AUTOMATED DERIVATION OF THE ADJOINT OF HIGH-LEVEL TRANSIENT FINITE ELEMENT PROGRAMS*

P. E. FARRELL†, D. A. HAM‡, S. W. FUNKE§, AND M. E. ROGNES¶

Abstract. In this paper we demonstrate a new technique for deriving discrete adjoint and tangent linear models of a finite element model. The technique is significantly more efficient and automatic than standard algorithmic differentiation techniques. The approach relies on a high-level symbolic representation of the forward problem. In contrast to developing a model directly in Fortran or C++, high-level systems allow the developer to express the variational problems to be solved in near-mathematical notation. As such, these systems have a key advantage: since the mathematical structure of the problem is preserved, they are more amenable to automated analysis and manipulation. The framework introduced here is implemented in a freely available software package named dolfin-adjoint, based on the FEniCS Project. Our approach to automated adjoint derivation relies on run-time annotation of the temporal structure of the model and employs the FEniCS finite element form compiler to automatically generate the low-level code for the derived models. This approach requires only trivial changes to a large class of forward models, including complicated time-dependent nonlinear models. The adjoint model automatically employs optimal checkpointing schemes to mitigate storage requirements for nonlinear models, without any user management or intervention. Furthermore, both the tangent linear and adjoint models naturally work in parallel, without any need to differentiate through calls to MPI or to parse OpenMP directives. The generality, applicability, and efficiency of the approach are demonstrated with examples from a wide range of scientific applications.

Key words. FEniCS project, libadjoint, dolfin-adjoint, adjoint, tangent linear, code generation

AMS subject classifications. 65N30, 68N20, 49M29

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1. Introduction. Adjoint models are key ingredients in many algorithms of computational science, such as parameter identification [40], sensitivity analysis [6], data assimilation [28], optimal control [30], adaptive observations [42], predictability analysis [36], and error estimation [4]. While deriving the adjoint model associated with a linear stationary forward model is straightforward, the development and implementation of adjoint models for nonlinear or time-dependent forward models is notoriously difficult for several reasons. First, each nonlinear operator of the forward model must be differentiated, which can be difficult for complex models. Second, the

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control flow of the adjoint model runs backwards, from the final time to the initial
time, and requires access to the solution variables computed during the forward run
if the forward problem is nonlinear. Since it is generally impractical for physically
relevant simulations to store all variables during the forward run, the adjoint model
developer must implement some checkpointing scheme that balances recomputation
and storage [17]. The control flow of such a checkpointing scheme must alternate be-
tween the solution of forward variables and that of adjoint variables and is thus highly
nontrivial to implement by hand on a large and complex code. For parallel computa-
tions, these difficulties are magnified by the fact that the control flow of parallel
communications reverses in the adjoint solve: forward sends become adjoint receives,
and forward receives become adjoint sends [46].

The traditional approach to model development is to implement the forward code
by hand in a low-level language (typically Fortran or C++). While this allows
the programmer a high degree of control over each memory access and floating point
operation, implementing these codes usually takes a large amount of time, and the
mathematical structure of the problem to be solved is irretrievably interwoven with
implementation details of how the solution is to be achieved. Then the adjoint code is
produced, either by hand or with the assistance of an algorithmic differentiation (AD)
tool (Figure 1.1). Such AD tools take as input a forward model written in a low-level
language and derive the associated discrete adjoint model, through some combination
of source-to-source transformations and operator overloading. However, this process
requires expert knowledge of both the tool and the model to be differentiated [39,
p. xii]. The root cause of the difficulty which AD tools have is that they operate on
low-level code in which implementation details and mathematics are inseparable and
therefore must both be differentiated: AD tools must concern themselves with matters
such as memory allocations, pointer analyses, I/O, and parallel communications (e.g.,
MPI or OpenMP).

A variant of this approach is to selectively apply AD to small sections of the model,
and then to connect and arrange these differentiated routines by hand to assemble the
discrete adjoint equations [15, 9, 38]. This approach attempts to reintroduce as much
as possible of the distinction between mathematics and implementation; however, it
requires even more expertise than a naïve black-box application of AD.

AD treats a model as a sequence of elementary instructions, where an instruction
is typically a native operation of the programming language such as addition, mul-
multiplication, or exponentiation. Instead, we consider a new, higher-level abstraction
for developing discrete adjoint models: to treat the model as a sequence of equation
solves. This offers an alternative approach to the development of discrete adjoint
models and is implemented in an open-source software library called libadjoint (see
section 2).
When libadjoint is applied to a low-level forward code, the developer must annotate the forward model. This involves embedding calls to the libadjoint library that record the temporal structure of the equations as they are solved. The recorded information is analogous to a tape in AD, but at a higher level of abstraction. Using this information, libadjoint can symbolically manipulate the annotated system to derive the structure of the adjoint equations or the tangent linear equations. If the adjoint developer further supplies callback functions for each operator that features in the annotation and any necessary derivatives, libadjoint can assemble each adjoint or tangent linear equation as required. The library therefore relieves the developer of deriving the adjoint equations, managing the complex life cycles of forward and adjoint variables, and implementing a checkpointing scheme. With this strategy, the task of developing the adjoint model (which requires significant expertise) is replaced with the tasks of describing and modularizing it (which are usually much more straightforward).

The aim of this work is to apply libadjoint to automatically derive the adjoint of models written in a high-level finite element system. In such systems, the discrete variational formulation is expressed in code which closely mimics mathematical notation. The low-level details of finite element assembly and numerical linear algebra are delegated to the system itself. For example, in the Sundance C++ library, the assembly is achieved by runtime Fréchet differentiation of the specified variational form [35]. In the FEniCS environment, the variational form is passed to a dedicated finite element form compiler, which generates low-level code for its assembly [26, 33, 34]. By exploiting optimizations that are impractical to perform by hand, such systems can generate very efficient implementations [27, 41, 37]. Moreover, a major advantage of this clean separation between mathematical intention and computer implementation is that it enables the automatic mathematical analysis and manipulation of the variational form. The absence of this separation in low-level models inhibits automated analysis at the level of variational forms. In addition, high-level systems can often provide automated variational form differentiation/linearization capabilities [2, 1, 35, 43], which can be of particular interest for adjoint models.

The main contribution of this paper is a new framework for the automated derivation of the discrete adjoint and tangent linear models of forward models implemented in the FEniCS software environment. The strategy is illustrated in Figure 1.2. In FEniCS model code, each equation solve may be naturally expressed as a single function call. This matches the basic abstraction of libadjoint exactly and there-
fore makes the integration of FEniCS and libadjoint particularly straightforward. Consequently, for a large class of forward models implemented in FEniCS, libadjoint can automatically derive the discrete adjoint model with only minor additions to the code. The necessary derivative terms are automatically computed using the form linearization capabilities of UFL [2, 1]. The adjoint equations derived by libadjoint are themselves valid FEniCS input, and the low-level adjoint code is generated using the same finite element form compiler as the forward model. As a result, the derived adjoint models approach optimal efficiency, automatically employ optimal checkpointing schemes, and inherit parallel support from the forward models.

This paper is organized as follows. In section 2, we describe the fundamental abstraction underlying libadjoint and give a brief review of libadjoint. In section 3, we outline the implementation of transient finite element forward models in the FEniCS framework. The integration of FEniCS and libadjoint is implemented in a new software framework called dolfin-adjoint, which is described in section 4. The advantages and limitations of the approach are discussed in section 5. Numerical examples drawn from a wide range of scientific applications are presented in section 6, before we make some concluding remarks in section 7.

2. The fundamental abstraction of libadjoint. In this section, we detail the basic abstraction upon which libadjoint is based, which is to treat the model as a sequence of equation solves. This abstraction applies to both stationary and time-dependent systems of partial differential equations, and to both linear and nonlinear systems.

