Adept C++ Software Library: User Guide

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If you have any queries about *Adept* that are not answered by this document or by the information on the *Adept* web site (http://www.met.reading.ac.uk/clouds/adept/) then please email me at r.j.hogan@ecmwf.int.

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Chapter 1

Introduction

1.1 What is Adept?

Adept (Automatic Differentiation using Expression Templates) is a C++ software library that enables algorithms to be automatically differentiated. Since version 2.0* it also provides array classes that can be used in array expressions. These two capabilities are fully integrated such that array expressions can be differentiated efficiently, but the array capability may also be used on its own.

The automatic-differentiation capability uses an operator overloading approach, so very little code modification is required. Differentiation can be performed in forward mode (the "tangent-linear" computation), reverse mode (the "adjoint" computation), or the full Jacobian matrix can be computed. This behaviour is common to several other libraries, namely ADOL-C (Griewank et al., 1996), CppAD (Bell, 2007) and Sacado (Gay, 2005), but the use of expression templates, an efficient way to store the differential information and several other optimizations mean that reverse-mode differentiation tends to be significantly faster and use less memory. In fact, *Adept* is also usually only a little slower than an adjoint code you might write by hand, but immeasurably faster in terms of user time; adjoint coding is very time consuming and error-prone. For technical details of how it works, benchmark results and further discussion of the factors affecting its speed when applied to a particular code, see Hogan (2014).

Expression templates also underpin a number of libraries that provide the capability to perform mathematical operations on entire arrays (Veldhuizen, 1995). Unfortunately, if *Adept* version 1.x and such an array library are used together, then the speed advantages of expression templates are lost, if indeed the libraries work together at all. Since version 2.0, *Adept* provides array classes that overcome this problem: its automatic differentiation and array capabilities are underpinned by a single unified expression template framework so that array expressions may be differentiated very efficiently. However, it should be stressed that *Adept* is useful as a fully functional array library even if you don't wish to use its automatic differentiation capability. *Adept* uses BLAS and LAPACK for matrix operations.

This user guide describes how to apply the *Adept* software library to your code, and many of the examples map on to those in the test directory of the *Adept* software package. Section 1.2 outlines how to install *Adept* on your system and how to compile your own code to use it. Chapter 2 describes how to use the automatic differentiation capability of the library, while chapter 3 describes how to use its array capability. Chapter 4 then describes general aspects such as exception handling, configuration options and license terms.

1.2 Installing Adept and compiling your code to use it

Adept is compatible with any C++98 compliant compiler, although most of the testing has been specifically on Linux with the GNU C++ compiler. The code is built with the help of a configure shell script generated by GNU autotools. If you are on a non-Unix system (e.g. Windows) and cannot use shell scripts, see section 1.2.2.

^{*}Note that the version 1.9.x series serve as beta releases for version 2.0 of Adept.

1.2.1 Unix-like platforms

On a Unix-like system, do the following:

- 1. Ensure you have a BLAS library installed, necessary for matrix multiplication. For the best performance in matrix operations it is recommended that you install an optimized package such as OpenBLAS[†] or ATLAS[‡]. If you have multiple BLAS libraries available on your system you can specify the one you want by calling the configure script below with --with-blas=openblas or similar.
- 2. Optionally install the LAPACK library, necessary for matrix inversion and solving linear systems of equations. If you do not install this then *Adept* will still compile but the functions inv and solve will fail at run time. Note that LAPACK relies on the underlying BLAS library for its speed.
- 3. The test and benchmarking programs can make use of additional libraries if available. If you also install any of the automatic differentiation tools ADOL-C, CppAD and/or Sacado then the benchmarking test program can compare them to *Adept*. One of the test programs uses the minimization algorithm from the GNU Scientific Library, if available, so you may wish to install that too.
- 4. Unpack the package (tar xvfz adept-2.x.tar.gz on Linux) and cd to the directory adept-2.x.
- 5. Configure the build using the configure script. The most basic method is to just run

```
./configure
```

More likely you will wish to compile with a higher level of optimization than the default (which is -02), achieved by setting the environment variable CXXFLAGS. You may also wish to specify the root directory of the installation, say to /foo. These may be done by running instead

```
./configure CXXFLAGS="-g -O3" --prefix=/foo
```

The -g option to CXXFLAGS ensures debugging information is stored. If a library you wish to use is installed in a non-system directory, say under /foo, then specify the locations as follows:

```
./configure CPPFLAGS="-I/foo/include" LDFLAGS="-L/foo/lib -Wl,-rpath,/foo/lib"
```

where the <code>-rpath</code> business is needed in order that the <code>Adept</code> shared library knows where to look for the libraries it is dependent on. If you have them then for the benchmarking program you can also add the non-system location of ADOL-C, CppAD and Sacado libraries with additional <code>-I</code> and <code>-L</code> arguments, but note that the <code>-rpath</code> argument is not needed in that case. You can see the more general options available by running <code>./configure --help</code>; for example, you can turn-off OpenMP parallelization in the computation of Jacobian matrices using <code>--disable-openmp</code>. See also section 4.5 for ways to make more fundamental changes to the configuration of <code>Adept</code>. The output from the <code>configure</code> script provides information on aspects of how <code>Adept</code> and the test programs will be built.

6. Build *Adept* by running

make

This will create the static and shared libraries in adept/.libs.

7. Install the header files and the static and shared libraries by running

```
make install
```

If this is to be installed to a system directory, you will need to log in as the super-user first, or run sudo make install on depending on your system.

8. Build the example and benchmarking programs by running

[†]OpenBLAS is available from http://www.openblas.net/.

[‡]ATLAS is available from http://math-atlas.sourceforge.net/.

make check

Note that this may be done without first installing the *Adept* library to a system directory. This compiles the following programs in the test directory:

- test_misc: the trivial example from Hogan (2014);
- test_adept: compares the results of numerical and automatic differentiation;
- test_with_without_ad: does the same but compiling the same source code both with and without automatic differentiation (see test/Makefile for how this is done),
- test_radiances: demonstrates the interfacing of *Adept* with code that provides its own Jacobian;
- test_gsl_interface: implementation of a simple minimization problem using the L-BFGS minimizer in the GSL library;
- test_checkpoint: demonstration of checkpointing, a useful technique for large codes;
- test_thread_safe: demonstration of the use of multiple OpenMP threads, each with its own instance of an *Adept* stack;
- test_no_lib: demonstrates the use of the adept_source.h header file that means there is no need to link to the *Adept* library in order to create an executable.
- test_arrays: tests many of the array capabilities described in chapter 3.
- test_fixed_arrays: tests the similar capabilities of arrays with fixed dimensions (known at compile time).

In the benchmark directory it compiles autodiff_benchmark for comparing the speed of the differentiation of two advection algorithms using *Adept*, ADOL-C, CppAD and Sacado (or whichever subset of these tools you have on your system). It also compiles animate for visualizing at a terminal what the algorithms are doing. Further information on running these programs can be found in the README files in the relevant directories. Note that despite the implication, "make check" does not automatically run any of the programs it makes to check they function correctly.

To compile source files that use the *Adept* library, you need to make sure that adept.h and adept_arrays.h are in your include path. If they are located in a directory that is not in the default include path, add something like -I/home/fred/include to the compiler command line. At the linking stage, add -ladept to the command line to tell the linker to look for the libadept.a static library, or equivalent shared library. If this file is in a non-standard location, also add something like -L/home/fred/lib -Wl, -rpath, /home/fred/lib before the -ladept argument to specify its location. Section 2.6.3 provdes an example Makefile for compiling code that uses the *Adept* library. Read on to see how you can compile an *Adept* application *without* needing to link to a library.

1.2.2 Non-Unix platforms, and compiling *Adept* applications without linking to an external library

WARNING: THIS FUNCTIONALITY IS BROKEN IN VERSION 1.9 BUT WILL BE FIXED WITH VERSION 2.0

Most of the difficulty in maintaining software that can compile on multiple platforms arises from the different ways of compiling software libraries, and the need to test on compilers that may be proprietary. Unfortunately I don't have the time to maintain versions of *Adept* that build specifically on Microsoft Windows or other non-Unix platforms. However, *Adept* is not a large library, so I have provided a very simple way to build an *Adept* application *without* the need to link to a pre-compiled *Adept* library. In one of your source files and one only, add this near the top:

Typically you would include this in the source file containing the main function. This header file is simply a concatenation of the *Adept* library source files, so when you compile a file that includes it, you compile in all the functionally of the *Adept* library. All other source files in your application should include the adept.h or adept_arrays.h header file as normal. When you link all your object files together to make an executable, the *Adept* functionality will be built in, even though you did not link to an external *Adept* library. A demonstration of this is in the test/test_no_lib.cpp source file, which needs to be compiled together with test/algorithm.cpp to make an executable. It is hoped that this feature will make it easy to use *Adept* on non-Unix platforms, although of course this feature works just as well on Unix-like platforms as well. If you want to use OpenBLAS on such platforms then you will still need to install that library in the normal way.

A further point to note is that, under the terms of the license, it is permitted to copy all the *Adept* include files, including adept_source.h, into an include directory in your software package and use it from there in both binary and source-code releases of your software. This means that users do not need to install *Adept* separately before they use your software. However, if you do this then remember that your use of these files must comply with the terms of the Apache License, Version 2.0; see section 4.7 for details.

Chapter 2

Using Adept for automatic differentiation

2.1 Introduction

This chapter describes how to use *Adept* to differentiate your code. For simplicity, none of the examples use array functionality described in the next chapter. *Adept* provides the following automatic-differentiation functionality:

Full Jacobian matrix Given the non-linear function $\mathbf{y} = f(\mathbf{x})$ relating vector \mathbf{y} to vector \mathbf{x} coded in C or C++, after a little code modification Adept can compute the Jacobian matrix $\mathbf{H} = \partial \mathbf{y}/\mathbf{x}$, where the element at row i and column j of \mathbf{H} is $H_{i,j} = \partial y_i/\partial x_j$. This matrix will be computed much more rapidly and accurately than if you simply recompute the function multiple times, each time perturbing a different element of \mathbf{x} by a small amount. The Jacobian matrix is used in the Gauss-Newton and Levenberg-Marquardt minimization algorithms.

Reverse-mode differentiation This is a key component in optimization problems where a non-linear function needs to be minimized but the state vector \mathbf{x} is too large for it to make sense to compute the full Jacobian matrix. Atmospheric data assimilation is the canonical example in the field of meteorology. Given a nonlinear function $J(\mathbf{x})$ relating the scalar to be minimized J to vector \mathbf{x} , Adept will compute the vector of adjoints $\partial J/\partial \mathbf{x}$. Moreover, for a component of the code that may be expressed as a multi-dimensional non-linear function $\mathbf{y} = f(\mathbf{x})$, Adept can compute $\partial J/\partial \mathbf{x}$ if it is provided with the vector of input adjoints $\partial J/\partial \mathbf{y}$. In this case, $\partial J/\partial \mathbf{x}$ is equal to the matrix-vector product $\mathbf{H}^T \partial J/\partial \mathbf{y}$, but it is computed here without computing the full Jacobian matrix \mathbf{H} . The vector $\partial J/\partial \mathbf{x}$ may then be used in a quasi-Newton minimization scheme (e.g., Liu and Nocedal, 1989).

Forward-mode differentiation Given the non-linear function $\mathbf{y} = f(\mathbf{x})$ and a vector of perturbations $\delta \mathbf{x}$, Adept will compute the corresponding vector $\delta \mathbf{y}$ arising from a linearization of the function f. Formally, $\delta \mathbf{y}$ is equal to the matrix-vector product $\mathbf{H}\delta \mathbf{x}$, but it is computed here without computing the full Jacobian matrix \mathbf{H} . Note that Adept is designed for the reverse case, so might not be as fast or economical in memory in the forward mode as libraries written especially for that purpose (although Hogan, 2014, showed that it was competitive).

Adept can automatically differentiate the following operators and functions:

- The standard binary mathematical operators +, -, \star and /.
- The assignment versions of these operators: +=, -=, *= and /=.
- The unary mathematical functions sqrt, exp, log, log10, sin, cos, tan, asin, acos, atan, sinh, cosh, tanh, abs, asinh, acosh, atanh, expm1, log1p, cbrt, erf, erfc, exp2 and log2.
- The binary functions pow, min and max.

Variables to take part in expressions to be differentiated have a special "active" type; such variables can take part in comparison operations ==, !=, >, <, >= and <=, as well as the diagnostic functions isfinite, isinf and isnan.

2.2. Code preparation 8

Note that at present Adept is missing some functionality that you may require:

• Differentiation is first-order only: it cannot directly compute higher-order derivatives such as the Hessian matrix (although one of the Frequently Asked Questions in section 4.6 describes how *Adept* can help compute the Hessian of a certain category of algorithms).

- It has limited support for complex numbers; no support for mathematical functions of complex numbers, and expressions involving operations (addition, subtraction, multiplication and division) on complex numbers are not optimized.
- It can be applied to C and C++ only; *Adept* could not be written in Fortran since the language provides no template capability.

It is hoped that future versions will remedy these limitations (and maybe even a future version of Fortran will support templates).

Section 2.2 describes how to prepare your code for automatic differentiation, and section 2.3 describes how to perform forward- and reverse-mode automatic differentiation on this code. Section 2.4 describes how to compute Jacobian matrices. Section 2.5 provides a detailed description of how to interface an algorithm implemented using *Adept* with a third-party minimization library. Section 2.6 describes how to call a function both with and without automatic differentiation from within the same program. Section 2.7 describes how to interface to software modules that compute their own Jacobians. Section 2.8 describes the user-oriented member functions of the Stack class that contains the differential information and section 2.9 describes the member functions of the "active" double-precision type adouble.

