ADIFOR Working Note #2: Using ADIFOR to Compute Dense and Sparse Jacobians

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Using ADIFOR to Compute Dense and Sparse Jacobians

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Abstract. ADIFOR is a source translator that, given a collection of Fortran subroutines for the computation of a “function,” produces Fortran code for the computation of the derivatives of this function. More specifically, ADIFOR produces code to compute the matrix-matrix product $JS$, where $J$ is the Jacobian of the “function” with respect to the user-defined independent variables, and $S$ is the composition of the derivative objects corresponding to the independent variables. This interface is flexible; by setting $S = x$, one can compute the matrix-vector product $Jx$, or by setting $S = I$, one can compute the whole Jacobian $J$. Other initializations of $S$ allow one to exploit a known sparsity structure of $J$. This paper illustrates the proper initialization of ADIFOR-generated derivative codes and the exploitation of a known sparsity structure of $J$.

1 Introduction

ADIFOR, (Automatic Differentiation in Fortran) is a tool for the automatic generation of derivative codes from user-supplied Fortran subroutines. That is, given a collection of Fortran subroutines for the computation of a “function,” ADIFOR produces Fortran code for the computation of the derivatives of this function. ADIFOR differs from previous approaches to automatic differentiation (see [13] for a recent survey) in several ways:

Generality: The “function” can be composed of many subroutines, and these subroutines may communicate via parameter lists and/or common blocks. In general, almost all of Fortran-77 is supported.

Portability: ADIFOR produces vanilla Fortran-77 code.

Efficiency: While primarily based on the so-called forward mode of automatic differentiation, ADIFOR uses the so-called reverse mode to process assignment statements with composite right-hand sides [9,10]. In addition to saving storage, this approach significantly enhances performance.

Parallelism and Vectorization: The code produced by ADIFOR respects the data-flow structure of the original program. That is, if the code vectorizes and parallelizes well, so does the ADIFOR-generated derivative code. If anything, the derivative code offers more scope for

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vectorization and parallelization because of the addition of another ‘data parallel’ dimension in
derivative objects.

**Extensibility:** The fact that ADIFOR employs a consistent subroutine-naming scheme allows the
user to supply his or her own derivative routines. In this fashion, the user can exploit domain-
specific knowledge, utilize vendor-supplied libraries, and speed up computational bottlenecks.

**Ease of Use:** ADIFOR requires the user to supply the Fortran source code for the subroutine
representing the function to be differentiated and for all lower-level subroutines. The user then
selects the variables (either in parameter lists or in common blocks) that correspond to the
independent and dependent variables. By using the powerful interprocedural analysis tools
of the ParaScope programming environment [3], ADIFOR then automatically determines
which other variables throughout the program must have derivative information associated
with them.

**Interactive Interface:** An X-windows interface for ADIFOR (called xadifor) is also provided.

Xadifor makes it easy for the user to set up the problem and to rerun ADIFOR if changes
in the code for the target function require a new translation.

ADIFOR is applied to the code of the subroutine that corresponds to the subroutine we wish to
differentiate (foo, say), and to all subroutines called directly or indirectly from foo. Let us assume
that foo describes a function $f : [x, w] \mapsto [y, z]$ and that we are interested in the derivatives $\frac{\partial y}{\partial x}$,
that is, the input variable $w$ is treated as constant, and the output variable $z$ is irrelevant. If this
is the case, we call $x$ the **independent variable** and $y$ the **dependent variable**. We are aware
of the fact that the terms “dependent,” “independent,” “variable,” and “parameters” are used in
many different contexts, yet we found that this terminology corresponds best to our mathematical
idea of derivatives, since we will compute derivatives of the “dependent” variables with respect to
the “independent” ones.

We require the user to tell ADIFOR the names of the independent variables and the names of
the dependent variables. In many codes, dependent and independent variables may share storage.
For example, on entry to foo, array A may be initialized to what we consider mathematically to be
the value of the independent variable $x$, and during the course of executing foo, $y$ will be written
into A. This poses no problem for ADIFOR. It produces a subroutine named gffoo$\langle n \rangle$ (where
$\langle n \rangle$ is some number encoding which variables were dependent and independent), which computes
the first derivatives of the function computed by foo, as well as foo itself.

To propagate derivative information in the forward mode, we have to associate derivative objects
with the independent variables, the dependent variables, and all those program variables whose
value depends (directly or indirectly) on an independent variable and that influence the value of a
dependent variable. That is, if $x$ is independent, $y$ is dependent, and $z$ depends on $x$ and $y$ depends
on $z$, then $z$ also needs a derivative object. A variable with which we associate a derivative object is
called an **active variable**, any other variable is a **passive variable**. Dependent and independent
variables are always active, and **integer** variables are always passive.

The user need not specify as passive or active variables local to foo or parameters or local
variables in routines called by foo. Using the powerful interprocedural analysis tools available
in the ParaScope environment [3], we can determine all active variables from a definition of the
independent and dependent ones. This allows for a simple user interface that corresponds as much as possible to the mathematical intuition underlying \texttt{foo}.

The derivative codes produced by \texttt{ADIFOR} have a gradient object associated with every active variable. The convention is to associate a gradient \texttt{g$<$var$>$} of leading dimension \texttt{lg$<$var$>$} with variable \texttt{<var>}$. The calling sequence of \texttt{g$<$foo$<$n$>$} is derived from that of \texttt{foo} by inserting an argument \texttt{g$p$} denoting the length of the gradient vectors as the first argument, and then copying the calling sequence of \texttt{foo}, inserting \texttt{g$<$var$>$} and \texttt{lg$<$var$>$} after every active variable \texttt{<var>}$. Passive variables are left untouched.