2.1. Mathematical framework. We consider systems of discretized partial differential equations expressed in the fundamental abstract form

\[ A(u)u = b(u), \]

where \( u \) is the vector of all prognostic variables, \( b(u) \) is the source term, and \( A(u) \) is the entire discretization matrix. In the time-dependent case, \( u \) is a block-structured vector containing all the values of the unknowns at all the time levels, \( A \) is a matrix with a lower-triangular block structure containing all of the operators featuring in the forward model, and \( b \) is a block-structured vector containing all of the right-hand-side terms for all of the equations solved in the forward model. The block-lower-triangular structure of \( A \) is a consequence of the forward propagation of information through time: later values depend on earlier values, but not vice versa.

It is to be emphasised that writing the model in the format of (2.1) does not imply that the whole of \( A \) is ever assembled at once, or the whole of \( u \) stored in memory. For instance, the forward solver will typically assemble one block-row of \( A \), solve it for a block-component of \( u \), forget as much as possible, and step forward in time.

Let \( m \) be some parameter upon which the forward equations depend. For example, \( m \) could be a boundary condition, initial condition, or coefficient appearing in the equations. The tangent linear model associated with (2.1) is then given by

\[ (A + G - R) \frac{du}{dm} = -\frac{\partial F}{\partial m}, \]

where

\[ G \equiv \frac{\partial A}{\partial u} u, \]

\[ R \equiv \frac{\partial b}{\partial u}, \]
and

\[ F \equiv A(u)u - b(u). \]

The unknown in (2.2) is the \( du/dm \) matrix, the Jacobian of the solution \( u \) with respect to the parameters \( m \). The \( G \) matrix of (2.3) arises because of the nonlinear dependency of the operator \( A \). \( A \) is a matrix (i.e., a rank-2 tensor), so differentiating it with respect to \( u \) yields a rank-3 tensor; the following contraction with \( u \) over the middle index reduces the rank again to 2. More precisely, in index notation we have

\[ G_{ik} \equiv \sum_j \frac{\partial A_{ij}}{\partial u_k} u_j. \]

The \( R \) matrix of (2.4) is the Jacobian of the right-hand-side \( b \) with respect to the solution \( u \). Written in index notation,

\[ R_{ij} \equiv \frac{\partial b_i}{\partial u_j}. \]

Let \( J \) be some functional of the solution \( u \). \( J \) is a function that takes in the system state and returns a single scalar diagnostic. For example, in aeronautical design, \( J \) may be the drag coefficient associated with a wing; in meteorology, \( J \) may be the weighted misfit between observations of the atmosphere and model results. The adjoint model associated with (2.1) is given by

\[ (A + G - R)^* z = \frac{\partial J}{\partial u}, \]

where \( z \) is the adjoint solution associated with \( J \) and * denotes the conjugate transpose.

To make matters more concrete, consider the following example. Suppose that the forward model approximately solves the time-dependent viscous Burgers’ equation,

\[ \frac{\partial u}{\partial t} + u \cdot \nabla u - \nabla^2 u = f, \]

for the velocity \( u \), subject to some suitable boundary conditions and a supplied initial condition \( u(0) = g \) with source term \( f \). For simplicity, suppose that the model linearizes the nonlinear advective term around the solution of the previous timestep (other choices are also possible). Discretizing with the Galerkin finite element method in space and the forward Euler method in time yields the timestep iteration

\[ u_{n+1} \leftarrow (M - \Delta t V(u_n) - \Delta t D)u_n + \Delta t f_n, \]

where \( n \) is the timelevel, \( \Delta t \) is the timestep, \( M \) is the mass matrix, \( D \) is the diffusion matrix, and \( V(u) \) is the advection matrix assembled at a given velocity \( u \). For brevity, define

\[ T(\cdot) = \Delta t V(\cdot) + \Delta t D - M. \]

System (2.10) can be cast into the form of (2.1) by writing \( N \) timestep iterations as

\[ \begin{pmatrix} I \\ T(u_0) \\ T(u_1) \\ \vdots \\ T(u_{N-1}) \end{pmatrix} \begin{pmatrix} u_0 \\ u_1 \\ u_2 \\ \vdots \\ u_N \end{pmatrix} = \begin{pmatrix} g \\ \Delta t f_0 \\ \Delta t f_1 \\ \vdots \\ \Delta t f_{N-1} \end{pmatrix}. \]
Using (2.8), the adjoint system is given by

\[
\begin{pmatrix}
I^* & (T(u_0) + \frac{\partial T(u_0)}{\partial u_0} u_0)^* \\
M^* & (T(u_1) + \frac{\partial T(u_1)}{\partial u_1} u_1)^* \\
& \ddots \\
& & \ddots \\
& & & M^* \\
\end{pmatrix}
\begin{pmatrix}
z_0 \\
z_1 \\
\vdots \\
z_N \\
\end{pmatrix} = \frac{\partial J}{\partial u},
\]

where the contraction of the derivative of a matrix with a vector is defined in (2.6).

The adjoint system reverses the temporal flow of information: where the forward and tangent linear models are block-lower-triangular, the adjoint model is block-upper-triangular, as visible in (2.13). The adjoint system is therefore typically solved by backward substitution: the adjoint variable associated with the end of time is solved for first, and then the solution proceeds backwards in time.

At its heart, libadjoint takes models cast in the form of (2.1), and derives, assembles, and solves the tangent linear (2.2) and adjoint (2.8) systems block-row by block-row.

### 2.2. Libadjoint

Applying libadjoint to a model breaks down into two main tasks. The automation of these two tasks by dolfin-adjoint, the software package presented in this paper, is described in section 4.

The first task, referred to as **annotation**, is to describe the forward code in the form of the fundamental abstraction (2.1). By describing the forward code in this form, libadjoint automates the reasoning necessary to derive the adjoint (2.8) and tangent linear (2.2) systems. As each equation is solved, the necessary semantic information about that equation is recorded. In particular, each equation records the variable solved for, the operators (matrices) that feature in the equation, and any dependencies these operators have on previous variables. This annotation is effected by making calls to library functions offered by the libadjoint application programming interface (API). In a low-level code, these calls must be inserted manually by the model developer, as the high-level semantic structure will have been obscured in the process of implementing the model. The annotation enables libadjoint to symbolically derive the structure of the discrete adjoint and tangent linear systems; however, without further information it cannot assemble the actual adjoint or tangent linear equations. The operators in the annotation are mere abstract handles.

The second task is to supply libadjoint with function callbacks for the operators that feature in the annotation. If these operators depend on previously computed variables, their derivatives must also be supplied; the code for these derivative callbacks may be written by hand or may be generated with an AD tool. With these callbacks, the derived equations may be automatically assembled. By modularizing the forward model in this manner, libadjoint can drive the assembly of the forward, tangent linear, and adjoint models, by calling the appropriate callbacks in the correct sequence. In a low-level code, these callbacks must be written manually. This can be a significant burden if the original forward code is poorly modularized.

The use of libadjoint offers several advantages. It can be applied to models for which black-box AD is intractable, and gains the speed and efficiency benefits of applying AD judiciously [15, 9, 38]. Using libadjoint makes development systematic: each incremental step in its application may be rigorously verified. As libadjoint internally
The adjoint of high-level finite element programs derives a symbolic representation of the discrete adjoint equations, it can compute when a forward or adjoint variable is no longer necessary, and thus the model developer is relieved of the management of variable deallocation. Furthermore, libadjoint can automatically check the consistency of the adjoint computed with the original forward model; this check greatly improves the maintainability of adjoint codes, as developers can be immediately notified when a change to the forward model is not mirrored in the adjoint model. Finally, as libadjoint has sufficient information to reassemble the forward equations, it is possible to implement checkpointing schemes entirely within the library itself. Checkpointing is a crucial feature for the efficient implementation of the adjoint of a time-dependent nonlinear model, but its implementation can be prohibitively difficult. The availability of optimal checkpointing schemes within libadjoint is a significant advantage.