2.2 Code preparation

If you have used ADOL-C, CppAD or Sacado then you will already be familiar with what is involved in applying an operator-overloading automatic differentiation package to your code. The user interface to *Adept* differs from these only in the detail. It is assumed that you have an algorithm written in C or C++ that you wish to differentiate. This section deals with the modifications needed to your code, while section 2.3 describes the small additional amount of code you need to write to differentiate it.

In all source files containing code to be differentiated, you need to include the adept.h header file and import the adouble type from the adept namespace. Assuming your code uses double precision, you then search and replace double with the "active" equivalent adouble, but doing this only for those variables whose values depend on the independent input variables. Under the hood this type is an alias for Active<double>. The single-precision equivalent is afloat, an alias for Active<float>. Active and passive variables of single and double precision may be used together in the same expressions, but note that by default all differentiation is done in double precision.

If you wish to enable your code to be easily recompiled to use different precisions, then you may alternatively use the generic Real type from the adept namespace with its active equivalent aReal (an alias for Active<Real>). Section 4.5 describes how to redefine Real to represent single, double or quadruple precision. Automatic differentiation will be performed using the same precision as Real, but but be aware that if this is defined to be the same as a single-precision float, accumulation of round-off error can make the accuracy of derivatives insufficient for minimization algorithms. The examples in the remainder of this chapter use only double precision.

Consider the following contrived algorithm from Hogan (2014) that takes two inputs and returns one output:

```
double algorithm(const double x[2]) {
  double y = 4.0;
  double s = 2.0*x[0] + 3.0*x[1]*x[1];
  y *= sin(s);
  return y;
}
```

The modified code would look like this:

```
#include <adept.h>
using adept::adouble;
```

```
adouble algorithm(const adouble x[2]) {
  adouble y = 4.0;
  adouble s = 2.0*x[0] + 3.0*x[1]*x[1];
  y *= sin(s);
  return y;
}
```

Changes like this need to be done in all source files that form part of an algorithm to be differentiated.

If you need to access the real number underlying an adouble variable a, for example in order to use it as an argument to the fprintf function, then use a.value() or adept::value(a). Any mathematical operations performed on this real number will not be differentiated.

You may use adouble as the template argument of a Standard Template Library (STL) vector type (i.e. std::vector<adouble>), or indeed any container where you access individual elements one by one. For types allowing mathematical operations on the whole object, such as the STL complex and valarray types, you will find that although you can multiply one std::complex<adouble> or std::valarray<adouble> object by another, mathematical functions (exp, sin etc.) will not work when applied to whole objects, and neither will some simple operations such as multiplying these types by an ordinary (non-active) double variable. Moreover, the performance is not great because expressions cannot be fully optimized when in these containers. Therefore If you need array functionality then you should use the features described in chapter 3. It is hoped that a future version of *Adept* will include its own complex type.

2.3 Applying reverse-mode differentiation

Suppose you wanted to create a version of algorithm that returned not only the result but also the gradient of the result with respect to its inputs, you would do this:

```
#include <adept.h>
double algorithm_and_gradient(
                   const double x_val[2], // Input values
                   adept::Stack stack,
                                         // Where the derivative information is stored
                                        // Import adouble from adept
 using adept::adouble;
 adouble x[2] = {x_val[0], x_val[1]};  // Initialize active input variables
                            // Start recording
 stack.new_recording();
                                    // Call version overloaded for adouble args
 adouble y = algorithm(x);
                                       // Defines y as the objective function
                                      // Run the adjoint algorithm
// Store the fire
 y.set_gradient(1.0);
 stack.compute_adjoint();
 dy_dx[0] = x[0].get_gradient();
dy_dx[1] = x[1].get_gradient();
                                       // Store the first gradient
                                         // Store the second gradient
                                         // Return the result of the simple computation
 return y.value();
```

The component parts of this function are in a specific order, and if this order is violated then the code will not run correctly. The steps are now described.

2.3.1 Set-up stack to record derivative information

```
adept::Stack stack;
```

The Stack object is where the differential version of the algorithm will be stored. When initialized, it makes itself accessible to subsequent statements via a global variable, but using thread-local storage to ensure thread safety. It must be initialized before the first adouble object is instantiated and it must not go out of scope until the last adouble object is destructed. This is because adouble objects register themselves with the currently active stack, and deregister themselves when they are destroyed; if the same stack is not active throughout the lifetime of such adouble objects then the code will crash with a segmentation fault.

In the example here, the Stack object is local to the scope of the function. If another Stack object had been initialized by the calling function and so was active at the point of entry to the function, then the local Stack object

would throw an adept::stack_already_active exception (see Test 3 described at test/README in the *Adept* package if you want to use multiple Stack objects in the same program). A disadvantage of local Stack objects is that the memory it uses must be reallocated each time the function is called. This can be overcome in several ways:

- Declare the Stack object to be static, which means that it will persist between function calls. This has the disadvantage that you won't be able to use other Stack objects in the program without deactivating this one first (see Test 3 in the *Adept* package, referred to above, for how to do this).
- Initialize Stack at a higher level in the program. If you need access to the stack, you may either pass a reference to it to functions such as algorithm_and_gradient, or alternatively you can use the adept::active_stack() function to return a pointer to the currently active stack object.
- Put it in a class so that it is accessible to member functions; this approach is demonstrated in section 2.5.

2.3.2 Initialize independent variables and start recording

```
adouble x[2] = {x_val[0], x_val[1]};
stack.new_recording();
```

The first line here simply copies the input values to the algorithm into adouble variables. These are the *inde*pendent variables, but note that there is no obligation for these to be stored as one array (as in CppAD), and for forward- and reverse-mode automatic differentiation you do not need to tell *Adept* explicitly via a function call which variables are the independent ones. The next line clears all differential statements from the stack so that it is ready for a new recording of differential information. Note that the first line here actually stores two differential statements, $\delta x [0] = 0$ and $\delta x [1] = 0$, which are immediately cleared by the new_recording function call. To avoid the small overhead of storing redundant information on the stack, we could replace the first line with

```
x[0].set_value(x_val[0]);
x[1].set_value(x_val[1]);
or
adept::set_values(x, 2, x_val);
```

which have the effect of setting the values of x without storing the equivalent differential statements.

Previous users of *Adept* version 0.9 should note that since version 1.0, the new_recording function replaces the start function call, which had to be put *before* the independent variables were initialized. The problem with this was that the independent variables had to be initialized with the set_value or set_values functions, otherwise the gradients coming out of the automatic differentiation would all be zero. Since it was easy to forget this, new_recording was introduced to allow the independent variables to be assigned in the normal way using the assignment operator (=). But don't just replace start in your version-0.9-compatible code with new_recording; the latter must appear *after* the independent variables have been initialized.

2.3.3 Perform calculations to be differentiated

```
adouble y = algorithm(x);
```

The algorithm is called, and behind the scenes the equivalent differential statement for every mathematical statement is stored in the stack. The result of the forward calculation is stored in y, known as a dependent variable. This example has one dependent variable, but any number is allowed, and they could be returned in another way, e.g. by passing a non-constant array to algorithm that is filled with the final values when the function returns.

2.3.4 Perform reverse-mode differentiation

```
y.set_gradient(1.0);
stack.compute_adjoint();
```

The first line sets the initial gradient (or adjoint) of y. In this example, we want the output gradients to be the derivatives of y with respect to each of the independent variables; to achieve this, the initial gradient of y must be unity.

More generally, if y was only an intermediate value in the computation of objective function J, then for the outputs of the function to be the derivatives of J with respect to each of the independent variables, we would need to set the gradient of y to $\partial J/\partial y$. In the case of multiple intermediate values, a separate call to set_gradient is needed for each intermediate value. If y was an array of length n then the gradient of each element could be set to the values in a double array y_ad using

```
adept::set_gradients(y, n, y_ad);
```

The compute_adjoint() member function of stack performs the adjoint calculation, sweeping in reverse through the differential statements stored on the stack. Note that this must be preceded by at least one set_gradient or set_gradients call, since the first such call initializes the list of gradients for compute_adjoint() to act on. Otherwise, compute_adjoint() will throw a gradients_not_initialized exception.

2.3.5 Extract the final gradients

```
dy_dx[0] = x[0].get_gradient();
dy_dx[1] = x[1].get_gradient();
```

These lines simply extract the gradients of the objective function with respect to the two independent variables. Alternatively we could have extracted them simultaneously using

```
adept::get_gradients(x, 2, dy_dx);
```

To do forward-mode differentiation in this example would involve setting the initial gradients of x instead of y, calling the member function compute_tangent_linear() instead of compute_adjoint(), and extracting the final gradients from y instead of x.

2.4 Computing Jacobian matrices

Until now we have considered a function with two inputs and one output. Consider the following more general function whose declaration is

```
void algorithm2(int n, const adouble* x, int m, adouble* y);
```

where x points to the n independent (input) variables and y points to the m dependent (output) variables. The following function would return the full Jacobian matrix:

```
#include <vector>
#include <adept.h>
void algorithm2_jacobian(
                    int n,
                                            // Number of input values
                    const double* x_val, // Input values
                    int m.
                                            // Number of output values
                    double* y_val,
                                            // Output values
                    double* jac) {
                                           // Output Jacobian matrix
  using adept::adouble;
                                            // Import Stack and adouble from adept
  adept::Stack stack;
                                            // Where the derivative information is stored
 std::vector<adouble> x(n); // Vector of active input variable:set_values(&x[0], n, x_val); // Initialize adouble inputs
                                           // Vector of active input variables
  adept.new_recording();
                                            // Start recording
  std::vector<adouble> y(m);
                                            // Create vector of active output variables
  algorithm2(n, &x[0], m, &y[0]);
                                            // Run algorithm
  stack.independent(&x[0], n);
                                            // Identify independent variables
  stack.dependent(&y[0], m);
                                             // Identify dependent variables
  stack.jacobian(jac);
                                             // Compute & store Jacobian in jac
```

Note that:

• The independent member function of stack is used to identify the independent variables, i.e. the variables that the derivatives in the Jacobian matrix will be with respect to. In this example there are n independent variables located together in memory and so can be identified all at once. Multiple calls are possible to identify further independent variables. To identify a single independent variable, call independent with just one argument, the independent variable (not as a pointer).

- The dependent member function of stack identifies the dependent variables, and its usage is identical to independent.
- The memory provided to store the Jacobian matrix (pointed to by jac) must be a one-dimensional array of size m×n, where m is the number of dependent variables and n is the number of independent variables.
- The resulting matrix is stored in the sense of the index representing the dependent variables varying fastest (column-major order).
- Internally, the Jacobian calculation is performed by multiple forward or reverse passes, whichever would be faster (dependent on the numbers of independent and dependent variables).
- The use of std::vector<adouble> rather than new adouble[n] ensures no memory leaks in the case of an exception being thrown, since the memory associated with x and y will be automatically deallocated when they go out of scope.

2.5 Real-world usage: interfacing Adept to a minimization library

Suppose we want to find the vector \mathbf{x} that minimizes an objective function $J(\mathbf{x})$ that consists of a large algorithm coded using the Adept library and encapsulated within a C++ class. In this section we illustrate how it may be interfaced to a third-party minimization algorithm with a C-style interface, specifically the free one in the GNU Scientific Library. The full working version of this example, using the N-dimensional Rosenbrock banana function as the function to be minimized, is "Test 4" in the test directory of the Adept software package. The interface to the algorithm is as follows:

```
#include <vector>
#include <adept.h>
using adept::adouble;
class State {
 public:
   // Construct a state with n state variables
   State(int n) { active_x_.resize(n); x_.resize(n); }
   // Minimize the function, returning true if minimization successful, false otherwise
   bool minimize();
   // Get copy of state variables after minimization
   void x(std::vector<double>& x_out) const;
   // For input state variables x, compute the function J(x) and return it
   double calc_function_value(const double* x);
   // For input state variables x, compute function and put its gradient in dJ_dx
   double calc_function_value_and_gradient(const double* x, double* dJ_dx);
   // Return the size of the state vector
   unsigned int nx() const { return active_x_.size(); }
 protected:
   // Active version: the algorithm is contained in the definition of this function
   adouble calc_function_value(const adouble* x);
   // DATA
   adept::Stack stack_;
                                    // Adept stack object
   std::vector<adouble> active_x_; // Active state variables
};
```

The algorithm itself is contained in the definition of calc_function_value(const adouble*), which is implemented using adouble variables (following the rules in section 2.2). However, the public interface to the class uses only standard double types, so the use of Adept is hidden to users of the class. Of course, a complicated algorithm may be implemented in terms of multiple classes that do exchange data via adouble objects. We will be using a quasi-Newton minimization algorithm that calls the algorithm many times with trial vectors \mathbf{x} , and for each call may request not only the value of the function, but also its gradient with respect to \mathbf{x} . Thus the public interface provides calc_function_value(const double*) and calc_function_value_and_gradient, which could be implemented as follows:

```
double State::calc_function_value(const double* x) {
  for (unsigned int i = 0; i < nx(); ++i) active_x_[i] = x[i];
  stack_.new_recording();
  return value(calc_function_value(&active_x_[0]));
}

double State::calc_function_value_and_gradient(const double* x, double* dJ_dx) {
  for (unsigned int i = 0; i < nx(); ++i) active_x_[i] = x[i];
  stack_.new_recording();
  adouble J = calc_function_value(&active_x_[0]);
  J.set_gradient(1.0);
  stack_.compute_adjoint();
  adept::get_gradients(&active_x_[0], nx(), dJ_dx);
  return value(J);
}</pre>
```

The first function simply copies the double inputs into an adouble vector and runs the version of calc_function_value for adouble arguments. Obviously there is an inefficiency here in that gradients are recorded that are then not used, and this function would be typically 2.5–3 times slower than an implementation of the algorithm that did not store gradients. Section 2.6 describes three ways to overcome this problem. The second function above implements reverse-mode automatic differentiation as described in section 2.3.