In its simplest form, the functionality of \texttt{ADIFOR} can be summarized as follows:

\begin{quote}
In general, if $x(1:n)$ are the independent variables, and $y(1:m)$ the dependent ones, then $g$x is a $g$p $\times$ $n$ matrix ($lg$x $\geq$ $g$p), and $g$y is a $g$p $\times$ $m$ matrix ($lg$y $\geq$ m). The functionality of $g$foo is: Given input values $x$ and $g$x, this subroutine computes $y = foo(x)$, and $g$y = $(foo'(x)g$x$T)^T$.
\end{quote}

In this paper, we shall not concern ourselves with the way code is generated or with the input provided to \texttt{ADIFOR}. For these details, the reader is referred to [2]. Even though the \texttt{ADIFOR} interface conceptually never changes, the actual initialization of \texttt{ADIFOR} code may vary depending on context. We focus instead on the proper and efficient use of \texttt{ADIFOR}-generated codes through detailed examination of the following cases:

- Dense Jacobian, one independent, one dependent variable
- Dense Jacobian, multiple independent, multiple dependent variables
- Sparse Jacobian, one independent, one dependent variable
- Sparse Jacobian, two independent variables, one dependent variable
- Partially separable functions

In most of these cases, a “variable” denotes an array; thus, we shall be dealing with vector-valued functions.

2 Case 1: Dense Jacobian, one independent, one dependent variable

Our first example is adapted from Problem C2 in the STDTST set of test problems for stiff ODE solvers [7] and was brought to our attention by George Corliss. The routine \texttt{FCN2} computes the right-hand side of a system of ordinary differential equations $y' = yp = f(x, y)$ by calling a subordinate routine \texttt{FCI}.
C File: FCN2.f

SUBROUTINE FCN2(M,X,Y,YP)
INTEGER N
DOUBLE PRECISION X, Y(M), YP(M)
INTEGER ID, IWT
DOUBLE PRECISION W(20)
COMMON /STCOM5/W, IWT, N, ID

CALL FCN(X,Y,YP)
RETURN
END

C File: FCN.f

SUBROUTINE FCN(X,Y,YP)

C ROUTINE TO EVALUATE THE DERIVATIVE F(X,Y) CORRESPONDING TO THE
C DIFFERENTIAL EQUATION:
CDY/DX = F(X,Y).
C THE ROUTINE STORES THE VECTOR OF DERIVATIVES IN YP(*). THE
C DIFFERENTIAL EQUATION IS SCALED BY THE WEIGHT VECTOR W(*)
C IF THIS OPTION HAS BEEN SELECTED (IF SO IT IS SIGNALLED
C BY THE FLAG IWT).

DOUBLE PRECISION X, Y(20), YP(20)
INTEGER ID, IWT, N
DOUBLE PRECISION W(20)
COMMON /STCOM5/W, IWT, N, ID
DOUBLE PRECISION SUM, CPARM(4), TEMP(20)
INTEGER I, IID
DATA CPARM/1.D-1, 1.D0, 1.D1, 2.D1/

IF (IWT.LT.0) GO TO 40
DO 20 I = 1, N
   TEMP(I) = Y(I)
   Y(I) = Y(I)*W(I)
20 CONTINUE
40 IID = MOD(IID,10)

C ADAPTED FROM PROBLEM C2
YP(I) = -Y(I) + 2.0D0
SUM = Y(I)*Y(I)
DO 50 I = 2, N
   YP(I) = -10.0D0*Y(I) + CPARM(IID-1)*(2**I)*SUM
   SUM = SUM + Y(I)*Y(I)
50 CONTINUE
IF (IWT.LT.0) GO TO 680
DO 660 I = 1, N
   YP(I) = YP(I)/W(I)
   Y(I) = YTEMP(I)
660 CONTINUE
680 CONTINUE
RETURN
END