3. The FEniCS system. The FEniCS Project is a collection of software components for automating the solution of differential equations [32, 31]. These components include the Unified Form Language (UFL) [2, 1], the FEniCS Form Compiler (FFC) [26], and DOLFIN [33, 34]. In the following, we only briefly outline the FEniCS pipeline and refer the reader to the aforementioned references for more information.

One of the key features of the FEniCS components is the use of code generation, and in particular domain-specific code generation, for finite element variational formulations: the user specifies the discrete variational problem to be solved in the domain-specific language UFL, the syntax of which mimics and encodes the mathematical formulation of the problem. Based on this high-level formulation, a special-purpose finite element form compiler generates optimized low-level C++ code for the evaluation of local element tensors. The generated code is then used by DOLFIN to perform the global assembly and numerical solution. DOLFIN also provides the underlying data structures such as meshes, function spaces, boundary conditions, and function values. DOLFIN provides both a C++ and a Python interface. For the C++ interface, the UFL specification and the form compilation must take place offline, and the generated code is explicitly included by the user. In the Python interface, the functionality is seamlessly integrated by way of runtime just-in-time compilation.

The approach taken by dolfin-adjoint is only applicable to models written using the Python interface, as only the Python interface has runtime access to the symbolic description of the forward model in UFL format.

DOLFIN abstracts the spatial discretization problem but not the temporal discretization problem. Transient DOLFIN-based solvers typically consist of a handwritten temporal loop in which one or more discrete variational (finite element) problems are solved with DOLFIN in each iteration. An overview of common operations relevant to dolfin-adjoint is given in Table 3.1.

<table>
<thead>
<tr>
<th>DOLFIN function signature</th>
<th>Short description</th>
</tr>
</thead>
<tbody>
<tr>
<td>solve(lhs == rhs, u, bcs)</td>
<td>Solve variational problem</td>
</tr>
<tr>
<td>u.assign(u_0)</td>
<td>Copy function u_0 to u</td>
</tr>
<tr>
<td>assemble(a)</td>
<td>Assemble variational form</td>
</tr>
<tr>
<td>bc.apply(A)</td>
<td>Apply a boundary condition to a matrix</td>
</tr>
<tr>
<td>solve(A, x, b)</td>
<td>Solve linear system</td>
</tr>
<tr>
<td>project(u, V)</td>
<td>Project u onto the function space V</td>
</tr>
</tbody>
</table>
from dolfin import *
from dolfin_adjoint import *
n = 30
mesh = UnitInterval(n)
V = FunctionSpace(mesh, "CG", 2)
ic = project(Expression("sin(2*pi*x[0])"), V)
u = Function(ic, name="Velocity")
u_next = Function(V, name="NextVelocity")
v = TestFunction(V)
u = Constant(0.0001)
timestep = Constant(1.0/n)
F = ((u_next - u)/timestep*v
 + u_next*grad(u_next)*v + nu*grad(u_next)*grad(v))*dx
bc = DirichletBC(V, 0.0, "on_boundary")
t = 0.0; end = 0.2
while (t <= end):
    solve(F == 0, u_next, bc)
u.assign(u_next)
t += float(timestep)

adj_inc_timestep()

Fig. 3.1. DOLFIN code for a simple discretization of the Burgers equation (2.9) with dolfin-
adjoint annotations. The dolfin-adjoint module overloads the existing solve and assign functions
(indicated in red), and allows the user to specify names of Functions for convenience. The only
change to the code body is the introduction of a call to adj_inc_timestep to indicate to libadjoint
that a new timestep is commencing (indicated in blue).

At the highest level of abstraction, a DOLFIN model developer can define the
variational problem for each timestep in terms of the unknown function, the left- and
right-hand-side forms, and boundary conditions and then call the DOLFIN solve
function. If the variational problem is linear (represented by a left-hand-side bilinear
form and a right-hand-side linear form, \( a = L \)), the linear system of equations is
assembled and solved under the hood. If the variational problem is nonlinear (rep-resented by a left-hand-side rank-1 form and a zero right-hand side, \( F = 0 \)), a Newton
iteration is invoked in which the Jacobian of the variational form \( F \) is derived auto-
matically, and the linear system in each Newton iteration assembled and solved until
the iteration has converged. The derivation of the Jacobian employs the algorithmic
differentiation algorithms of UFL: because the form is represented in a high-level
abstraction, the symbolic differentiation of the form is straightforward [1, sections
17.5.2 and 17.7]. When moving from one timestep to the next, the assign function is
typically used to update the previous function with the new value. A sample solver
demonstrating this type of usage, solving the nonlinear Burgers' equation, is listed in
Figure 3.1.

However, DOLFIN also supports more prescriptive programming models, in which
explicit calls are employed to assemble matrices and solve the resulting linear systems.
If the variational problem is linear and the left-hand side is constant, the assembly of
the stiffness matrix may occur outside the temporal loop, and the matrix be reused.
The solution of the linear systems may also be further controlled by specifying direct
LU or Krylov solvers, or even matrix-free solvers.
DOLFIN’s abstraction of a transient problem as an explicit sequence of variational problems exactly matches the fundamental abstraction of libadjoint; this matching of abstractions is the basis of the work presented here.

4. Applying libadjoint to DOLFIN. In this section, we present the internal details of how dolfin-adjoint integrates DOLFIN with libadjoint. This includes the annotation of the forward model execution, the recording of any necessary values, and the generation and registration of callback functions. All of these processes happen automatically, without any intervention by the model developer.

4.1. Annotations. The basic mechanism of automatically annotating DOLFIN models employed here is to overload the DOLFIN functions that change the values of variables. The overloaded versions annotate the event and then pass control to the original DOLFIN functions. All of the functions listed in Table 3.1 are annotated.

Figure 3.1 presents a Burgers’ equation model modified for use with dolfin-adjoint. The modifications and overloaded functions are highlighted. This demonstrates that only minimal source changes are required.

In a low-level model, the information that libadjoint needs to record is not explicitly represented as data in the code; therefore, annotation has to be at least partially done by hand, as the programmer must supply the information instead. By contrast, in the Python interface to DOLFIN, the equation to be assembled is explicitly represented as data at runtime; DOLFIN’s `solve` function takes as input the variational form of the equation to be solved, and so all information necessary for the annotation is available during the `solve` function. This makes the automatic annotation possible.

By contrast, when using the C++ interface to DOLFIN, the code generation happens offline, and the equations are not explicitly represented as data at runtime; it is for this reason that the automated runtime derivation of the adjoint model in the manner described here is not possible. Therefore, dolfin-adjoint only supports the use of the Python interface to DOLFIN.

When called, the adjoint-aware `solve` function inspects the left- and right-hand sides of the provided equation to determine the information needed by libadjoint. This includes the variable being solved for, the operators which feature in the equation, and their dependencies on values that were previously computed. UFL supports the interrogation of forms to automatically extract the necessary information. This information is then registered with libadjoint using the relevant API calls. When an overloaded call completes, dolfin-adjoint will record the resulting value if this is required for later use in the adjoint computation.

For nonlinear solves expressed in the form $F(u) = 0$, where $F$ is a rank-1 (vector) form, the adjoint code annotates the equivalent equation

\begin{equation}
Iu = Iu - F(u),
\end{equation}

where $I$ is the identity matrix associated with the function space of $u$, so that it can be cast into the form of a row of (2.1). While it would be possible to annotate each linear solve in the Newton iteration, this would be inefficient: with this method, the adjoint run would have to rewind through each iteration of the nonlinear solver, whereas by annotating in the manner of (4.1), only one linearized solve is necessary. This is akin to the method suggested in [14], which computes the derivative of a Newton iteration in one step by linearizing about the computed forward solution.