The minimize member function could be implemented using GSL as follows:

```
#include <iostream>
#include <qsl/qsl_multimin.h>
bool State::minimize() {
  // Minimizer settings
  const double initial_step_size = 0.01;
  const double line_search_tolerance = 1.0e-4;
  const double converged_gradient_norm = 1.0e-3;
  // Use the "limited-memory BFGS" quasi-Newton minimizer
  const gsl_multimin_fdfminimizer_type* minimizer_type
    = gsl_multimin_fdfminimizer_vector_bfgs2;
  // Declare and populate structure containing function pointers
  gsl_multimin_function_fdf my_function;
  my_function.n = nx();
  my_function.f = my_function_value;
  my_function.df = my_function_gradient;
  my_function.fdf = my_function_value_and_gradient;
  my_function.params = reinterpret_cast<void*>(this);
  // Set initial state variables using GSL's vector type
  qsl_vector *x;
  x = gsl_vector_alloc(nx());
  for (unsigned int i = 0; i < nx(); ++i) qsl_vector_set(x, i, 1.0);
  // Configure the minimizer
  gsl_multimin_fdfminimizer* minimizer
    = gsl_multimin_fdfminimizer_alloc(minimizer_type, nx());
  gsl_multimin_fdfminimizer_set(minimizer, &my_function, x,
                                initial_step_size, line_search_tolerance);
  // Begin loop
  size_t iter = 0;
  int status;
  do {
    ++iter;
    // Perform one iteration
    status = gsl_multimin_fdfminimizer_iterate(minimizer);
    // Quit loop if iteration failed
    if (status != GSL_SUCCESS) break;
    // Test for convergence
```

The GSL interface requires three functions to be defined, each of which takes a vector of state variables \mathbf{x} as input: my_function_value, which returns the value of the function; my_function_gradient, which returns the gradient of the function with respect to \mathbf{x} ; and my_function_value_and_gradient, which returns the value and the gradient of the function. These functions are provided to GSL as function pointers (see above), but since GSL is a C library, we need to use the 'extern "C"' specifier in their definition. Thus the function definitions would be:

When the <code>gsl_multimin_fdfminimizer_iterate</code> function is called, it chooses a search direction and performs several calls of these functions to approximately minimize the function along this search direction. The <code>this</code> pointer (i.e. the pointer to the <code>State</code> object), which was provided to the <code>my_function</code> structure in the definition of the <code>minimize</code> function above, is provided as the second argument to each of the three functions above. Unlike in C, in C++ this pointer needs to be cast back to a pointer to a <code>State</code> type, hence the use of <code>reinterpret_cast</code>.

That's it! A call to minimize should successfully minimize well behaved differentiable multi-dimensional functions. It should be straightforward to adapt the above to work with other minimization libraries.

2.6 Calling an algorithm with and without automatic differentiation from the same program

The calc_function_value(const double*) member function defined in section 2.5 is sub-optimal in that it simply calls the calc_function_value(const adouble*) member function, which not only computes the value of the function, it also records the derivative information of all the operations involved. This information

is then ignored. This overhead makes the function typically 2.5–3 times slower than it needs to be, although sometimes (specifically for loops containing no trancendental functions) the difference between an algorithm coded in terms of doubles and the same algorithm coded in terms of adoubles can exceed a factor of 10 (Hogan, 2014). The impact on the computational speed of the entire minimization process depends on how many requests are made for the function value only as opposed to the gradient of the function, and can be significant. We require a way to avoid the overhead of *Adept* computing the derivative information for calls to calc_function_value(const double*), without having to maintain two versions of the algorithm, one coded in terms of doubles and the other in terms of adoubles. The three ways to achieve this are now described.

2.6.1 Function templates

The simplest approach is to use a function template for those functions that take active arguments, as demonstrated in the following example:

```
#include <adept.h>
class State {
  public:
    ...
    template <typename xdouble>
        xdouble calc_function_value(const xdouble* x);
    ...
};

// Example function definition that must be in a header file included
// by any source file that calls calc_function_value
template <typename xdouble>
inline
xdouble State::calc_function_value(const xdouble* x) {
    xdouble y = 4.0;
    xdouble s = 2.0*x[0] + 3.0*x[1]*x[1];
    y *= sin(s);
    return y;
}
```

This takes the example from section 2.2 and replaces adouble by the template type xdouble. Thus, calc_function_value can be called with either double or adouble arguments, and the compiler will compile inline the inactive or active version accordingly. Note that the function template need not be a member function of a class.

This technique is good if only a small amount of code needs to be differentiated, but for large models the use of inlining is likely to lead to duplication of compiled code leading to large executables and long compile times. The following two approaches do not have this drawback and are suitable for large codes.

2.6.2 Pausable recording

WARNING: THIS FUNCTIONALITY IS BROKEN IN VERSION 1.9 BUT WILL BE FIXED WITH VERSION 2.0

The second method involves compiling the entire code with the ADEPT_RECORDING_PAUSABLE preprocessor variable defined, which can be done by adding an argument <code>-DADEPT_RECORDING_PAUSABLE</code> to the compler command line. This modifies the behaviour of mathematical operations performed on <code>adouble</code> variables: instead of performing the operation and then storing the derivative information, it performs the operation and then only stores the derivative information if the *Adept* stack is not in the "paused" state. We then use the following member function definition instead of the one in section 2.5:

```
double State::calc_function_value(const double* x) {
   stack_.pause_recording();
   for (unsigned int i = 0; i < nx(); ++i) active_x_[i] = x[i];
   double J = value(calc_function_value(&active_x_[0]));
   stack_.continue_recording();
   return J;</pre>
```

By pausing the recording for all operations on adouble objects, most of the overhead of storing derivative information is removed. The extra run-time check to see whether the stack is in the paused state, which is carried out by mathematical operations involving adouble objects, generally adds a small overhead. However, in algorithms where most of the number crunching occurs in loops containing no trancendental functions, even if the stack is in the paused state, the presence of the check can prevent the compiler from agressively optimizing the loop. In that instance the third method may be preferable.

2.6.3 Multiple object files per source file

}

The third method involves compiling each source file containing functions with adouble arguments twice. The first time, the code is compiled normally to produce an object file containing compiled functions including automatic differentiation. The second time, the code is compiled with the <code>-DADEPT_NO_AUTOMATIC_DIFFERENTIATION</code> flag on the compiler command line. This instructs the <code>adept.h</code> header file to turn off automatic differentiation by defining the <code>adouble</code> type to be an alias of the <code>double</code> type. This way, a second set of object files are created containing overloaded versions of the same functions as the first set but this time without automatic differentiation. These object files can be compiled together to form one executable. In the example presented in section 2.5, the <code>calc_function_value</code> function would be one that would be compiled twice in this way, once to provide the <code>calc_function_value</code> (const <code>adouble*</code>) version and the other to provide the <code>calc_function_value</code> (const <code>double*</code>) version. Note that any functions that do not include <code>adouble</code> arguments must be compiled only once, because otherwise the linker will complain about multiple versions of the same function.

The following shows a Makefile from a hypothetical project that compiles two source files (algorithm1.cpp and algorithm2.cpp) twice and a third (main.cpp) once:

```
# Specify compiler and flags
CPP = g++
CPPFLAGS = -Wall - O3 - q
# Normal object files to be created
OBJECTS = algorithm1.o algorithm2.o main.o
# Object files created with no automatic differentiation
NO_AD_OBJECTS = algorithm1_noad.o algorithm2_noad.o
# Program name
PROGRAM = my_program
# Include-file location
INCLUDES = -I/usr/local/include
# Library location and name, plus the math library
LIBS = -L/usr/local/lib -lm -ladept
# Rule to build the program (typing "make" will use this rule)
$(PROGRAM): $(OBJECTS) $(NO_AD_OBJECTS)
       $(CPP) $(CPPFLAGS) $(OBJECTS) $(NO_AD_OBJECTS) $(LIBS) -0 $(PROGRAM)
# Rule to build a normal object file (used to compile all objects in OBJECTS)
       $(CPP) $(CPPFLAGS) $(INCLUDES) -c $<
# Rule to build a no-automatic-differentiation object (used to compile ones in NO_AD_OBJECTS)
%_noad.o: %.cpp
       $(CPP) $(CPPFLAGS) $(INCLUDES) -DADEPT_NO_AUTOMATIC_DIFFERENTIATION -c $< -o $@
```

There is a further modification required with this approach, which arises because if a header file declares both the double and adouble versions of a function, then when compiled with -DADEPT_NO_AUTOMATIC_DIFFERENTIATION it appears to the compiler that the same function is declared twice, leading to a compile-time error. This can be overcome by using the preprocessor to hide the adouble version if the code is compiled with this flag, as follows (using the example from section 2.5):

```
#include <adept.h>
class State {
  public:
    ...
```

```
double calc_function_value(const double* x);
protected:
#ifndef ADEPT_NO_AUTOMATIC_DIFFERENTIATION
   adouble calc_function_value(const adouble* x);
#endif
   ...
};
```

A final nuance is that if the code contains an adouble object x, then x.value() will work fine in the compilation when x is indeed of type adouble, but in the compilation when it is set to a simple double variable, the value() member function will not be found. Hence it is better to use adept::value(x), which returns a double regardless of the type of x, and works regardless of whether the code was compiled with or without the -DADEPT_NO_AUTOMATIC_DIFFERENTIATION flag.

2.7 Interfacing with software containing hand-coded Jacobians

Often a complicated algorithm will include multiple components. Components of the code written in C or C++ for which the source is available are straightforward to convert to using *Adept*, following the rules in section 2.2. For components written in Fortran, this is not possible, but if such components have their own hand-coded Jacobian then it is possible to interface *Adept* to them. More generally, in certain situations automatic differentiation is much slower than hand-coding (see the Lax-Wendroff example in Hogan, 2014) and we may wish to hand-code certain critical parts. In general the Jacobian matrix is quite expensive to compute, so this interfacing strategy makes most sense if the component of the algorithm has a small number of inputs or a small number of outputs. A full working version of the following example is given as "Test 3" in the test directory of the *Adept* package.

Consider the example of a radiative transfer model for simulating satellite microwave radiances at two wavelengths, I and J, which takes as input the surface temperature T_s and the vertical profile of atmospheric temperature T_s from a numerical weather forecast model. Such a model would be used in a data assimilation system to assimilate the temperature information from the satellite observations into the weather forecast model. In addition to returning the radiances, the model returns the gradient $\partial I/\partial T_s$ and the gradients $\partial I/\partial T_i$ for all height layers i between 1 and n, and likewise for radiance J. The interface to the radiative transfer model is the following:

The calling function needs to allocate 2*n elements for the temperature Jacobian dradiance_dtemperature to be stored, and the stored Jacobian will be oriented such that the radiance index varies fastest.

Adept needs to be told how to relate the radiance perturbations δI and δJ , to perturbations in the input variables, δT_s and δT_i (for all layers i). Mathematically, we wish the following relationship to be stored within the Adept stack:

$$\delta I = \frac{\partial I}{\partial T_s} \delta T_s + \sum_{i=1}^n \frac{\partial I}{\partial T_i} \delta T_i. \tag{2.1}$$

This is achieved with the following wrapper function, which has adouble inputs and outputs and therefore can be called from within other parts of the algorithm that are coded in terms of adouble objects:

```
for (int i = 0; i < n; ++i) t[i] = value(temperature[i]);</pre>
// Declare variables to hold the inactive outputs and their Jacobians
double r[2];
double dr_dst[2];
std::vector<double> dr_dt(2*n);
// Call the non-Adept function
simulate_radiances(n, st, &t[0], &r[0], dr_dst, &dr_dt[0]);
// Copy the results into the active variables, but use set_value in order
// not to write any equivalent differential statement to the Adept stack
radiance[0].set_value(r[0]);
radiance[1].set_value(r[1]);
// Loop over the two radiances and add the differential statements to the Adept stack
for (int i = 0; i < 2; ++i) {</pre>
  // Add the first term on the right-hand-side of Equation 1 in the text
 radiance[i].add_derivative_dependence(surface_temperature, dr_dst[i]);
  // Now append the second term on the right-hand-side of Equation 1. The third argument
  // "n" of the following function says that there are n terms to be summed, and the fourth
  // argument "2" says to take only every second element of the Jacobian dr_dt, since the
  // derivatives with respect to the two radiances have been interlaced. If the fourth
  // argument is omitted then relevant Jacobian elements will be assumed to be contiguous
  // in memory.
  radiance[i].append_derivative_dependence(temperature, &dr_dt[i], n, 2);
```

In this example, the form of add_derivative_dependence for one variable on the right-hand-side of the derivative expression has been used, and the form of append_derivative_dependence for an array of variables on the right-hand-side has been used. As described in section 2.9, both functions have forms that take single variables and arrays as arguments. Note also that the use of std::vector<double> rather than new double[n] ensures that if simulate_radiances throws an exception, the memory allocated to hold dr_dt will be freed correctly.

2.8 Member functions of the Stack class

This section describes the user-oriented member functions of the Stack class. Some functions have arguments with default values; if these arguments are omitted then the default values will be used. Some of these functions throw *Adept* exceptions, defined in section 4.4.

- Stack (bool activate_immediately = true) The constructor for the Stack class. Normally Stack objects are constructed with no arguments, which means that the object will attempt to make itself the currently active stack by placing a pointer to itself into a global variable. If another Stack object is currently active, then the present one will be fully constructed, left in the unactivated state, and an stack_already_active exception will be thrown. If a Stack object is constructed with an argument "false", it will be started in an unactivated state, and a subsequent call to its member function activate will be needed to use it.
- void new_recording() Clears all the information on the stack in order that a new recording can be started. Specifically this function clears all the differential statements, the list of independent and dependent variables (used in computing Jacobian matrices) and the list of gradients used by the compute_tangent_linear and compute_adjoint functions. Note that this function leaves the memory allocated to reduce the overhead of reallocation in the new recordings.
- bool pause_recording() Stops recording differential information every time an adouble statement is executed. This is useful if within a single program an algorithm needs to be run both with and without automatic differentiation. This option is only effective within compilation units compiled with ADEPT_RECORDING_PAUSABLE defined; if it is, the function returns true, otherwise it returns false. Further information on using this and the following function are provided in section 2.6.2.

