$
 (19.3)

If  $h_{j,1} \neq 0$  is such that

$$[0,1]h_{i,1}^k f^{[k]}(t_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j) \subseteq \boldsymbol{u}_j, \tag{19.4}$$

and

$$h_j = \operatorname{sign}(h_{j,0}) \cdot \min\{|h_{j,0}|, |h_{j,1}|\},$$
 (19.5)

then there exists a unique solution  $y(t; t_j, y_j)$  to (19.1) for all  $t \in t_j + [0, 1]h_j$  and all  $y_j \in y_j$ . Moreover,

$$y(t; t_j, y_j) \in \widetilde{\boldsymbol{y}}_j$$
 for all  $t \in t_j + [0, 1]h_j$  and all  $y_j \in \boldsymbol{y}_j$ .

#### 19.2 The HOE class

The class implementing the HOE method is

```
204 \langle \text{ class HOE } 204 \rangle \equiv
       class HOE {
       public:
          HOE(int n);
          void compAprioriEnclosure (const interval &t0, const iVector &y0,
               bool & info);
          void acceptSolution();
          \langle set functions HOE 221 \rangle
          \langle \text{ get functions HOE } 222 \rangle
          \simHOE();
       private:
          Apriori *apriori_trial, *apriori;
          Control *control;
          AD\_ODE *tayl\_coeff;
          double h, h_next, h_trial, t_trial;
          int order_trial;
          iVector term, p, u, v;
          const interval one;
                                     /* one = [0,1] */
          interval comp\_beta(const iVector \&v, const iVector \&u, int k);
       };
```

This code is used in chunk 226.

#### 19.3 Implementation of the HOE method

#### 19.3.1 Computing $p_j$

We have to enclose  $\mathcal{P}_j$  in (19.2). On the first step, if  $t_0$  is not a representable machine number, an interval  $t_0$  containing  $t_0$  would be given. We assume that, in general,  $t_i \in t_j$ . Then

$$t \in \mathbf{t}_j + [0, 1]h_{j,0}$$
 and  $t - t_j \in \mathbf{t}_j - \mathbf{t}_j + [0, 1]h_{j,0}$ .

The width of an interval  $\boldsymbol{a}$  is  $w(\boldsymbol{a}) = \overline{\boldsymbol{a}} - \underline{\boldsymbol{a}}$ . (Width of an interval vector is defined componentwise.) Denote  $\boldsymbol{t}_j^* = (\boldsymbol{t_j} - \boldsymbol{t_j})/h_{j,0} + [0,1]$ . Then

$$t_j^* = \begin{cases} (t_j - t_j)/h_{j,0} + [0,1] & \text{if } w(t_j) > 0, \\ [0,1] & \text{if } w(t_j) = 0. \end{cases}$$

Hence  $t - t_j \in \boldsymbol{t}_j^* \, h_{j,0}$ , and we can bound  $\mathcal{P}_j$  as

$$\mathcal{P}_j \subseteq \sum_{i=0}^{k-1} oldsymbol{t}_j^{*i} \, h_{j,0}^i f^{[i]}(oldsymbol{t}_j, oldsymbol{y}_j).$$

We enclose first  $h_{j,0}^i f^{[i]}(\boldsymbol{t}_j, \boldsymbol{y}_j)$  for  $i = 1, \dots, k-1$ . Then, we use an interval form of Horner's rule to compute

$$p_{j} := \boldsymbol{y}_{j} + \boldsymbol{t}_{j}^{*} \Big( h_{j,0} f^{[1]}(\boldsymbol{t}_{j}, \boldsymbol{y}_{j}) + \cdots + \boldsymbol{t}_{j}^{*} \Big( h_{j,0}^{k-2} f^{[k-2]}(\boldsymbol{t}_{j}, \boldsymbol{y}_{j}) + \boldsymbol{t}_{j}^{*} h_{j,0}^{k-1} f^{[k-1]}(\boldsymbol{t}_{j}, \boldsymbol{y}_{j}) \Big) \cdots \Big).$$
206  $\langle \text{compute } \boldsymbol{p}_{j} | 206 \rangle \equiv /*$ 

$$\boldsymbol{t} \boldsymbol{-} 0 = \boldsymbol{t}_{j}$$

$$\boldsymbol{y} 0 \supseteq \boldsymbol{y}_{j}$$

$$h \boldsymbol{-} trial = h_{j,0}$$

$$order \boldsymbol{-} trial = k$$

$$tayl \boldsymbol{-} coeff \text{ contains enclosures on } h_{j,0}^{i} f^{[i]}(\boldsymbol{t}_{j}, \boldsymbol{y}_{j}) \text{ for } i = 0, \dots, k-1$$

```
tayl_coeff contains enclosures on h_{j,0}^*f^{(i)}(m{t}_j,m{y}_j) for i=0,\dots,k-1 t\_enc\supseteq m{t}_j^*=(m{t}_j-m{t}_j)/h_{j,0}+[0,1] p\supseteq m{p}_j */
```

```
tayl\_coeff \neg set(t0, y0, h\_trial, order\_trial - 1); \\ tayl\_coeff \neg compTerms(); \\ \textbf{interval} \ t\_enc = (t0 - t0)/h\_trial + one; \\ tayl\_coeff \neg getTerm(p, order\_trial - 1); \\ \textbf{for (int } i = order\_trial - 2; \ i \geq 0; \ i--) \ \{ \\ scaleV(p, t\_enc); \\ tayl\_coeff \neg getTerm(term, i); \\ addViVi(p, term); \\ \}
```

This code is used in chunk 217.

#### 19.3.2 Computing $u_j$ and $\tilde{y}_j$

If  $u_j$  is symmetric with no component equal to [0,0] then (19.3) holds. Denote

$$tol_j = rtol \cdot ||\boldsymbol{y}_j|| + atol.$$

Throughout this manuscript, we shall assume the infinity norm, unless stated otherwise. For an interval vector a,

$$\|\boldsymbol{a}\| = \max_{i} \{ |\underline{\boldsymbol{a}}_{i}|, |\overline{\boldsymbol{a}}_{i}| \}.$$

Let  $u_j$  be the *n*-vector with each component  $h_{j,0}[-\operatorname{tol}_j/2,\operatorname{tol}_j/2]$ . Then, we form

$$\widetilde{\boldsymbol{y}}_i = \boldsymbol{p}_i + \boldsymbol{u}_j.$$

$$\begin{array}{c} y0 \supseteq \boldsymbol{y}_{j} \\ control \neg atol = \mathrm{atol} \\ control \neg tol = \mathrm{rtol} \\ h \_trial = h_{j,0} \\ p \supseteq \boldsymbol{p}_{j} \\ \hline tol \ge \mathrm{rtol} \cdot \|\boldsymbol{y}_{j}\| + \mathrm{atol} \\ u \supseteq \boldsymbol{u}_{j} \\ apriori\_trial \supseteq \widetilde{\boldsymbol{y}}_{j} = \boldsymbol{p}_{j} + \boldsymbol{u}_{j}. \\ \\ \boldsymbol{\psi}_{j} = \boldsymbol{v}_{j} \\ control \neg trial \supseteq \boldsymbol{v}_{j} \\ control \neg trial \supseteq \boldsymbol{v}_{j} \\ control \neg trial \supseteq \boldsymbol{v}_{j} \\ control \neg trial \vdash tontrol \neg tol + control \neg tol; \\ control \neg tol + control \neg tol + control \neg tol; \\ control \neg tol + control \neg tol + control \neg tol; \\ control \neg tol + control \neg tol + control \neg tol; \\ control \neg tol + control \neg tol + control \neg tol + control \neg tol; \\ control \neg tol + control \neg tol$$

#### Motivation for the choice of $u_j$

Denote

$$z_j = f^{[k]}(t_j + [0, 1]h_{j,0}, \widetilde{\boldsymbol{y}}_j).$$

We can consider  $|h_{j,1}|^k \|\mathbf{w}(\mathbf{z}_j)\|$  as an estimate of the *local excess* at  $\mathbf{t}_{j+1}$  that we introduce on the step from  $\mathbf{t}_j$  to  $\mathbf{t}_{j+1}$ .

The magnitude of an interval a is  $|a| = \max\{|\underline{a}|, |\overline{a}|\}$ . Magnitude of an interval vector is defined componentwise. From (19.4), (19.5), and the choice for  $u_j$ ,

$$|h_{j,1}|^k \operatorname{w}(\boldsymbol{z}_j) \leq \operatorname{w}([0,1]h_{j,1}^k \boldsymbol{z}_j) = |h_{j,1}|^k |\boldsymbol{z}_j|$$
  
$$\leq \operatorname{w}(\boldsymbol{u}_j) = h_{j,0} (\operatorname{tol}_j, \operatorname{tol}_j, \cdots, \operatorname{tol}_j)^T.$$

Hence.

$$|h_{j,1}|^k \|\mathbf{w}(\boldsymbol{z}_j)\| \le h_{j,0} \cdot \text{tol}_j.$$

If  $h_{j,1} = h_{j,0}$ , we have a local excess per unit step (LEPUS) control. If  $h_{j,1} < h_{j,0}$ , then usually  $h_{j,1}$  is not much smaller than  $h_{j,0}$ , and we have almost LEPUS.

#### 19.3.3 Computing a stepsize

We enclose first

$$v_j = h_{j,0}^k f^{[k]}(t_j + [0,1]h_{j,0}, \widetilde{y}_j).$$

210 
$$\langle$$
 compute stepsize 210  $\rangle$   $\equiv$  /\*

$$t\theta = t_{j}$$
 $apriori\_trial \supseteq \widetilde{\boldsymbol{y}}_{j}$ 
 $h\_trial = h_{j,0}$ 

 $order\_trial = k$ 

tayl\_coeff contains enclosures on 
$$h_{j,0}^i f^{[i]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)$$
 for  $i = 0, \ldots, k$ 

 $v \supseteq h_{j,0}^k f^{[k]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)$ 

 $*/\\tayl\_coeff \neg set(t0 + one * h\_trial, apriori\_trial \neg y, h\_trial, order\_trial);$ 

 $tayl\_coeff \neg comp Terms();$  $tayl\_coeff \neg get Term(v, order\_trial);$ 

See also chunks 212 and 214.

This code is used in chunk 217.

Now we consider two cases:  $\underline{t}_j = \overline{t}_j$  and  $\underline{t}_j < \overline{t}_j$ .

#### The case $\underline{t}_j = \overline{t}_j$

Let  $\gamma > 0$  be the largest number such that

$$[0,1]h_{j,1}^{k}f^{[k]}(\boldsymbol{t}_{j}+[0,1]h_{j,0},\widetilde{\boldsymbol{y}}_{j}) = [0,1](\gamma h_{j,0}^{k})f^{[k]}(\boldsymbol{t}_{j}+[0,1]h_{j,0},\widetilde{\boldsymbol{y}}_{j})$$

$$= [0,\gamma]\boldsymbol{v}_{j}$$

$$\subseteq \boldsymbol{u}_{j}.$$

Such a  $\gamma$  exists since  $u_j$  is symmetric. Then, we write  $\beta = \gamma^{1/k}$  and set

$$h_j = \begin{cases} \beta h_{j,0} & \text{if } \beta < 1\\ h_{j,0} & \text{if } \beta \ge 1. \end{cases}$$

In practice, we compute  $\beta \ni \beta > 0$ , with  $\underline{\beta} \ge 0$  and find

$$h_{j} = \begin{cases} \downarrow(\underline{\beta} h_{j,0}) & \text{if } \underline{\beta} < 1, \ h_{j,0} > 0 \\ \uparrow(\underline{\beta} h_{j,0}) & \text{if } \underline{\beta} < 1, \ h_{j,0} < 0 \\ h_{j,0} & \text{if } \underline{\beta} \ge 1. \end{cases}$$

$$(19.6)$$

Here  $\uparrow$  denotes rounding up, and  $\downarrow$  denotes rounding down.

212  $\langle \text{compute stepsize 210} \rangle + \equiv /*$ 

$$\frac{u \supseteq \mathbf{u}_{j}}{v \supseteq \mathbf{v}_{j} = h_{j,0}^{k} f^{[k]}(\mathbf{t}_{j} + [0,1]h_{j,0}, \widetilde{\mathbf{y}}_{j})}{h = h_{j}}$$

\*/
interval  $beta = comp\_beta(v, u, order\_trial);$   $assert(inf(beta) \ge 0);$ if (inf(beta) < 1) {
 if  $(h\_trial > 0)$   $round\_down();$  else  $round\_up();$   $h = inf(beta) * h\_trial;$ }
else  $h = h\_trial;$ 

#### The case $\underline{t}_i < \overline{t}_j$

Similar to (19.4), we try to find (the largest)  $|h_{j,1}| \leq |h_{j,0}|$  such that

$$(t - t_j)^k f^{[k]}(t_j + [0, 1]h_{j,0} \subseteq (\boldsymbol{t}_j - \boldsymbol{t}_j + [0, 1]h_{j,1})^k f^{[k]}(\boldsymbol{t}_j + [0, 1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)$$

$$\subseteq \boldsymbol{u}_j.$$
(19.7)

Normally  $|h_{j,0}| \gg |t_j - t_j|$ , and therefore,

$$(\boldsymbol{t}_{j} - \boldsymbol{t}_{j} + [0, 1]h_{j,0})^{k} \approx [0, 1]h_{j,0}^{k}.$$

Hence, we may consider setting  $h_j$  as in (19.6). However, (19.7) may not hold. To find  $h_j$  such that it is likely to hold, we set

$$h_i = 0.9 h_{i,0}$$

and check if

$$\begin{split} \left( (\boldsymbol{t}_{j} - \boldsymbol{t}_{j}) + [0, 1]h_{j} \right)^{k} f^{[k]} (\boldsymbol{t}_{j} + [0, 1]h_{j,0}, \widetilde{\boldsymbol{y}}_{j}) \\ &= \left( \frac{\boldsymbol{t}_{j} - \boldsymbol{t}_{j} + [0, 1]h_{j}}{h_{j,0}} \right)^{k} h_{j,0}^{k} f^{[k]} (\boldsymbol{t}_{j} + [0, 1]h_{j,0}, \widetilde{\boldsymbol{y}}_{j}) \\ &= \left( \frac{\boldsymbol{t}_{j} - \boldsymbol{t}_{j} + [0, 1]h_{j}}{h_{j,0}} \right)^{k} \boldsymbol{v}_{j} \subseteq \boldsymbol{u}_{j}. \end{split}$$

If the above inclusion test fails, we reduce  $h_j$  and repeat as shown below.

```
214 \langle compute stepsize 210 \rangle + \equiv
            if (inf(t\theta) < sup(t\theta)) {
                                                     v \supseteq \boldsymbol{v}_j
                                                    h = h_i
                                            h_trial = h_{i,0}
                                          t\theta = t_i
                               control \rightarrow hmin = h_{min}
                                                    tt \supseteq \boldsymbol{t}_i - \boldsymbol{t}_i
                                              t\_enc \supseteq rac{oldsymbol{t}_j - oldsymbol{t}_j + [0,1]h_j}{h_{j,0}}
                                                   v \supseteq \left(\frac{\boldsymbol{t}_j - \boldsymbol{t}_j + [0, 1]h_j}{h_{j,0}}\right)^k h_{j,0}^k f^{[k]}(\boldsymbol{t}_j + [0, 1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)
                 interval tt = t\theta - t\theta;
                 while (fabs(h) > control \rightarrow hmin) {
                     t\_enc = (tt + one * h)/h\_trial;
                     scale V(v, pow(t\_enc, order\_trial));
                     if (subseteq(v, u)) break;
                     h = 0.9 * h;
            }
```

#### 19.3.4 Forming the time interval

Once  $h_j$  is found, we need to determine the next representable integration point and the machine interval over which the a priori bounds hold.