In the cases where a developer of DOLFIN models preassembles forms into tensors using the `assemble` function, only low-level matrices and vectors are given as inputs to
the \texttt{solve} call; the semantic information about the forms is no longer available. This problem is resolved by supplying an overloaded \texttt{assemble} function which associates the form to be assembled with the assembled tensor. When the matrix-vector version of \texttt{solve} is called, it uses this association to recover the forms involved in the equation, and annotates as described above.

The correctness of the adjoint relies on the correctness of the annotation. If the annotation does not exactly record the structure of the forward model, the gradient computed using the adjoint will be inconsistent. For example, the annotation would become inconsistent if the model accessed the underlying \texttt{.vector()} of a \texttt{Function} and changed the values by hand; this would not be recorded by libadjoint. This restriction is mitigated somewhat by the fact that the replay feature of libadjoint can be used to check the correctness of the annotation. As will be discussed in section 5.1, libadjoint has sufficient information to replay the forward equations. This can be used to automatically check the correctness of the annotation, by replaying the annotation and comparing each value to that computed during the original model run.

4.2. Callbacks. In the dolfin-adjoint code, the registration of callbacks occurs at the same time as the annotation, in the overloaded function calls. For each operator in the equations registered, Python functions are generated by dolfin-adjoint at runtime, and these are associated with the operator through the relevant libadjoint API calls. Depending on the precise details of the operator, libadjoint’s requirements vary: if the operator is on the main diagonal of $A$ in (2.1), the function returns the form itself (or its transpose, in the adjoint case), while if the operator is not on the main diagonal, the function returns the action of the form on a given input vector (or its transpose action). In either case, if the form depends on previously computed solutions, the derivative of the form with respect to these variables must be generated, along with the associated transposes. For the computation of these transposes and derivatives, we rely on the relevant features of the UFL in which the form is expressed, in particular on its powerful AD capabilities [1, sections 17.5.2 and 17.7].

The function callbacks also take references to information other than the form, so that the exact conditions of the forward solve can be recovered by dolfin-adjoint as necessary. In particular, time-dependent boundary conditions and forcing terms are implemented in DOLFIN via \texttt{Expression} classes which take in the current time as a parameter. Before a function is defined, the current value of every parameter is recorded, and the function then includes this record as part of its lexical closure. When the function is called, it restores each parameter value to the value it had when it was created. Similarly, functions take references to any Dirichlet boundary conditions that are applied to the equation as part of their lexical closure, so that the associated homogeneous Dirichlet boundary conditions may be applied to the corresponding adjoint or tangent linear equations.

Again, because of the fact that the equation to be solved is represented as data at runtime, the definition of the callbacks can happen entirely automatically. Following the registration of these callbacks, libadjoint can compose the appropriate terms at will to assemble the adjoint or tangent linear system corresponding to (2.1).

4.3. The dolfin-adjoint user interface. The highest-level interface is the function \texttt{compute gradient}: given a \texttt{Functional} $J$ and a \texttt{Parameter} $m$, it computes the gradient $\frac{dJ}{dm}$ using the adjoint solution. Example usage is given in Figure 4.1.

If the user wishes to directly access the adjoint or tangent linear solutions, the functions \texttt{compute adjoint} and \texttt{compute tlm} are available. These functions iterate over the adjoint and tangent linear solutions, with the tangent linear solutions
J = Functional(0.5*inner(u, u)*dx*dt[FINISH_TIME])

ic_param = InitialConditionParameter("Velocity")
dJdic = compute_gradient(J, ic_param)
print norm(dJdic)
plot(dJdic, interactive=True)

Fig. 4.1. Sample dolfin-adjoint user code complementing the Burgers model presented in Figure 3.1: The adjoint is generated and used to compute the gradient of the functional $J = \frac{1}{2} \int_{\Omega} u(T) \cdot u(T) dx$ with respect to the initial condition $u(0)$.

### Table 4.1

<table>
<thead>
<tr>
<th>dolfin-adjoint statement</th>
<th>Short description</th>
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<tbody>
<tr>
<td>J = Functional(0.5*inner(u, u)<em>dx</em>dt[FINISH_TIME])</td>
<td>Functional $J = \frac{1}{2} \int_{\Omega} u(T) \cdot u(T) dx$</td>
</tr>
<tr>
<td>J = Functional(0.5*inner(u, u)<em>dx</em>dt)</td>
<td>Functional $J = \frac{1}{2} \int_{\Omega(T)} u(t) \cdot u(t) dt$</td>
</tr>
<tr>
<td>m = InitialConditionParameter(&quot;Velocity&quot;)</td>
<td>Control variable for the initial velocity</td>
</tr>
<tr>
<td>compute_gradient(J, m)</td>
<td>Computes the gradient $dJ/du_0$</td>
</tr>
<tr>
<td>compute_adjoint(J)</td>
<td>Generator for the adjoint solutions</td>
</tr>
<tr>
<td>compute_tlm(m)</td>
<td>Generator for the tangent linear solutions</td>
</tr>
</tbody>
</table>

advancing in time and the adjoint solutions going backwards. A list of important dolfin-adjoint statements is given in Table 4.1.

5. Discussion.

5.1. Checkpointing. The adjoint and tangent linear models are linearizations of the forward model. If the forward model is nonlinear, then the solution computed by the forward model must be available during the execution of the linearized models: the adjoint and tangent linear models depend on the forward solution. In the tangent linear case, this is not a major burden: the tangent linear system is block-lower-triangular, like the nonlinear forward model, and so each tangent linear equation can be solved immediately after the associated forward equation, with no extra storage of the forward solutions necessary. However, as the adjoint system is solved backwards in time, the forward solution (or the ability to recompute it) must be available for the entire length of the forward and adjoint solves.

In large simulations, it quickly becomes impractical to store the entire forward solution through time at once. The alternative to storing the forward solutions is to recompute them when necessary from checkpoints stored during the forward run; however, a naïve recomputation scheme would greatly increase the computational burden of the problem to be solved. Therefore, some balance of storage and recomputation is necessary. This problem has been extensively studied in the algorithmic differentiation literature [17, 24, 55, 50], and several algorithmic differentiation tools support the automatic implementation of such a checkpointing scheme. Of particular note is the algorithm of Griewank and coworkers, which achieves logarithmic growth of the storage requirements and recomputation costs [17, 18] and is provably optimal for the case in which the number of timesteps to be performed is known in advance [19].

Despite these theoretical advances, implementing checkpointing schemes by hand is a major challenge. The complexity of programming them inhibits their widespread use. Many adjoint models do not implement checkpointing schemes [53] or implement
suboptimal checkpointing schemes. In order to implement a checkpointing scheme, the model control flow must jump between adjoint solves and forward solves, which is difficult to achieve if the model has a large or complex state. Instead, models may rely on temporal interpolation schemes to approximate the forward state. This choice introduces approximation errors into the adjoint equations, and the adjoint will no longer supply a gradient consistent with the discrete forward model [51].

The main function of the annotation is to enable libadjoint to derive the associated adjoint and tangent linear models. However, libadjoint can also use the annotation without manipulation to assemble and solve the original forward model. While this may not be of direct utility to the model developer, the ability to replay the forward model means that checkpointing schemes may be implemented entirely within libadjoint itself. When the developer requests the assembly of an adjoint equation from libadjoint, libadjoint will detect any dependencies of that adjoint equation that are not available, and will automatically recompute the relevant forward solutions from the appropriate checkpoint, without any further intervention by the model developer. libadjoint uses the revolve library [18] to identify the optimal placement of checkpoints, subject to the specified disk and memory storage limits. If the number of timesteps is known a priori, libadjoint uses revolve’s optimal offline checkpointing algorithm [17]; if this is not possible, libadjoint uses revolve’s online checkpointing algorithm [50].