- bool continue_recording() Instruct a stack that may have previously been put in a paused state to now continue recording differential information as normal. This option is only effective within compilation units compiled with ADEPT_RECORDING_PAUSABLE defined; if it is, the function returns true, otherwise it returns false.
- bool is_recording() Returns false if recording has been paused with pause_recording() and the code has been compiled with ADEPT_RECORDING_PAUSABLE defined. Otherwise returns true.
- void compute_tangent_linear() Perform a tangent-linear calculation (forward-mode differentiation) using
 the stored differential statements. Before calling this function you need call the adouble::set_gradient
 or set_gradients function (see section 2.9) on the independent variables to set the initial gradients, otherwise the function will throw a gradients_not_initialized exception. This function is synonymous
 with forward().
- void compute_adjoint() Perform an adjoint calculation (reverse-mode differentiation) using the stored
 differential statements. Before calling this function you need call the adouble::set_gradient or
 set_gradients function on the dependent variables to set the initial gradients, otherwise the function
 will throw a gradients_not_initialized exception. This function is synonymous with reverse().
- void independent (const adouble& x) Before computing Jacobian matrices, you need to identify the independent and dependent variables, which correspond to the columns and rows of he Jacobian, respectively. This function adds x to the list of independent variables. If it is the nth variable identified in this way, the nth column of the Jacobian will correspond to derivatives with respect to x.
- void dependent (const adoubles y) Add y to the list of dependent variables. If it is the mth variable identified in this way, the mth row of the Jacobian will correspond to derivatives of y with respect to each of the independent variables.
- void independent (const adouble* x_ptr, size_t n) Add n independent variables to the list, which must be stored consecutively in memory starting at the memory pointed to by x_ptr.
- void dependent (const adouble* y_ptr, size_t n) Add n dependent variables to the list, which must be stored consecutively in memory starting at the memory pointed to by y_ptr.
- void jacobian(double* jacobian_out) Compute the Jacobian matrix, i.e., the gradient of the m dependent variables (identified with the dependent (...) function) with respect to the n independent variables (identified with independent (...). The result is returned in the memory pointed to by jacobian_out, which must have been allocated to hold $m \times n$ values. The result is stored in column-major order, i.e., the m diemension of the matrix varies fastest. If no dependents or independents have been identified, then the function will throw a dependents_or_independents_not_identified exception. In practice, this function calls jacobian_forward if $n \le m$ and jacobian_reverse if n > m.
- **void jacobian_forward(double* jacobian_out)** Compute the Jacobian matrix by executing n forward passes through the stored list of differential statements; this is typically faster than **jacobian_reverse** for $n \le m$.
- void jacobian_reverse (double* jacobian_out) Compute the Jacobian matrix by executing m reverse passes through the stored list of differential statements; this is typically faster than jacobian_forward for n > m.
- void clear_gradients() Clear the gradients set with the set_gradient member function of the adouble class. This enables multiple adjoint and/or tangent-linear calculations to be performed with the same recording.
- **void clear_independents()** Clear the list of independent variables, enabling a new Jacobian matrix to be computed from the same recording but for a different set of independent variables.

- **void clear_dependents()** Clear the list of dependent variables, enabling a new Jacobian matrix to be computed from the same recording but for a different set of dependent variables.
- size_t n_independents () Return the number of independent variables that have been identified.
- size_t n_dependents() Return the number of dependent variables that have been identified.
- size_t n_statements() Return the number of differential statements in the recording.
- size_t n_operations() Return the total number of operations in the recording, i.e the total number of terms on the right-hand-side of all the differential statements.
- size_t max_gradients() Return the number of working gradients that need to be stored in order to perform a forward or reverse pass.
- size_t memory() Return the number of bytes currently used to store the differential statements and the working gradients. Note that this does not include memory allocated but not currently used.
- size_t n_gradients_registered() Each time an adouble object is created, it is allocated a unique index that is used to identify its gradient in the recorded differential statements. When the object is destructed, its index is freed for reuse. This function returns the number of gradients currently registered, equal to the number of adouble objects currently created.
- void print_status(std::ostream& os = std::cout) Print the current status of the Stack object, such
 as number of statements and operations stored and allocated, to the stream specified by os, or standard
 output if this function is called with no arguments. Sending the Stack object to the stream using the "<<"
 operator results in the same behaviour.</pre>
- void print_statements (std::ostream& os = std::cout) Print the list of differential statements to the specified stream (or standard output if not specified). Each line corresponds to a separate statement, for example "d[3] = 1.2*d[1] + 3.4*d[2]".
- bool print_gradients (std::ostream& os = std::cout) Print the vector of gradients to the specified stream (or standard output if not specified). This function returns false if no set_gradient function has been called to set the first gradient and initialize the vector, and true otherwise. To diagnose what compute_tangent_linear and compute_adjoint are doing, it can be useful to call print_gradients immediately before and after.
- void activate() Activate the Stack object by copying its this pointer to a global variable that will be
 accessed by subsequent operations involving adouble objects. If another Stack is already active, a
 stack_already_active exception will be thrown. To check whether this is the case before calling
 activate(), check that the active_stack() function (described below) returns 0.
- void deactivate() Deactivate the Stack object by checking whether the global variable holding the pointer to the currently active Stack is equal to this, and if it is, setting it to 0.
- bool is active () Returns true if the Stack object is the currently active one, false otherwise.
- void start () This function was present in version 0.9 to activate a Stack object, since in that version they were not constructed in an activated state. This function has now been deprecated and will always throw a feature_not_available exception.
- int max_jacobian_threads () Return the maximum number of OpenMP threads available for Jacobian calculations. The number will be 1 if either the library was or the current source code is compiled without OpenMP support (i.e. without the -fopenmp compiler and linker flag). (Introduced in *Adept* version 1.1.)

- int set_max_jacobian_threads (int n) Set the maximum number of threads to be used in Jacobian calculations to n, if possible. A value of 1 indicates that OpenMP will not be used, while a value of 0 indicates that the maximum available will be used. Returns the maximum that will be used, which may be fewer than requested, e.g. 1 if the *Adept* library was compiled without OpenMP support. (Introduced in *Adept* version 1.1.)
- void preallocate_statements (int n) If you know in advance roughly how many differential statements will be stored by an algorithm then you may be able to speed-up the first use of the stack by preallocating the memory needed to store them. More memory will still be allocated if needed, but this should reduce the number of allocations and copies.
- **void preallocate_operations (int n)** Likewise, if you know in advance roughly how many operations will be stored then you can speed-up the first use of the stack with this member function.

The following non-member functions are provided in the adept namespace:

- adept::Stack* active_stack() Returns a pointer to the currently active Stack object, or 0 if there is none.
- bool is_thread_unsafe() Returns true if your code has been compiled with ADEPT_STACK_THREAD_UNSAFE, false otherwise.

2.9 Member functions of the adouble object

This section describes the user-oriented member functions of the adouble class. Some functions have arguments with default values; if these arguments are omitted then the default values will be used. Some of these functions throw *Adept* exceptions, defined in section 4.4.

double value() Return the underlying double value.

- void set_value (double x) Set the value of the adouble object to x, without storing the equivalent differential statement in the currently active stack.
- void set_gradient(const double& gradient) Set the gradient corresponding to this adouble variable.
 The first call of this function (for any adouble variable) after a new recording is made also initializes the vector of working gradients. This function should be called for one or more adouble objects after a recording has been made but before a call to Stack::compute_tangent_linear() or Stack::compute_adjoint().
- void get_gradient(double& gradient) Set gradient to the value of the gradient corresponding to this adouble object. This function is used to extract the result after a call to
 Stack::compute_tangent_linear() or Stack::compute_adjoint(). If the set_gradient function
 was not called since the last recording was made, this function will throw a gradients_not_initialized
 exception. The function can also throw a gradient_out_of_range exception if new adouble objects were
 created since the first set_gradient function was called.
- void add_derivative_dependence (const adouble& r, const adouble& g) Add a differential statement to the currently active stack of the form $\delta \mathbf{1} = \mathbf{g} \times \delta \mathbf{r}$, where 1 is the adouble object from which this function is called. This function is needed to interface to software containing hand-coded Jacobians, as described in section 2.7; in this case \mathbf{g} is the gradient $\partial \mathbf{l}/\partial \mathbf{r}$ obtained from such software.
- void append_derivative_dependence (const adouble& r, const adouble& g) Assuming that the same adouble object has just had its add_derivative_dependence member function called, this function appends $+ \mathbf{g} \times \delta \mathbf{r}$ to the most recent differential statement on the stack. If the calling adouble object is different, then a wrong_gradient exception will be thrown. Note that multiple append_derivative_dependence calls can be made in succession.

void add_derivative_dependence(const adouble* r, const double* g,

size_t n = 1, size_t m_stride = 1) Add a differential statement to the currently active stack of the form $\delta l = \sum_{i=0}^{n-1} m[i] \times \delta r[i]$, where 1 is the adouble object from which this function is called. If the g_stride argument is provided, then the index to the g array will be $i \times g$ _stride rather than i. This is useful if the Jacobian provided is oriented such that the relevant gradients for 1 are not spaced consecutively.

void append_derivative_dependence(const adouble* rhs, const double* g,

```
size_t n = 1, size_t q_stride = 1)
```

Assuming that the same adouble object has just called the add_derivative_dependence function, this function appends $+\sum_{i=0}^{\mathbf{n}-1}\mathbf{m}[i]\times\delta\mathbf{r}[i]$ to the most recent differential statement on the stack. If the calling adouble object is different, then a wrong_gradient exception will be thrown. The g_stride argument behaves the same way as in the previous function described.

The following non-member functions are provided in the adept namespace:

- double value (const adouble& x) Returns the underlying value of x as a double. This is useful to enable x to be used in fprintf function calls. It is generally better to use adept::value(x) rather than x.value(), because the former also works if you compile the code with the ADEPT_NO_AUTOMATIC_DIFFERENTIATION flag set, as discussed in section 2.6.3.
- void set_values(adouble* x, size_t n, const double* x_val) Set the value of the n adouble objects starting at x to the values in x_val, without storing the equivalent differential statement in the currently active stack.
- void set_gradients(adouble* x, size_t n, const double* gradients) Set the gradients corresponding to the n adouble objects starting at x to the n doubles starting at gradients. This has the same effect as calling the set_gradient member function of each adouble object in turn, but is more concise.
- void get_gradients(const adouble* y, size_t n, double* gradients) Copy the gradient of the n adouble objects starting at y into the n doubles starting at gradients. This has the same effect as calling the get_gradient member function of each adouble object in turn, but is more concise. This function can throw a gradient_out_of_range exception if new adouble objects were created since the first set_gradients function or set_gradient member function was called.

Chapter 3

Using Adept's array functionality

3.1 Introduction

The design of *Adept*'s array capability and many of the functions is inspired to a significant extent by the built-in array support in Fortran 90 (and later), and a lesser extent by Matlab, although implemented in the "C++ way", e.g. default row-major order with all array indices starting from zero. Future additions to the array capability in *Adept* will attempt to reproduce built-in Fortran array functions if available*. This design makes *Adept* a good choice if you have Fortran code that you wish to convert to C++. *Adept* provides the following array functionality:

Multi-dimensional arrays. Standard dynamically sized arrays can have an arbitrary number of dimensions (although indexing and slicing is supported only up to 7), and may refer to non-contiguous areas of memory. See section 3.2.

Mathematical operators and functions. *Adept* supports array expressions containing the standard mathematical operators +, -, * and /, as well as their assignment versions +=, -=, *= and /=. When applied to arrays, they work "element-wise", applying the same operation to every element of the arrays. *Adept* also supports array operations on all the mathematical functions listed in section 2.1. The following operators and functions return boolean array expressions: ==, !=, >, <, >= and <=, isfinite, isinf and isnan. See section 3.3.

Array slicing. There are many ways to produce an array that references a subset of another array, and therefore can be used as an Ivalue in a statement. Arrays can be indexed with scalar integers, a contiguous range of integers, a strided range of integers or an arbitrary list of integers. This is facilitated with "__" (a double underscore) and "end", such that A(__, end-1) returns a vector pointing to the penultimate column of matrix A. The member function subset produces an array pointing to a contiguous subset of the original array, while diag_vector and diag_matrix produce arrays pointing to the diagonal of the original array. T produces an array pointing to the transpose of the original array. See section 3.4.

Passing arrays to and from functions. *Adept* uses a reference-counting approach to implement the storage of array data, enabling multiple array objects to point to the same data, or parts of it in the case of array slices. This makes it straightforward to pass arrays to and from functions without having to perform a deep copy. See section 3.5.

Array reduction operations. The functions sum, mean, product, minval, maxval and norm2 perform reduction operations that return an array of lower rank to the expression they are applied to. The functions all and any do the same but for boolean expressions. count returns the number of true elements in a boolean expression. See section 3.6.

^{*}This decision may puzzle some readers, since Fortran is a dirty word to many C++ users due to the limitations of the FORTRAN 77 language. Many of these limitations were overcome in Fortran 90, whose array functionality in particular is rather well designed. Indeed, the pioneering "Blitz++" C++ array library (Veldhuizen, 1995) also reproduces many Fortran array functions. All references to Fortran in this document imply the 1990 (or later) standard.