Most software for the numerical solution of stiff systems of ODEs requires the user to supply a subroutine for the Jacobian of $f$ with respect to $y$. Such a subroutine can easily be generated by ADIFOR. For the purposes of automatic differentiation, the vector $Y$ is the independent variable, and the vector $YP$ is the dependent variable. Then ADIFOR produces

```fortran
subroutine g$fcn2s6(g$p$, m, x, y, g$y$, ldg$y$, yp, g$yp$, ldg$yp$)
C
C      ADIFOR: runtime gradient index
      integer g$p$
C      ADIFOR: translation time gradient index
      integer g$p$max$
      parameter (g$p$max$ = 20)
C      ADIFOR: gradient iteration index
      integer g$i$
C
      integer ldg$y$
      integer ldg$yp$
      integer n
      double precision x, y(m), yp(m)
      integer id, iwt
      double precision w(20)
      common /stcom5/ w, iwt, n, id
C
C      ADIFOR: gradient declarations
      double precision g$y(ldg$y, m), g$yp(ldg$yp, m)
      if (g$p$ .gt. g$p$max$) then
         print *, "Parameter g$p$ is greater than g$p$max."
         stop
      endif
      call g$fcn86(g$p$, x, y, g$y$, ldg$y$, yp, g$yp$, ldg$yp$)
      return
   end
```

```fortran
subroutine g$fcn86(g$p$, x, y, g$y$, ldg$y$, yp, g$yp$, ldg$yp$)
C
C      ADIFOR: runtime gradient index
```
integer g$p\$
C ADFOR: translation time gradient index
integer g$pmax$
parameter (g$pmax$ = 20)
C ADFOR: gradient iteration index
integer g$i$
C
integer ldg$y$
integer ldg$yp$
C ROUTINE TO EVALUATE THE DERIVATIVE F(X,Y) CORRESPONDING TO THE
C DIFFERENTIAL EQUATION;
C DY/DX = F(X,Y).
C THE ROUTINE STORES THE VECTOR OF DERIVATIVES IN YP(*). THE
C DIFFERENTIAL EQUATION IS SCALED BY THE WEIGHT VECTOR W(*)
C IF THIS OPTION HAS BEEN SELECTED (IF SO IT IS SIGNALED
C BY THE FLAG IWT).
double precision x, y(20), yp(20)
integer id, iwt, n
do double precision w(20)
common /stcom5/ w, iwt, n, id
do double precision sum, cparm(4), ytemp(20)
integer i, id
data cparm /1.d-1, 1.d0, 1.d1, 2.d1/
C
C ADFOR: gradient declarations
do double precision g$y(ldg$y, 20), g$yp(ldg$yp, 20)
do double precision g$sum(g$pmax$), g$ytemp(g$pmax$, 20)
if (g$p gt. g$pmax$) then
  print *, "Parameter g$p is greater than g$pmax."
  stop
endif
if (iwt .lt. 0) then
  goto 40
endif
do 99999, i = 1, n
C ytemp(i) = y(i)
do g$i = 1, g$p
  g$ytemp(g$i, i) = g$y(g$i, i)
endo
do ytemp(i) = y(i)
C y(i) = y(i) * w(i)
do g$i = 1, g$p
  g$y(g$i, i) = w(i) * g$y(g$i, i)
endo
y(i) = y(i) * w(i)
20 continue
99999 continue
40  $\text{id} = \text{mod}($\text{id}, 10$)
C  ADAPTED FROM PROBLEM C2
C  $y_p(1) = -y(1) + 2.0d0$
  do $g\text{Si$} = 1$, $g\text{Sp$}
  g\text{Sy}(g\text{Si$}, 1) = -g\text{Sy}(g\text{Si$}, 1)$
  enddo
  $y_p(1) = -y(1) + 2.0d0$
C  sum = $y(1) * y(1)$
  do $g\text{Si$} = 1$, $g\text{Sp$}
  g\text{Sum}(g\text{Si$}) = y(1) * g\text{Sy}(g\text{Si$}, 1) + y(1) * g\text{Sy}(g\text{Si$}, 1)$
  enddo
  sum = $y(1) * y(1)$
  do 99998, i = 2, n
C  $y_p(i) = -10.0d0 * i * y(i) + \text{cparm}($\text{uid}, -1$) * (2 ** i) * \text{sum}$
  do $g\text{Si$} = 1$, $g\text{Sp$}
  g\text{Sy}(g\text{Si$}, i) = \text{cparm}($\text{uid}, -1$) * (2 ** i) * g\text{Sum}(g\text{Si$}) + -1$
  *0.0d0 * i * g\text{Sy}(g\text{Si$}, i)$
  enddo
  $y_p(i) = -10.0d0 * i * y(i) + \text{cparm}($\text{uid}, -1$) * (2 ** i) * \text{sum}$
C  sum = sum + $y(i) * y(i)$
  do $g\text{Si$} = 1$, $g\text{Sp$}
  g\text{Sum}(g\text{Si$}) = g\text{Sum}(g\text{Si$}) + y(i) * g\text{Sy}(g\text{Si$}, i) + y(i) * g\text{Sy}$
  *(g\text{Si$}, i)$
  enddo
  sum = sum + $y(i) * y(i)$
50  continue
99998 continue
  if ($\text{Iwt}, \text{L}, 0$) then
    goto 680
  endif
  do 99997, i = 1, n
C  $y_p(i) = y_p(i) / w(i)$
  do $g\text{Si$} = 1$, $g\text{Sp$}
  g\text{Sy}(g\text{Si$}, i) = (1 / w(i)) * g\text{Sy}(g\text{Si$}, i)$
  enddo
  $y_p(i) = y_p(i) / w(i)$
C  $y(i) = \text{ytemp}(i)$
  do $g\text{Si$} = 1$, $g\text{Sp$}
  g\text{Sy}(g\text{Si$}, i) = g\text{Sytemp}(g\text{Si$}, i)$
  enddo
  $y(i) = \text{ytemp}(i)$
660 continue
99997 continue
680 continue
return
end

In accordance with the general policy outlined in § 1, the derivative objects $g\text{Sy}$ and $g\text{Sy}_p$ are
declared as matrices with 20 columns (since both \( y \) and \( yp \) were declared as vectors of length 20) and leading dimension \( 1dgsy \) and \( 1dgspy \), respectively. The parameter \( gsp \) denotes the actual length of the gradient objects in a call to \( gsfcn2$6 \). Since Fortran 77 does not allow dynamic memory allocation, derivative objects for local variables are statically allocated with leading dimension \( pmax \), whose value was selected by the user during the invocation of ADIFOR. A variable and its associated derivative object are treated in the same fashion; that is, if \( x \) is a function parameter, so is \( gsx \). Derivative objects corresponding to locally declared variables or variables in common blocks are declared locally or in common blocks as well.