1. If  $h_j > 0$ , we compute  $t_{j+1} = \downarrow (\underline{t}_j + h_j) \quad \text{and} \quad T_j = [\underline{t}_j, t_{j+1}].$ 

2. If  $h_i < 0$ , we compute

$$t_{j+1} = \uparrow (\overline{t}_j + h_j)$$
 and  $T_j = [t_{j+1}, \overline{t}_j].$ 

#### Proposition 19.1.

(i) 
$$T_j \subseteq t_j + [0, 1]h_{j,0}$$

(ii) If 
$$|h_j| \geq w(t_j)$$
, then  $t_j \subseteq T_j$ 

**Proof.**  $h_{j,0} > 0$ . Then  $0 < h_j \le h_{j,0}$  and

$$T_j = [\underline{\boldsymbol{t}}_j, t_{j+1}] \subseteq [\underline{\boldsymbol{t}}_j, \underline{\boldsymbol{t}}_j + h_j] \subseteq [\underline{\boldsymbol{t}}_j, \underline{\boldsymbol{t}}_j + h_{j,0}] = \underline{\boldsymbol{t}}_j + [0, 1]h_{j,0}$$
  
$$\subseteq \boldsymbol{t}_j + [0, 1]h_{j,0}.$$

If 
$$h_j \ge w(t_j) = \overline{t}_j - \underline{t}_j$$
, then  $\overline{t}_j \le \downarrow (\underline{t}_j + h_j) \le \underline{t}_j + h_j$ , and

$$[\underline{\boldsymbol{t}}_j, \overline{\boldsymbol{t}}_j] \subseteq [\underline{\boldsymbol{t}}_j, \downarrow (\underline{\boldsymbol{t}}_j + h_j)] = \boldsymbol{T}_j.$$

$$h_{j,0} < 0$$
. Then  $0 > h_j \ge h_{j,0}$  and

$$\begin{aligned} \boldsymbol{T}_j &= [t_{j+1}, \overline{\boldsymbol{t}}_j] \subseteq [\overline{\boldsymbol{t}}_j + h_j, \overline{\boldsymbol{t}}_j] \subseteq [\overline{\boldsymbol{t}}_j + h_{j,0}, \overline{\boldsymbol{t}}_j] = \overline{\boldsymbol{t}}_j + [0, 1] h_{j,0} \\ &\subseteq \boldsymbol{t}_j + [0, 1] h_{j,0}. \end{aligned}$$

If 
$$-h_j \geq \overline{t}_j - \underline{t}_j$$
, then  $\underline{t}_j \geq \uparrow (\overline{t}_j + h_j) \geq \overline{t}_j + h_j$ , and

$$t_j = [\underline{t}_j, \overline{t}_j] \subseteq [\uparrow (\underline{t}_j + h_j), \overline{t}_j] = T_j.$$

215  $\langle$  form time interval 215  $\rangle \equiv /*$ 

$$h = h_j$$
$$t\theta = \mathbf{t}_j$$

$$t$$
\_trial =  $t_{j+1}$   
 $apriori$ \_trial $\rightarrow t = T_i$ 

double td; if (h > 0) {  $td = inf(t\theta)$ ;  $round\_down()$ ;  $t\_trial = td + h$ ;

```
apriori\_trial\_t = \mathbf{interval}(td, t\_trial); } else { td = sup(t\theta); round\_up(); t\_trial = td + h; apriori\_trial\_t = \mathbf{interval}(t\_trial, td); } assert(subseteq(t\theta, apriori\_trial\_t));
```

This code is used in chunk 217.

#### 19.3.5 Selecting a trial stepsize for the next step

We can select for the next step  $h_{j+1,0} = \beta h_{j,0}$ . This is reasonable since, if  $|h_{j+1,0}| > |h_{j,0}|$ , we have computed an a priori enclosure with  $h_{j,0}$ , but we could have possibly done this with a stepsize between  $h_{j,0}$  and  $h_{j+1,0}$ . That is, we assume that we might be successful on the next step with  $|h_{j+1,0}| = \beta |h_{j,0}| > |h_{j,0}|$ . Similar considerations apply when  $|h_{j+1,0}| < |h_{j,0}|$ . Hence, we select

$$h_{i+1,0} = \beta h_{i,0}$$

for the next step.

216  $\langle$  select stepsize 216  $\rangle \equiv /*$ 

$$h\_trial = h_{j,0}$$

$$beta \supseteq \beta$$

$$h\_next = h_{j+1,0}$$

$$*/$$
 $h\_next = inf(beta) * h\_trial;$ 

This code is used in chunk 217.

#### 19.3.6 Computing a priori bounds

First, we check if the stepsize is not very small. Then, we compute  $p_j$ ,  $\tilde{y}_j$ , and  $h_j$ . If  $|h_j| \leq h_{\min}$ , we cannot validate existence and uniqueness and return false. Otherwise, we form  $T_j$  and select  $h_{j+1,0}$  for the next step.

217 \( \text{validate existence and uniqueness } 217 \rangle \) = \( \text{void HOE: } \cdot \const \) const into

void HOE::compAprioriEnclosure (const interval & $t\theta$ , const iVector & $y\theta$ , bool &info)

```
{
              /*
                                            t\theta = t_j
                                            y\theta \supseteq \boldsymbol{y}_i
                                     h_{trial} = h_{i,0}
                             order\_trial = k
                       apriori\_trial \rightarrow t \supseteq T_i
                      apriori\_trial 
eg y \supseteq \widetilde{oldsymbol{y}}_j
                               tayl_coeff contains enclosures on h_{j,0}^i f^{[i]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j) for i = 0, \dots, k
                                    h_trial = h_{i+1,0}
        if (fabs(h\_trial) \leq control \neg hmin) {
            info = false;
            return;
         \langle \text{ compute } \boldsymbol{p}_j | 206 \rangle;
\langle \text{ compute } \boldsymbol{u}_j | \text{ and } \widetilde{\boldsymbol{y}}_j | 207 \rangle;
        \langle \text{ compute stepsize } 210 \rangle;
        if (fabs(h) \leq control \rightarrow hmin) {
            info = false;
            return;
        (form time interval 215);
         ⟨ select stepsize 216 ⟩;
        info = true;
This code is used in chunk 227.
```

#### 19.4 Other functions

#### 19.4.1 Constructor and destructor

```
219 \langle \text{constructor-destructor HOE } 219 \rangle \equiv \\ \text{HOE}::\text{HOE}(\text{int } n) \\ : one(\text{interval}(0,1)) \{\\ sizeV(term,n);\\ sizeV(p,n);\\ sizeV(u,n);\\ sizeV(v,n);\\ apriori\_trial = \text{new Apriori}(n);\\ apriori = \text{new Apriori}(n);\\ assert(apriori \land apriori\_trial);\\ tayl\_coeff = 0;
```

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```
control = 0;
         HOE :: \sim HOE()
            delete apriori;
            delete apriori_trial;
      This code is used in chunk 227.
                  Accept a solution
      To accept a trial enclosure, we store it in the *apriori object.
220 \langle accept solution (HOE) 220 \rangle \equiv
         void HOE::acceptSolution() {
                       apriori_trial contains \widetilde{\boldsymbol{y}}_j and \boldsymbol{T}_j
                              apriori contains \widetilde{\boldsymbol{y}}_{j-1} and \boldsymbol{T}_{j-1}
                              h_trial = h_{i,0}
                              h-next = h_{j+1,0}
                             apriori contains \widetilde{\boldsymbol{y}}_i and \boldsymbol{T}_j
                              h_trial = h_{j+1,0}
                   */
            apriori \rightarrow t = apriori \_trial \rightarrow t;
            assign V (apriori \rightarrow y, apriori trial \rightarrow y);
            h\_trial = h\_next;
      This code is used in chunk 227.
      19.4.3 Set functions
221 \langle set functions HOE 221 \rangle \equiv
         void set(Control *ctrl, AD *ad)
            control = ctrl;
            tayl\_coeff = ad \neg tayl\_coeff\_ode;
         void init(const\ interval\ \&t\theta, const\ iVector\ \&y\theta) {
            apriori \rightarrow init(t0, y0);
         void setTrialStepsize(double h\theta) {
            h\_trial = h\theta;
```

```
void setTrialOrder(int order0) {
           order\_trial = order\theta;
     This code is used in chunk 204.
     19.4.4 Get functions
222 \langle \text{ get functions HOE } 222 \rangle \equiv
        double getStepsize() const {
           return h;
        double getTrialStepsize() const {
           return h_trial;
        const interval \&getT() const {
           return apriori→t;
        interval getTrialT() const {
           return apriori_trial→t;
        const iVector &getApriori() const {
           return apriori \rightarrow y;
        const iVector & getTrialApriori() const {
           return apriori_trial→y;
     See also chunk 223.
     This code is used in chunk 204.
     Obtain error term
     We obtain h_{i,0}^i f^{[i]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_i) for given i=0,1,\ldots,k by
223 \langle \text{ get functions HOE } 222 \rangle + \equiv
                    e \supseteq h_{j,0}^i f^{[i]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)
        void getErrorTerm(iVector &e, int i) const {
           tayl\_coeff \neg getTerm(e, i);
     19.4.5 Enclosing \beta
```

The function  $comp\_beta$  returns an enclosure of  $\beta$ .

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```
224 \langle \text{ compute } \beta \text{ 224} \rangle \equiv
                  gamma = \gamma, the largest representable \gamma > 0 such that \gamma v_j \subseteq u_j
                      beta \supseteq \beta = \gamma^{1/k}
             */
       interval HOE::comp\_beta(const iVector \&v, const iVector \&u, int k)
          double gamma = \mathbf{v\_blas} :: compH(v, u);
          interval i-gamma = gamma;
          interval i_pw = interval(1.0)/interval(double(k));
          interval beta = pow(i\_gamma, i\_pw);
          return beta;
     This code is used in chunk 227.
     Files
226 \langle hoe.h 226 \rangle \equiv
     \#ifndef HOE_H
     #define HOE_H
       namespace vnodelp {
          ⟨ class HOE 204 ⟩
     #endif
227 \langle hoe.cc 227 \rangle \equiv
     #include <cmath>
     #include <cassert>
     #include <algorithm>
     #include "vnodeinterval.h"
     #include "vnoderound.h"
     #include "basiclinalg.h"
     #include "intvfuncs.h"
     #include "control.h"
     #include "solution.h"
     #include "allad.h"
     #include "hoe.h"
       using namespace v_blas;
       namespace vnodelp {
          (constructor-destructor HOE 219)
          (validate existence and uniqueness 217)
          (accept solution (HOE) 220)
          \langle \text{ compute } \beta \text{ 224} \rangle
       }
```

### Computing tight bounds on the solution

To compute tight bounds on the solution, we employ the interval Hermite-Obreschkoff (IHO) method developed in [23]. It consists of two phases: a predictor and a corrector. We present the relevant theory first and then describe its implementation.

#### 20.1 Theory background

#### 20.1.1 Predictor

For  $y_j \in \boldsymbol{y}_j \subseteq \widetilde{\boldsymbol{y}}_j$ , we have

$$y(t_{j+1}; t_j, y_j) \in y_j + \sum_{i=1}^q h_j^i f^{[i]}(t_j, y_j) + h_j^{q+1} f^{[q+1]}(\boldsymbol{T}_j, \widetilde{\boldsymbol{y}}_j), \tag{20.1}$$

where  $h_j = t_{j+1} - t_j$  and  $t_j$ ,  $t_{j+1} \in \boldsymbol{T}_j$ . Let  $J(f^{[i]}; \boldsymbol{y}_j)$  be the Jacobian of  $f^{[i]}$  evaluated at  $\boldsymbol{y}_j$  and denote

$$U_{j+1} = I + \sum_{i=1}^{q} h_j^i J(f^{[i]}; y_j).$$

These Jacobians are computed by generating TCs for the solution of the associated variational equation

$$Y' = \frac{\partial f}{\partial y}Y, \quad Y(t_j) = I,$$

where I is the  $n \times n$  identity matrix.

We assume that at  $t_j$  the solution  $y(t_j; t_0, y_0)$  is contained in

$$egin{aligned} m{y}_j \ &\{u_j + S_j lpha + A_j r \mid lpha \in m{lpha}, \ r \in m{r}_j\}, \quad ext{and} \ &\{u_j + S_j lpha + Q_j r \mid lpha \in m{lpha}, \ r \in m{r}_{\mathrm{QR},j}\}, \end{aligned}$$

where  $u_j \in \boldsymbol{y}_i$ .

If we apply the mean-value theorem to the  $f^{[i]}$  in (20.1), we obtain that for any

$$y_{j} \in \{u_{j} + S_{j}\alpha + A_{j}r \mid \alpha \in \boldsymbol{\alpha}, r \in \boldsymbol{r}_{j}\}$$

$$\cap \{u_{j} + S_{j}\alpha + Q_{j}r \mid \alpha \in \boldsymbol{\alpha}, r \in \boldsymbol{r}_{QR,j}\}$$

$$\cap \boldsymbol{y}_{j},$$

$$y(t_{j+1}; t_{j}, y_{j}) \in u_{j} + \sum_{i=1}^{q} h_{j}^{i} f^{[i]}(t_{j}, u_{j}) + h_{j}^{q+1} f^{[q+1]}(\boldsymbol{T}_{j}, \widetilde{\boldsymbol{y}}_{j})$$

$$+ (\boldsymbol{U}_{j+1} S_{j}) \boldsymbol{\alpha} + \{(\boldsymbol{U}_{j+1} A_{j}) \boldsymbol{r}_{j} \cap (\boldsymbol{U}_{j+1} Q_{j}) \boldsymbol{r}_{QR, j} \}.$$

$$(20.2)$$

For brevity, denote

$$\begin{split} \widehat{u}_{j+1} &= u_j + \sum_{i=1}^q h_j^i f^{[i]}(t_j, u_j) \\ \boldsymbol{z}_{j+1} &= h_j^{q+1} f^{[q+1]}(\boldsymbol{T}_j, \widetilde{\boldsymbol{y}}_j), \quad \text{and} \\ \boldsymbol{x}_{j+1} &= (\boldsymbol{U}_{j+1} S_j) \boldsymbol{\alpha} + \left\{ (\boldsymbol{U}_{j+1} A_j) \boldsymbol{r}_j \cap (\boldsymbol{U}_{j+1} Q_j) \boldsymbol{r}_{\text{QR}, j} \right\}. \end{split}$$

Then

$$y(t_{j+1};t_0,y_0) \in \boldsymbol{y}_{j+1}^* := (\widehat{u}_{j+1} + \boldsymbol{z}_{j+1} + \boldsymbol{x}_{j+1}) \cap \widetilde{\boldsymbol{y}}_j.$$

#### 20.1.2 Corrector

In the corrector, we compute  $y_{j+1} \subseteq y_{j+1}^*$ . Usually,  $y_{j+1}$  is much tighter than  $y_{j+1}^*$ .

Let

$$y_j = y(t_j; t_0, y_0)$$
 and  $y_{j+1} = y(t_{j+1}; t_0, y_0)$ .

Denote  $k = p + q + 1 \ (p, q \ge 0)$  and

$$c_i^{q,p} = \frac{q! (q+p-i)!}{(p+q)! (q-i)!} \qquad (q,p, \text{ and } i \ge 0).$$
 (20.3)

Denote also

$$\gamma_{p,q} = \frac{q!p!}{(p+q)!} \tag{20.4}$$

Then [23]

$$\sum_{i=0}^{q} (-1)^{i} c_{i}^{q,p} h_{j}^{i} f^{[i]}(t_{j+1}, y_{j+1}) \in \sum_{i=0}^{p} c_{i}^{p,q} h_{j}^{i} f^{[i]}(t_{j}, y_{j}) + (-1)^{q} \gamma_{p,q} h_{j}^{p+q+1} f^{[p+q+1]}(\boldsymbol{T}_{j}, \widetilde{\boldsymbol{y}}_{j}).$$
(20.5)

Denote

$$y_{i+1}^* = m(y_{i+1}^*), \tag{20.6}$$

$$\boldsymbol{B}_{j+1} = \sum_{i=0}^{q} (-1)^{i} c_{i}^{q,p} h_{j}^{i} J(f^{[i]}; \boldsymbol{y}_{j+1}^{*}), \tag{20.7}$$

$$\mathbf{F}_{j} = \sum_{i=0}^{p} c_{i}^{p,q} h_{j}^{i} J(f^{[i]}; \mathbf{y}_{j}), \tag{20.8}$$

$$C_{i+1} = m(\mathbf{B}_{i+1}), \tag{20.9}$$

$$S_{j+1} = (C_{j+1}^{-1} F_j) S_j, (20.10)$$

$$\mathbf{A}_{j+1} = (C_{j+1}^{-1} \mathbf{F}_j) A_j, \tag{20.11}$$

$$Q_{j+1} = (C_{j+1}^{-1} F_j) Q_j, (20.12)$$

$$e_{j+1} = (-1)^q \gamma_{p,q} h_j^{p+q+1} f^{[p+q+1]}(\mathbf{T}_j, \widetilde{\mathbf{y}}_j), \tag{20.13}$$

$$g_{j+1} = \sum_{i=0}^{p} c_i^{p,q} h_j^i f^{[i]}(u_j) - \sum_{i=0}^{q} (-1)^i c_i^{q,p} h_j^i f^{[i]}(y_{j+1}^*), \tag{20.14}$$

$$d_{j+1} = g_{j+1} + e_{j+1}$$
, and (20.15)

$$\boldsymbol{w}_{j+1} = C_{j+1}^{-1} \boldsymbol{d}_{j+1} + (I - C_{j+1}^{-1} \boldsymbol{B}_{j+1}) (\boldsymbol{y}_{j+1}^* - y_{j+1}^*). \tag{20.16}$$

(For sufficiently small  $h_j$ , we can enclose the inverse of  $C_{j+1}$ .) Since

$$y_{j+1}, y_{j+1}^* \in \mathbf{y}_{j+1}^* \text{ and } y_j, u_j \in \mathbf{y}_j,$$

we can apply the mean-value theorem to the two sums in (20.5), and using the above notation derive that

$$y_{j+1} \in y_{j+1}^* + S_{j+1}\alpha + A_{j+1}r_j + w_{j+1}$$
 and (20.17)

$$y_{j+1} \in y_{j+1}^* + S_{j+1}\alpha + Q_{j+1}r_{QR,j} + w_{j+1}.$$
 (20.18)

Denoting

$$s_{i+1} = (A_{i+1}r_i) \cap (Q_{i+1}r_{\text{OR},i}),$$

we have

$$y(t_{j+1}; t_0, y_0) \in \boldsymbol{y}_{j+1} := (y_{j+1}^* + \boldsymbol{S}_{j+1} \boldsymbol{\alpha} + \boldsymbol{s}_{j+1} + \boldsymbol{w}_{j+1}) \cap \boldsymbol{y}_{j+1}^*.$$

#### 20.1.3 Computing a solution representation

We have to determine  $u_{j+1}$ ,  $S_{j+1}$ ,  $A_{j+1}$ ,  $Q_{j+1}$ ,  $r_{j+1}$ , and  $r_{QR,j+1}$  for the next step.