3.2. The Array class 24

Conditional operations. Two convenient ways are provided to perform an operation on an array depending on the result of a boolean expression: where and find. The statement A. where (B>0) =C assigns elements of C to elements of A whenever the corresponding element of B is greater than zero. For vectors only, the same result could be obtained with A (find (B>0)) = C (find (B>0)). See section 3.7.

Fixed-size arrays. Adept provides a fixed-size array class with dimensions (up to seven) that are known at compile time. The functionality is very similar to standard dynamic arrays.

Special square matrices. *Adept* uses specific classes for symmetric, triangular and band-diagonal matrices, the latter of which use compressed storage and include diagonal and tridiagonal matrices. Certain operations such as matrix multiplication and solving linear equations are optimized especially for these objects. See section 3.9.

Matrix multiplication. Matrix multiplication can be applied to one- and two-dimensional arrays using the matmul function, of for extra syntactic sugar, the "**" pseudo-operator. *Adept* uses whatever BLAS (Basic Linear Algebra Subroutines) support is available on your system, including optimized versions for symmetric and band-diagonal matrices. See section 3.10.

Linear algebra. *Adept* uses the LAPACK library to invert matrices and solve linear systems of equations. See section 3.11.

Array bounds and alias checking. *Adept* checks at compile time that terms in an array expression accord in rank, and at run time that they accord in the size of each dimension. Run-time alias checking is performed to determine if any objects on the right-hand-side of a statement overlap in memory with the left-hand-side of the statement, making a temporary copy of the right-hand-side if they do. This can be overridden with the noalias function. See section 3.12.

3.2 The Array class

The bread and butter of array operations is provided by the Array class template (in the adept namespace along with all other public types and classes), which has the following declaration:

```
namespace adept {
  template <int Rank, typename Type = Real, bool IsActive = false>
  class Array;
}
```

The first template argument provides the number of dimensions of the array and may be 1 or greater, although indexing and slicing is only supported up to 7 dimensions. The second argument is the numerical type being stored and can be any simple integer or real number, including bool. The default type is adept::Real, which is the default floating-point type the *Adept* library has been compiled to use for computing derivatives, and is usually double. The final argument states whether the array is "active", i.e. whether it participates in the differentiation of an algorithm.

A number of typedefs are provided for the most common types of array: Vector, Matrix and Array3D provide inactive arrays of type Real and rank 1–3. The corresponding active types are aVector, aMatrix and aArray3D. Vectors of other numeric types are boolVector, intVector, floatVector, afloatVector, and similarly for matrices and 3D arrays of these types. If you wanted shortcuts for 4-dimensional active and passive arrays, they could be defined using:

```
typedef adept::Array<4> Array4D;
typedef adept::Array<4, adept::Real, true> aArray4D;
```

An Array can be constructed in numerous ways:

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```
Matrix M(dim);  // This works for an array with an arbitrary number of dimensions
Vector v = M(__,0); // Link to a existing array, in this case the first column of M
Vector v(M(__,0)); // Has exactly the same effect as the previous example
Matrix N = log(M); // Initialize with the size and values of a mathematical expression
```

Note that in the remaining code examples it will be assumed that using namespace adept has already been called. When new memory is needed, the Array object creates a Storage object that contains the memory needed, and stores pointers to both the Storage object and the start of the data. By default the data are accessed in C-style row-major order (i.e. the final index corresponds to the array dimension that varies most rapidly in memory). However, this is flexible since in addition to storing the length of each of its n dimensions, a rank-n Array also stores n "offsets" that define the separation of elements in memory in each dimension. Thus, a 3-by-4 matrix with row-major storage would store offsets of (4,1). The same size matrix would use column-major storage simply by storing offsets of (1,3). To make new arrays use column-major storage, call the following function:

```
set_array_row_major_order(false);
```

Note that this does not change the storage of any existing objects. Note also that when array expressions are evaluated, the data are requested in row-major order, so the use of column-major arrays will incur a performance penalty.

It can be seen from one of the constructors above (the example involving a Vector) that an Array can be configured to "link" to part of an existing Array, and modifications to the numbers in one will be seen by the other. This is a very useful feature as it allows slices of an array to be passed to functions and modified; see section 3.4. Note that the array or sub-array being linked to must be of the same rank, type and activeness as the linking array. Internally, linking is achieved by both the arrays pointing to the same Storage object, which itself contains a reference count of the number of arrays pointing to it. When an Array is destructed the reference count is reduced by one and only if it falls to zero will the data get deallocated.

After it has been constructed, an Array can be resized, relinked or cleared completely as follows:

```
M.resize(5,2);  // Works up to 7 dimensions
Index dim[2]={5,2};
M.resize(dim);  // Works for any number of dimensions
v.link(M(end-1,__));  // Size of v set to that of the argument and link to data
v >>= M(end-1,__);  // Convenient syntax for linking, similar to Fortran's "->"
M.clear();  // Returns array to original empty state
```

The member functions resize and clear unlink from any existing data, while link can only be used if the array is already in an empty state. Note that if you assign one array to another (e.g. N=M), then they must be of the same size; if they are not then you should clear the left-hand-side first.

The Array class implements a number of member functions for inquiring about its properties:

size() Returns the total number of elements, i.e. the product of the lengths of each of the dimensions.

dimension (i) Returns the length of dimension i.

offset (i) Returns the separation in memory of elements along dimension \mathtt{i} .

gradient_index() For active arrays, returns the gradient index of the first element of the array, which is always positive; for inactive arrays it returns a negative number.

empty() Returns true if the array is in the empty state, or false otherwise.

An Array may be filled using the << operator for the first element followed by either the << or , operators for subsequent elements:

For multidimensional arrays, elements are filled such that the final dimension ticks over fastest (regardless of whether the array uses row-major storage internally), and new rows are started when a row is complete. Moreover, other arrays can be part of the list of elements, provided that they fit in. In this context, a rank-1 array is treated as a row vector. An index_out_of_bounds exception is thrown if an array is overfilled, while an empty_array exception is thrown if an attempt is made to fill an empty array.

If you compile your code with C++11 features enabled then you can use the "initializer list" feature to fill arrays using the C-like curly bracket syntax:

Another convenient property of this syntax is that temporary arrays with explicit values can be used in expressions:

```
v = w * Vector{3.0, 4.2, 5.1};
```

When interfacing with other libraries, direct access to the data is often required. The Array class provides the following member functions:

data () Returns a pointer to the first element in the array, i.e. the element found by indexing all the dimensions of the array with zero. It is up to the caller to understand the layout of the data in memory and not to stray outside. Remember that an array may be strided and the stride may even be negative so that the data returned from increasing indices are actually from earlier memory addresses. Note that a double-precision active array is not stored as an array of adouble objects, but as an array of double data and a single gradient index for the first element. Thus the pointer returned by data() will point to the underlying inactive data. In contexts where the Array object is const, a const pointer will be returned.

const_data() It is sometimes convenient to specify explicitly that read-only access is required, in which case you can use const_data() to return a const pointer to the first element in the array.

3.3 Operators and mathematical functions

The operators and mathematical functions listed in section 2.1 have been overloaded so that they work exactly as you would expect. Consider this example:

The penultimate illustrates that all elements of an Array can be set to the same value, although note that this will only work if the array is not in the empty state. The final line illustrates how terms with different rank, type and activeness can participate in the same expression. Scalars and arrays can participate in the same expression on the right-hand-side of a statement provided that the arrays have the same size as the array on the left-hand-side. Objects of different type (in this case single and double precision) can be combined in a mathematical operation, and the type of that operation will be the larger (higher precision) of the two types. If active and inactive objects participate in an expression then the left-hand-side must also be active. Expression templates ensure that no temporary arrays need to be created to store the output of intermediate parts of the expression. The functions max and min behave just like binary operators (such as + and *) in this regard, as shown by the following:

```
c = max(a,b);  // Element-wise comparison of a and b
c = min(a,3.0);  // Return minimum of each element of a and 3
```

3.4. Array slicing 27

The examples so far have floating-point results, but some operators (e.g. "==") and some functions (e.g. isinf) take floating-point arguments and return a boolean. The *Adept* versions take floating-point array expressions as arguments and return bool expressions of the same rank and size. Finally, the *Adept* versions of the operators !, | | and && take a bool expression as arguments and return a bool expression of the same size and rank.

3.4 Array slicing

This section concerns the many ways that sub-parts of an Array can be extracted to produce an object that can be used as an Ivalue; that is, if the object is modified then it will modify part of the original Array. It should be stressed that none of these methods results in any rearrangement of data in memory, so they should be efficient.

The first way this can be done is via the function-call and member-access operators (i.e. operator() and operator[], respectively) of the Array. In the case of the function-call operator, the same number of arguments as the rank of the array must be provided, where each argument states how its corresponding dimension should be treated. The nature of the resulting object depends on the type of all of the arguments in a way that is similar to how Fortran arrays behave, although note that array indices always start at 0. The four different behaviours are as follows:

Extract single value. If every argument is an integer scalar or scalar expression, then a reference to a single element of the array will be extracted. If an argument is an integer expression containing end, then end will be interpretted to be the index to the final element of that dimension (a feature borrowed from Matlab). If the array is active then the returned object will be of a special "active reference" type that can be used as an Ivalue and ensures that any expressions making use of this element can be differentiated. Now for some examples:

```
aMatrix A(4,3);

aReal x = A(1,1); // Copy element at second row and second column into x

A(end-1,1) \star= 2.0; // Double the element in the penultimate column and 2nd row of A

A(3) = 4.0; // Error: number of indices does not match number of dimensions
```

Extract regular subarray. If every argument is either (i) an integer scalar or scalar expression, or (ii) a regular range of indices, and there is at least one of (ii), then an Array object will be returned of the same type and activeness as the original. However, for each argument of type (i), the rank of the returned array will be one less than that of the original. There are three ways to express a regular range of indices: "__" represents all indices of a particular dimension, range (a, b) represents a contiguous range of indices between a and b (equivalent to a:b in Fortran and Matlab), and stride(a, b, c) represents a regular range of indices between a and b with spacing c (equivalent to a:b:c in Fortran and a:c:b in Matlab). Note that a, b and c may be scalar expressions containing end, but c must not be zero although it can be negative to indicate a reversed ordering. The rank of the returned array is known at compile time; thus if range arguments are found at run-time to contain only one element (e.g. range(1,1)) then the dimension being referred to will be not be removed in the returned array but will remain as a singleton dimension. This behaviour is the same as indexing an array dimension with 1:1 in Fortran. Now for some examples:

```
v(range(1,end-1)) // Subset of vector v that excludes 1st & last points A(0,stride(end,0,-1)) // First row of A as a vector treated in reverse order A(range(0,0),stride(0,0,1)) // A 1-by-1 matrix containing the first element of A
```

Extract irregular subarray. If an array is indexed as in either of the two methods above, except that one or more dimensions is instead indexed using a rank-1 Array of integers, then the result is a special "indexed-array" type that stores how each dimension is indexed. If it then participates either on the left- or right-hand-side of a mathematical expression then when an element is requested, the indices will be queried to map the request to obtain the correct element from the original array. This is much less efficient than using regular ranges of indices as above. It also means that if an indexed array is passed to a function expecting an object of type Array, then it will first be converted to an Array and any modifications performed within the function will not be passed back to the original array. For example:

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```
intVector index(3);
index << 2, 3, 5;
Array A(4,4);
A(0,index) = 2.0; // Set irregularly spaced elements of the first row of A</pre>
```

Slice leading dimension. In C, an element is extracted from a two-dimensional array using A[i][j], and A[i] returns a pointer to a single row of A, where i and j are integers. To enable similar functionality, if A is an *Adept* matrix then A[i] indexes the leading dimension by integer i returning an array of rank one less than the original. This is equivalent to A(i, ...). Furthermore, A[i][j] will return an individual element as in C, but it should be stressed that A(i, j) is more efficient since it does not involve the creation of intermediate arrays.

There are a few other ways to produce Ivalues that consist of a subset or a reordering of an array. They are implemented as member functions of the Array class, in order to distinguish from non-member functions that produce a copy of the data and therefore cannot be usefully used as Ivalues. For example, A.T() and transpose(A) both return the transpose of matrix A, but the former is faster since it does not make a copy of the original data, while the latter is more flexible since it can be applied to array expressions (e.g. transpose(A*B)). The member functions available are:

- subset (int ibegin0, int iend0, ...) This function returns a contiguous subset of an array as an array of the same rank that points to the original data. It takes twice as many arguments as the array has dimensions, with each pair of arguments representing the indices to the first and last element to include from a particular dimension. Exactly the same result can be obtained using range but the subset form is more concise. For example, for a matrix M, M. subset (1, 5, 3, 10) is equivalent to M(range(1, 5), range(3, 10)).
- **T**() This function returns the transpose of a rank-2 array (a matrix). The returned array points to the same data but with its dimensions reversed. A compile-time error occurs if this function is used on an array with rank other than 2. Currently *Adept* doesn't allow the transpose of a rank-1 array (a vector), since vectors are not intended to have an intrinsic orientation. When orientation matters, such as in matrix multiplication, the intended orientation may be inferred from the context or specified explicitly.
- permute (int i0, int i1, ...) This function is the generalization of the transpose for multi-dimensional arrays: it returns an array of the same rank as the original but with the dimensions rearranged according to the arguments. There must be the same number of arguments as there are dimensions, and each dimension (starting at 0) must be provided once only. The returned array is linked to the original; the permutation is achieved simply by rearranging the list of dimensions and the list of "offsets" (the separation in memory of elements along each dimension individually).
- **diag matrix ()** When this function is applied to a rank-1 Array of length n, it returns an n-by-n diagonal matrix (specifically a DiagMatrix; see section 3.9) that points to the data from the rank-1 array along its diagonal.
- diag_vector() When this function is applied to a rank-2 Array with equally sized dimensions, it returns a rank-1 array pointing to the data along its diagonals. An invalid_operation exception is thrown if applied to a non-square matrix, and a compile-time error if applied to an array of rank other than 2.
- **diag_vector(i)** When applied to a square rank-2 n-by-n Array, this returns a rank-1 array of length n abs(i) pointing to the ith superdiagonal of the square matrix, or the -ith subdiagonal if i is negative. An invalid_exception exception occurs if applied to a non-square matrix, and a compile-time error if applied to an array of rank other than 2.
- submatrix.on_diagonal (ibegin, iend) When applied to a square rank-2 array, this function returns a square matrix that shares part of the diagonal of the original matrix. Thus A.submatrix_on_diagonal (int ibegin, int iend) is equivalent to A(range(ibegin, iend), range(ibegin, iend)). Its purpose is to provide a subsetting facility for symmetric, triangular and band-diagonal matrices (see section 3.9) for which general array indexing is not available. If applied to a non-square matrix, an invalid_operation exception will be thrown.