Subroutine \( gsfcn2$6 \) relates to the Jacobian

\[
J_{yp} = \begin{pmatrix}
\frac{\partial y_1}{\partial y_1} & \cdots & \frac{\partial y_1}{\partial y_m} \\
\vdots & \ddots & \vdots \\
\frac{\partial y_m}{\partial y_1} & \cdots & \frac{\partial y_m}{\partial y_m}
\end{pmatrix}
\]

as follows: Given input values for \( gsp, m, x, y, gsy, 1dgsy, \) and \( ldgsy \), the routine \( gsfcn2$6 \) computes both \( yp \) and \( gsyyp \), where

\[
gsyyp(1:gsp, 1:m) = (J_{yp}(gsp(1:gsp, 1:m)^T))^T.
\]

The superscript \( T \) denotes matrix transposition. The user must allocate \( gsyyp \) and \( gsy \) with leading dimensions \( 1dgsyyp \) and \( 1dgsy \) that are at least \( gsp \). While the implicit transposition may seem awkward at first, this is the only way to handle assumed-size arrays (like \texttt{real a(*)}) in subroutine calls.

Assume that \( m \) and \( gsp \) are 20 and that \( 1dgsyyp \) and \( 1dgsy \) are at least 20. Then we can compute the derivative matrix \( J_{yp} \) simply by initializing \( gsy \) to the identity:

***************
* Approach 1 *
***************
\begin{verbatim}
DG 10 I = 1, M
   DO 5 J = 1, M
      GSY(I,J) = 0.0D
   5 CONTINUE
   GSY(I,1) = 1.0D0
10 CONTINUE
   call gsfcn2$6(20, m, x, y, gsy, ldgsy, yp, gsyyp, ldgsyyp)
\end{verbatim}

On exit from \( gsfcn2$6 \), the variable \( gsyyp \) contains the transpose of the Jacobian \( J_{yp} \).

Alternatively, we could have computed the Jacobian one column at a time:

***************
* Approach 2 *
***************
\begin{verbatim}
DG 10 I = 1, M
   *
   * initialize first row of GSY to i-th unit vector
\end{verbatim}
* DO $J = 1, M$
  $GSY(1,J) = 0.0D$
  CONTINUE
  $GSY(1,1) = 1.0D$
* call ADIFOR-generated derivative code
* call g$func2$6($l$, $m$, $x$, $y$, $g$,$y$, $lg$,$y$, $yp$, $g$,$yp$, $ldg$,$yp$)
* store ith column of the Jacobian in ith row of Jactrans array
* DO $15 J = 1,M$
  JACTRANS(I,J) = G$YP(I,J)$
  CONTINUE

Even though $g$,$yp(i,j)$ as computed in Approach 1 equals jactrans(i,j) computed in Approach 2, the second method is significantly less efficient. This inefficiency arises from the fact that the value of yp itself is computed once in the first approach, but m times in the second approach. Thus, it is usually best to compute as large a slice of the Jacobian as memory restrictions will allow.

3 Case 2: Dense Jacobian, multiple independent and multiple dependent variables

The second example involves a code that models adiabatic flow [16], a commonly used module in chemical engineering. This code models the separation of a pressurized mixture of hydrocarbons into liquid and vapor components in a distillation column, where pressure (and, as a result, temperature) decrease. This example was communicated to us by Larry Biegler.

In its original version, the top-level subroutine

```
subroutine aifl(kf)
   integer kf
```

has only one argument. All other information is passed in common blocks. For demonstration purposes, we changed the interface slightly to

```
subroutine aifl(kf,feed,pressure,liquid,vapor)
   integer kf
   real feed(*), pressure(*), liquid(*), vapor(*)
```

copying the values passed in those arguments into the proper common blocks in aifl. As our first example, assume that we are interested in \( \frac{\partial \text{liquid}}{\partial \text{feed}} \) and \( \frac{\partial \text{vapor}}{\partial \text{feed}} \). In this case, ADIFOR generates

\[ \frac{\partial \text{liquid}}{\partial \text{feed}} + \frac{\partial \text{vapor}}{\partial \text{feed}} = 1 \]

\(^1\) Actually, it is sufficient to compute one or the other, since, because of conservation laws, \( \frac{\partial \text{liquid}}{\partial \text{feed}} + \frac{\partial \text{vapor}}{\partial \text{feed}} \) equals the identity matrix.
subroutine g$aifl$26(g$p, kf, feed, g$feed, ldg$feed, pressure,
$    liquid, g$liquid, ldg$liquid,
$    vapor, g$vapor, ldg$vapor)
integer g$p, kf, ldg$feed, ldg$liquid, ldg$vapor
real feed(*), g$feed(ldg$feed, *), pressure(*),
$    liquid(*), g$liquid(ldg$liquid, *),
$    vapor(*), g$vapor(ldg$vapor, *)

In our example, the feed was a mixture of the hydrocarbons N-butane, N-pentane, 1-butene, cis-2-butene, trans-2-butene, and propylene, so the length of feed, liquid, and vapor was six, with feed(1) corresponding to the N-butane feed, and so on. So if we set g$p=6 and initialize g$feed to a 6 × 6 identity matrix, then on exit g$liquid(i,j) contains
\[
\frac{\partial (\text{component } j \text{ in liquid})}{\partial (\text{component } i \text{ in feed})}
\]
which predicts by what amount the liquid portion of substance $j$ will change, if the feed of component $i$ changes.