Let

$$u_{j+1} = m(y_{j+1}), (20.19)$$

$$S_{j+1} = m(\mathbf{S}_{j+1}), \tag{20.20}$$

$$\mathbf{v}_{i+1} = y_{i+1}^* - u_{i+1} + (\mathbf{S}_{i+1} - S_{i+1})\alpha + \mathbf{w}_{i+1}. \tag{20.21}$$

$$A_{i+1} = m(A_{i+1}), (20.22)$$

$$\mathbf{r}_{j+1} = (A_{j+1}^{-1} \mathbf{A}_{j+1}) \mathbf{r}_j + A_{j+1}^{-1} \mathbf{v}_{j+1}, \tag{20.23}$$

 $Q_{j+1}$  the orthonormal matrix described in Subsection 20.1.4, and (20.24)

$$\mathbf{r}_{QR,j+1} = (Q_{j+1}^{-1}\mathbf{Q}_{j+1})\mathbf{r}_{QR,j} + Q_{j+1}^{-1}\mathbf{v}_{j+1}.$$
(20.25)

Using (20.19-20.25) in (20.17-20.18), we derive that

$$y_{j+1} \in \{u_{j+1} + S_{j+1}\alpha + A_{j+1}r \mid \alpha \in \alpha, r \in r_{j+1}\}$$
 and  $y_{j+1} \in \{u_{j+1} + S_{j+1}\alpha + Q_{j+1}r \mid \alpha \in \alpha, r \in r_{QR,j+1}\}.$ 

We assumed above that we can enclose  $A_{j+1}^{-1}$ . If we cannot enclose  $A_{j+1}^{-1}$ , or if

$$Q_{j+1}\boldsymbol{r}_{\mathrm{QR},j+1} \subseteq A_{j+1}\boldsymbol{r}_{j+1},$$

we set

$$A_{j+1} = Q_{j+1}$$
 and  $r_{j+1} = r_{QR,j+1}$ .

# 20.1.4 Computing $Q_{j+1}$

Let

$$\widetilde{A}_{j+1} = m(\mathbf{Q}_{j+1})$$
 and  $D = \operatorname{diag}(w(\mathbf{r}_{QR,j})).$ 

Let also  $P_{j+1}$  be a permutation matrix such that the columns of  $\widetilde{A}_{j+1}DP_{j+1}$  are sorted in non-increasing order in the Euclidean norm. Then, we perform the QR factorization of  $\widetilde{A}_{j+1}DP_{j+1} = Q_{j+1}R_{j+1}$  and use  $Q_{j+1}$  in our method.

# 20.2 Implementation

#### 20.2.1 The IHO class

```
235 ⟨class IHO 235⟩ ≡
class IHO {
public:
IHO(int n);
⟨set and get functions 277⟩
void compCoeffs();
void compTightEnclosure(interval &t_next);
void acceptSolution();
virtual ~IHO();
```

```
private:
         void compCpq(\mathbf{int}\ p,\mathbf{int}\ q);
         void compCqp(\mathbf{int}\ p,\mathbf{int}\ q);
         interval compErrorConstant(int p, int q);
       private:
          unsigned int p, q, order\_trial;
         interval h_trial;
         pVector y_pred_point;
         iVector y, y_pred, globalExcess, temp, temp2, x, u_next, predictor_excess,
              corrector\_excess, z, w, gj, term, d, s;
         iMatrix Fj, M, Cinv, G, B, S, A, Q, U, V, Ainv;
         pMatrix C, A_point;
         MatrixInverse *matrix_inverse;
         Solution *solution, *trial_solution;
         interval *C_pq, *C_qp;
         HOE *hoe;
          AD *ad:
         Control *control;
         interval errorConstant;
       };
    This code is used in chunk 292.
              Computing a tight enclosure
    20.2.2
    The main functions is
236 \langle compute tight enclosure 236 \rangle \equiv
       void IHO::compTightEnclosure(interval &t_next)
          (initialize IHO method 238)
          \langle \text{ predictor: compute } \boldsymbol{y}_{i+1}^* | 242 \rangle
```

This code is used in chunk 293.

 $\langle \operatorname{set} \boldsymbol{t}_{j+1} | 264 \rangle$ 

 $\langle \text{ corrector: compute } oldsymbol{y}_{j+1} \ \ {}_{247} 
angle$ 

(find solution representation for next step 265)

#### 20.2.3 Initialization

#### Stepsize

We assume we have enclosures on the solution at  $t_j \in \mathbf{t}_j$  and now wish to compute enclosures at  $t_{j+1} \in \mathbf{t}_{j+1}$ . Hence, we have for the stepsize

$$h_j = t_{j+1} - t_j \in \mathbf{h}_j = \mathbf{t}_{j+1} - \mathbf{t}_j.$$

```
238 \langle initialize IHO method 238 \rangle \equiv /*
t\_next = t_{j+1}
solution t = t_{j}
h\_trial \supseteq h_{j} = t_{j+1} - t_{j}
*/
h\_trial = t\_next - solution t;
See also chunks 240 and 241.
This code is used in chunk 236.
```

#### Order and method coefficients

The order is in  $order\_trial$ . Initially,  $order\_trial = 0$ . If  $control \neg order \neq order\_trial$ , we set  $order\_trial = control \neg order$  and compute the coefficients (20.3) and (20.4) of the method. If the value for the order is k, we require that p + q + 1 = k and  $p \leq q$ . We set

```
p = |(k-1)/2| and q = \lceil (k-1)/2 \rceil.
239 \langle compute IHO method coefficients 239\rangle \equiv
        void IHO::compCoeffs()
          if (order\_trial \neq control \neg order) {
                      /* deal with order */
             order\_trial = control \neg order;
             double pq = (order\_trial - 1)/2.0;
             p = \mathbf{int}(floor((pq)));
             q = \mathbf{int}(ceil((pq)));
             assert(p + q + 1 \equiv order\_trial);
                 /* reallocate memory if necessary */
             if (C_pq) delete [C_pq]
             C_pq = \mathbf{new} \ \mathbf{interval}[p+1];
             if (C_-qp) delete[] C_-qp;
             C_{-}qp = \mathbf{new interval}[q+1];
                 /* compute coefficients */
             compCpq(p,q);
             compCqp(p,q);
             errorConstant = compErrorConstant(p, q);
     This code is used in chunk 293.
```

240  $\langle$  initialize IHO method 238 $\rangle + \equiv$ 

compCoeffs();

#### Local excess

In the HOE method, we enclose

$$h_{j,0}^{q+1} f^{[q+1]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)$$

and

$$h_{j,0}^{p+q+1}f^{[p+q+1]}(\boldsymbol{t}_j+[0,1]h_{j,0},\widetilde{\boldsymbol{y}}_j)=h_{j,0}^kf^{[k]}(\boldsymbol{t}_j+[0,1]h_j,\widetilde{\boldsymbol{y}}_j).$$

In the IHO method, we have to enclose

$$h_j^{q+1}f^{[q+1]}(\boldsymbol{T}_j,\widetilde{\boldsymbol{y}}_j) \quad \text{and} \quad h_j^{p+q+1}f^{[p+q+1]}(\boldsymbol{T}_j,\widetilde{\boldsymbol{y}}_j),$$

where  $T_j \subseteq t_j + [0,1]h_{j,0}$ , cf. Proposition 19.1

We have

$$h_{j}^{q+1} f^{[q+1]}(\boldsymbol{T}_{j}, \widetilde{\boldsymbol{y}}_{j}) = \left(\frac{h_{j}}{h_{j,0}}\right)^{q+1} h_{j,0}^{q+1} f^{[q+1]}(\boldsymbol{T}_{j}, \widetilde{\boldsymbol{y}}_{j})$$

$$\subseteq \left(\frac{h_{j}}{h_{j,0}}\right)^{q+1} h_{j,0}^{q+1} f^{[q+1]}(\boldsymbol{t}_{j} + [0, h_{j,0}], \widetilde{\boldsymbol{y}}_{j}).$$

Similarly, we have

$$\begin{split} h_j^k f^{[k]}(\boldsymbol{T}_j, \widetilde{\boldsymbol{y}}_j) &= \left(\frac{h_j}{h_{j,0}}\right)^k h_{j,0}^k f^{[k]}(\boldsymbol{T}_j, \widetilde{\boldsymbol{y}}_j) \\ &\subseteq \left(\frac{h_j}{h_{j,0}}\right)^k h_{j,0}^k f^{[k]}(\boldsymbol{t}_j + [0, h_{j,0}], \widetilde{\boldsymbol{y}}_j). \end{split}$$

Hence, we just re-scale.

241  $\langle \text{ initialize IHO method } 238 \rangle + \equiv /*$ 

$$h_{trial} \ni h_i$$

 $tayl\_coeff \rightarrow getStepsize()$  returns  $h_{i,0}$ 

tayl\_coeff contains enclosures on  $h_{j,0}^i f^{[i]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)$ 

for 
$$i = 0, ..., k = p + q + 1$$

$$rescale \ni h_j/h_{j,0}$$

$$predictor\_excess \supseteq h_{j,0}^{q+1} f^{[q+1]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)$$

$$predictor\_excess \supseteq h_j^{q+1} f^{[q+1]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)$$

$$corrector\_excess \supseteq h_{j,0}^{p+q+1} f^{[p+q+1]}(\boldsymbol{t}_j + [0,1]h_{j,0}\widetilde{\boldsymbol{y}}_j)$$

$$corrector\_excess \supseteq h_j^{p+q+1} f^{[p+q+1]}(\boldsymbol{t}_j + [0,1]h_{j,0}, \widetilde{\boldsymbol{y}}_j)$$

\*/
interval rescale = h\_trial/ad-tayl\_coeff\_ode-getStepsize();

 $hoe \rightarrow getErrorTerm(predictor\_excess, q + 1);$ 

 $scale V(predictor\_excess, \mathbf{v\_bias} :: pow(rescale, q + 1));$ 

 $hoe \neg getErrorTerm(corrector\_excess, p + q + 1);$ 

scale V (corrector\_excess, pow(rescale, order\_trial));

# 20.2.4 Predictor

```
We enclose \hat{u}_{j+1}, U_{j+1}, x_{j+1}, and y_{j+1}^*. Below, t_j \in t_j.
242 \langle predictor: compute y_{i+1}^* 242 \rangle \equiv
            \langle \hat{u}_{j+1} = u_j + \sum_{i=1}^{q} h_i^i f^{[i]}(t_j, u_j) \ 243 \rangle
            \langle \boldsymbol{U}_{j+1} = I + \sum_{i=1}^{q} h_i^i J(f^{[i]}; \boldsymbol{y}_i) \ 244 \rangle
            \langle x_{j+1} = (U_{j+1}S_j)\alpha + \{(U_{j+1}A_j)r_j \cap (U_{j+1}Q_j)r_{QR,j}\} 245\rangle
            \langle \boldsymbol{y}_{j+1}^* = (\widehat{u}_{j+1} + \boldsymbol{z}_{j+1} + \boldsymbol{x}_{j+1}) \cap \widetilde{\boldsymbol{y}}_j \ 246 \rangle
        This code is used in chunk 236.
243 \ \langle \widehat{u}_{j+1} = u_j + \sum_{i=1}^q h_i^i f^{[i]}(t_j, u_j) \ 243 \rangle \equiv /*
                                solution \rightarrow t = t_i
                               solution \neg u = u_j
                                     h\_trial \supseteq h_j
                                               q order
                                 tayl_coeff contains enclosures on h_i^i f^{[i]}(t_i, u_i) for i = 0, \ldots, q
                                      u\_next \ni \widehat{u}_{j+1}
             assign V(temp, solution \neg u); /* temp stores u_j as an interval */
             ad \rightarrow tayl\_coeff\_ode \rightarrow set(solution \rightarrow t, temp, h\_trial, q);
             ad \rightarrow tayl\_coeff\_ode \rightarrow compTerms();
             ad \rightarrow tayl\_coeff\_ode \rightarrow sumTerms(u\_next, q);
        This code is used in chunk 242.
244 \ \langle \boldsymbol{U}_{j+1} = I + \sum_{i=1}^{q} h_i^i J(f^{[i]}; \boldsymbol{y}_i) \ 244 \rangle \equiv
                                     solution \rightarrow t = t_i
                                    solution \rightarrow y \supseteq \mathbf{y}_i
                                           h_{\perp}trial \supseteq h_i
                                                     q order
                               tayl\_coeff\_var contains enclosures on h_i^i J(f^{[i]}; y_i)
                                                           for i = 0, 1, ..., q
                                                   U\supseteq \boldsymbol{U}_{j+1}=I+\sum_{i=1}^q h_j^i\,J(f^{[i]};\boldsymbol{y}_j)
             ad \rightarrow tayl\_coeff\_var \rightarrow set(solution \rightarrow t, solution \rightarrow y, h\_trial, q);
             ad \rightarrow tayl\_coeff\_var \rightarrow comp\ Terms();
             ad \rightarrow tayl\_coeff\_var \rightarrow sum Terms(U, q);
        This code is used in chunk 242.
```

Now we evaluate

```
245 \langle x_{j+1} = (U_{j+1}S_j)\alpha + \{(U_{j+1}A_j)r_j \cap (U_{j+1}Q_j)r_{QR,j}\} 245 \rangle \equiv
                          solution contains u_i, S_i, \boldsymbol{\alpha}, A_i, Q_i, \boldsymbol{r}_i, \boldsymbol{r}_{\text{QR},i}, \boldsymbol{y}_i
                                   U \supseteq \boldsymbol{U}_{j+1}
                                  M \supset \boldsymbol{U}_{i+1} A_i
                              temp \supseteq (U_{j+1}A_j)r_j
                                  M \supseteq U_{i+1}Q_i
                            temp2 \supseteq (\boldsymbol{U}_{j+1}Q_j)\boldsymbol{r}_{\mathrm{QR},j}
                              temp \supseteq (\boldsymbol{U}_{j+1}A_j)\boldsymbol{r}_j \cap (\boldsymbol{U}_{j+1}Q_j)\boldsymbol{r}_{QR,j}
                                  M \supseteq \boldsymbol{U}_{i+1}S_i
                              tem2 \supseteq (\boldsymbol{U}_{j+1}S_j)\boldsymbol{\alpha}
                                    x \supseteq x_{j+1} = (U_{j+1}S_j)\alpha + \{(U_{j+1}A_j)r_j \cap (U_{j+1}Q_j)r_{QR,j}\}
          multMiMp(M, U, solution \rightarrow A);
          multMiVi(temp, M, solution \rightarrow r);
          multMiMp(M, U, solution \rightarrow Q);
          multMiVi(temp2, M, solution \neg rQR);
          bool finite\_temp = finite\_interval(temp);
          bool finite\_temp2 = finite\_interval(temp2);
          if (\neg finite\_temp \land \neg finite\_temp2) {
              control \rightarrow ind = failure;
             return;
          else {
             if (finite\_temp \land finite\_temp2) {
                 bool b = intersect(temp, temp2, temp);
                 assert(b);
             else {
                 if (\neg finite\_temp) assign V(temp, temp2);
                          /* else temp is finite and temp2 is not. We do not do anything. */
          }
          multMiMp(M, U, solution \rightarrow S);
          multMiVi(temp2, M, solution \neg alpha);
          addViVi(x, temp, temp2);
       This code is used in chunk 242.
```