Once these basic checkpointing parameters have been specified, the only change required to the model code is to add a call to the adjinc timestep function at the end of the forward time loop to indicate that the model timestep has been incremented; libadjoint’s use of revolve is fundamentally based on the timesteps of the model, but DOLFIN currently has no native concept of time or timestepping. From the perspective of the model developer, the fact that the checkpointing algorithm can be implemented entirely within libadjoint is attractive, as it allows for the adjoints of long forward runs to be achieved with optimal storage and recomputation costs for almost no extra developer effort.

5.2. Parallelism. The algorithmic differentiation of parallel programs implemented using OpenMP or MPI is a major research challenge [52, 46, 12]. Even when using an AD tool to generate the majority of the adjoint code, this challenge means that the adjoints of the communication routines are usually written by hand. This was the strategy used by the parallel adjoint MITgcm ocean model [21, 22].

In the libadjoint context, the annotation is orthogonal to the parallel implementation of the model: the fact that the matrices and solutions happen to be distributed over multiple processing units is independent of the dependency structure of the equations to be solved. However, the callbacks supplied by the developer must be parallel-aware: for example, the action callback of an operator must call the relevant parallel update routine, and the transpose action must call the adjoint of the parallel update routine. By itself, libadjoint does not remove the need for the adjoint of the parallel communication routines; the developer must still reverse the information flow of the communications to implement the transpose action callbacks.

By contrast, DOLFIN handles all of these parallel communication patterns automatically, even in the adjoint case. The high-level input to DOLFIN contains no parallel communication calls: DOLFIN derives the correct communication patterns at runtime. The adjoint equations to be solved are represented in the same UFL format as the forward model, and are passed to the same DOLFIN runtime system. The necessary parallel communication patterns for the adjoint equations are therefore
automatically derived in exactly the same way as the parallel communication patterns for the forward equations, for both the MPI and OpenMP cases [32, section 6.4]. This circumvents the need to adjoin the parallel communication calls. Indeed, there is no parallel-specific code in dolfin-adjoint: by operating at this high level of abstraction, the problem of the reversal of communication patterns in the adjoint model simply vanishes, and the adjoint model inherits the same parallel scalability properties as the forward model. The remarkably straightforward implementation of the parallel adjoint model is a major advantage of adopting the combination of a high-level finite element system and a high-level approach to deriving its adjoint.

5.3. Matrix-free models. In some scientific applications, the problems to be solved are so large that it is not practical to explicitly store the matrix associated with a single block-equation in the sense of (2.1). For example, such problems arise in the Stokes equations for mantle convection [11] and the Boltzmann transport equations for radiation transport [44]. In such situations, matrix-free solution algorithms, algorithms that never demand the whole matrix at once, are used instead. Instead of assembling a matrix $M$ and passing it to the linear solver routine, a function $f$ that computes the action $f: v \mapsto Mv$ of the matrix on a given vector $v$ is supplied.

In general, automatically differentiating programs that use matrix-free solvers is difficult. If an AD tool were to be applied in a na"ively black-box fashion, the analysis of the tool would need to trace the flow of information from the independent variables through the registration of the action function pointer into the inner loop of the matrix-free linear solver algorithm, and differentiate backwards through this chain. In the typical case, the linear solver algorithm is taken from an external library, which may be written in a different programming language and for which the source may not be immediately available. Due to this difficulty, the only previous research on algorithmically differentiating matrix-free solvers of which the authors are aware has been limited to very specific interfaces in PETSc that are confined to structured meshes [25].

However, the fact that the model takes a matrix-free approach is ultimately an implementation detail: it is a choice made in how the discrete problem is to be solved, but it does not change the discrete problem or its adjoint. The difficulties faced by applying an AD tool to models that use matrix-free solvers stem from the fact that the tool must untangle the mathematical structure of the problem from the details of how its solution is to be achieved; in the matrix-free case, this untangling is particularly difficult. However, with the high-level abstraction adopted in this work, such problems disappear: by operating at the level of equation solves, automatically deriving the discrete adjoint proceeds in exactly the same manner, and it does not matter whether those equation solves happen to assemble matrices or to use matrix-free solvers.

5.4. Limitations. The first major limitation of dolfin-adjoint is that all changes to object values must happen through the DOLFIN interface. DOLFIN permits the user to access and manipulate the raw memory addresses of function values: however, if the user modifies the underlying array, dolfin-adjoint currently does not identify that this has taken place and consequently cannot differentiate through the modification. Therefore, the derived adjoint and tangent linear models will be inconsistent with the implemented forward model. However, the replay feature described in section 4.1 mitigates this limitation somewhat: dolfin-adjoint can replay its annotation and compare it against the values recorded in the forward run, automatically identifying inconsistencies introduced in this way.

Another limitation of dolfin-adjoint is that it cannot fully automate the differ-
Differentiation of functionals with respect to mesh parameters, as is typically necessary in shape optimization. UFL is currently unable to symbolically differentiate equation operators with respect to the spatial coordinates; furthermore, differentiating through the mesh generation procedure is also necessary [13]. However, it is possible to directly use the automatically computed adjoint solutions for shape optimization via the shape calculus approach [48, 47], which circumvents the need for discretely differentiating through the mesh generation process.

6. Examples. In this section, we present three numerical examples drawn from a range of scientific applications, illustrating different discretizations and solution strategies. These examples were run with DOLFIN 1.0, libadjoint 0.9, and dolfin-adjoint 0.6. The software is freely available from http://dolfin-adjoint.org.

6.1. Cahn–Hilliard. As an initial example, the Cahn–Hilliard solver from the DOLFIN examples collection was adjoined using the techniques described above. The Cahn–Hilliard equation is a nonlinear parabolic fourth-order partial differential equation used to describe the separation of two components of a binary fluid [7, 8]:

\[ \frac{\partial c}{\partial t} - \nabla \cdot M \left( \nabla \left( \frac{df}{dc} - \epsilon^2 \nabla^2 c \right) \right) = 0 \quad \text{in } \Omega, \]

\[ M \left( \nabla \left( \frac{df}{dc} - \epsilon^2 \nabla^2 c \right) \right) = 0 \quad \text{on } \partial \Omega, \]

\[ M \epsilon^2 \nabla c \cdot \hat{n} = 0 \quad \text{on } \partial \Omega, \]

\[ c(t=0) = c_0 \quad \text{on } \Omega, \]

where \( c \) is the unknown concentration field, \( M \) and \( \epsilon \) are scalar parameters, \( \hat{n} \) is the unit normal, \( c_0 \) is the given initial condition, and \( f = 100c^2(1-c)^2 \).

In order to solve the problem using a standard Lagrange finite element basis, the fourth-order PDE is separated into two coupled second-order equations:

\[ \frac{\partial c}{\partial t} - \nabla \cdot M \nabla \mu = 0 \quad \text{in } \Omega, \]

\[ \mu - \frac{df}{dc} + \epsilon^2 \nabla^2 c = 0 \quad \text{in } \Omega, \]

where \( \mu \) is an additional prognostic variable. To solve the problem, the equations are cast into variational form and discretized with linear finite elements, and Crank–Nicolson timestepping is applied [10].

The example code was slightly modified for use with dolfin-adjoint. The subclassing of objects to implement the equation was replaced with calls to solve, and adjinc_timestep was added at the end of the timeloop as described in section 5.1. These modifications were trivial and involved changing less than ten lines of code.

The functional chosen was the Willmore functional integrated over time,

\[ W(c(t), \mu(t)) = \frac{1}{4\epsilon} \int_{t=0}^{t=T} \int_{\Omega} \left( \epsilon \nabla^2 c(t) - \frac{1}{\epsilon} \frac{df}{dc} \right)^2 \, dx \, dt \]

\[ = \frac{1}{4\epsilon} \int_{t=0}^{t=T} \int_{\Omega} \left( -\frac{1}{\epsilon} \mu(t) \right)^2 \, dx \, dt, \]

where \( T \) denotes the final time of the simulation. This functional is physically relevant, as it is intimately connected to the finite-time stability of transition solutions of the
Cahn–Hilliard equation [5]. The problem was solved on a mesh of $\Omega = [0,1]^2$ with 501,264 vertices (> 1 million degrees of freedom) and run for 50 timesteps with a timestep $\Delta t = 5 \times 10^{-6}$. The solution was computed on 24 cores using DOLFIN’s MPI support; both the forward and adjoint models ran in parallel with no further modification. The solutions of the concentration field at the initial and final times are shown in Figure 6.1.