Suppose that we also wish to treat the pressure at the various inlets as an independent variable, but (because of the conservation law) we decide not to declare “vapor” as a dependent variable, ADIFOR generates

subroutine g$aifl$14(g$p, kf, feed, g$feed, ldg$feed,
$    pressure, g$pressure, ldg$pressure,
$    liquid, g$liquid, ldg$liquid, vapor)

The initialization is a little more complicated this time. Assuming that we have 3 feeds (so pressure has three elements), the total number of independent variables is $6 + 3 = 9$. g$liquid measures the sensitivity of the 6 substances with respect to changes in the 9 independent variables. Thus,
\[
J_{\text{liquid}} = \begin{bmatrix}
\frac{\partial \text{liquid}}{\partial \text{pressure}} & \frac{\partial \text{liquid}}{\partial \text{feed}}
\end{bmatrix}
\]
is a 6 × 9 matrix. ADIFOR computes
\[
g\text{liquid} = \left( J_{\text{liquid}} \begin{bmatrix} g\text{feed}^T \\ g\text{pressure}^T \end{bmatrix} \right)^T.
\]
If we wish to compute the whole Jacobian $J$, then
\[
\begin{bmatrix} g\text{feed}^T \\ g\text{pressure}^T \end{bmatrix}
\]
must be initialized to a 9 × 9 identity matrix. Thus, g$feed$^T must contain the first six rows of a 9 × 9 identity matrix (since there are six variables in the feed), and g$pressure$^T must contain the
last three rows of a $9 \times 9$ identity matrix. This configuration is achieved by initializing

$$g_{\text{feed}} = \begin{pmatrix} 1 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 \end{pmatrix}, \quad \text{and} \quad g_{\text{pressure}} = \begin{pmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}.$$

4 Case 3: Sparse Jacobian, one independent, one dependent variable

From the previous discussion, ADIFOR may seem to be well suited for computing dense Jacobian matrices, but rather expensive for sparse Jacobians. A primary reason is that the forward mode of automatic differentiation upon which ADIFOR is mainly based (see [2]) requires roughly $g_{\text{p}}$ operations for every assignment statement in the original function. Thus, if we compute a Jacobian $J$ with $n$ columns by setting $g_{\text{p}} = n$, its computation will require roughly $n$ times as many operations as the original function evaluation, independent of whether $J$ is dense or sparse. However, it is well known [5,8] that the number of function evaluations that are required to compute an approximation to the Jacobian by finite differences can be much less than $n$ if $J$ is sparse. Fortunately, the same idea can be applied to greatly reduce the running time of ADIFOR-generated derivative code as well.

The idea is best understood with an example. Assume that we have a function

$$F = \begin{pmatrix} f_1 \\ f_2 \\ f_3 \\ f_4 \\ f_5 \end{pmatrix} : x \in \mathbb{R}^4 \mapsto y \in \mathbb{R}^5$$

whose Jacobian $J$ has the following structure (symbols denote nonzeros, and zeros are not shown):

$$J = \begin{pmatrix} \bigcirc & \bigcirc & \Diamond & \Diamond \\ \bigcirc & \bigcirc & \Delta & \Box \\ \bigcirc & \bigcirc & \Delta & \Box \\ \end{pmatrix}.$$

That is, the function $f_1$ depends only on $x_1$, $f_2$ depends only on $x_1$ and $x_4$, and so on. The key idea in sparse finite difference approximations is to identify so-called structurally orthogonal columns $j_i$ of $J$—that is, columns whose inner product is zero, independent of the value of $x$. In our example, columns 1 and 2 are structurally orthogonal, and so are columns 3 and 4. This means that the set
of functions that depend nontrivially on $x_1$, and the set of functions that depend nontrivially on $x_2$ are disjoint.

To exploit this structure, recall that **ADIFOR** (ignoring transposes) computes $J \cdot S$, where $S$ is a matrix with $g \# p$ columns. For our example, setting $S = I_{4 \times 4}$ will give us $J$ at roughly four times the cost of evaluating $F$, but if we exploit the structural orthogonality and set

$$S = \begin{pmatrix} 1 & 0 \\ 1 & 0 \\ 0 & 1 \\ 0 & 1 \end{pmatrix},$$

the running time for the **ADIFOR** code is roughly halved. *Note that the ADIFOR-generated code remains unchanged.*

As a more realistic example, we consider the swirling flow problem, part of the MINPACK-2 test problem collection [1]. Here we solve a nonlinear system of equations $F(x) = 0$ for $F : \mathbb{R}^n \to \mathbb{R}^n$. The swirling flow code has the form

```fortran
subroutine dswirl3(nxmax,x,fvec,fjac,ldfjac,job,eps,nint)
  integer nxmax, ldfjac, job, nint
  double precision x(*), fvec(*), fjac(ldfjac,*), eps
```

Like all codes in the MINPACK-2 test collection, it is set up to compute the function values (in **fvec**) and, if desired, the analytic first-order derivatives (in **fjac**) as well. The vectors **x** and **fvec** are of size $nxmax = 14 \times nint$. For example, for $nint = 4$, the Jacobian of $F$ is of size $nxmax = 56$ and has the structure shown in Figure 1.

![Figure 1: Structure of the swirling flow Jacobian, $n = 56$](image)

The derivative subroutine produced by **ADIFOR** is
If we initialize $g_0$ to a $5\times 56$ identity matrix and let $g_0 = 56$ and $f(14g_0)$ at least 56, then the exit from $g_0$ is a $56 \times 56$ identity matrix and $g_0 = 56$ and $f=56$. As it turns out, less than 7% of the total operations performed on the Jacobian involve non-trivial operations. On the other hand, by using a graph-coloring algorithm designed to identify structurally orthogonal columns (we used the one described in [1]), we can determine that the Jacobian can be grouped into 4 sets of structurally orthogonal columns independent of the size of the problem. In our example, columns 1, 10, 16, 31, and 37 were in the first group; columns 2, 17, 32, 43, and 48 were in the second group; and so on. We can then advantage of this fact by initializing the second column of $g_0$, such that it has 10 in rows 2, 17, 32, 43, and 48 and so on. The structure of $g_0$ thus initialized is shown in Figure 2 together with the resulting compressed Jacobian $g_0$, where the first column of $g_0$ is not initialized. Note that instead of $g_0 = 56$ we now get by with $g_0 = 14$, a sizeable reduction in cost.