Finally, we compute

```
246 \quad \langle \boldsymbol{y}_{j+1}^* = (\widehat{u}_{j+1} + \boldsymbol{z}_{j+1} + \boldsymbol{x}_{j+1}) \cap \widetilde{\boldsymbol{y}}_j \ \ 246 \rangle \equiv \quad /*
u\_next \ni \widehat{u}_{j+1}
predictor\_excess \supseteq \boldsymbol{z}_{j+1}
x \supseteq \boldsymbol{x}_{j+1}
hoe \neg getTrialApriori() \supseteq \widetilde{\boldsymbol{y}}_j
y\_pred \supseteq u_{j+1} + \boldsymbol{z}_{j+1}
y\_pred \supseteq \widehat{u}_{j+1} + \boldsymbol{z}_{j+1} + \boldsymbol{x}_{j+1}
y\_pred \supseteq (\widehat{u}_{j+1} + \boldsymbol{z}_{j+1} + \boldsymbol{x}_{j+1}) \cap \widetilde{\boldsymbol{y}}_j
\stackrel{*/}{addViVi(y\_pred, u\_next, predictor\_excess);}
addViVi(y\_pred, x);
bool \ b = intersect(y\_pred, hoe \neg getTrialApriori(), y\_pred);
assert(b);
This code is used in chunk 242.
```

#### 20.2.5 Corrector

The computation of  $y_{i+1}$  is given by

247 
$$\langle \text{ corrector: compute } \boldsymbol{y}_{j+1} | 247 \rangle \equiv$$

$$\langle \boldsymbol{F}_{j} = \sum_{i=0}^{p} c_{i}^{p,q} h_{j}^{i} J(f^{[i]}; \boldsymbol{y}_{j}) | 248 \rangle$$

$$\langle \boldsymbol{B}_{j+1} = \sum_{i=0}^{q} (-1)^{i} c_{i}^{q,p} h_{j}^{i} J(f^{[i]}; \boldsymbol{y}_{j+1}^{*}) | 249 \rangle$$

$$\langle \boldsymbol{C}_{j+1} = \mathbf{m}(\boldsymbol{B}_{j+1}) | 250 \rangle$$

$$\langle \boldsymbol{G}_{j+1} = \boldsymbol{C}_{j+1}^{-1} \boldsymbol{F}_{j} | 251 \rangle$$

$$\langle \boldsymbol{S}_{j+1} = \boldsymbol{G}_{j+1} \boldsymbol{S}_{j} | 252 \rangle$$

$$\langle \boldsymbol{A}_{j+1} = \boldsymbol{G}_{j+1} \boldsymbol{A}_{j} | 253 \rangle$$

$$\langle \boldsymbol{Q}_{j+1} = \boldsymbol{G}_{j+1} \boldsymbol{Q}_{j} | 254 \rangle$$

$$\langle \boldsymbol{e}_{j+1} = (-1)^{q} \gamma_{p,q} h_{j}^{p+q+1} f^{[p+q+1]}(\boldsymbol{T}_{j}, \boldsymbol{\tilde{y}}_{j}) | 255 \rangle$$

$$\langle \boldsymbol{g}_{j+1} = \sum_{i=0}^{p} c_{i}^{p,q} h_{j}^{i} f^{[i]}(\boldsymbol{u}_{j}) - \sum_{i=0}^{q} (-1)^{i} c_{i}^{q,p} h_{j}^{i} f^{[i]}(\boldsymbol{y}_{j+1}^{*}) | 256 \rangle$$

$$\langle \boldsymbol{d}_{j+1} = \boldsymbol{g}_{j+1} + \boldsymbol{e}_{j+1} | 260 \rangle$$

$$\langle \boldsymbol{w}_{j+1} = \boldsymbol{C}_{j+1}^{-1} \boldsymbol{d}_{j+1} + (\boldsymbol{I} - \boldsymbol{C}_{j+1}^{-1} \boldsymbol{B}_{j+1}) (\boldsymbol{y}_{j+1}^{*} - \boldsymbol{y}_{j+1}^{*}) | 261 \rangle$$

$$\langle \boldsymbol{s}_{j+1} = (\boldsymbol{A}_{j+1} \boldsymbol{r}_{j}) \cap (\boldsymbol{Q}_{j+1} \boldsymbol{r}_{QR,j}) | 262 \rangle$$

$$\langle \boldsymbol{y}_{j+1} = (\boldsymbol{y}_{j+1}^{*} + \boldsymbol{S}_{j+1} \boldsymbol{\alpha} + \boldsymbol{s}_{j+1} + \boldsymbol{w}_{j+1}) \cap \boldsymbol{y}_{j+1}^{*} | 263 \rangle$$

This code is used in chunk 236.

#### Computing $F_j$

The matrices  $h_j^i J(f^{[i]}; \boldsymbol{y}_j)$  for  $i = 1, \ldots, q$  have been already enclosed in the predictor. Since  $p \leq q$ , we have the terms that we need to enclose  $\boldsymbol{F}_j$  in (20.8). In the code below, we obtain the enclosure on  $h_j^i J(f^{[i]}; \boldsymbol{y}_j)$  in M and then scale M such that it contains  $c_i^{p,q} h_j^i J(f^{[i]}; \boldsymbol{y}_j)$ .

 $B \supseteq \boldsymbol{B}_{j+1}$ 

 $ad \rightarrow tayl\_coeff\_var \rightarrow set(t\_next, y\_pred, h\_trial, q);$ 

 $ad \rightarrow tayl\_coeff\_var \rightarrow comp Terms();$ 

for (int i = q;  $i \ge 1$ ; i - -) {

setM(B, 0.0);

```
ad \rightarrow tayl\_coeff\_var \rightarrow getTerm(M,i);
scaleM(M, C\_qp[i]);
addMiMi(B, M);
}
addId(B);
This code is used in chunk 247.
(C_{j+1} = m(\boldsymbol{B}_{j+1}) \ 250) \equiv /*
\boldsymbol{B} \supseteq \boldsymbol{B}_{j+1}
\boldsymbol{C} = C_{j+1} = m(\boldsymbol{B}_{j+1})
*/
midpoint(C, B);
This code is used in chunk 247.
```

The function encloseInverse below encloses the inverse of a point matrix. If this function fails, it returns false; otherwise, it returns true.

$$251 \ \langle \boldsymbol{G}_{j+1} = C_{j+1}^{-1} \boldsymbol{F}_{j} \ 251 \rangle \equiv /*$$

$$C = C_{j+1}$$

$$F_{j} \supseteq \boldsymbol{F}_{j}$$

$$Cinv \ni C_{j+1}^{-1} \text{ if } ok$$

$$G \supseteq \boldsymbol{G}_{j+1} = C_{j+1}^{-1} \boldsymbol{F}_{j}$$

$$*/$$

$$\boldsymbol{bool} \ ok = matrix\_inverse\_encloseMatrixInverse(Cinv, C);$$

$$\textbf{if} \ (\neg ok)$$

$$\{ \\ control\_ind = failure;$$

$$\# \textbf{ifdef VNODE\_DEBUG} \\ printMessage("Could\_not\_invert\_the\_C\_matrix.");$$

$$\# \textbf{endif} \\ \textbf{return};$$

$$\} \\ multMiMi(G, Cinv, F_{j});$$

$$This code is used in chunk 247.$$

$$252 \ \langle \boldsymbol{S}_{j+1} = \boldsymbol{G}_{j+1} \boldsymbol{S}_{j} \ 252 \rangle \equiv /*$$

$$solution\_S = S_{j}$$

$$G \supseteq \boldsymbol{G}_{j+1}$$

$$S \supseteq \boldsymbol{S}_{j+1} = \boldsymbol{G}_{j+1} \boldsymbol{S}_{j}$$

\*/ 
$$multMiMp(S, G, solution \neg S);$$
 This code is used in chunk 247.

253 
$$\langle \mathbf{A}_{j+1} = \mathbf{G}_{j+1} A_j | 253 \rangle \equiv /*$$

$$solution A = A_j$$

$$G \supseteq \mathbf{G}_{j+1}$$

$$A \supseteq \mathbf{A}_{j+1} = \mathbf{G}_{j+1} A_j$$

\*/  $multMiMp(A, G, solution \rightarrow A);$ This code is used in chunk 247.

$$254 \qquad \langle \mathbf{Q}_{j+1} = \mathbf{G}_{j+1}Q_j \ \ 254 \rangle \equiv \qquad /*$$

$$solution \neg Q = Q_j$$

$$G \supseteq \mathbf{G}_{j+1}$$

$$Q \supseteq \mathbf{Q}_{j+1} = \mathbf{G}_{j+1}Q_j$$

\*/  $multMiMp(Q, G, solution \rightarrow Q);$ This code is used in chunk 247.

# Computing $e_{j+1}$

$$\begin{aligned} & corrector\_excess \text{ encloses } \boldsymbol{h}_{j}^{p+q+1}f^{[p+q+1]}(\boldsymbol{T}_{j},\widetilde{\boldsymbol{y}}_{j}). \text{ We multiply it by } (-1)^{q}\gamma_{p,q}.\\ & 255 \ \ \langle \boldsymbol{e}_{j+1} = (-1)^{q}\gamma_{p,q}\,\boldsymbol{h}_{j}^{p+q+1}\,f^{[p+q+1]}(\boldsymbol{T}_{j},\widetilde{\boldsymbol{y}}_{j})\,\,255\,\rangle \equiv \qquad /*\\ & corrector\_excess \supseteq \boldsymbol{h}_{j}^{p+q+1}\,f^{[p+q+1]}(\boldsymbol{T}_{j},\widetilde{\boldsymbol{y}}_{j})\\ & \underline{errorConstant} \supseteq (-1)^{q}\gamma_{p,q}\\ & \underline{corrector\_excess} \supseteq (-1)^{q}\gamma_{p,q}\boldsymbol{h}_{j}^{p+q+1}\,f^{[p+q+1]}(\boldsymbol{T}_{j},\widetilde{\boldsymbol{y}}_{j}) \end{aligned}$$

\*/
scale V (corrector\_excess, errorConstant);
This code is used in chunk 247.

# Enclosing $g_{j+1}$

$$\begin{array}{l} 256 \ \, \langle \, g_{j+1} = \sum_{i=0}^p c_i^{p,q} h_j^i f^{[i]}(u_j) - \sum_{i=0}^q (-1)^i c_i^{q,p} h_j^i f^{[i]}(y_{j+1}^*) \,\, 256 \, \rangle \equiv \\ \ \, \langle \, g_{j+1}^{\rm f} = \sum_{i=0}^p c_i^{p,q} h_j^i f^{[i]}(u_j) \,\, 257 \, \rangle \\ \ \, \langle \, g_{j+1}^{\rm b} = \sum_{i=0}^q (-1)^i c_i^{q,p} h_j^i f^{[i]}(y_{j+1}^*) \,\, 258 \, \rangle \\ \ \, \langle \, g_{j+1} = g_{j+1}^{\rm f} - g_{j+1}^{\rm b} \,\, 259 \, \rangle \\ \ \, \text{This code is used in chunk 247.} \end{array}$$

\*/

```
We have the terms h_i^i f^{[i]}(u_j) for i = 1, \dots, q computed in the predictor.
257 \langle g_{j+1}^{\rm f} = \sum_{i=0}^{p} c_i^{p,q} h_j^i f^{[i]}(u_j) \; 257 \rangle \equiv /*
                            tayl_coeff contains h_i^i f^{[i]}(u_j) for i = 0, 1, \dots q
                                          p order
                                         qj = 0
                                     term \ni h_i^i f^{[i]}(u_j)
                                 C_pq[i] \ni c_i^{p,q}
                                    term \ni c_i^{p,q} h_j^i f^{[i]}(u_j)
                                        gj \ni g_{j+1}^{f} = \sum_{i=0}^{p} c_i^{p,q} h_j^i f^{[i]}(u_j)
            set V(gj, 0.0);
            for (int i = p; i \ge 0; i--) {
                ad \rightarrow tayl\_coeff\_ode \rightarrow getTerm(term, i);
                scale V(term, C\_pq[i]);
                addViVi(gj, term);
        This code is used in chunk 256.
258 \langle g_{j+1}^{b} = \sum_{i=0}^{q} (-1)^{i} c_{i}^{q,p} h_{i}^{i} f^{[i]}(y_{j+1}^{*}) 258 \rangle \equiv
                                       t_next = t_{i+1}
                                       y\_pred \supseteq \boldsymbol{y}_{i+1}^*
                                      h_trial \supseteq h_j
                                               q order
                                       temp2 = 0
                             y-pred-point = y_{j+1}^* = m(\boldsymbol{y}_{j+1}^*)
                                         temp interval vector storing y_{i+1}^*
                                  \mathit{tayL}\mathit{coeff} contains h_j^i f^{[i]}(y_{j+1}^*) after \mathit{compTerms} is called
                                        term \ni h_i^i f^{[i]}(y_{i+1}^*)
                                      C_{-qp}[i] \ni (-1)^i c_i^{q,p}
                                     term \ni (-1)^{i} c_{i}^{q,p} h_{j}^{i} f^{[i]}(y_{j+1}^{*})
temp2 \ni g_{j+1}^{b} = \sum_{i=0}^{q} (-1)^{i} c_{i}^{q,p} h_{j}^{i} f^{[i]}(y_{j+1}^{*})
```

```
\label{eq:midpoint} \begin{split} & \textit{midpoint}(\textit{y\_pred\_point}, \textit{y\_pred}); \\ & \textit{assignV}(\textit{temp}, \textit{y\_pred\_point}); \\ & \textit{ad\_tayl\_coeff\_ode\_set}(\textit{t\_next}, \textit{temp}, \textit{h\_trial}, \textit{q}); \\ & \textit{ad\_tayl\_coeff\_ode\_compTerms}(); \\ & \textit{setV}(\textit{temp2}, 0.0); \\ & \textit{for} \ (\textbf{int} \ i = q; \ i \geq 0; \ i--) \ \{ \\ & \textit{ad\_tayl\_coeff\_ode\_getTerm}(\textit{term}, i); \\ & \textit{scaleV}(\textit{term}, \textit{C\_qp}[i]); \\ & \textit{addViVi}(\textit{temp2}, \textit{term}); \\ \} \end{split}
```

This code is used in chunk 256.

$$259 \langle g_{j+1} = g_{j+1}^{f} - g_{j+1}^{b} 259 \rangle \equiv /*$$

$$gj \ni g_{j+1}^{f}$$

$$temp2 \ni g_{j+1}^{b}$$

$$gj \ni g_{j+1} = g_{j+1}^{f} - g_{j+1}^{b}$$

$$*/sub ViVi(gj, temp2);$$

This code is used in chunk 256.

# Computing $d_{j+1}$

This code is used in chunk 247.