To test the checkpointing implementation, libadjoint was configured to use the multistage checkpointing algorithm of revolve [49], with five checkpoints available in memory and ten on disk. As described in section 5.1, the use of this checkpointing algorithm is entirely transparent to the DOLFIN user.

To verify the correctness of the adjoint solution, the Taylor remainder convergence test was applied. Let $\hat{W}(c_0)$ be the Willmore functional considered as a pure function of the initial condition; i.e., to evaluate $\hat{W}(c_0)$, solve the PDE (6.1) with initial condition $c_0$ to compute $\mu(t)$, and then evaluate $W$ as in (6.4). The Taylor remainder convergence test is based on the observation that, given an arbitrary perturbation $\tilde{c}$ to the initial conditions $c_0$,

\begin{equation}
|\hat{W}(c_0 + h\tilde{c}) - \hat{W}(c_0)| \rightarrow 0 \quad \text{at } O(|h|),
\end{equation}

but that

\begin{equation}
|\hat{W}(c_0 + h\tilde{c}) - \hat{W}(c_0) - h\tilde{c}^T \nabla \hat{W}| \rightarrow 0 \quad \text{at } O(|h|^2),
\end{equation}

where the gradient $\nabla \hat{W}$ is computed using the adjoint solution $z$:

\begin{equation}
\nabla \hat{W} = -\left\langle z, \frac{\partial F}{\partial c_0} \right\rangle,
\end{equation}

where $F \equiv 0$ is the discrete system corresponding to (6.1) [20]. This test is extremely sensitive to even slight errors in the implementation of the adjoint, and it rigorously checks that the computed gradient is consistent with the discrete forward model. The perturbation $\tilde{c}$ was pseudorandomly generated, with each value uniformly distributed in $[0,1]$.

The results of the Taylor remainder convergence test can be seen in Table 6.1. As expected, the convergence orders (6.5) and (6.6) hold, indicating that the adjoint

\begin{figure}
\centering
\begin{subfigure}{0.45\textwidth}
\includegraphics[width=\textwidth]{fig61a}
\caption{$t = 0$.}
\end{subfigure}
\begin{subfigure}{0.45\textwidth}
\includegraphics[width=\textwidth]{fig61b}
\caption{$t = 2.5 \times 10^{-4}$.}
\end{subfigure}
\caption{The concentration $c$ of the Cahn–Hilliard problem at the initial and final times.}
\end{figure}
indicating that the functional gradient computed using the adjoint is correct. The Taylor remainders incorporating gradient information computed using the adjoint converges at second order, as expected, the Taylor model was verified, with the functional evaluated at the final time rather than integrated. The functional is computed using the tangent linear model. The Taylor remainders converge at second order, indicating that the tangent linear solution computed is correct. Similarly, the tangent linear model was verified, with the functional evaluated at the final time rather than integrated. The functional gradient in the direction of the perturbation \( \tilde{c} \) is computed using the tangent linear model

\[
\tilde{c}^T \nabla W = \left\langle \frac{\partial W}{\partial c_T}, \frac{\partial c_T}{\partial \tilde{c}_0} \tilde{c} \right\rangle,
\]

where \( \frac{\partial c_T}{\partial \tilde{c}_0} \) is the Jacobian matrix of the final solution with respect to the initial condition. The tangent linear model computes the term \( \frac{\partial c_T}{\partial \tilde{c}_0} \). For the tangent linear verification, the model was again run on 24 processors. The results of the Taylor remainder convergence test can be seen in Table 6.2. As expected, the theoretical convergence orders hold, indicating that the tangent linear solution computed is correct.

The efficiency of the adjoint implementation was benchmarked using a lower-resolution mesh with 40328 degrees of freedom, as follows. First, the unannotated model was run. Then, the forward model was run again, with annotation, to quantify the cost of annotating the forward model. Finally, the forward and adjoint models were run together. During the forward run, all variables were stored, and checkpointing was not used during the adjoint run, to isolate the intrinsic cost of assembling the adjoint system. For each measurement, five runs were performed on a single processor, and the minimum time taken for the computation was recorded.

For this configuration, the Newton solver typically employs five linear solves. As the adjoint replaces each Newton solve with one linear solve, a coarse estimate of the optimal performance ratio is 1.2. The numerical results can be seen in Table 6.3. The overhead of the annotation is less than 1%. This overhead will further reduce with increasing mesh resolution, as the cost of the annotation and symbolic manipulations

| Table 6.1 | The Taylor remainders for the Willmore functional \( \hat{W} \) evaluated at a perturbed initial condition \( \tilde{c}_0 \equiv c_0 + h\tilde{c} \), where the perturbation direction \( \tilde{c} \) is pseudorandomly generated. All calculations were performed on the fine mesh with more than one million degrees of freedom. As expected, the Taylor remainder incorporating gradient information computed using the adjoint converges at second order, indicating that the functional gradient computed using the adjoint is correct. |
| --- | --- | --- | --- |
| \( h \) | \( |\hat{W}(\tilde{c}_0) - \hat{W}(c_0)| \) | Order | \( |\hat{W}(\tilde{c}_0) - \hat{W}(c_0) - \tilde{c}_0^T \nabla \hat{W}| \) Order |
| 1 × 10^{-7} | 3.4826 × 10^{-6} | 3.0017 × 10^{-9} | 2.0013 |
| 5 × 10^{-8} | 1.7405 × 10^{-6} | 7.4976 × 10^{-10} | 2.0005 |
| 2.5 × 10^{-8} | 8.7099 × 10^{-7} | 1.8737 × 10^{-10} | 2.0004 |
| 1.25 × 10^{-8} | 4.3499 × 10^{-7} | 4.6829 × 10^{-11} | 2.0000 |
| 6.25 × 10^{-9} | 2.1749 × 10^{-7} | 1.1716 × 10^{-11} | 1.9999 |

| Table 6.2 | The Taylor remainders for the Willmore functional, with gradients computed using the tangent linear model. The Taylor remainders converge at second order, indicating that the tangent linear model is correct. |
| --- | --- | --- | --- |
| \( h \) | \( |\hat{W}(\tilde{c}_0) - \hat{W}(c_0)| \) | Order | \( |\hat{W}(\tilde{c}_0) - \hat{W}(c_0) - \tilde{c}_0^T \nabla \hat{W}| \) Order |
| 1 × 10^{-6} | 0.76441 | 0.03120 | 2.012 |
| 5 × 10^{-7} | 0.39007 | 0.00773 | 2.006 |
| 2.5 × 10^{-7} | 0.19698 | 2.001 |
| 1.25 × 10^{-8} | 0.04960 | 4.0085 × 10^{-4} | 2.003 |
| 6.25 × 10^{-9} | 0.04960 | 1.1987 × 10^{-4} | 2.001 |

solution computed is correct. Similarly, the tangent linear model was verified, with the functional evaluated at the final time rather than integrated. The functional gradient in the direction of the perturbation \( \tilde{c} \) is computed using the tangent linear model

\[
\tilde{c}^T \nabla W = \left\langle \frac{\partial W}{\partial c_T}, \frac{\partial c_T}{\partial \tilde{c}_0} \tilde{c} \right\rangle,
\]
Table 6.3
Timings for the Cahn–Hilliard adjoint. The efficiency of the adjoint approaches the theoretical ideal value of 1.2.