**Figure 2.** Left: Structure of $g_0$. Right: Structure of $g_0'$.
enddo

g$s(color(i),i) = 1
enddo

call g$dswrl3 (nocolors, nxmax, x, g$s, pmax,
+ fvec, g$fvec, pmax,
+ fjac, ldffc, 1, eps, nint)
c
job = 1 indicates that only the function value is to be computed in
dswrl3.
c	onzero(j,i) is TRUE if the (j,i) entry in the Jacobian is nonzero,
and FALSE otherwise.
do i = 1, n
do j = 1, n
if nonzero(j,i) then
  jac(j,i) = g$fvec(color(i),j)
else
  jac(j,i) = 0.0
endif
enddo
enddo

Computing the Jacobian with ADIFOR in this way performed at least as well as the analytic
MINPACK-2 Jacobian on both a SPARC-compatible Solbourne 5E/900 and a one-processor Cray
Y/MP.

5 Case 4: Sparse Jacobian, two independent variables, one dependent variable

The coating thickness problem, conveyed to us by Janet Rogers of the National Institute of
Standards and Technology, presents many alternatives for using ADIFOR-generated subroutines.
The code for this problem is (in abbreviated form) shown below:

```
SUBROUTINE fun(n,m,np,nq,
+ beta,xplusd,ldxp)
+ f,ldf)
c
SUBROUTINE fun(n,m,np,nq,
+ beta,xplusd,ldxp)
+ f,ldf)
c Subroutine Arguments
c = n number of observations
c = m number of columns in independent variable
c = np number of parameters
c = nq number of responses per observation
c = beta current values of parameters
c = xplusd current value of independent variable, i.e., x + delta
c = ldxp leading dimension of xplusd
c = f predicted function values

14
c $\equiv \ldf$ leading dimension of f

c Variable Declarations

INTEGER i, j, k, ldf, klxdp, m, n, np, nq, numpars
INTEGER ia, ib
DOUBLE PRECISION beta(np), f(ldf, nq), xplusd(ldxdp, m)

double precision par(20), fn(2)

do 10 k = 1, np
   par(k) = beta(k)
10 continue

do 100 i = 1, n
   do 20 j = 1, m
      par(np + j) = xplusd(i, j)
20 continue

compute function values (fn) given parameters (par)

call fn(par, fn)

   f(i, 1) = fn(1)
   f(i, 2) = fn(2)

100 continue
return
cend

subroutine fn(x, fn)
integer m, n, np
parameter (np=8, m=2, nq=2)
integer i
double precision x(np+m), fn(nq)
double precision beta(np), xplusd(m)

do 10 i = 1, np
   beta(i) = x(i)
10 continue

do 20 i = 1, m
   xplusd(i) = x(np+i)
20 continue

c compute first of multi-response observations

   fn(1) = beta(1)
   + beta(2)*xplusd(1)
+ + beta(3)*xplusd(2)  
+ + beta(4)*xplusd(1)*xplusd(2)

c compute second of multi-response observations

fn(2) = bet(5)  
+ + beta(6)*xplusd(1)  
+ + beta(7)*xplusd(2)  
+ + beta(8)*xplusd(1)*xplusd(2)

return  
end

The special format of this code is due to its embedding in the ODRPACK software for orthogonal
distance regression. We are interested in the derivatives of f with respect to the variables beta and
xplusd. We shall explore various ways to do this in some detail.

5.1 Approach 1 – Generate derivatives only for fn

The easiest approach is to generate the derivative code only for fn, since it is clear from
the code that f(i,1:2) depends only on beta(1:np) and xplusd(i,1:m). ADIFOR then produces

subroutine g$fnc$3(x,g$x,1dg$x,fn,g$fn,1dg$fn)
in-ger m, np, nq
parameter( np = 8, m = 2, nq = 2)
double precision x(np+m), fn(nq), g$x(1dg$x, np+m), g$fn(1dg$fn,nq)

If inside fn we replace the call to fn with a call to g$fnc$3, always initializing g$x to a 10 x 10
identity matrix before the call, then

\[ g^f_{n}(k,j) = \frac{\partial f(i,j)}{\partial \beta(k)} , k = 1, \ldots 8, j = 1, 2. \]

and

\[ g^f_{n}(k,j) = \frac{\partial f(i,j)}{\partial xplusd(i,k - np)} , k = 9, 10. \]

Closer inspection reveals that the 10 x 2 array g$fn always has the following structure (numbers
are used to identify nonzero elements):

\[
\begin{pmatrix}
1 & 0 \\
2 & 0 \\
3 & 0 \\
0 & 5 \\
0 & 6 \\
0 & 7 \\
0 & 8 \\
9 & 10 \\
11 & 12
\end{pmatrix}
\]
In other words, \texttt{fn(i,1)} depends only on \texttt{beta(1:4)}, and \texttt{fn(i,2)} depends only on \texttt{beta(5:8)}. Hence, we can compute a compressed version of \texttt{g$fn} at reduced cost by merging rows 1 and 5, 2 and 6, 3 and 7, and 5 and 8 of \texttt{g$fn}. Keeping in mind that \texttt{g$fn} is the transpose of the Jacobian, this is an especially simple case of the compression strategy outlined in the previous section. This is achieved by initializing

\[
g$x = \begin{pmatrix} 1 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\ \end{pmatrix}
\]

which results in

\[
g$fn = \begin{pmatrix} 1 & 5 \\ 2 & 6 \\ 3 & 7 \\ 4 & 8 \\ 9 & 10 \\ 11 & 12 \\ \end{pmatrix}
\]

All the nonzero values of the Jacobian are now computed at roughly 60% of the cost of the previous approach.

On a SPARC-compatible Solbourne S5E/900 with a clock resolution of 0.01 seconds, executing \texttt{fun} took 0.01 seconds, computing derivative values using \texttt{g$fnc} without compression took 0.06 seconds, and exploiting the structure of \texttt{g$fn} through the initialization of \texttt{g$x} shown above reduced that time to 0.03 seconds.

5.2 Approach 2 – Generate derivatives for \texttt{fun}

An alternative method of applying \texttt{ADIFOR} is to process subroutine \texttt{fun}. \texttt{ADIFOR} detects the interprocedural data dependence between \texttt{fun} and \texttt{fnc} and therefore generates \texttt{g$fun$176} as well as \texttt{g$fnc$3}, with \texttt{g$fnc$3} called properly within \texttt{g$fun$176}. We obtain

\[
\text{subroutine } \texttt{g$fun$176}(g$p,n,m,\texttt{np,\texttt{nq,beta,g$beta,ldg$beta}},
\]

\[
\text{integer } g$p, n, m, \texttt{np,\texttt{nq,ldg$beta,ldg$xplusrd,ldxpd,lddf,ldg$f,ldg$ldf}},
\]

\[
\text{double precision beta(np), g$beta(1dg$beta,\texttt{np}}),
\]

\[
\text{g$xplusrd(1dxpd,m), g$xplusrd(1dg$xplusrd,ldxpd,m),}
\]

\[
\text{f(1df,\texttt{np}), g$f(1dg$f,1df,\texttt{nq})}
\]