## Computing $w_{j+1}$

Now, we compute

```
_{261} \ \langle \boldsymbol{w}_{j+1} = C_{j+1}^{-1} \boldsymbol{d}_{j+1} + (I - C_{j+1}^{-1} \boldsymbol{B}_{j+1}) (\boldsymbol{y}_{j+1}^* - y_{j+1}^*) \ \text{261} \, \rangle \equiv
                                y\_pred\_point = y_{i+1}^*
                                           y\_pred \supseteq \boldsymbol{y}_{i+1}^*
                                                   B \supseteq \boldsymbol{B}_{i+1}
                                             Cinv \ni C_i^{-1}
                                                    d \supseteq \mathbf{d}_{j+1}
                                             term \supseteq \boldsymbol{y}_{j+1}^* - y_{j+1}^*
                                                 M \supseteq C_j^{-1} \boldsymbol{B}_{j+1}
                                                 M \supseteq I - C_i^{-1} \boldsymbol{B}_{j+1}
                                             temp \supseteq (I - C_{i+1}^{-1} \boldsymbol{B}_{j+1}) (\boldsymbol{y}_{i+1}^* - y_{i+1}^*)
                                                  w \supseteq \boldsymbol{w}_{j+1} = C_{i+1}^{-1} \boldsymbol{d}_{j+1} + (I - C_{i+1}^{-1} \boldsymbol{B}_{j+1}) (\boldsymbol{y}_{i+1}^* - y_{i+1}^*)
             subViVp(term, y_pred, y_pred_point);
             multMiMi(M, Cinv, B);
             subFromId(M);
             multMiVi(temp, M, term);
             multMiVi(w, Cinv, d);   /* w = Cinv * d + temp */
             addViVi(w, temp);
        This code is used in chunk 247.
262 \langle s_{j+1} = (A_{j+1}r_j) \cap (Q_{j+1}r_{QR,j}) \ 262 \rangle \equiv /*
                                       solution \neg r \supseteq r_j
                                                      A \supseteq \mathbf{A}_{i+1}
                                solution \neg rQR \supseteq r_{\text{OR},i}
                                                      Q \supseteq \mathbf{Q}_{i+1}
                                                temp \supseteq \boldsymbol{A}_{i+1}\boldsymbol{r}_i
                                                       s \supseteq oldsymbol{Q}_{j+1} oldsymbol{r}_{	ext{QR},j}
                                                       s \supseteq s_{i+1} = (\boldsymbol{A}_{i+1} \boldsymbol{r}_i) \cap (\boldsymbol{Q}_{i+1} \boldsymbol{r}_{QR,j})
             multMiVi(temp, A, solution \neg r);
             multMiVi(s, Q, solution \neg rQR);
             b = intersect(s, temp, s);
             assert(b);
        This code is used in chunk 247.
```

$$263 \ \langle \mathbf{y}_{j+1} = (y_{j+1}^* + \mathbf{S}_{j+1} \alpha + \mathbf{s}_{j+1} + \mathbf{w}_{j+1}) \cap \mathbf{y}_{j+1}^* \ 263 \rangle \equiv /*$$

$$trial\_solution\_alpha \supseteq \alpha$$

$$S \supseteq \mathbf{S}_{j+1}$$

$$\mathbf{y}\_pred \supseteq \mathbf{y}_{j+1}^*$$

$$\mathbf{y}\_pred\_point = \mathbf{y}_{j+1}^*$$

$$s \supseteq \mathbf{s}_{j+1}$$

$$w \supseteq \mathbf{w}_{j+1}$$

$$globalExcess \supseteq \mathbf{s}_{j+1} + \mathbf{w}_{j+1}$$

$$temp \supseteq \mathbf{S}_{j+1}\alpha$$

$$temp \supseteq \mathbf{S}_{j+1}\alpha + \mathbf{s}_{j+1} + \mathbf{w}_{j+1}$$

$$temp \supseteq \mathbf{y}_{j+1}^* + \mathbf{S}_{j+1}\alpha + \mathbf{s}_{j+1} + \mathbf{w}_{j+1}$$

$$temp \supseteq \mathbf{y}_{j+1}^* + \mathbf{S}_{j+1}\alpha + \mathbf{s}_{j+1} + \mathbf{w}_{j+1}$$

$$trial\_solution\_y \supseteq \mathbf{y}_{j+1} = (\mathbf{y}_{j+1}^* + \mathbf{S}_{j+1}\alpha + \mathbf{x}_{j+1}) \cap \mathbf{y}_{j+1}^*$$

$$^*/$$

$$addViVi(globalExcess, \mathbf{s}, \mathbf{w});$$

$$multMiVi(temp, \mathbf{S}, trial\_solution\_alpha);$$

$$addViVi(temp, \mathbf{S}, trial\_solution\_alpha);$$

$$addViVi(temp, \mathbf{g}, \mathbf{b})$$

$$addViVi(temp, \mathbf{y}\_pred\_point);$$

$$b = intersect(trial\_solution\_y, \mathbf{y}\_pred, temp);$$

$$assert(b);$$
This code is used in chunk 247.

$$264 \ \langle \mathbf{set} \ \mathbf{t}_{j+1} \ 264 \rangle \equiv trial\_solution\_t = t\_next;$$
This code is used in chunk 236.

#### 20.2.6 Enclosure representation

```
265 \langle find solution representation for next step 265 \rangle \equiv \langle u_{j+1} = \mathrm{m}(\boldsymbol{y}_{j+1}) | 266 \rangle

\langle S_{j+1} = \mathrm{m}(\boldsymbol{S}_{j+1}) | 267 \rangle

\langle \boldsymbol{v}_{j+1} = \boldsymbol{y}_{j+1}^* - u_{j+1} + (\boldsymbol{S}_{j+1} - \boldsymbol{S}_{j+1})\boldsymbol{\alpha} + \boldsymbol{w}_{j+1} | 268 \rangle

\langle A_{j+1} = \mathrm{m}(\boldsymbol{A}_{j+1}) | 269 \rangle

\langle \boldsymbol{r}_{j+1} = (A_{j+1}^{-1}\boldsymbol{A}_{j+1})\boldsymbol{r}_j + A_{j+1}^{-1}\boldsymbol{v}_{j+1} | 270 \rangle

\langle compute Q_{j+1} | 273 \rangle

\langle \boldsymbol{r}_{\mathrm{QR},j+1} = (Q_{j+1}^{-1}\boldsymbol{Q}_{j+1})\boldsymbol{r}_{\mathrm{QR},j} + Q_{j+1}^{-1}\boldsymbol{v}_{j+1} | 271 \rangle

\langle reset if needed 272 \rangle

This code is used in chunk 236.
```

```
266 \langle u_{j+1} = m(y_{j+1}) \ 266 \rangle \equiv
                       trial\_solution \neg y \supseteq \mathbf{y}_{j+1}trial\_solution \neg u = u_{j+1}
           midpoint(trial\_solution \neg u, trial\_solution \neg y);
       This code is used in chunk 265.
267 \langle S_{j+1} = m(S_{j+1}) \ 267 \rangle \equiv /*
                          trial\_solution \neg S = S_{j+1} = m(S_{j+1})
           midpoint(trial\_solution \neg S, S);
       This code is used in chunk 265.
268 \langle v_{j+1} = y_{j+1}^* - u_{j+1} + (S_{j+1} - S_{j+1})\alpha + w_{j+1} | 268 \rangle \equiv
                                      y\_pred\_point = y_{i+1}^*
                                 trial\_solution \neg u = u_{j+1}
                                                      S \supseteq \boldsymbol{S}_{i+1}
                                 trial\_solution \neg S = S_{i+1}
                           trial\_solution \neg alpha \supseteq \alpha
                                                      S \supseteq S_{i+1} - S_{i+1}
                                                       z \supseteq (S_{j+1} - S_{j+1})\alpha
                                                       z \supseteq (S_{j+1} - S_{j+1})\alpha + w_{j+1}
                                                       z \supseteq -u_{j+1} + (S_{j+1} - S_{j+1})\alpha + w_{j+1}
                                                      z \supseteq v_{j+1} = y_{j+1}^* - u_{j+1} + (S_{j+1} - S_{j+1})\alpha + w_{j+1}
           subMiMp(S, trial\_solution \neg S);
           multMiVi(z, S, trial\_solution \rightarrow alpha);
           addViVi(z, w);
           sub ViVp(z, trial\_solution \neg u);
           addViVp(z, y\_pred\_point);
       This code is used in chunk 265.
269 \langle A_{j+1} = m(A_{j+1}) \ 269 \rangle \equiv
                                                A \supseteq \mathbf{A}_{j+1}
                          trial\_solution \rightarrow A = A_{i+1}
```

```
midpoint(trial\_solution \neg A, A);
         This code is used in chunk 265.
_{270} \langle \boldsymbol{r}_{j+1} = (A_{j+1}^{-1} \boldsymbol{A}_{j+1}) \boldsymbol{r}_{j} + A_{j+1}^{-1} \boldsymbol{v}_{j+1} | 270 \rangle \equiv
                                 trial\_solution \rightarrow A = A_{j+1}
                                                            A \supseteq \mathbf{A}_{j+1}
                                                             z \supseteq \boldsymbol{v}_{i+1}
                                            solution \neg r \supseteq r_i
                                                      Ainv \ni A_{i+1}^{-1} \text{ if } ok
                                                      temp \supseteq A_{j+1}^{-1} \boldsymbol{v}_{j+1}
                                                          M \supseteq A_{j+1}^{-1} \boldsymbol{A}_{j+1}
                                  trial\_solution \neg r \supseteq (A_{j+1}^{-1} \boldsymbol{A}_{j+1}) \boldsymbol{r}_j
                                  trial\_solution \neg r \supseteq r_{j+1} = (A_{j+1}^{-1} A_{j+1}) r_j + A_{j+1}^{-1} v_{j+1}
              ok = matrix\_inverse \neg encloseMatrixInverse (Ainv, trial\_solution \neg A);
             if (ok) {
                  multMiVi(temp, Ainv, z);
                  multMiMi(M, Ainv, A);
                  multMiVi(trial\_solution \neg r, M, solution \neg r);
                  addViVi(trial\_solution \neg r, temp);
         This code is used in chunk 265.
271 \ \langle \boldsymbol{r}_{\text{QR},j+1} = (Q_{i+1}^{-1}\boldsymbol{Q}_{i+1})\boldsymbol{r}_{\text{QR},j} + Q_{i+1}^{-1}\boldsymbol{v}_{j+1} \ 271 \rangle \equiv
                                       trial\_solution \neg Q = Q_{j+1}
                                                                   z \supseteq v_{i+1}
                                           solution \neg rQR \supseteq r_{QR,j}
                                                                 Q \supseteq \mathbf{Q}_{j+1}
                                                           Ainv \ni Q_{i+1}^{-1} \text{ if } ok
                                                           temp \supseteq Q_{i+1}^{-1} \boldsymbol{v}_{i+1}
                                                                M \supseteq Q_{i+1}^{-1} \boldsymbol{Q}_{i+1}
                                 \mathit{trial\_solution} \neg \mathit{rQR} \supseteq (Q_{j+1}^{-1} \boldsymbol{Q}_{j+1}) \boldsymbol{r}_{\text{QR},j}
                                 trial\_solution \neg rQR \supseteq r_{\text{QR},j+1} = (Q_{j+1}^{-1}Q_{j+1})r_{\text{QR},j} + Q_{j+1}^{-1}v_{j+1}
                       */
```

```
b = matrix\_inverse \neg orthogonalInverse (Ainv, trial\_solution \neg Q);
         if (b \equiv false) {
            control \neg ind = failure;
      #ifdef VNODE_DEBUG
            printMessage("Could_not_invert_the_Q_matrix.");
      #endif
            return;
         multMiVi(temp, Ainv, z);
         multMiMi(M, Ainv, Q);
         multMiVi(trial\_solution \neg rQR, M, solution \neg rQR);
         addViVi(trial\_solution \neg rQR, temp);
      This code is used in chunk 265.
272 \langle \text{ reset if needed } 272 \rangle \equiv
                                                 trial\_solution \neg Q = Q_{j+1}
                                             trial\_solution \neg rQR \supseteq r_{QR, j+1}
                                                 trial\_solution \rightarrow A = A_{i+1}
                                                  trial\_solution \neg r \supseteq r_{j+1}
                                                              temp \supseteq Q_{j+1} r_{QR,j+1}
                                                             temp2 \supseteq A_{i+1} \boldsymbol{r}_{i+1}
                       if ( subseteq(temp, temp2) \lor \neg ok )
                                                 trial\_solution \neg A = Q_{j+1}
                                                  trial\_solution \neg r \supseteq r_{\text{QR},j+1}
         multMpVi(temp, trial\_solution \rightarrow Q, trial\_solution \rightarrow rQR);
         multMpVi(temp2, trial\_solution \rightarrow A, trial\_solution \rightarrow r);
         if (subseteq(temp, temp2) \lor \neg ok) {
            assignM(trial\_solution \rightarrow A, trial\_solution \rightarrow Q);
            assign V (trial\_solution \neg r, trial\_solution \neg rQR);
      This code is used in chunk 265.
```

## Computing $Q_{i+1}$

```
273 \langle \text{ compute } Q_{j+1} | 273 \rangle \equiv
                                     Q \supseteq \boldsymbol{Q}_{j+1}
                              A\_point = m(\boldsymbol{Q}_{i+1})
                              A\_point = m(\mathbf{Q}_{i+1}) \operatorname{diag} w(\mathbf{r}_{QR,j})
                                      C = A-point with columns sorted in non-increasing order in \|\cdot\|_2
                                      C = Q_{i+1}R_{i+1}
                     trial\_solution \neg Q = Q_{i+1}
              */
        midpoint(A\_point, Q);
        int n = sizeM(Q);
        for (int i = 0; i < n; i +++)
           for (int j = 0; j < n; j ++)
              A\_point[i][j] *= \mathbf{v\_bias} :: width(solution \neg rQR[j]);
        if (\neg(inf\_normM(A\_point) < \mathbf{numeric\_limits} \langle \mathbf{double} \rangle :: max())) {
      #ifdef VNODE_DEBUG
           printMessage("The computed enclosures are too wide");
      #endif
           control \rightarrow ind = failure;
           return;
        sortColumns(C, A\_point);
        b = computeQR(trial\_solution \neg Q, C);
        assert(b);
     This code is used in chunk 265.
     20.2.7
                Constructor
274 \langle \text{ constructor IHO } 274 \rangle \equiv
        IHO::IHO(int n)
           order\_trial = 0;
           solution = new Solution(n);
           trial\_solution = new Solution(n);
           matrix\_inverse = new MatrixInverse(n);
           C_{pq} = C_{qp} = 0;
           size V(y, n);
           size V(y\_pred, n);
           size V(globalExcess, n);
           size V(y\_pred\_point, n);
           size V(temp, n);
           size V(temp2, n);
           size V(x, n);
```

```
size V(u\_next, n);
          sizeV(predictor\_excess, n);
          sizeV(corrector\_excess, n);
           size V(z, n);
           size V(w, n);
          size V(gj, n);
           sizeV(term, n);
           size V(d, n);
          size V(s, n);
          sizeM(Fj, n);
           sizeM(M, n);
           sizeM(Cinv, n);
          sizeM(G, n);
          sizeM(B, n);
          sizeM(S, n);
          sizeM(A, n);
          sizeM(Q, n);
          sizeM(U, n);
          sizeM(V, n);
           sizeM(Ainv, n);
          sizeM(C, n);
          sizeM(A\_point, n);
     This code is used in chunk 293.
     20.2.8 Destructor
275 \langle destructor IHO 275 \rangle \equiv
        IHO::\simIHO()
          delete matrix_inverse;
          delete trial_solution;
          delete solution;
          \mathbf{delete}[] C_{\bullet}pq;
          \mathbf{delete}[] \ C_{\bullet}qp;
     This code is used in chunk 293.
     20.2.9 Accepting a solution
276 \langle accept solution (IHO) 276 \rangle \equiv
        void IHO :: acceptSolution() {
          solution \rightarrow t = trial\_solution \rightarrow t;
           assign V (solution \neg y, trial\_solution \neg y);
           assign V (solution \neg u, trial\_solution \neg u);
```

```
assignM(solution \neg S, trial\_solution \neg S);
            assign V (solution \neg alpha, trial\_solution \neg alpha);
            assignM(solution \rightarrow A, trial\_solution \rightarrow A);
            assign V (solution \neg r, trial\_solution \neg r);
            assignM(solution \neg Q, trial\_solution \neg Q);
            assign V (solution \neg rQR, trial\_solution \neg rQR);
     This code is used in chunk 293.
     20.2.10 Set and get functions
277 \langle set and get functions 277 \rangle \equiv
        void set(Control *c, HOE *hoem, AD *ad0)
            assert(c \land hoem \land ad\theta);
            control = c;
           hoe = hoem;
            ad = ad\theta;
        void getTightEnclosure(iVector &y) const
           y = solution \rightarrow y;
        void init(const interval \&t, const iVector \&y)
            trial\_solution \rightarrow init(t, y);
           solution \neg init(t, y);
        const iVector &getGlobalExcess() const
           return globalExcess;
     This code is used in chunk 235.
      20.2.11 Constants
      Computing c_i^{p,q}
      We show how to compute
             c_i^{p,q} = \frac{p!}{(p+q)!} \frac{(q+p-i)!}{(p-i)!}.
     From
            c_{i-1}^{p,q} = \frac{p!}{(p+q)!} \frac{(q+p-i+1)!}{(p-i+1)!} = \frac{p!}{(p+q)!} \frac{(q+p-i)!}{(p-i)!} \frac{q+p-i+1}{p-i+1}
                  = \frac{q+p-i+1}{p-i+1}c_i^{p,q}
```

we compute

This code is used in chunk 293.