<table>
<thead>
<tr>
<th></th>
<th>Runtime (s)</th>
<th>Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forward model</td>
<td>103.93</td>
<td></td>
</tr>
<tr>
<td>Forward model + annotation</td>
<td>104.24</td>
<td>1.002</td>
</tr>
<tr>
<td>Forward model + annotation + adjoint model</td>
<td>127.07</td>
<td>1.22</td>
</tr>
</tbody>
</table>

To derive the adjoint are independent of mesh size, while the costs of assembly and solves do scale with mesh size. The adjoint model takes approximately 1.22 times the cost of the forward model. This ratio compares very well with the theoretical estimate: the adjoint implementation achieves almost optimal performance.

6.2. Stokes. As described in section 5.3, the approach presented in this paper is capable of deriving the adjoint for models that use matrix-free solvers. To demonstrate this capability, a matrix-free variant of the mantle convection model presented in [54] was adjoined:

\[
-\nabla \cdot \sigma - \nabla p = (Ra T)e, \\
\nabla \cdot u = 0, \\
\frac{\partial T}{\partial t} + u \cdot \nabla T - \nabla^2 T = 0,
\]

(6.9)

where \( \sigma \) is the deviatoric stress tensor, \( p \) is the pressure, \( Ra \) is the thermal Rayleigh number, \( T \) is the temperature, \( u \) is the velocity, and \( e \) is a unit vector in the direction of gravity (here the \(-x_2\)-direction). For the configuration reported below, the Rayleigh number \( Ra \) was set to \( 10^6 \), which yields a vigorously convective system. For the Stokes equations, no-slip conditions are applied on the top and bottom boundaries, and a stress-free condition is applied on the remainder of the boundary. For the temperature equation, Dirichlet conditions are applied on the top and bottom boundaries, while homogeneous Neumann conditions are applied on the left and right boundaries. The initial temperature field is set based on an analytical expression derived from boundary layer theory; for full details, see [54].

The Stokes equations were discretized using the \( P_2 \times P_1 \) Taylor–Hood element, while the advection equation for the temperature field was discretized using the \( P_{1DG} \) element. The domain \( \Omega = [0, 2] \times [0, 1] \) was discretized with 40 elements in the \( x \)- and \( y \)-directions, leading to a degree of freedom count of 24403. The solution of the Stokes equations was achieved in a matrix-free manner using DOLFIN’s interfaces to the PETSc matrix-free solvers [3]. Both the forward and adjoint problems were parallelized using DOLFIN’s OpenMP support and were run on eight cores.

The functional taken was the Nusselt number of the temperature evaluated at the final time

\[
Nu(T) = \int_{\Gamma_{\text{top}}} \frac{\partial T}{\partial x_2} \, dx_2 / \int_{\Gamma_{\text{bottom}}} T \, dx_2,
\]

(6.10)

which measures the efficiency of the convection by comparing the total heat transferred to that transferred by thermal conduction alone. The adjoint was computed using 30 checkpoints on disk and 30 checkpoints in memory. The results are illustrated in Figure 6.2.
Fig. 6.2. (a) The initial condition for temperature of the mantle convection simulation. (b) The temperature field after 200 timesteps. Plumes are clearly visible. (c) The gradient of the Nusselt number with respect to the temperature initial condition, computed using the adjoint.
Table 6.4

The Taylor remainder \( R \equiv \hat{\text{Nu}}(T_0 + \frac{h}{2}\tilde{T}) - \hat{\text{Nu}}(T_0 - \frac{h}{2}\tilde{T}) \) for the Nusselt functional \( \hat{\text{Nu}} \) evaluated at a perturbed initial condition, where the perturbation direction \( \tilde{T} \) is the vector of all ones. All calculations were performed on the mesh with 24403 degrees of freedom. The third-order convergence of the Taylor remainders indicates that the adjoint is correct.

| \( h \)          | \( |R| \)     | Order | \( |R - h\tilde{T}^T \nabla \hat{\text{Nu}}| \) | Order |
|------------------|--------------|-------|---------------------------------|-------|
| \( 7.5 \times 10^{-3} \) | 0.14568      |       | \( 3.6260 \times 10^{-2} \)     |       |
| \( 3.75 \times 10^{-3} \)  | 0.05905      | 1.302 | \( 4.3465 \times 10^{-3} \)     | 3.06  |
| \( 1.875 \times 10^{-3} \) | 0.02787      | 1.083 | \( 5.1659 \times 10^{-4} \)     | 3.07  |
| \( 9.375 \times 10^{-4} \)  | 0.01373      | 1.021 | \( 5.2311 \times 10^{-5} \)     | 3.30  |

Table 6.5

Timings for the Stokes mantle convection adjoint. The efficiency of the adjoint exceeds the theoretical ideal value of 2, as the adjoint linear solves happen to converge faster.

<table>
<thead>
<tr>
<th></th>
<th>Runtime (s)</th>
<th>Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forward model</td>
<td>96.03</td>
<td></td>
</tr>
<tr>
<td>Forward model + annotation</td>
<td>96.05</td>
<td>1.0</td>
</tr>
<tr>
<td>Forward model + annotation + adjoint model</td>
<td>178.75</td>
<td>1.86</td>
</tr>
</tbody>
</table>

The adjoint was verified using a higher-order analogue of the Taylor remainder convergence test described in section 6.1. Let \( \hat{\text{Nu}} \) be the Nusselt number considered as a pure function of the initial condition for temperature; i.e., to evaluate \( \hat{\text{Nu}}(T_0) \), solve the PDE (6.9) with initial condition \( T_0 \) to compute the final temperature \( T \), and then evaluate \( \text{Nu}(T) \) as in (6.10). The test is based on the observation that, given an arbitrary perturbation \( \tilde{T} \) to the initial conditions \( T_0 \),

\[
\hat{\text{Nu}} \left( T_0 + \frac{h}{2}\tilde{T} \right) - \hat{\text{Nu}} \left( T_0 - \frac{h}{2}\tilde{T} \right) \to 0 \quad \text{at } O(|h|),
\]

but that

\[
\hat{\text{Nu}} \left( T_0 + \frac{h}{2}\tilde{T} \right) - \hat{\text{Nu}} \left( T_0 - \frac{h}{2}\tilde{T} \right) - h\tilde{T}^T \nabla \hat{\text{Nu}} \to 0 \quad \text{at } O(|h|^3),
\]

where the gradient \( \nabla \hat{\text{Nu}} \) is computed using the adjoint solution \( z \):

\[
\nabla \hat{\text{Nu}} = - \left< z, \frac{\partial F}{\partial T_0} \right>,
\]

where \( F \equiv 0 \) is the discrete system corresponding to (6.9). The higher-order version of the Taylor remainder convergence test was used because it reaches the asymptotic region more rapidly, which is useful for strongly nonlinear problems such as this. The results of the Taylor remainder convergence test can be seen in Table 6.4. As expected, the convergence orders (6.11) and (6.12) hold, indicating that the adjoint solution computed is correct.

The efficiency of the adjoint implementation was benchmarked using the same procedure as described in section 6.1, again using a lower-resolution configuration of 1603 degrees of freedom. The results can be seen in Table 6.5. The performance overhead of solving the linear systems matrix-free is significant, and these timings should not be taken as representative of the potential performance of the modeling.
system. The overhead of the annotation is extremely small, within the distribution of timings of the unannotated run. The adjoint model takes approximately 86% of the cost of the forward model. In this case, the forward model performs two Picard iterations per timestep, each of which induce a corresponding linear solve in the adjoint equations. Therefore, a naive estimate of the ideal theoretical efficiency is 2. On investigation, the proximate cause of the adjoint run being cheaper than the forward run was that the matrix-free linear solvers happened to converge more quickly during the adjoint run.

6.3. Viscoelasticity. Most biological tissue responds in a viscoelastic, rather than purely elastic, manner. As a final example, we consider a nontrivial discretization of a viscoelastic model for the deformation and stress development in the upper part of the spinal cord under pressure induced by the pulsating flow of cerebrospinal fluid [16].