3.5 Passing arrays to and from functions

When writing functions taking array arguments, there are three different ways to do it depending on the extent to which the function needs to be able to modify the array. In the case of constant array arguments, a constant reference should be used; for example:

This function works with all three types of argument. The last example illustrates that when an inactive rank-1 expression is passed to the function, it is evaluated and the result placed in a temporary vector that is passed to the function.

At the other extreme, we may wish to create a function that modifies an array argument, including the possibility of changing its size; for example:

In this case, due to the C++ rule that a non-constant reference cannot bind to a temporary object, the function can only take a *non-temporary* Vector as an argument. This is fair enough; it would not make sense to resize the subset of an array, or an expression. However, it is very common to want to pass a subset of an array to a function and for the function to modify the values of the array, but not to resize it. In *Adept* this is achieved as follows:

Even though the Vector has been passed by value, the Vector copy constructor performs a "shallow copy", which means that little more than the array dimensions and a pointer to the data are copied. Therefore, in the first two examples above the vector v inside the function points to data in w, and can therefore modify w. By contrast, when an expression is passed to the function, a new Vector is created to hold the result of the expression, and when this is modified inside the function it does not affect the data in the calling routine.

The fact that Array copy constructors perform shallow copies also improves the efficiency of functions that return arrays such as the following:

```
Matrix square(const Matrix& in) {
   Matrix out = in*in; // Create an matrix containing the result of in*in
   return out;
}
Matrix A(100,100); // Allocate memory for "A"
Matrix B = square(A); // Copy constructor: shallow copy of "out" into "B"
```

At the return statement, matrix out is received by the copy constructor of matrix B, so a shallow copy is performed. This means that the description of matrix out is copied to B, including a pointer to Storage object containing both the data and a count of the number of references to it; this counter is increased by one. Matrix out is then destructed, and the counter is immediately reduced by one. The net result is that B has "stolen" the data in the matrix from out without it having been copied, thus avoiding unnecessary allocation of memory on the heap followed by copying and deallocation.

The shallow-copy implementation leads to behaviour that users may not be expecting. If an array is initialized from another array in either of the following two ways:

```
Matrix M(3,4);
Matrix A(M); // Call copy constructor
Matrix B = M; // Call copy constructor
```

then the result is that A, B and M share the same data, rather than a copy being made. To make a deep copy, it is necessary to do the following:

```
Matrix M(3,4);
Matrix A;  // Create empty matrix
A = M;  // Call assignment operator for deep copy
```

This is annoying, but the alternative is that there would be no clean way to pass a subset of an array to a function that then modifies its values. The same behaviour is implemented in the Blitz++ array class (Veldhuizen, 1995).

It should be noted that with the introduction of "move semantics" in the C++11 standard, the it is possible to detect when an array returned from a function is about to be destructed, and therefore invoke a move constructor that implements a shallow copy. This negates one of the two reasons from making the copy constructor execute only a shallow copy. But it does not help in passing array subsets to functions, unless two versions of every function were created, one accepting an Ivalue reference (Array&) and the other accepting an rvalue reference (Array&A), which is hardly practical.

If you compile your code with C++11 features enabled then move semantics can sometimes make assignment more efficient. Consider code calling the square function above:

```
Matrix A(10,10), B(10,10);
B = square(A); // Move assignment operator performs shallow copy
Matrix C(B); // B and C now share the same data
B = square(A); // Move assignment operator performs deep copy
```

Both assignments are to temporary objects about to be destructed, so the move assignment operator is called. This operator checks how many references there are to the data in $\mathbb B$. In the first case there is only one reference, so the data in $\mathbb B$ can safely be discarded and a shallow copy (a "move") of the data in the temporary is performed. In the second case there are two references, so a deep copy must be performed in order that $\mathbb C$ sees the change in $\mathbb B$.

3.6 Array reduction operations

A family of functions return a result that is reduced in rank compared to their argument, and operate in the same way as Fortran functions of the same name. Consider the sum function, which can be used either to sum all the elements in an array expression and return a scalar, or to sum elements along the dimension specified in the second argument and return an array whose rank is one less than the first argument:

```
Array A(3,4);

Real x = sum(A); // Sum all elements of matrix A

Vector v = sum(A,1); // Sum along the row dimension returning a vector of length 3
```

Functions that are used in the same way are mean, product, minval, maxval and norm2 (the square-root of the sum of the squares of each element). Note the difference between maxval and max: the behaviour of max is outlined in section 3.3. Three further functions operate in the same way but on boolean arrays: all returns true only if all elements are true, any returns true if any element is true (and false otherwise), while count returns the number of true elements. Each of these can work on an individual dimension as with sum and friends.

A further function, dot_product (a, b), takes two arguments that must be rank-1 arrays of the same length and returns the dot product. This is essentially the same as sum(a*b).

3.7 Conditional operations

There are two main ways to perform an operation on an array depending on the result of a boolean expression. The first is similar to the Fortran where construct:

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```
Array A(3,4);

Array B(3,4);

A.where(B > 0.0) = 2.0 * B; // Only assign to A if B > 0

A.where(!isnan(B)) = either_or(-B, 0.0); // Read from either one expression or the other
```

In the first example, A is only assigned if a condition is met, and therefore A must be of the same size and rank of the boolean expression. In the second example A is filled with elements from the first argument of either_or if the boolean expression is true, or from the second argument otherwise; if A is empty then it will be resized to the size of the boolean expression. In both cases, the expressions on the right-hand-side may be scalars or array expressions of the same size as the boolean expression. Equivalent expressions are possible replacing the assignment operator with the +=, -=, *= and /= operators, in which case A must already be the same size as the boolean expression.

An alternative approach that works with only vectors uses the find function. This is similar to the equivalent Matlab function and returns an IndexVector (a vector of integers of sufficient precision to index an array) containing indices to the true elements of the vector:

```
Vector v(10), w(10);
v(find(v > 5.0)) = 3.0;
IndexVector index = find(v > 5.0);
v(index) = 2.0 * u(index);
```

This will work if no true elements are found: find will return an empty array, and when v is indexed by an empty vector, no action will be taken. In general, find is less efficient than where.

3.8 Fixed-size arrays

The size of the Array class is dynamic, which is somewhat sub-optimal for small arrays whose dimensions are known at compile time. *Adept* provides an alternative class template for an array whose size is known at compile time and whose data are stored on the stack. It has the following declaration:

```
namespace adept {
  template <typename Type, bool IsActive, int Dim0, int Dim1 = 0, ...>
  class FixedArray;
}
```

The type (e.g. double) and activeness are specified by the first two template arguments, while the remaining template arguments provide the size of the dimensions, up to 7. Only as many sizes need to be specified as there are dimensions. A user working with arrays of a particular size could use typedef to provide convenient names; for example:

```
typedef FixedArray<double, false, 4> Vector4;
typedef FixedArray<double, false, 4, 4> Matrix44;
typedef FixedArray<double, true, 4> aVector4;
typedef FixedArray<double, true, 4, 4> aMatrix44;
```

In the adept namespace, *Adept* defines Vector2, Vector3, Matrix22, Matrix33 and their active counterparts. Fixed arrays have all the same capabilities as dynamic arrays, with a few exceptions:

- Since their size is fixed, there are no member functions resize, clear or in-place_transpose.
- Since for the lifetime of the object it is associated with data on the stack, it cannot link to other data. This means that there is no member function link, and also if it is passed by value to a function then the contents of the array will be copied, rather than the behaviour of the Array class where the receiving function links to the original data.

All the same slicing operations are available as discussed in section 3.4, and they return the same types when applied to fixed arrays as they do when applied to dynamic arrays. Thus most operations return an Array object that links to a subset of the data within the FixedArray object.

3.9 Special square matrices

Adept offers several special types of square matrix that can participate in array expressions. They are more efficient than Arrays in certain operations such as matrix multiplication and assignment, but less efficient in operations such as accessing individual elements. All use an internal storage scheme compatible with BLAS (Basic Linear Algebra Subprograms). All are specializations of the SpecialMatrix class template, which has the following declaration:

```
namespace adept {
  template <typename Type, class Engine, bool IsActive = false>
  class SpecialMatrix;
}
```

The first template argument is the numerical type, the second provides the functionality specific to the type of matrix being simulated, and the third states whether the matrix participates in the differentiation of an algorithm. The specific types of special matrix are as follows:

Square matrices. SquareMatrix provides a dense square matrix of type Real with aSquareMatrix its active counterpart. Its functionality is similar to a rank-2 Array, except that its dimensions are always equal and the data along its fastest varying dimension are always contiguous in memory, which may make it faster than Array in some instances.

Symmetric matrices. SymmMatrix provides a symmetric matrix of type Real, and aSymmMatrix is its active equivalent. Internally this type uses row-major unpacked storage with the data held in the lower triangle of the array and zeros in the upper triangle (equivalent to column-major storage with data in the upper triangle). If the oposite configuration is required then it is available by specifying different template arguments to the SpecialMatrix class template. Note that with normal access methods, the storage scheme is opaque to the user; for example, S(1,2)=2.0 and S(2,1)=2.0 have the same effect.

Triangular matrices. LowerMatrix and UpperMatrix (and their active equivalents prefixed by "a") provide triangular matrices of type Real. Internally they use row-major unpacked storage, although column-major storage is available by specifying different template arguments to the SpecialMatrix class template.

Band diagonal matrices. DiagMatrix, TridiagMatrix and PentadiagMatrix provide diagonal, tridiagonal and pentadiagonal Real matrices, respectively (with their active equivalents prefixed by "a"). Internally, row-major BLAS-type band storage is used such that an n-by-n tridiagonal matrix stores 3n rather than n^2 elements. Adept supports arbitrary numbers of sub-diagonals and super-diagonals, accessible by specifying different template arguments to the SpecialMatrix class template.

A SpecialMatrix can be constructed and resized as for Arrays (see section 3.2), with the following additions:

```
SymmMatrix S(4); // Initialize a 4-by-4 symmetric matrix
S.resize(5); // Resize to a 5-by-5 matrix
```

These are applicable to all types of SpecialMatrix.

In terms of array indexing and slicing, the member functions T, diag and diag_submatrix described in section 3.4 are all available, but if you index a SpecialMatrix with S(a,b) then a and b must be scalars or scalar expressions. For triangular or band-diagonal matrices, if the requested element is one of the zero parts of the matrix then it can only be used as an rvalue in an expression. If you wish to extract arbitrary subarrays from a SpecialMatrix then it must first be converted to a Matrix:

```
SymmMatrix S(6);
intVector index(3);
index << 2, 3, 5;
Matrix M = Matrix(S)(index, stride(0,4,2));</pre>
```

3.10 Matrix multiplication

Matrix multiplication may be invoked in two equivalent ways: using the matmul function or the "**" pseudo-operator. Following Fortran, the two arguments may be either rank-1 or rank-2, but at least one argument must be

of rank-2. The orientation of any rank-1 argument is inferred from whether it is the first or second argument, as shown here:

```
Matrix A(3,5), B(5,3), C;
Vector v(5), w;
C = matmul(A,B); // Matrix-matrix multiplication: return a 3x3 matrix
w = matmul(v,B); // Interpret v as a row vector: return a vector of length 3
w = matmul(A,v); // Interpret v as a column vector: return a vector of length 3
```

In this way it is never necessary to transpose a vector; the appropriate orientation to use is inferred from the context. You may find it clearer to use "**" for matrix multiplication as illustrated here:

The "**" pseudo-operator has been implemented in *Adept* by overloading the dereference operator such that "*A" returns a special type when applied to array expressions, and overloading the multiply operator to perform matrix multiplication when one of these types is on the right-hand-side. This means that ** has the same precedence as ordinary multiplication, and both will be applied in order of left to right. Thus, in the first example above, matrix-matrix multiplication is performed followed by matrix-vector multiplication. The second example shows how to make this more efficient with parentheses to specify that the rightmost matrix multiplication should be applied first, leading to two matrix-vector multiplications. The final example shows an expression that would fail at runtime with an inner_dimension_mismatch exception due to the matrix multiplication being applied to matrices whose inner dimensions do not match.

In order to get the best performance, *Adept* does not use expression templates for matrix multiplication but rather calls the appropriate level-2 BLAS function for matrix-vector multiplication and level-3 BLAS function for matrix-matrix multiplication. For matrix multiplication involving active vectors and matrices, *Adept* first uses BLAS to perform the matrix multiplication and then stores the equivalent differential statements. There are therefore a few factors that users should be aware of in order to get the best performance:

• If an array expression rather than an array is provided as an argument to matrix multiplication, it will first be converted to an Array of the same rank. Therefore, if the same expression is used more than once in a sequence of matrix multiplications, better performance will be obtained by precomputing the array expression and storing it in a temporary matrix:

```
Matrix A(5,5), B(5,5), C(5,5), D(5,5)
// Slow implementation:
C = transpose(2.0*A*B) ** (2.0*A*B);
D = (2.0*A*B) ** C;
// Faster implementation:
{
    Matrix tmp = 2.0*A*B;
    C = tmp.T() ** tmp;
    D = tmp ** C;
} // "tmp" goes out of scope here
```

• If the left-hand argument of a matrix multiplication is a symmetric, triangular or band matrix then a specialist BLAS function will be used that is faster than the one for general dense matrices. *Adept* may not be able to tell if the result of an array expression is symmetric, triangular or has a band structure, and so may not call the most efficient BLAS function. The user can help as follows:

 $^{^{\}dagger}A$ drawback of the $\star\star$ interface with the orientation of vector arguments being inferred is that in an expression like $A\star\star v\star\star B$ (where A and B are matrices and v is a vector), v is interpreted as a column vector in $A\star\star v$, which returns a column vector result, but this result is then implicitly transposed when it is used as the left-hand argument of the matrix multiplication with B. Moreover, the order of precedence affects the result, since this expression will not give the same answer as $A\star\star(v\star\star B)$.

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• BLAS requires that the fastest-varying dimension of input matrices are contiguous and increasing. This is always the case for the special square matrices described in section 3.9, but not necessarily for a Matrix or an aMatrix, which are particular cases of the general Array type. If the fastest-varying dimension of such a matrix is not contiguous and increasing then *Adept* will copy it to a temporary matrix before invoking matrix multiplications, as in the following example:

```
Matrix A(5,5), B, C(5,5);

B.link(A(\_, stride(end,1,-1)); // Fastest varying dim is contiguous but decreasing

C = A ** A; // Matrix multiplication applied directly with A

C = B ** B; // Adept will copy B to a temporary matrix before multiplication
```

A final member function to mention in this section is in_place_transpose(), which is only applicable to matrices. It transposes the matrix by swapping the dimensions and the offsets to each dimension, but leaving the actual data untouched. This means that a matrix with row-major storage will be changed to column-major, and vice versa.

3.11 Linear algebra

Adept provides the functions solve and inv to solve systems of linear equations and to invert a matrix, respectively, which themselves call the most appropriate function from LAPACK.[‡]

3.12 Bounds and alias checking

When encountering an array or active expression, *Adept* performs several checks to test the validity of the expression both at compile time and at runtime:

Activeness check. An expression in which an active expression is assigned to an inactive array will fail to compile.

Rank check. An expression will fail to compile if the rank of the array on the left-hand-side of the "=" operator (or the operators "+=", "*=", etc.) does not match the rank of the array expression on the right-hand-side. However, a scalar (rank-0) expression can be assigned to an array of any rank; its value will be assigned to all elements of the array. Compile-time rank checks are also performed for each binary operation (binary operators such as "+" and binary functions such as pow) making up an array expression: compilation will fail if the two arguments do not have the same rank and neither is of rank 0.

Dimension check. When a binary operation is applied to two array expressions of rank n then Adept checks at run-time that each of the n dimensions has the same length. Otherwise, a size_mismatch exception is thrown.

Alias check. *Adept* checks to see whether the memory referenced in the array object on the left-hand-side of a statement overlaps with the memory referenced by any of the objects on the right-hand-side, as in this example of a shift-right operation:

[‡]Statements involving solve and inv cannot yet be automatically differentiated.

```
Vector v(6);
v(range(1,end)) = v(range(0,end-1));
```

In order to prevent the right-hand-side changing during the operation, *Adept* copies the expression on the right-hand-side to a temporary array and then assigns the left-hand-side array to this temporary, which is equivalent to the following

```
{
    Vector tmp;
    tmp = v(range(0,end-1));
    v(range(1,end)) = tmp;
} // tmp goes out of scope here
```

However, for speed *Adept* does not check to see whether individual memory locations are shared, which means that for certain strided operations this copying to a temporary array is unnecessary. Nor is it necessary if elements of an array will be accessed in exactly the same order on the left-hand-side as the right-hand-side. If the user is sure that alias checking is not necessary then he or she can override alias checking for part or all of an array expression using the noalias function, as follows:

```
v(stride(1,end,2)) = noalias(v(stride(0,end-1,2))); // No overlap between RHS and LHS 
 <math>v = 1.0 + noalias(exp(v)); // LHS \& RHS accessed in same order
```

Bounds check. If the preprocessor variable ADEPT_BOUNDS_CHECKING is defined then additional run-time checks will be performed when an array is indexed or sliced using the methods described in section 3.4; if an index is ount of bounds then a index_out_of_bounds exception will be thrown. This makes indexing and slicing of arrays slower so would normally only be used for debugging.

Chapter 4

General considerations

4.1 Setting and checking the global configuration

The following non-member functions are provided in the adept namespace:

- std::string version() Returns a string containing the version number of the Adept library (e.g. "1.9.8").
- **std::string compiler_version()** Returns a string containing the compiler name and version used to compile the *Adept* library.
- std::string compiler_flags() Returns a string containing the compiler flags used when compiling the *Adept* library.
- **std::string configuration()** Returns a multi-line string listing numerous aspects of the way *Adept* has been configured.
- int set_max_blas_threads(int n) Set the maximum number of threads used for matrix operations by the BLAS library, or zero to use the upper limit on your system. The number returned is the number actually used.
- int num_blas_threads(int n) Return the maximum number of threads available for matrix operations by the BLAS library.

4.2 Parallelizing *Adept* programs

Adept currently has limited built-in support for parallelization. If the algorithms that you wish to differentiate are individually small enough to be treated by a single processor core, and you wish to differentiate multiple algorithms independently (or the same algorithm but with multiple sets of inputs) then parallelization is straightforward. This is because the global variable containing a pointer to the Adept stack uses thread-local storage. This means that if a process spawns multiple threads (e.g. using OpenMP or Pthreads) then each thread can declare one adept::Stack object and all adouble operations will result in statements being stored on the stack object specific to that thread. The Adept package contains a test program test_thread_safe that demonstrates this approach in OpenMP.

If your problem is larger and you wish to use parallelism to speed-up the differentiation of a single large algorithm then the build-in support is more limited. Provided your program and the *Adept* library were compiled with OpenMP enabled (which is the default for the *Adept* library if your compiler supports OpenMP), the computation of Jacobian matrices will be parallelized. By default, the maximum number of concurrent threads will be equal to the number of available cores, but this can be overridden with the set_max_jacobian_threads member function of the Stack class. Note that the opportunity for speed-up depends on the size of your Jacobian matrix: for an $m \times n$ matrix, the number of independent passes through the stored data is $\min(m,n)$ and each thread treats ADEPT_MULTIPASS_SIZE of them (see section 4.5.2), so the maximum number of threads that can be exploited is $\min(m,n)/\text{ADEPT_MULTIPASS_SIZE}$. Again, the test_thread_safe program can demonstrate the parallelization

of Jacobian calculations. Note, however, that if the jacobian function is called from within an OpenMP thread (e.g. if the program already uses OpenMP with each thread containing its own adept::Stack object), then the program is likely not to be able to spawn more threads to assist with the Jacobian calculation.

If you need Jacobian matrices then the ability to parallelize the calculation of them is useful since this tends to be more computationally costly than recording the original algorithm. If you only require the tangent-linear or adjoint calculations (equivalent to a Jacobian calculation with n = 1 or m = 1, respectively), then unfortunately you are stuck with single threading. It is intended that a future version of *Adept* will enable all aspects of differentiating an algorithm to be parallelized with either or both of OpenMP and MPI.

If your BLAS library has support for parallelization then be aware that the performance may be poor if other parts of the program are parallelized. This occurs with OpenBLAS, which uses Pthreads, if you also use parallelized Jacobian calculations, which use OpenMP. In this instance you can turn off parallelization of array operations with the set_max_blas_threads(1) function in the adept namespace. The number of available threads for array operations is returned by the max_blas_threads() function. Alternatively, you can use the OPENBLAS_NUM_THREADS environment variable to control the number of threads used by OpenBLAS, and the OMP_NUM_THREADS environment variable to control the number used in Jacobian calculations.

4.3 Tips for the best performance

- If you are working with single-threaded code, or in a multi-threaded program but with only one thread using a Stack object, then you can get slightly faster code by compiling all of your code with -DADEPT_STACK_THREAD_UNSAFE. This uses a standard (i.e. non-thread-local) global variable to point to the currently active stack object, which is slightly faster to access.
- If you compile with the -g option to store debugging symbols, your object files and executable will be much larger because every mathematical statement in the file will have the name of its associated templated type stored in the file, and these names can be long. Once you have debugged your code, you may wish to omit debugging symbols from production versions of the executable. There appears to be no performance penalty associated with the debugging symbols, at least with the GNU C++ compiler.
- A high compiler optimization setting is recommended to inline the function calls associated with mathematical expressions. On the GNU C++ compiler, the -o3 setting is recommended.
- By default the Jacobian functions are compiled to process a strip of rows or columns of the Jacobian matrix at once. The optimum width of the strip depends on your platform, and you may wish to change it. To make the Jacobian functions process *n* rows or columns at once, recompile the *Adept* library with —DADEPT_MULTIPASS_SIZE=*n*.
- If you suspect memory usage is a problem, you may investigate the memory used by *Adept* by simply sending your Stack object to a stream, e.g. "std::cout << stack". You may also use the memory() member function, which returns the total number of bytes used. Further details of similar functions is given in section 2.8.

4.4 Exceptions thrown by the Adept library

Some functions in the *Adept* library can throw exceptions, and the exceptions that can be thrown are typically derived from either adept::autodiff_exception or adept::array_exception. These classes are derived from adept::exception, which is itself derived from std::exception. Most indicate an error in the users code, usually associated with calling *Adept* functions in the wrong order.

An overly comprehensive exception-catching implementation that takes different actions depending on whether a specific *Adept* exception, an exception related to automatic differentiation, a general *Adept* exception, or a non-*Adept* exception is thrown, could have the following form:

```
try {
  adept::Stack stack;
  // ... Code using the Adept library goes here ...
catch (adept::stack_already_active& e) {
  // Catch a specific Adept exception
  std::cerr << "Error: " << e.what() << std::endl;
  // ... any further actions go here ...
catch (adept::autodiff_exception& e) {
  // Catch any Adept exception related to automatic differentiation not yet caught
  std::cerr << "Error: " << e.what() << std::endl;
  // ... any further actions go here ...
catch (adept::exception& e) {
  // Catch any other Adept exception not yet caught
 std::cerr << "Error: " << e.what() << std::endl;
  // ... any further actions go here ...
catch (...) {
  // Catch any exceptions not yet caught
  std::cerr << "An error occurred" << std::endl;</pre>
  // ... any further actions go here ...
```

All exceptions implement the what () member function, which returns a const char* containing an error message.

4.4.1 General exceptions

The following exceptions are not specific to arrays or automatic differentiation and inherit directly from adept::exception::

feature_not_available This exception is thrown by deprecated functions, such as Stack::start(). It is also thrown by functions that are not available because a certain library is not being used, such as inv if LAPACK was not available at compile time.

4.4.2 Automatic-differentiation exceptions

The following exceptions relate to automatic differentiation (the functionality described in chapter 2), and all are in the adept namespace:

gradient_out_of_range This exception can be thrown by the adouble::get_gradient member function if the index to its gradient is larger than the number of gradients stored. This can happen if the adouble object was created after the first adouble::set_gradient call since the last Stack::new_recording call. The first adouble::set_gradient call signals to the *Adept* stack that the main algorithm has completed and so memory can be allocated to store the gradients ready for a forward or reverse pass through the differential statements. If further adouble objects are created then they may have a gradient index that is out of range of the memory allocated.

gradients_not_initialized This exception can be thrown by functions that require the list of working gradients to have been initialized (particularly the functions Stack::compute_tangent_linear and Stack::compute_adjoint). This initialization occurs when adouble::set_gradient is called.

stack_already_active This exception is thrown when an attempt is made to make a particular Stack object "active", but there already is an active stack in this thread. This can be thrown by the Stack constructor or the Stack: activate member function.

dependents_or_independents_not_identified This exception is thrown when an attempt is made to compute a Jacobian but the independents and/or dependents have not been identified.

- wrong_gradient This exception is thrown by the adouble::append_derivative_dependence if the adouble object that it is called from is not the same as that of the most recent adouble::add_derivative_dependence.
- non_finite_gradient This exception is thrown if the users code is compiled with the preprocessor variable ADEPT_TRACK_NON_FINITE_GRADIENTS defined, and a mathematical operation is carried out for which the derivative is not finite. This is useful to locate the source of non-finite derivatives coming out of an algorithm.

4.4.3 Array exceptions

The following exceptions relate to arrays (the functionality described in chapter 3), and all are in the adept namespace:

- **size_mismatch** A mathematical operation taking two arguments has been applied to array expressions that are not of the same size. The same exception is thrown if an array expression is applied to an array of a different size.
- **inner_dimension_mismatch** Matrix multiplication has been attempted with arrays whose inner dimensions don't agree.
- **empty_array** An empty array has been used in an operation when a non-empty array is required; for example, if an attempt is made to link an array to an empty array (see section 3.2 for more information on linking).
- invalid_dimension Attempt to create an array with a negative dimension.
- index_out_of_bounds An element or range of elements has been requested from an array but one of the indices provided is out of range; for a dimension of length n, the index is not in the range 0 to n-1. Note that bounds checking is only applied if the preprocessor variable ADEPT_BOUNDS_CHECKING is defined.
- matrix_ill_conditioned An attempt has been made to factorize an ill-conditioned matrix (either via solve or inv).

4.5 Configuring the behaviour of Adept

The behaviour of the Adept library can be changed by defining one or more of the Adept preprocessor variables. This can be done either by editing the adept/base.h file and uncommenting the relevant #define lines, or by compiling your code with -Dxxx compiler options (replacing xxx by the relevant preprocessor variable. There are two types of preprocessor variable: the first types only apply to the compilation of user code, while the second types require the Adept library to be recompiled.

4.5.1 Modifications not requiring a library recompile

The preprocessor variables that apply only to user code and do not require the *Adept* library to be recompiled are as follows:

ADEPT_STACK_THREAD_UNSAFE If this variable is defined, the currently active stack is stored as a global variable but is not defined to be "thread-local". This is slightly faster, but means that you cannot use multi-threaded code with separate threads holding their own active Stack object. Note that although defining this variable does not require a library recompile, all source files that make up a single executable must be compiled with this option (or all not be).

- ADEPT_RECORDING_PAUSABLE This option enables an algorithm to be run both with and without automatic differentiation from within the same program via the functions Stack::pause_recording() and Stack::continue_recording(). Note that although defining this variable does not require a library recompile, all source files that make up a single executable must be compiled with this option (or all not be). Further details on this option are provided in section 2.6.2.
- **ADEPT_NO_AUTOMATIC_DIFFERENTIATION** This option turns off automatic differentiation by treating adouble objects as double. It is useful if you want to compile one source file twice to produce versions with and without automatic differentiation. Further details on this option are provided in section 2.6.3.
- **ADEPT_TRACK_NON_FINITE_GRADIENTS** Often when an algorithm is first converted to use an operator-overloading automatic differentiation library, the gradients come out as Not-a-Number or Infinity. The reason is often that the algorithm contains operations for which the derivative is not finite (e.g. \sqrt{a} for a=0), or constructions where a non-finite value is produced but subsequently made finite (e.g. $\exp(-1.0/a)$ for a=0). Usually the algorithm can be recoded to avoid these problems, if the location of the problematic operations can be identified. By defining this preprocessor variable, a non_finite_gradient exception will be thrown if any operation results in a non-finite derivative. Running the program within a debugger (and ensuring that the exception is not caught within the program) enables the offending line to be identified.
- **ADEPT_INITIAL_STACK_LENGTH** This preprocessor variable is set to an integer, and is used as the default initial amount of memory allocated for the recording, in terms of the number of statements and operations.
- **ADEPT_REMOVE_NULL_STATEMENTS** If many variables in your code are likely to be zero then redundant operations will be added to the list of differential statements. For example, the assignment $a = b \times c$ with active variables b and c both being zero results in the differential statement $\delta a = 0 \times \delta b + 0 \times \delta c$. This preprocessor variable checks for zeros and removes terms on the right-hand-side of differential statements if it finds them. In this case it would put $\delta a = 0$ on the stack instead. This option slows down the recording stage, but speeds up the subsequent use of the recorded stack for adjoint and Jacobian calculations. The speed up of the latter is only likely to exceed the slow down of the former if your code contains many zeros. For most codes, this option causes a net slow down.
- adouble objects are only used in the return values for functions, then defining this preprocessor variable will lead to slightly faster code, because it will be assumed that when a copy constructor is called, the index to its gradient can simply be copied because the object being copied will shortly be destructed (otherwise communication with the Stack object is required to unregister one and immediately register the other). You need to be sure that the code being compiled with this option does not invoke the copy constructor in any other circumstances. Specifically, it must not include either of these constructions: "adouble x = y;" or "adouble x (y);", where y is an adouble object. If it does, then strange errors will occur.
- **ADEPT_BOUNDS_CHECKING** If this variable is defined, check that all array indices are within the bounds of the array throwing an index_out_of_bounds exception if necessary. If this variable is not defined then these checks are not performed, which is faster but means that attempts to access arrays out of bounds will result either of corruption of other memory used by the process, or a segmentation fault.

4.5.2 Modifications requiring a library recompile

The preprocessor variables that require the *Adept* library to be recompiled are as follows. Note that if these variables are used they must be the same when compiling both the library and the user code. This is safest to implement by editing section 2 of the adept/base.h header file.

ADEPT_FLOATING_POINT_TYPE If you want to compile *Adept* to use a precision other than double for the Real type, and hence for automatic differentiation, then define this preprocessor variable to be the floating-point type required, e.g. float or long double. To use from the compiler command-line, use the argument -DADEPT_FLOATING_POINT_TYPE=float or -DADEPT_FLOATING_POINT_TYPE="long double".

- **ADEPT_STACK_STORAGE_STL** Use the C++ standard template library vector or valarray classes for storing the recording and the list of gradients, rather than dynamically allocated arrays. In practice, this tends to slow down the code.
- **ADEPT_MULTIPASS_SIZE** This is set to an integer, invariably a power of two, specifying the number of rows or columns of a Jacobian that are calculated at once. The optimum value depends on the platform and the capability of the compiler to optimize loops whose length is known at compile time.
- ADEPT_MULTIPASS_SIZE_ZERO_CHECK This is also set to an integer; if it is greater than ADEPT_MULTIPASS_SIZE, then the Stack::jacobian_reverse function checks gradients are non-zero before using them in a multiplication.
- ADEPT_THREAD_LOCAL This can be used to specify the way that thread-local storage is declared by your compiler. Thread-local storage is used to ensure that the *Adept* library is thread-safe. By default this variable is not defined initially, and then later in adept/base.h it is set to __declspec(thread) on Microsoft Visual C++, an empty declaration on Mac (since thread-local storage is not available on many Mac platforms) and __thread otherwise (appropriate for at least the GCC, Intel, Sun and IBM compilers). To override the default behaviour, define this variable yourself in adept/base.h.

4.6 Frequently asked questions

Why are all the gradients coming out of the automatic differentiation zero? You have almost certainly omitted or misplaced the call of the adept::Stack member function "new_recording()". It should be placed after the independent variables in the algorithm have been initialized, but before any subsequent calculations are performed on these variables. If it is omitted or placed before the point where the independent variables are initialized, the differential statements corresponding to this initialization (which are all of the form $\delta x = 0$), will be placed in the list of differential statements and will unhelpfully set to zero all your gradients right at the start of a forward pass (resulting from a call to forward()) or set them to zero right at the end of a reverse pass (resulting from a call to reverse()).

Why are the gradients coming out of the automatic differentiation NaN or Inf (even though the value is correct)?

This can occur if the algorithm contains operations for which the derivative is not finite (e.g. \sqrt{a} for a=0), or constructions where a non-finite value is produced but subsequently made finite (e.g. $\exp(-1.0/a)$ for a=0). Usually the algorithm can be recoded to avoid these problems, if the location of the problematic operations can be identified. The simplest way to locate the offending statement is to recompile your code with the -g option and the ADEPT_TRACK_NON_FINITE_GRADIENTS preprocessor variable set (see section 4.5.1). Run the program within a debugger and a non_finite_gradient exception will be thrown, which if not caught within the program will enable you to locate the line in your code where the problem originated. You may need to turn optimizations off (compile with -00) for the line identification to be accurate. Another approach is to add the following in a C++ source file:

```
#include <fenv.h>
int _feenableexcept_status = feenableexcept(FE_INVALID|FE_DIVBYZERO|FE_OVERFLOW);
```

This will cause a floating point exception to be thrown when a NaN or Inf is generated, which can again be located in a debugger.

Why are the gradients coming out of the automatic differentiation wrong? Before suspecting a bug in *Adept*, note that round-off error can lead to incorrect gradients even in hand-coded differential code. Consider the following:

```
int main() {
   Stack stack;
   adouble a = 1.0e-26, b;
   stack.new_recording();
   b = sin(a) / a;
```

```
b.set_gradient(1.0);
stack.compute_adjoint();
std::cout << "a=" << a << ", b=" << b << ", db/da=" << a.get_gradient() << "\n";
}</pre>
```

We know that near a=0 we should have b=1 and the gradient should be 0. But running the program above will give a gradient of 1.71799e+10. If you hand-code the gradient, i.e.

```
double A = 1.0e-26;
double dB_dA = cos(A)/A - sin(A) / (A*A);
```

you will you will also get the wrong gradient. You can see that the answer is the difference of two very large numbers and so subject to round-off error. This example is therefore not a bug of *Adept*, but a limitation of finite-precision machines. To check this, try compiling your code using either the ADOL-C or CppAD automatic differentiation tools; I have always found these tools to give exactly the same gradient as *Adept*. Unfortunately, round-off error can build up over many operations to give the wrong result, so there may not be a simple solution in your case.

Can Adept reuse a stored tape for multiple runs of the same algorithm but with different inputs? No. Adept does not store the full algorithm in its stack (as ADOL-C does in its tapes, for example), only the derivative information. So from the stack alone you cannot rerun the function with different inputs. However, rerunning the algorithm including recomputing the derivative information is fast using Adept, and is still faster than libraries that store enough information in their tapes to enable a tape to be reused with different inputs. It should be stressed that for any algorithm that includes different paths of execution ("if" statements) based on the values of the inputs, such a tape would need to be rerecorded anyway. This includes any algorithm containing a look-up table.

Why does my code crash with a segmentation fault? This means it is trying to access a memory address not belonging to your program, and the first thing to do is to run your program in a debugger to find out at what point in your code this occurs. If it is in the adept::aReal constructor (note that aReal is synonymous with adouble), then it is very likely that you have tried to initiate an adept::adouble object before initiating an adept::Stack object. As described in section 2.3.1, there are good reasons why you need to initialize the adept::Stack object first.

How can I interface *Adept* **with a matrix library such as Eigen?** Unfortunately the use of expression templates in *Adept* means that it does not work optimally (if it works at all) with third-party matrix libraries that use expression templates. This is the reason why Adept 2.0 combines array functionality with automatic differentiation in a single expression-template framework.

Do you have plans to enable Adept to produce Hessian matrices? Not in the near future; refining the array functionality is a higher priority at the moment. However, if your objective function $J(\mathbf{x})$ (also known as a cost function or penalty function) has a quadratic dependence on each of the elements of $\mathbf{y}(\mathbf{x})$, where \mathbf{y} is a nonlinear function of the independent variables \mathbf{x} , then the Hessian matrix $\nabla^2_{\mathbf{x}}J$ can be computed from the Jacobian matrix $\mathbf{H} = \partial \mathbf{y}/\partial \mathbf{x}$. This is the essence of the Gauss-Newton and Levenberg-Marquardt algorithms. Consider the optimization problem of finding the parameters \mathbf{x} of nonlinear model $\mathbf{y}(\mathbf{x})$ that provides the closest match to a set of "observations" \mathbf{y}^o in a least-squares sense. For maximum generality we add constraints that penalize differences between \mathbf{x} and a set of a priori values \mathbf{x}^a , as well as a regularization term. In this case the objective function could be written as

$$J(\mathbf{x}) = [\mathbf{y}(\mathbf{x}) - \mathbf{y}^o]^{\mathsf{T}} \mathbf{R}^{-1} [\mathbf{y}(\mathbf{x}) - \mathbf{y}^o] + [\mathbf{x} - \mathbf{x}^a]^{\mathsf{T}} \mathbf{B}^{-1} [\mathbf{x} - \mathbf{x}^a] + \mathbf{x}^{\mathsf{T}} \mathbf{T} \mathbf{x}.$$
(4.1)

Here, all vectors are treated as column vectors, \mathbf{R} is the error covariance matrix of the observations, \mathbf{B} is the error covariance matrix of the *a priori* values, and \mathbf{T} is a Twomey-Tikhonov matrix that penalizes either spatial gradients or curvature in \mathbf{x} . The Hessian matrix is then given by

$$\nabla_{\mathbf{x}}^{2} J = \mathbf{H}^{\mathrm{T}} \mathbf{R}^{-1} \mathbf{H} + \mathbf{B}^{-1} + \mathbf{T}, \tag{4.2}$$

which can be coded up using Adept to compute **H**. Each term on the right-hand-side of (4.2) has its corresponding term in (4.1), so it is easy to work out what the Hessian would look like if only a subset of the terms in (4.1) were present.

Why doesn't the ternary operator work? Some compilers will fail to compile the following function:

```
adept::adouble piecewise(adept::adouble x) {
  return x < 1.0 ? x*x : 2.0*x-1.0;
}</pre>
```

The reason is that these compilers require that the two possible outcomes of the ternary operator have the same type, but due to the use of expression templates, the types of these mathematical expressions are actually different. The ternary operator cannot be overloaded to allow such arguments. The solution is to explicitly convert the outcomes to adouble:

```
adept::adouble piecewise(adept::adouble x) {
  return x < 1.0 ? adept::adouble(x*x) : adept::adouble(2.0*x-1.0);
}</pre>
```

Why is my executable so huge? Probably you are including debugging symbols by compiling with the -g option. Expression templates need long strings to describe them, so this extra content can increase the size of object files and executables by a factor of ten. This does not slow down execution, but for production code you may wish to compile without debugging symbols.

4.7 Copyright and license for *Adept* software

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In addition to the legally binding terms of the license, it is *requested* that:

- You cite Hogan (2014) in publications describing algorithms and software that make use of the *Adept* library. While not not a condition of the license, this is good honest practice in science and engineering.
- If you make modifications to the *Adept* library that might be useful to others, you release your modifications under the terms of the Apache License, Version 2.0, so that they are available to others and could also be merged into a future official version of *Adept*. If you do not state the license applied to your modifications then by default they will be under the terms of the Apache License. You will retain copyright of your modifications, but if your modifications are written in the course of employment then under almost all circumstances (including employment by a University) it is your employer who holds the copyright. Therefore you should obtain permission from them to release your modifications under the Apache License.

Note that other source files in the *Adept* package used for demonstrating and benchmarking *Adept* are released under the GNU all-permissive license*, which is specified at the top of all files it applies to.

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Adept version 1.0 was released under the terms of the GNU General Public License (GPL) and so could not be released as part of a larger work unless the entire work was released under the conditions of the GPL. It is hoped that the switch to the Apache License will facilitate wider use of Adept.

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