# Computing $(-1)^i c_i^{q,p}$

When computing  $c_i^{q,p}$ , we multiply them by  $(-1)^i$ , that is  $(-1)^i c_i^{q,p}$ .

$$c_{i-1}^{q,p} = \frac{q!}{(q+p)!} \frac{(p+q-i+1)!}{(q-i+1)!} = \frac{q!}{(q+p)!} \frac{(p+q-i)!}{(q-i)!} \frac{p+q-i+1}{q-i+1}$$
$$= \frac{p+q-i+1}{q-i+1} c_i^{q,p}$$

we compute

$$(-1)^i c_i^{q,p} = (-1) \frac{q-i+1}{p+q-i+1} \left( (-1)^{i-1} c_{i-1}^{q,p} \right), \text{ where } c_0^{q,p} = 1.$$

$$\begin{array}{l} 280 \quad \langle \, (-1)^i c_i^{q,p} = (-1)^i \frac{q!}{(p+q)!} \frac{(q+p-i)!}{(q-i)!} \, \, 280 \, \rangle \equiv \\ \mathbf{void} \ \mathbf{IHO} :: comp Cqp \, (\mathbf{int} \ p, \mathbf{int} \ q) \\ \{ \quad /* \\ \qquad \qquad C_- qp \, [i] \ni (-1)^i c_i^{q,p} \quad \text{ for } i=1,\ldots,q \\ \\ \overset{*/}{\mathbf{int}} \ tmp = q+p+1; \\ C_- qp \, [0] = 1.0; \\ \mathbf{for} \ (\mathbf{int} \ i=1; \ i \leq q; \ i++) \\ C_- qp \, [i] = (-C_- qp \, [i-1] * \mathbf{double} (q-i+1)) / \mathbf{double} (tmp-i); \\ \} \end{array}$$

This code is used in chunk 293.

#### Even number

This function returns *true* if its arguments  $(k \ge 0)$  is even and *false* otherwise. We check if the last bit of k is 1.

```
281 \langle check if a number is even 281 \rangle \equiv inline bool isEven(unsigned\ int\ k) \{ if (k \equiv 0)\ return\ true; if (k \& \#01)\ return\ false; return true; \}
```

This code is used in chunk 293.

# Computing $(-1)^q q! p! / (p+q)!$

For given p and q, compErrorConst computes  $(-1)^q q! p! / (p+q)!$ . We use the relation

$$\frac{q!p!}{(p+q)!} = \frac{p!}{(q+1)(q+2)\cdots(p+q)} = \frac{1}{q+1}\frac{2}{q+2}\cdots\frac{p}{q+p}.$$

```
282 \langle \operatorname{compute} (-1)^q q! p! / (p+q)! \ 282 \rangle \equiv /*
\operatorname{err\_const} \supseteq (-1)^q q! p! / (p+q)!
*/
\operatorname{interval} \ \mathbf{IHO} :: \operatorname{compErrorConstant}(\mathbf{int} \ p, \mathbf{int} \ q)
\{ \\ \operatorname{interval} \ \operatorname{err\_const}(1.0); \\ \operatorname{for} \ (\mathbf{int} \ i=1; \ i \leq p; \ i++) \ \{ \\ \operatorname{err\_const} \ *= \mathbf{double}(i); \\ \operatorname{err\_const} \ /= (q+i); \\ \} \\ \operatorname{if} \ (\neg isEven(q)) \ \operatorname{err\_const} = -\operatorname{err\_const}; \\ \operatorname{return} \ \operatorname{err\_const};
```

This code is used in chunk 293.

#### 20.2.12 Sorting columns of a matrix

Given an  $n \times n$  point matrix A, we sort its columns such that, in the 2-norm, they are in non-increasing order. In sortColumns the columns of the input matrix A are sorted, and the result is stored in B.

```
284 \langle sort columns 284\rangle \equiv
       void sortColumns(pMatrix \&B, const pMatrix \&A)
          \langle compute column norms of A 286\rangle
           (check if sorting is needed 287)
          (perform sorting 289)
     This code is used in chunk 291.
       First, we create a structure to store the index and the norm of the column corre-
     sponding to this index.
285 \langle \text{index-norm structure 285} \rangle \equiv
       struct index_norm {
          int index;
          double norm;
     This code is used in chunk 291.
       We store in an array b in b[i]. norm the 2-norm of column i of A.
286 \langle compute column norms of A 286 \rangle \equiv
       int n = \mathbf{v\_blas} :: sizeM(B);
       index\_norm *b = new index\_norm[n];
       pVector tmp;
       \mathbf{v}_{\mathbf{blas}} :: size V(tmp, n);
       for (int j = 0; j < n; j +++) {
          b[j].index = j;
          getColumn(tmp, A, j);
          b[j].norm = \mathbf{v\_blas} :: norm2(tmp);
     This code is used in chunk 284.
287 \langle check if sorting is needed 287\rangle \equiv
       bool sorting\_needed = false;
       for (int i = 0; i < n - 1; i + +) {
          if (b[i].norm < b[i+1].norm) {
             sorting\_needed = true;
             break;
     This code is used in chunk 284.
       We use the standard gsort function. We need a compare function for it.
288 \langle compare function 288 \rangle \equiv
```

```
inline int v_compare (const void *a1, const void *b1)
         index\_norm *a = (index\_norm *) a1;
         index\_norm *b = (index\_norm *) b1;
         if (a \rightarrow norm > b \rightarrow norm) return -1;
         if (a \rightarrow norm < b \rightarrow norm) return 1;
         return 0:
    This code is used in chunk 291.
       We sort the b array based on b[i]. norm. Then the jth column of B is the b[j]. index
    column of A.
289 \langle \text{ perform sorting } 289 \rangle \equiv
       if (sorting_needed) std::qsort((void *) b, n, sizeof(index_norm), v_compare);
       for (int j = 0; j < n; j ++)
          getColumn(tmp, A, b[j].index);
          setColumn(B, tmp, j);
       \mathbf{delete}[] b;
    This code is used in chunk 284.
    Files
291
       \langle \text{sortcolumns.cc} 291 \rangle \equiv
     #include <cstdlib>
     #include <cassert>
     #include "vnodeinterval.h"
     #include "basiclinalg.h"
       namespace vnodelp {
          using namespace v_bias;
          using namespace v_blas;
          (index-norm structure 285)
          (compare function 288)
          ⟨sort columns 284⟩
292 \langle iho.h 292 \rangle \equiv
     #ifndef IHO_H
     #define IHO_H
     #include <cassert>
     #include <cmath>
     #include "ad_ode.h"
     #include "ad_var.h"
```

```
#include "hoe.h"
     #include "intvfuncs.h"
     #include "basiclinalg.h"
     #include "matrixinverse.h"
       namespace vnodelp {
          using namespace v_bias;
          using namespace v_blas;
          ⟨ class IHO 235 ⟩
     #endif
293 \langle iho.cc 293 \rangle \equiv
     #include <cmath>
     #include "vnodeinterval.h"
     #include "basiclinalg.h"
     #include "solution.h"
     #include "control.h"
     #include "allad.h"
     #include "matrixinverse.h"
     #include "iho.h"
     #include "debug.h"
     #include "miscfuns.h"
       namespace vnodelp {
       using namespace v_bias;
       using namespace v_blas;
       extern void sortColumns(pMatrix \&B, const pMatrix \&A);
       extern bool computeQR(\mathbf{pMatrix} \& B, \mathbf{const} \ \mathbf{pMatrix} \ \& A);
       ⟨ constructor IHO 274 ⟩
       ⟨ destructor IHO 275⟩
       (check if a number is even 281)
       \langle c_i^{p,q} = \frac{p!}{(p+q)!} \frac{(q+p-i)!}{(p-i)!} 279 \rangle
       \langle\, (-1)^i c_i^{q,p} = (-1)^i \tfrac{q!}{(p+q)!} \tfrac{(q+p-i)!}{(q-i)!} \,\, 280\, \rangle
       \langle \text{ compute } (-1)^q q! p! / (p+q)! \ 282 \rangle
       (accept solution (IHO) 276)
        (compute tight enclosure 236)
       ⟨ compute IHO method coefficients 239 ⟩}
```

# Chapter 21

# The VNODE class

# 21.1 Declaration

```
295 \langle class VNODE 295 \rangle \equiv
       typedef enum {
          on, off
       } stepAction;
       class VNODE \{
       public:
         int steps;
          VNODE(AD *ad);
          void integrate(interval \&t\theta, iVector \&y\theta, const interval \&t\_end);
          ⟨set VNODE parameters 331⟩
          ⟨get functions (VNODE) 329⟩
          \simVNODE();
       private:
          void acceptSolution(interval \&t\theta, iVector \&y\theta);
          double compHstart(\mathbf{const}\ \mathbf{interval}\ \&t\theta, \mathbf{const}\ \mathbf{iVector}\ \&y\theta);
       private:
          int direction;
          interval t_trial, Tj, h_accepted;
          iVector temp;
          double h_start, h_min;
          pVector tp;
          Control *control;
         HOE *hoe;
         IHO *iho;
          AD *ad;
     This code is used in chunk 333.
```

# 21.2 The integrator function

The integrator function consists of the following steps:

- 1. check input correctness and determine the direction of the integration
- 2. initialize
- 3. validate existence and uniqueness and select stepsize
- 4. check if the end point is reached
- 5. compute a tight enclosure
- 6. decide how to proceed

#### This code is used in chunk 334.

#### 21.2.1 Input correctness

```
control-ind must not be failure

298 ⟨check input correctness 298⟩ ≡
    if (control-ind ≡ failure) {
        vnodeMessage("Previous_call_to_integrate()_failed");
        vnodeMessage("Call_setFirstEntry()_before_calling_integrate");
        return;
    }
}
```

See also chunks 299, 300, 301, 302, 303, and 304.

This code is used in chunk 296.

```
The initial condition and the final points must be representable machine intervals.
299 \langle check input correctness 298\rangle + \equiv
        if (\neg \mathbf{v}\_\mathbf{bias} :: finite\_interval(t\theta) \lor \neg \mathbf{v}\_\mathbf{bias} :: finite\_interval(t\_end)) {
            vnodeMessage("t0\(\text{\lambda}\)and\(\text{\lambda}\)t\(\text{\lambda}\)end\(\text{\lambda}\)must\(\text{\lambda}\)be\(\text{\lambda}\)finite");
            control \rightarrow ind = failure;
           return;
        for (unsigned int i = 0; i < size V(y\theta); i++)
           if (\neg \mathbf{v}\_\mathbf{bias} :: finite\_interval(y\theta[i])) {
              vnodeMessage("y0\u00cdmust\u00cdcontain\u00cdfinite\u00cdintervals");
              control \rightarrow ind = failure;
              return;
        If the integrator is re-entered with the same end point, t_end, as the t_end from
      the most recent call, then it should return.
300 \langle check input correctness 298\rangle + \equiv
        t\_trial = t\theta;
        if (t\_trial \equiv t\_end \land control \neg ind \neq first\_entry)  {
            vnodeMessage("Set_different_t_end");
           return;
        At least one of the tolerances must be positive.
301 \langle check input correctness 298\rangle + \equiv
        if (control \neg atol \leq 0 \land control \neg rtol \leq 0) {
            vnodeMessage ("Set_nonnegative_tolerances_and_at_least\
                 □one□positive□tolerance");
           return;
         }
        The order must be between 3 and qetMaxOrder().
302 \langle check input correctness 298\rangle + \equiv
        if (control \neg order < 3 \mid control \neg order > getMaxOrder()) {
            vnodeMessage("Set_order_>=_3_and_order_<=_MAX_ORDER");</pre>
            control \rightarrow ind = failure;
           return;
         }
        The value for the minimum stepsize must be non-negative
303 \langle check input correctness 298\rangle + \equiv
        if (control \rightarrow hmin < 0) {
            vnodeMessage("Set \_minimum \_stepsize \_>= \_0");
           return;
```

The initial and final time intervals may not intersect.

```
304 \langle \text{check input correctness 298} \rangle +\equiv  if (\neg \mathbf{v\_bias} :: disjoint(t0, t\_end)) { control \neg ind = failure; vnodeMessage("t0\_and\_t\_end\_must\_be\_disjoint"); return; }
```

#### 21.2.2 Determine direction

The user provides  $t_0$  and  $t_{end}$ . The solver decides if the integration is from left to right or right to left.

If a first entry into *integrate*, we initialize *direction* with 0. We also save the direction from the most recent integration.

```
305 ⟨determine direction 305⟩ ≡
    if (control¬ind ≡ first\_entry) direction = 0;
    int last_direction = direction;
    See also chunks 306 and 307.
    This code is used in chunk 296.
```

- 1. If  $\overline{t}_0 < \underline{t}_{end}$ , the integration is in positive direction.
- 2. If  $\underline{t}_0 > \overline{t}_{end}$ , the integration is in negative direction.

```
306 \langle \text{ determine direction } 305 \rangle + \equiv direction = 1;

if (\mathbf{v\_bias} :: sup(t\_end) < \mathbf{v\_bias} :: inf(t0))

direction = -1;
```

If the direction of the integration is reversed without calling first setFirstEntry(), then print a message and return.

```
307 ⟨determine direction 305⟩ +≡

if (last_direction ≠ 0 ∧ last_direction ≡ -direction) {

vnodeMessage("Integration_direction_changed_without_r\
esetting.\n""_\L\L\L\Call_setFirstEntry()_before_integrating.");

control¬ind = failure;
return;
}
```

## 21.2.3 Initialization

Initializing an integration includes the following parts discussed below.

```
308 \langle initialize integration 308 \rangle \equiv steps = 0; \langle find minimum stepsize 309 \rangle \langle find initial stepsize 310 \rangle \langle set IHO method 312 \rangle \langle set HOE method 311 \rangle This code is used in chunk 296.
```

#### Find minimum stepsize

- If  $control \rightarrow hmin \equiv 0$ , the solver computes minimum stepsize.
- If  $contol \neg ind \equiv success$ , this is a re-entry to the integrator; it computes minimum stepsize.
- If control- $hmin > 0 \land contol$ - $ind ≡ first\_entry$ , the user has specified minimum stepsize. We still have to check if this value is not smaller than what the solver finds as a minimum stepsize.

```
309 ⟨ find minimum stepsize 309⟩ ≡
    h_min = control→hmin;
if (control→hmin ≡ 0 ∨ control→ind ≡ success)
    control→hmin = compHmin(t0, t_end);
if (control→hmin > 0 ∧ control→ind ≡ first_entry) {
    double h = compHmin(t0, t_end);
    if (control→hmin < h) {
        control→ind = failure;
        vnodeMessage("Set_larger_value_for_hmin");
        return;
    }
    assert(control→hmin > 0);
}
```

# Find initial stepsize

If  $control \rightarrow ind \equiv first\_entry$ , the solver computes initial stepsize.

```
310 \langle find initial stepsize 310 \rangle \equiv

if (control \neg ind \equiv first\_entry) {

h\_start = compHstart(t0, y0);
}

if (h\_start < control \neg hmin) {

control \neg ind = failure;

vnodeMessage ("Minimum_stepsize_reached.");

return;
}
```

```
if (direction \equiv -1) {
   if (control \rightarrow ind \equiv first\_entry) h\_start = -h\_start;
```

This code is used in chunk 308.