The standard linear solid viscoelastic model equations can be phrased [45] as:

\[ A_0^1 \frac{\partial}{\partial t} \sigma_0 + A_0^0 \sigma_0 - \nabla v + \gamma = 0, \]

\[ A_1^1 \frac{\partial}{\partial t} \sigma_1 - \nabla v + \gamma = 0, \]

\[ \nabla \cdot (\sigma_0 + \sigma_1) = 0, \]

\[ \text{skw}(\sigma_0 + \sigma_1) = 0 \]

for \((t; x, y, z) \in (0, T] \times \Omega\). Here, \(A_0^1, A_0^0, A_1^1\) are fourth-order compliance tensors, the divergence and gradient are taken rowwise, and \(\text{skw}\) denotes the skew-symmetric component of a tensor field. The total stress tensor \(\sigma\) is the sum of the Maxwell and elastic contributions. In the isotropic case, each of the compliance tensors \(A_0^0, A_1^0, A_1^1\) reduces to a two-parameter map:

\[ A_j^i = (C_j^i)^{-1}, \quad C_j^i \varepsilon = \mu_j^i \varepsilon + \lambda_j^i \text{tr}(\varepsilon) I, \]

where \(\mu_j^i, \lambda_j^i\) are positive Lamé parameters. The system is closed by initial conditions for the Maxwell stress, essential boundary conditions for the velocity, and traction boundary conditions for the total stress \(\sigma \cdot \hat{n}\), where \(\hat{n}\) is the outward normal on the domain boundary. The cord was kept fixed at the top and bottom, and the parameters were set to \(\mu_0^0 = 37.466, \lambda_0^0 = 10^4, \mu_0^1 = 4.158, \lambda_0^1 = 10^3, \mu_1^1 = 2.39, \lambda_1^1 = 10^3\) (kPa). The traction boundary condition was set to

\[ \sigma \cdot \hat{n} = -p\hat{n}, \]

where \(p\) is a periodically varying pressure modeled as \(p(t; x, y, z) = a \sin(2 \pi t)(171 - 78)^{-1}(z - 78)\) (kPa), where \(a = 0.05\) is the amplitude of the pressure.

The discretization of (6.14) is performed using the (locking-free) scheme introduced in [45], allowing for direct approximation of the stresses while enforcing the symmetry of the total stress weakly. The temporal discretization is carried out via a two-step TR-BDF2 scheme, that is, a Crank–Nicolson step followed by a two-step backward difference scheme, while the spatial mixed finite element discretization is based on seeking approximations \(\sigma_0(t), \sigma_1(t), v(t), \gamma(t)\) in the space

\[ Z = BDM_3^1 \times BDM_3^1 \times P_{0DG}^3 \times P_{0DG}^3. \]
where BDM$_1$ denotes the lowest-order $H$(div)-conforming Brezzi–Douglas–Marini elements and P$_{0DG}$ denotes piecewise constants. See [45] for more details.

Abnormal stress conditions in the interior of the spinal cord may be of biomedical interest [29, 23]. In particular, we focus on the contribution of the Maxwell stress tensor in the horizontal plane at the final time $T$:

$$J = \int_{\Omega} (\sigma(T) \cdot e_z)^2 \, dx,$$

where $e_z$ denotes a unit vector in the $z$-direction.

The problem was solved on a tetrahedral mesh generated from patient-specific imaging data, yielding a total of 879204 degrees of freedom, for $t \in [0, 1.25]$ with a timestep of $\Delta t = 0.01$. As the system is linear, the matrices corresponding to each of the steps in the TR-BDF$_2$ scheme were preassembled outside of the timeloop, and their LU factorizations were cached. This optimization is recognized by dolfin-adjoint, which then applies the analogous factorization strategy to the corresponding adjoint solve. The Maxwell stress in the horizontal plane at $t = 1.25$ is illustrated in Figure 6.3, and norms of the forward and adjoint stress tensors are illustrated in Figure 6.4.

To verify the correctness of the adjoint solution, the Taylor remainder convergence test was applied. Let $\hat{J}(a)$ be the Maxwell stress functional considered as a pure function of the amplitude of the applied pressure; i.e., to evaluate $\hat{J}(a)$, solve the PDE (6.14) with pressure amplitude $a$ to compute $\sigma_0(T)$, and then evaluate $J$ as in (6.17). The results of the Taylor remainder convergence test can be seen in Table 6.6. As expected, the convergence orders (6.5) and (6.6) hold, indicating that the adjoint solution computed is correct.

The efficiency of the adjoint implementation was benchmarked using the same procedure as described in section 6.1, again using a lower-resolution configuration of 86976 degrees of freedom. The results can be seen in Table 6.7. As the problem is
Fig. 6.4. (a) $L^2$-norm squared of the Maxwell stress in the horizontal plane versus time. (b) The $L^2$-norm of the adjoint Maxwell stress tensor $z_0$ and the adjoint elastic stress tensor $z_1$ versus time.

Table 6.6
The Taylor remainders for the functional given by (6.17). All calculations were performed on the mesh with 879204 degrees of freedom. The convergence of the Taylor remainders indicates that the adjoint is correct.

| $\delta a$   | $|\hat{J}(a + \delta a) - \hat{J}(a)|$ | Order | $|\hat{J}(a + \delta a) - \hat{J}(a) - \nabla \hat{J} \cdot \delta a|$ | Order |
|--------------|----------------------------------|-------|-----------------------------------|-------|
| 0.05         | $9.1012 \times 10^{-3}$          |       | $3.0337 \times 10^{-3}$           |       |
| 0.025        | $3.7921 \times 10^{-3}$          | 1.263 | $7.58417 \times 10^{-4}$          | 2.0000|
| 0.0125       | $1.7064 \times 10^{-3}$          | 1.152 | $1.8959 \times 10^{-4}$          | 2.0000|
| $6.25 \times 10^{-3}$ | $8.0583 \times 10^{-4}$          | 1.0824 | $4.7397 \times 10^{-5}$          | 2.0001|
| $3.125 \times 10^{-3}$ | $3.9106 \times 10^{-4}$          | 1.0430 | $1.1848 \times 10^{-5}$          | 2.0001|

Table 6.7
Timings for the viscoelasticity adjoint. The efficiency of the adjoint approaches the theoretical ideal value of 2.

<table>
<thead>
<tr>
<th></th>
<th>Runtime (s)</th>
<th>Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forward model</td>
<td>119.93</td>
<td></td>
</tr>
<tr>
<td>Forward model + annotation</td>
<td>120.24</td>
<td>1.002</td>
</tr>
<tr>
<td>Forward model + annotation + adjoint model</td>
<td>243.99</td>
<td>2.029</td>
</tr>
</tbody>
</table>

linear, the theoretical estimate of the ideal efficiency ratio is 2, which is approximately achieved by the implementation presented here.

7. Conclusion. Naumann’s recent book on algorithmic differentiation states [39, p. xii]:

[T]he automatic generation of optimal (in terms of robustness and efficiency) adjoint versions of large-scale simulation code is one of the great open challenges in the field of High-Performance Scientific Computing.

The framework presented here, dolfin-adjoint, provides a robust and efficient mechanism for automatically deriving adjoint and tangent linear models of a wide variety of finite element models implemented in the Python interface to the DOLFIN library. Only minimal changes are required to adapt such a forward model for use with dolfin-adjoint. The adjoint model draws on the advantages of libadjoint to deliver optimal checkpointing strategies, and inherits the seamless parallelism of the FEniCS frame-
work. The numerical results obtained demonstrate optimal efficiency in the adjoint model.

The approach employed in dolfin-adjoint is analogous to but fundamentally different from that adopted by algorithmic differentiation: by operating at a higher level of abstraction, a much greater degree of automation and efficiency has been achieved. The framework presented here enables adjoint models to be derived automatically, reliably, and robustly, relieving the model developers of the adjoint development task.

REFERENCES