#### Setting the HOE method

We set in the HOE method the value for the initial stepsize, order, control structure, and an AD object.

```
311 \langle set HOE method 311 \rangle \equiv
           hoe \rightarrow setTrialStepsize(h\_start);
           hoe \rightarrow setTrialOrder(control \rightarrow order);
           hoe \rightarrow set(control, ad);
           hoe \rightarrow init(t\theta, y\theta);
       This code is used in chunk 308.
```

#### Setting the IHO method

In the IHO method, we set the control object, the HOE method, and an AD object. We also compute the coefficients of the method and set the initial point.

```
312 \langle set IHO method 312 \rangle \equiv
          iho \rightarrow set(control, hoe, ad);
          iho \neg compCoeffs();
          iho \rightarrow init(t\theta, y\theta);
       This code is used in chunk 308.
```

#### 21.2.4 Methods involved in the initialization

#### Minimum stepsize

We determine the minimum magnitude,  $h_{\min}$ , for the stepsize allowed as follows:

```
\hat{t} = \max\{|\boldsymbol{t}_0|, |\boldsymbol{t}_{\text{end}}|\},
h_{\min} = \max \{ \triangle (\operatorname{next}(\widehat{t}) - \widehat{t}), w(\boldsymbol{t}_{\text{end}}) \}.
```

```
Here next(a) returns the next representable machine number to the right of a.
314 \langle \text{ compute } h_{\min} | 314 \rangle \equiv
            double compHmin(const interval &t0, const interval &t_end)
                                     t\theta = t_0
                                t_{-}end = t_{\mathrm{end}}
                                        \overline{t} = \max\{|\boldsymbol{t}_0|, |\boldsymbol{t}_{\mathrm{end}}|\}
                               t_{-}next = next machine number to the right of t
                                        t = \triangle(\operatorname{next}(\widehat{t}) - \widehat{t})
                                        t = \max \left\{ \triangle \left( \operatorname{next}(\widehat{t}) - \widehat{t} \right), \operatorname{w}(\boldsymbol{t}_{\text{end}}) \right\}
```

```
double t = \mathbf{std} :: max(\mathbf{v\_bias} :: mag(t\theta), \mathbf{v\_bias} :: mag(t\_end));
              double t\_next = nextafter(t, t + 1);
              v_bias :: round_up();
              t = t\_next - t;
              t = \mathbf{std} :: max(t, \mathbf{v\_bias} :: width(t\_end));
              return t;
       This code is used in chunk 334.
       Initial stepsize
       On the first step we compute
               h_{0,0} = \left(\frac{\text{tol}}{\|(k+1)f^{[k+1]}(\boldsymbol{y}_0)\|}\right)^{1/k},
       (see [26]) where
               tol = rtol \cdot ||\boldsymbol{y}_0|| + atol.
315 \langle compute initial stepsize 315 \rangle \equiv
          double VNODE::compHstart(const interval &t\theta, const iVector &y\theta)
                                           y\theta = \boldsymbol{y}_0
                                             k order
                          tayl_coeff_ode contains f^{[i]}(\boldsymbol{t}_0,\boldsymbol{y}_0) for i=0,1,\ldots,k+1
                                                       after compTerm is called
                                       temp \supseteq f^{[k+1]}(\boldsymbol{t}_0, \boldsymbol{y}_0)
                                           tol = tol = rtol \cdot ||\boldsymbol{y}_0|| + atol
                                             t = (k+1)||f^{[k+1]}(\boldsymbol{t}_0, \boldsymbol{y}_0)||
                                             t = \frac{\operatorname{tol}}{(k+1)\|f^{[k+1]}(\boldsymbol{t}_0,\boldsymbol{y}_0)\|}
                                             t = \left(\frac{\operatorname{tol}}{(k+1)\|f^{[k+1]}(\boldsymbol{t}_0, \boldsymbol{y}_0)\|}\right)^{1/k}
              int k = control \neg order;
              ad \rightarrow tayl\_coeff\_ode \rightarrow set(t0, y0, 1, k + 1);
              ad \rightarrow tayl\_coeff\_ode \rightarrow compTerms();
              ad \rightarrow tayl\_coeff\_ode \rightarrow getTerm(temp, k + 1);
              double tol = control \neg rtol * inf\_norm V(y0) + control \neg atol;
              double t = (k+1) * inf\_norm V(temp);
```

```
 \begin{array}{l} \langle \operatorname{check} t \ 316 \rangle \\ t = tol/t; \\ t = \operatorname{std} :: pow(t, 1.0/k); \\ \operatorname{return} t; \\ \} \end{array}
```

This code is used in chunk 334.

The t that we compute above is normally t > 0. To prevent the case that it might be zero, we set t to be the largest of  $control \neg atol$  and  $control \neg rtol$ .

```
316 \langle \text{check } t \text{ 316} \rangle \equiv  if (t \equiv 0) t = \text{std} :: max(control \rightarrow atol, control \rightarrow rtol); This code is used in chunk 315.
```

## 21.2.5 Validate existence and uniqueness

We try to validate existence and uniqueness with  $t_j$  and  $y_j$ . If validate returns false, we set control-ind = failure.

```
317 \langle \text{validate and select stepsize } 317 \rangle \equiv /*
t0 = t_j
y0 = y_j
*/
bool info;
hoe \sim compAprioriEnclosure(t0, y0, info);
if (info \equiv false) \ \{
control \sim ind = failure;
vnodeMessage("Could_not_validate_solution");
return;
```

#### 21.2.6 Check last step

This code is used in chunk 296.

If validate returns successfully, we obtain an interval  $[t_j, t_{j+1}]$ , or  $[t_{j+1}, t_j]$  if the integration is backwards, over which we have verified existence and uniqueness. Denote this interval by  $T_j$ .

We want to prevent the solver from taking a very small last stepsize. To accomplish this, we adapt the idea for taking a last step from [13]. We consider three cases:

- 1.  $t_{\text{end}} \subseteq T_j$
- 2.  $\mathbf{t}_{\mathrm{end}} \cap \mathbf{T}_j = \emptyset$
- 3.  $t_{j+1} \in t_{\text{end}}$

```
318 \langle check if last step 318 \rangle \equiv
                                                 /*
                          Tj \supseteq T_j
                       tend = t_{end}
                */
          Tj = hoe \rightarrow getTrialT();
         if (\mathbf{v\_bias}:: subseteq(t\_end, Tj)) {
             (case 1 320)
         else {
            if (\mathbf{v\_bias} :: disjoint(t\_end, Tj)) {
                \langle \text{ case 2 } 321 \rangle
            else {
                interval tmp;
                if (direction \equiv 1) \ tmp = \mathbf{v\_bias} :: sup(Tj);
                else tmp = inf(Tj); /* tmp \ni t_{j+1} */
                if (\mathbf{v\_bias} :: subseteq(tmp, t\_end)) {
                   \langle \text{ case 3 323} \rangle
                else assert(0);
```

This code is used in chunk 296.

Case 1:  $t_{end} \subseteq T_j$ . We set  $t_{j+1} = t_{end}$ ; Figure 21.1.

 $320 \langle case 1 320 \rangle \equiv$  $t\_trial = t\_end;$ /\* t\_end is inside Tj \*/ This code is used in chunk 318.

Case 2:  $t_{end} \cap T_i = \emptyset$ . Denote

$$\Theta = |\boldsymbol{t}_{\mathrm{end}} - t_j| = \max\{|\underline{\boldsymbol{t}}_{\mathrm{end}} - t_j|, |\overline{\boldsymbol{t}}_{\mathrm{end}} - t_j|\}.$$

- (i) If  $w(T_j) < \Theta/2$ , then the next point we chose is the already selected  $t_{j+1}$ .
- (ii) Otherwise,  $w(T_j) \ge \Theta/2$ ; see Figure 21.2.

We compute

$$T_{j}^{*} = \begin{cases} T_{j} \cap (t_{j} + [0, \Theta/2]) & \text{if } t_{j} < t_{j+1} \\ T_{j} \cap (t_{j} - [0, \Theta/2]) & \text{if } t_{j} > t_{j+1} \end{cases}$$
(21.1)

and select for the next step

$$t_{j+1} \leftarrow \begin{cases} \overline{T}_{j+1}^* & \text{if } t_j < t_{j+1} \\ \underline{T}_{j+1}^* & \text{if } t_j > t_{j+1} \end{cases}$$
 (21.2)

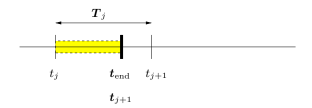


Figure 21.1. The case  $t_{end} \subseteq T_j$ . We set  $t_{j+1} = t_{tend}$ .

Case 3:  $t_{j+1} \in t_{end}$ . We expect this situation to occur very rarely. We set

$$t_{j+1} \leftarrow \begin{cases} \underline{t}_{\text{end}} & \text{if } t_j < t_{j+1} \\ \overline{t}_{\text{tend}} & \text{if } t_j > t_{j+1}. \end{cases}$$

```
323 \langle \text{case 3 323} \rangle \equiv

if (direction \equiv 1) t\_trial = \mathbf{v\_bias} :: inf(t\_end);

else t\_trial = \mathbf{v\_bias} :: sup(t\_end);

This code is used in chunk 318.
```

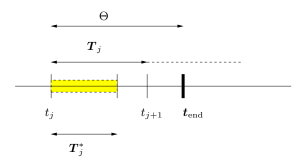


Figure 21.2. When close to  $t_{end}$ , we take the "middle" as the next integration point.

#### 21.2.7 Compute a tight enclosure

```
To compute a tight enclosure at t_{j+1}, we call 324 \langle compute enclosure 324\rangle \equiv iho \neg compTightEnclosure(t\_trial); This code is used in chunk 296.
```

#### 21.2.8 Decide

We decide how to proceed as shown below.

- 1. If  $control \neg ind \equiv first\_entry$  we set  $control \neg ind \equiv success$ .
- 2. If  $control \neg ind \equiv success$ , we consider two cases.
  - (a) If  $control interrupt \equiv no$ , we continue the integration.
  - (b) If  $control \rightarrow interrupt \equiv before\_accept$ , we return from the integration.
- 3. If  $control \neg ind \equiv failure$ , we return from the integration.

```
325 \langle \operatorname{decide} 325 \rangle \equiv

bool ret = false;

switch (\operatorname{control} - \operatorname{ind}) {

case \operatorname{first\_entry}:

\operatorname{control} - \operatorname{ind} = \operatorname{success};

case \operatorname{success}:

if (\operatorname{control} - \operatorname{interrupt} \equiv \operatorname{before\_accept}) ret = \operatorname{true};

break;

case \operatorname{failure}: return;
}

\operatorname{acceptSolution}(t0, y0);

if (\operatorname{ret}) return;

This code is used in chunk 296.
```

#### Accept solution

We accept the solution computed in the *hoe* and *iho* objects. We also update t0 and y0.

```
326 ⟨VNODE accept solution 326⟩ ≡
void VNODE :: acceptSolution(interval &tθ, iVector &yθ)

{
hoe¬acceptSolution();
iho¬acceptSolution();
h_accepted = t_trial - tθ;
tθ = t_trial;
iho¬getTightEnclosure(yθ);
steps++;
}
This code is used in chunk 334.
```

#### 21.3 Constructor/destructor

```
327 \langle \text{constructor (VNODE) } 327 \rangle \equiv
        VNODE::VNODE(AD *a)
           steps = 0;
           direction = 0;
          int n = a \rightarrow size;
           sizeV(temp, n);
           size V(tp, n);
           hoe = \mathbf{new} \ \mathbf{HOE}(n);
           iho = \mathbf{new} \ \mathbf{IHO}(n);
           control = new Control;
           assert(a);
           ad = a;
     This code is used in chunk 334.
328 \langle destructor (VNODE) 328 \rangle \equiv
        VNODE :: \sim VNODE()
           delete control;
           delete iho;
           delete hoe;
     This code is used in chunk 334.
```

#### 21.4 Get functions

To check if an integration is successful, we call

```
329 \langle \text{ get functions (VNODE) } 329 \rangle \equiv
       unsigned int getMaxOrder() const
         return ad→getMaxOrder();
       bool successful() const
         if (control \neg ind \equiv success \lor control \neg ind \equiv first\_entry) return true;
         return false;
     See also chunk 330.
    This code is used in chunk 295.
330 \langle \text{get functions (VNODE)} 329 \rangle + \equiv
       double getStepsize() const {
                                            /* Returns the last stepsize taken. */
         return midpoint (h_accepted);
       const iVector &getAprioriEncl() const {
             /* Obtains a reference to the a priori enclosure */
         return hoe→getApriori();
       const interval & getT() const { /* Returns T_i. */
         return hoe \neg getT();
       double getGlobalExcess() {
                                          /* Returns an estimate of the global excess. */
         width(tp, iho \neg getGlobalExcess());
         return inf\_norm V(tp);
       double getGlobalExcess(int i) {
             /* Returns an estimate of the global excess in component i. */
         width(tp, iho \neg getGlobalExcess());
         return tp[i];
       int getNoSteps() const {
             /* Returns the number of accepted steps during an integration. */
         return steps;
```

#### 21.5 Set parameters

We set integration parameters by the following functions.

```
331 \langle set VNODE parameters 331 \rangle \equiv
```

```
void setTols(\mathbf{double}\ a, \mathbf{double}\ r = 0) {
     /* Sets atol and rtol. By default, rtol is set to 0 */
  control \neg atol = a;
  control \neg rtol = r;
void setOrder(int p) { /* Sets order. */
  control \neg order = p;
void setHmin(double h) {
                                  /* Sets a value for the minimum stepsize. */
  control \rightarrow hmin = h;
void setOneStep(stepAction action) {
      /* Indicates an interrupt after each computed solution. */
  if (action \equiv on) control \neg interrupt = before\_accept;
  else
           /* off */
     control \neg interrupt = no;
                              /* Indicates first entry */
void setFirstEntry() {
  control \neg ind = first\_entry;
```

#### This code is used in chunk 295.

#### 21.6 **Files**

#### 21.6.1Interface

```
333 \langle \text{vnodeint.h} \quad 333 \rangle \equiv
    #ifndef VNODEINT_H
    #define VNODEINT_H
    #include <algorithm>
    #include "miscfuns.h"
    #include "vnodeinterval.h"
    #include "vnoderound.h"
    #include "vector_matrix.h"
    #include "ad_ode.h"
    #include "ad_var.h"
    #include "allad.h"
    #include "solution.h"
    #include "control.h"
    #include "matrixinverse.h"
    #include "iho.h"
    #include "hoe.h"
    #include "vtiming.h"
    #include "debug.h"
      namespace vnodelp {
```

```
\langle class VNODE 295 \rangle
    #endif
    21.6.2 Implementation
334 \langle integ.cc 334 \rangle \equiv
    #include <cmath>
    #include <algorithm>
    #include <ostream>
       using namespace std;
    #include "vnodeinterval.h"
    #include "vector_matrix.h"
    #include "matrixinverse.h"
    #include "basiclinalg.h"
    #include "vnodeint.h"
      namespace vnodelp {
         (constructor (VNODE) 327)
         ⟨ destructor (VNODE) 328⟩
         \langle \text{ compute } h_{\min} | 314 \rangle
         (compute initial stepsize 315)
         (integrator 296)
         ⟨VNODE accept solution 326⟩
```

#### 21.7 Interface to the VNODE-LP Package

```
The interface to the VNODE-LP package is stored in
335 \langle \text{vnode.h} \quad 335 \rangle \equiv
    \#ifndef VNODE_H
    #define VNODE_H
    #include <algorithm>
    #include "vnodeinterval.h"
    #include "vnoderound.h"
    #include "vector_matrix.h"
    #include "ad_ode.h"
    #include "ad_var.h"
    #include "allad.h"
    #include "vnodeint.h"
    #include "solution.h"
    #include "control.h"
    #include "iho.h"
    #include "hoe.h"
    #include "matrixinverse.h"
    #include "basiclinalg.h"
```

```
#include "fadbad_ad.h"
#include "fadbad_advar.h"
#include "fadbadad.h"
#endif
```

# Part V **AD Implementation**

#### Chapter 22

# Using FADBAD++

Currently, VNODE employs the FADBAD++ package [29].

#### 22.1 Computing ODE Taylor coefficients

#### 22.1.1 FadbadODE class

```
339 \langle class FadbadODE 339 \rangle \equiv
       typedef T(interval) Tinterval;
       typedef void(*Tfunction)(int n, Tinterval *yp, const Tinterval
            *y, Tinterval t, void *param);
       class FadbadODE : public AD_ODE {
       public:
         FadbadODE(int n, Tfunction, void *param = 0);
         void set(\mathbf{const}\ \mathbf{interval}\ \&t\theta,\mathbf{const}\ \mathbf{iVector}\ \&y\theta,\mathbf{const}\ \mathbf{interval}\ \&h,\mathbf{int}
              k);
         void compTerms();
         void sumTerms(iVector &sum, int m);
         void getTerm(iVector & term, int i) const;
         interval getStepsize() const;
         void eval(void *param);
         \simFadbadODE();
       private:
          Tfunction fcn;
         Tinterval *y\_coeff, *f\_coeff, t;
         int size;
         int order;
         interval stepsize;
    This code is used in chunk 349.
```

#### 22.1.2 Function description

In the constructor, we allocate the necessary memory, set the ODE problem, and generate the computational graph by calling fcn.

```
340 \langle \text{constructor (FadbadODE) } 340 \rangle \equiv
       FadbadODE::FadbadODE(int n, Tfunction f, void *param)
       : AD_ODE() {
          size = n;
          y\_coeff = \mathbf{new} \ \mathbf{Tinterval}[2*n];
          f\_coeff = y\_coeff + n;
          fcn = f;
          fcn(size, f\_coeff, y\_coeff, t, param);
     This code is used in chunk 350.
341 \langle FadbadODE destructor 341\rangle \equiv
       FadbadODE :: \sim FadbadODE()
          delete y\_coeff;
     This code is used in chunk 350.
342 (initialize Taylor coefficients (FadbadODE) 342) \equiv
       void FadbadODE::set(const\ interval\ \&t\theta, const\ iVector\ \&y\theta, const
                 interval &h, int k)
          t[0] = t\theta;
          t[1] = h;
          stepsize = h;
          order = k;
          for (int eqn = 0; eqn < size; eqn ++) y\_coeff[eqn][0] = y0[eqn];
     This code is used in chunk 350.
       The Taylor coefficients for the right side of y' = f(t, y) are evaluated by eval and
     stored in f-coeff. Then, the ith coefficient for y is h/i\times (the (i-1)st coefficient for f).
343 \langle compute Taylor coefficients (FadbadODE) 343 \rangle \equiv
       void FadbadODE :: compTerms()
          for (int eqn = 0; eqn < size; eqn +++)
            f_coeff [eqn].reset(); /* reset previously created terms */
          for (int coeff = 1; coeff \leq order; coeff ++)
                                                               /* compute coefficients */
            for (int eqn = 0; eqn < size; eqn ++) {
               f_{-}coeff[eqn].eval(coeff-1);
```

```
y\_coeff\left[\mathit{eqn}\right][\mathit{coeff}\right] = \mathit{stepsize} * f\_\mathit{coeff}\left[\mathit{eqn}\right][\mathit{coeff} - 1] / \mathbf{double}(\mathit{coeff});
     This code is used in chunk 350.
344
        \langle \text{sum terms (FadbadODE) } 344 \rangle \equiv
        void FadbadODE::sumTerms(iVector &sum,int m)
           interval s;
           for (int eqn = 0; eqn < size; eqn +++) {
              for (int coeff = m; coeff \ge 0; coeff --) s += y\_coeff[eqn][coeff];
              sum[eqn] = s;
        }
     This code is used in chunk 350.
        \langle \text{get term (FadbadODE) } 345 \rangle \equiv
345
        void FadbadODE::getTerm(iVector &term, int i) const
           for (int eqn = 0; eqn < size; eqn ++) term[eqn] = y\_coeff[eqn][i];
     This code is used in chunk 350.
346 \langle obtain stepsize 346 \rangle \equiv
        interval FadbadODE::getStepsize() const
           {\bf return}\ step size;
     This code is used in chunk 350.
        To evaluate the function and rebuild the computational graph, we do
347 \langle rebuild computational graph 347 \rangle \equiv
        void FadbadODE :: eval(void *param)
           \mathit{fcn}\left(\mathit{size}\,,\mathit{f\_coeff}\,,\,\mathit{y\_coeff}\,,t,\mathit{param}\,\right);
     This code is used in chunk 350.
      22.1.3 Files
349 \langle fadbad_ad.h 349 \rangle \equiv
      #ifndef Fadbad_ODE
      #define Fadbad_ODE
```

```
#include "vnodeinterval.h"
    #include "basiclinalg.h"
    #include "ad_ode.h"
    #include "ffadiff.h"
    #include "fadbad_intv.inc"
      namespace vnodelp {
         (class FadbadODE 339)
    #endif
350 \langle fadbad_ad.cc 350 \rangle \equiv
    #include "fadbad_ad.h"
      namespace vnodelp {
         \langle \text{constructor (FadbadODE)} 340 \rangle
         FadbadODE destructor 341
         (initialize Taylor coefficients (FadbadODE) 342)
         (compute Taylor coefficients (FadbadODE) 343)
         sum terms (FadbadODE) 344
         get term (FadbadODE) 345 \
         (obtain stepsize 346)
         (rebuild computational graph 347)
```

# 22.2 Computing Taylor coefficients for the variational equation

#### 22.2.1 FadbadVarODE class

```
352 \langle class FadbadVarODE 352 \rangle \equiv
       typedef T\langle F\langle interval \rangle \rangle TFinterval;
       typedef void(*TFfunction)(int n, TFinterval *yp, const TFinterval
           *y, TFinterval tf, void *param);
       class FadbadVarODE : public AD_VAR {
       public:
         FadbadVarODE(int n, TFfunction f, void *param = 0);
         void set (const interval & t\theta, const iVector & y\theta, const interval & h, int
             k);
         void compTerms();
         void sumTerms(iMatrix \& sum, int m);
         void getTerm(iMatrix &term, int i) const;
         void eval(void *param) { fcn(size, tf_out, tf_in, tf, param); }
         \simFadbadVarODE();
       private:
         TFinterval *tf_in, *tf_out, tf;
```

```
TFfunction fcn;
         int size;
         int order;
         interval stepsize;
       };
    This code is used in chunk 360.
    22.2.2 Function description
353 \langle constructor (FadbadVarODE) 353 \rangle \equiv
       FadbadVarODE::FadbadVarODE(int n, TFfunction f, void *param)
       {
          size = n;
          tf_in = new TFinterval[2 * n];
          tf\_out = tf\_in + n;
         fcn = f;
         fcn(size, tf\_out, tf\_in, tf, param);
    This code is used in chunk 361.
354 \langle destructor (FadbadVarODE) 354 \rangle \equiv
       FadbadVarODE :: \sim FadbadVarODE()
          delete[] tf_in;
    This code is used in chunk 361.
355 \langle initialize Taylor coefficients (FadbadVarODE) 355 \rangle \equiv
       void FadbadVarODE::set(const interval &t\theta, const iVector &y\theta, const
                 interval &h, int k)
          stepsize = h;
          order = k;
          tf[0].x() = t\theta;
          tf[1].x() = h;
         for (int eqn = 0; eqn < size; eqn +++) {
            tf_in[eqn][0] = y\theta[eqn];
    This code is used in chunk 361.
356 \langle compute Taylor coefficients (FadbadVarODE) 356 \rangle \equiv
       void FadbadVarODE :: compTerms()
```

```
for (int eqn = 0; eqn < size; eqn ++) tf\_out[eqn].reset();
         for (int eqn = 0; eqn < size; eqn ++) tf_in[eqn][0].diff(eqn, size);
         for (int coeff = 0; coeff < order; coeff ++) {
            for (int eqn = 0; eqn < size; eqn +++) {
              tf\_out[eqn].eval(coeff);
              tf\_in[eqn][coeff + 1] = stepsize * (tf\_out[eqn][coeff]/\mathbf{double}(coeff + 1));
    This code is used in chunk 361.
357 \langle \text{sum terms (FadbadVarODE) } 357 \rangle \equiv
       void FadbadVarODE :: sum Terms (iMatrix &Sum, int k)
         for (int row = 0; row < size; row ++)
            for (int col = 0; col < size; col ++) {
              interval s = 0.0;
              for (int coeff = k; coeff \ge 1; coeff --) s += tf_i in[row][coeff].d(col);
              Sum[row][col] = s;
         for (int row = 0; row < size; row ++) Sum[row][row] += 1.0;
    This code is used in chunk 361.
358
       \langle \text{get term (FadbadVarODE) } 358 \rangle \equiv
       void FadbadVarODE:: getTerm(iMatrix & Term, int i) const
         for (int row = 0; row < size; row ++)
            for (int col = 0; col < size; col ++) Term[row][col] = tf\_in[row][i].d(col);
    This code is used in chunk 361.
    22.3
               Files
360 \langle fadbad_advar.h | 360 \rangle \equiv
    #ifndef Fadbad_Var_ODE
    #define Fadbad_Var_ODE
    #include "vnodeinterval.h"
    #include "vector_matrix.h"
    #include "ad_var.h"
     #include "ffadiff.h"
     #include "fadbad_intv.inc"
       namespace vnodelp {
         using namespace v_bias;
         using namespace v_blas;
```

```
⟨ class FadbadVarODE 352 ⟩
    #endif
361 \langle fadbad_advar.cc 361 \rangle \equiv
    #include "fadbad_advar.h"
      namespace vnodelp {
         (constructor (FadbadVarODE) 353)
         ⟨ destructor (FadbadVarODE) 354 ⟩
         (initialize Taylor coefficients (FadbadVarODE) 355)
         (compute Taylor coefficients (FadbadVarODE) 356)
         ⟨sum terms (FadbadVarODE) 357⟩
         (get term (FadbadVarODE) 358)
      }
    22.4
              Encapsulated FADBAD++ AD
    Now we encapsulate all AD using FADBAD++ in
362 \langle \text{encapsulated FADBAD} + \text{AD } 362 \rangle \equiv
      class FADBAD_AD : public AD {
      public:
             FADBAD_AD(int n, Tfunction f, TFfunction tf)
        : AD(n, new FadbadODE(n, f), new FadbadVarODE(n, tf)),
             max\_order(MaxLength-2) \{ \}
        FADBAD_AD(int n, Tfunction f, TFfunction tf, void *p)
        : AD(n, new FadbadODE(n, f, p), new FadbadVarODE(n, tf, p)),
             max\_order(MaxLength - 2) \{ \}
         virtual int getMaxOrder() const {
           return max_order;
      private: const int max_order;
      };
    This code is used in chunk 363.
      and store the new class in
363 \langle fadbadad.h 363 \rangle \equiv
    #ifndef FADBADAD_H
    \#\mathbf{define}\ \mathtt{FADBADAD\_H}
    #include "fadbad_ad.h"
    #include "fadbad_advar.h"
    #include "allad.h"
      namespace vnodelp {
```

#### Appendix A

# **Miscellaneous Functions**

#### A.1 Vector output

```
We output a vector to the standard output by  \begin{array}{l} 366 \  \, \langle \, \mathrm{print} \,\, \mathrm{vector} \,\, 366 \, \rangle \equiv \\ \# \mathrm{include} \,\, \langle \, \mathrm{iostream} \rangle \\ \mathrm{using} \,\, \mathrm{namespace} \,\, \mathrm{std}; \\ \mathrm{template} \, \langle \, \mathrm{class} \,\, \mathbf{T} \, \rangle \,\, \mathrm{void} \,\, print \, Vector(\, \mathrm{const} \,\, \mathbf{T} \,\, \&v, \, \mathrm{const} \,\, \mathrm{char} \,\, *s = 0) \\ \big\{ \\ \mathrm{if} \,\, (s) \,\, cout \,\, \ll \, s \,\, \ll \,\, \parallel_{\square = \square} \,\, \parallel \,\, \ll \,\, endl; \\ \mathrm{for} \,\, (\, \mathrm{unsigned} \,\, \mathrm{int} \,\, i = 0; \,\, i < \, \mathbf{v} \,\, \mathrm{blas} \, :: \, size \, V(v); \,\, i + +) \,\,\, cout \,\, \ll \,\, v[i] \,\, \ll \,\, endl; \\ cout \,\, \ll \,\, endl; \\ \big\} \\ \mathrm{This} \,\, \mathrm{code} \,\, \mathrm{is} \,\, \mathrm{used} \,\, \mathrm{in} \,\, \mathrm{chunk} \,\, 137. \end{aligned}
```

#### A.2 Check if an interval is finite

```
367 ⟨check finite 367⟩ ≡
    namespace v_bias {
        inline bool finite_interval(const interval &a)
        {
            return (isfinite(inf(a)) ∧ isfinite(sup(a)));
        }
    }
    See also chunk 368.
    This code is used in chunk 374.

We check if an interval vector contains finite intervals by
368 ⟨check finite 367⟩ +≡
        namespace v_blas {
```

```
\label{eq:const_ivector} \begin{aligned} & \textbf{inline bool} \ \textit{finite\_interval}(\textbf{const iVector} \ \&a) \\ & \{ & \\ & \textbf{for (unsigned int } i=0; \ i < a.size(); \ i++) \\ & \textbf{if } (\neg \textbf{v\_bias} :: \textit{finite\_interval}(a[i])) \ \textbf{return} \ \textit{false}; \\ & \textbf{return} \ \textit{true}; \\ & \} \end{aligned}
```

#### A.3 Message printing

```
We would like to print various messages.
369 \pmod{\text{message printing } 369} \equiv
     #ifdef VNODE_DEBUG
     \#define printMessage(s)
          cerr \ll "\n_{\sqcup}***_{\sqcup}" \ll \__FILE\_\_":" \ll \__LINE\_\_ \ll "_{\sqcup\sqcup\sqcup}" \ll s \ll endl;
     \# \mathbf{else}
     \#define printMessage(s) (0)
     #endif
     #ifdef VNODE_DEBUG
     \#define exitOnError(s)
         printMessage(s);
         exit(-1);
     \#else
     \#define exitOnError(s) (0)
     #endif
     This code is used in chunk 372.
370 \langle VNODE-LP \text{ message } 370 \rangle \equiv
       void vnodeMessage(const char *s)
         This code is used in chunk 373.
```

#### A.4 Check intersection

```
371 ⟨check vector intersection 371⟩ ≡
void checkIntersection(const iVector &a, const iVector &b)
{
v_bias::interval ai, bi;
```

```
for (unsigned int i = 0; i < vnodelp :: sizeV(a); i++) {
           ai = a[i];
           bi = b[i];
           if (\mathbf{vnodelp} :: disjoint(ai, bi))
              cout \ll "i_{\square} = " \ll i \ll " = " \ll endl \ll ai \ll endl \ll bi;
    This code is used in chunk 373.
    Files
372 \langle debug.h 372 \rangle \equiv
     #ifndef DEBUG_H
     #define DEBUG_H
     #include "basiclinalg.h"
       (message printing 369)
       using namespace v_blas;
       void checkIntersection(const iVector &a, const iVector &b);
     #endif
373 \langle debug.cc 373 \rangle \equiv
     #include <ostream>
     #include <stdarg.h>
     #include <stdio.h>
     #include <stdlib.h>
     #include <iostream>
       using namespace std;
     #include "vnodeinterval.h"
     #include "basiclinalg.h"
       using namespace std; namespace vnodelp {
         using namespace v_bias;
         using namespace v_blas;
         ⟨ check vector intersection 371 ⟩
         ⟨VNODE-LP message 370⟩
374 \langle \mathtt{miscfuns.h} \ 374 \rangle \equiv
     #ifndef MISCFUNS_H
     \#define MISCFUNS_H
     #include <cmath>
     #include "basiclinalg.h"
       using namespace std; (check finite 367) namespace vnodelp {
              void vnodeMessage(const char *s);
     #endif
```

#### A.5 Timing

The getTime function returns the current user time. The getTotalTime subtracts the end time from the start time and returns the result.

```
375 \langle \text{vtiming.h} \quad 375 \rangle \equiv
    #ifndef VTIMING_H
    #define VTIMING_H
       double getTime();
       double getTotalTime(double start_time, double end_time);
    #endif
376 \langle \text{vtiming.cc} 376 \rangle \equiv
    #include <cassert>
    #include <sys/times.h>
    #include <unistd.h>
    #include <ctime>
    #include "vnodeinterval.h"
    #include "vnoderound.h"
       using namespace std;
       static struct tms Tms;
       double getTime()
         times(\& Tms);
         v_bias::round_nearest();
         long int ClockTcks = sysconf(\_SC\_CLK\_TCK);
         return (Tms.tms_utime)/(double(ClockTcks));
       double getTotalTime(double start_time, double end_time)
         assert(start\_time \leq end\_time);
         \mathbf{v}_{\mathbf{bias}} :: round\_nearest();
         return (end\_time - start\_time);
```

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 \langle \boldsymbol{S}_{j+1} = \boldsymbol{G}_{j+1} S_j | 252 \rangle Used in chunk 247.
 \langle \boldsymbol{y}_{j+1}^* = (\widehat{u}_{j+1} + \boldsymbol{z}_{j+1} + \boldsymbol{x}_{j+1}) \cap \widetilde{\boldsymbol{y}}_j 246 \rangle Used in chunk 242.
\langle \boldsymbol{B}_{j+1} = \sum_{i=0}^{q} (-1)^i c_i^{q,p} h_j^i J(f^{[i]}; \boldsymbol{y}_{j+1}^*) 249 \rangle Used in chunk 247.
\langle \boldsymbol{F}_j = \sum_{i=0}^p c_i^{p,q} h_i^i J(f^{[i]}; \boldsymbol{y}_j) 248 \rangle Used in chunk 247.
\langle \boldsymbol{U}_{j+1} = \boldsymbol{I} + \sum_{i=1}^{q} h_i^i J(f^{[i]}; \boldsymbol{y}_i) 244\rangle Used in chunk 242.
\langle \, \boldsymbol{d}_{j+1} = g_{j+1} + \boldsymbol{e}_{j+1} \, 260 \, \rangle Used in chunk 247. \langle \, \boldsymbol{e}_{j+1} = (-1)^q \gamma_{p,q} \, \boldsymbol{h}_j^{p+q+1} \, f^{[p+q+1]}(\boldsymbol{T}_j, \widetilde{\boldsymbol{y}}_j) \, 255 \, \rangle Used in chunk 247.
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 \langle v_{j+1} = y_{j+1}^* - u_{j+1} + (S_{j+1} - S_{j+1})\alpha + w_{j+1} 268 \rangle Used in chunk 265.
\langle \boldsymbol{w}_{j+1} = C_{j+1}^{-1} \boldsymbol{d}_{j+1} + (I - C_{j+1}^{-1} \boldsymbol{B}_{j+1}) (\boldsymbol{y}_{j+1}^* - \boldsymbol{y}_{j+1}^*) 261 \rangle Used in chunk 247. \langle \boldsymbol{y}_{j+1} = (\boldsymbol{y}_{j+1}^* + \boldsymbol{S}_{j+1} \boldsymbol{\alpha} + \boldsymbol{s}_{j+1} + \boldsymbol{w}_{j+1}) \cap \boldsymbol{y}_{j+1}^* 263 \rangle Used in chunk 247.
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