Now we have three-dimensional derivative objects, which somewhat complicates the initialization of \texttt{g$xplusrd} and the interpretation of the results in \texttt{g$f}. However, this is not too difficult if we keep in mind that we wish to initialize
\[
\begin{pmatrix}
g$\beta^T \\
g$xplusd^T
\end{pmatrix}
\]
to an identity matrix. The number of elements in \texttt{xplusd} is \(n \times m\), and the number of elements in \texttt{beta} is \texttt{np}. For the coating thickness problem, \(n=63\), \(m=2\), and \texttt{np}=8. Hence, the identity matrix should be \(134 \times 134\). This is also the value we shall use for \texttt{g$p}$. Initialization of \texttt{g$\beta} follows the scheme outlined in § 3; that is, the first 8 rows should be an \(8 \times 8\) identity matrix, and the remaining 126 rows should be initialized to zero. How to initialize \texttt{g$xplusd} is less readily apparent, for it is not immediately obvious how to form a \(126 \times 126\) identity matrix from a three-dimensional structure. However, if one looks at the way Fortran stores two-dimensional structures in memory, a simple scheme for storing the Jacobian develops. In Fortran, element \((j,i)\) in an \(n \times m\) array is stored as if it were element \(n \times (i-1) + j\) of a one-dimensional array. Thus, we can apply this technique to map the 126 columns of the Jacobian that should be initialized to the identity onto \texttt{g$xplusd}. Specifically, element \((np+k,j,i)\) is initialized to 1 if and only if \(k = 63 \times (i-1) + j\). The following code segment accomplishes this initialization.

c n=63, m=2, np=8
\[
g$p$ = np + m*n
do 44 i = 1, np
do 144 j = 1, g$p$
g$\beta(j,i) = 0.0$
144 continue
g$\beta(i,i) = 1.0$
44 continue
do 45 i = 1, 4
do 145 j = 1, n
do 245 k = 1, g$p$
g$xplusd(k,j,i) = 0.0
245 continue
g$xplusd(np*((i-1)*n)+j,j,i) = 1.0
145 continue
45 continue

When initialized in this manner, \texttt{ADIFOR} computes
\[
g$f = \left( J_f = \begin{pmatrix} \frac{\partial f}{\partial \beta} & \frac{\partial f}{\partial xplusd} \end{pmatrix} \right)^T.
\]
However, the performance of this approach is poor, since we totally ignore the sparsity structure of the Jacobian. As a result, the computation of \(J_f\) takes 0.77 seconds on a Solbourne 5E/900. A better way to find the Jacobian of \(f\) using \texttt{g$fun$} is to take note of the structures used by \texttt{fun}. From this, it becomes obvious that \(\frac{\partial f}{\partial xplusd}\) is nonzero only when \(i = k\). As a consequence, we may change the
\[
gsp = np + m \times n \\
g\timesplusd(np+((i-1) \times m)+j,j,i) = 1.0
\]

to the much simpler
\[
gsp = np + m \\
g\timesplusd(np+i,j,i) = 1.0
\]

with the understanding that \( g\timesf(np+i,j,k) (i = 1..m) \) represents \( \frac{\partial f[i,k]}{\partial \timesplusd[j,i]} \). This is equivalent to initializing
\[
g\timesbeta = \\
\begin{pmatrix}
1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\end{pmatrix}
\]
\[
, \text{ and } g\timesplusd[n] = \\
\begin{pmatrix}
0 & 0 \\
0 & 0 \\
0 & 0 \\
0 & 0 \\
0 & 0 \\
0 & 0 \\
1 & 0 \\
0 & 1 \\
\end{pmatrix}
\]

This implementation is much more efficient than that described in the preceding paragraph and more closely mimics the behavior of the original subroutine \texttt{fun}. As a consequence, the time required to execute \texttt{g\timesfun\$176} using this initialization is 0.07 seconds.

As discussed in § 5.1, only half of the derivatives of \texttt{f} with respect to \texttt{beta} are nonzero. Specifically, \( \frac{\partial f[k]}{\partial \timesbeta[i]} \) is nonzero for \( j = 1.4 \) and zero for \( j = 5.8 \), while \( \frac{\partial f[k]}{\partial \timesplusd[i,j]} \) is zero for \( j = 1.4 \) and nonzero for \( j = 5.8 \). This information can be used to further compress the Jacobian. The initialization
\[
g\timesbeta = \\
\begin{pmatrix}
1 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 1 & 0 & 0 & 0 & 1 & 0 \\
0 & 0 & 0 & 1 & 0 & 0 & 0 & 1 \\
0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\end{pmatrix}
\]
\[
, \text{ and } g\timesplusd[n] = \\
\begin{pmatrix}
0 & 0 \\
0 & 0 \\
0 & 0 \\
0 & 0 \\
0 & 0 \\
1 & 0 \\
0 & 1 \\
\end{pmatrix}
\]

compresses the Jacobian into only 6 columns. Columns 1 through 4 represent the nonzero derivatives of \texttt{f} with respect to \texttt{beta}, while columns 5 and 6 correspond to the derivatives of \texttt{f[i,j]} with respect to \texttt{xplusd[i,1..2]}, as above. This initialization may be accomplished with the following code fragment.
c n=63, m=2, np=8
  halfnp = 4
  g$p$ = 4 + m
  do 44 i = 1, halfnp
     do 144 j = 1, g$p$
        g$beta(j,i) = 0.0
        g$beta(j,i+halfnp) = 0.0
     144 continue
     g$beta(i,i) = 1.0
     g$beta(i,i+halfnp) = 1.0
  44 continue
  do 45 i = 1, m
     do 145 j = 1, n
        do 245 k = 1, g$p$
           g$plusd(k,j,i) = 0.0
        245 continue
        g$plusd(halfnp+i,j,i) = 1.0
  145 continue
  45 continue

This approach is efficient, capable of computing all derivatives in 0.03 seconds. However, it has the disadvantage that the initialization routine might have to be changed if fnC or np is altered.

6 Computing Gradients of Partially Separable Functions

A particular class of functions that arises often in optimization contexts is that of the so-called partially separable functions [6,11,12,14,15]. That is, we have a function $f: \mathbb{R}^n \rightarrow \mathbb{R}$ which can be expressed as

$$f(x) = \sum_{i=1}^{n} f_i(x).$$

Usually each $f_i$ depends on only a few (say, $n_i$) of the $x$'s, and one can take advantage of this fact in computing the (sparse) Hessian of $f$.

As was pointed out to us by Andreas Griewank, this structure can also be used advantageously in computing the (usually dense) gradient $\nabla f$ of $f$.

Assume that the code for computation of $f$ looks as follows:

```
subroutine f(n,x,fval)
  integer n
  real x(n), fval, temp

  fval = 0

  call f1(n,x,temp)
  fval = fval + temp
```

20
call fnb(n,x,temp)
fval = fval + temp

return
end

If we submit f to ADIFOR, it generates

subroutine g$fn(n,x,g$x,ldg$x,fval,g$fval,ldg$fval).

To compute $\nabla f$, the first (and only) row of the Jacobian of f, we set g$p= n$ and initialize g$x$ to a $n \times n$ identity matrix. Hence, the cost of computing $\nabla f$ is of the order of $n$ times the function evaluation.

As an alternative, we realize that with $f : \mathbb{R}^n \rightarrow \mathbb{R}^{nb}$ defined as

$$g = \begin{pmatrix} f_1 \\ \vdots \\ f_{nb} \end{pmatrix},$$

we have the identities

$$f(x) = e^T g(x),$$
and hence

$$\nabla f(x) = e^T J_g,$$

where $e$ is the vector of all ones, and $J_g$ is the Jacobian of $g$. We can get the gradient of $f$ by computing $J_g$ and adding up its rows. The corresponding code fragment for computing $f$ is

subroutine f(n,x,fval)
integer n
real x(n)

integer nf, i
parameter (nf = <whatever>)
real gval(nf)

call g(n,x,gval)

fval = 0
do i = 1,nb
   fval = fval + gval(i)
enddo

return
end

It may not appear that we have gained anything, since $J_g$ is $nf \times n$: if we initialize g$x$ in

subroutine g$g$(g$p,n,x,g$x,ldg$x,g$val,g$gval,ldg$gval)
to an \( n \times n \) identity matrix, then the computation of \( J_g \) still takes about \( n \) times as long as the computation of \( g \) (or \( f \)).

The key observation is that the Jacobian \( J_g \) is likely to be sparse, since

\[
J_g = \begin{pmatrix}
(\nabla f_1)^T \\
\vdots \\
(\nabla f_n)^T
\end{pmatrix},
\]

and each of the \( f_i \)'s depends only on \( n_i \) of the \( x \)'s. By using the graph coloring techniques described in Section 4, we can compute \( J_g \) at a cost that is proportional to the number of columns in the compressed \( J_g \), and then add up its (sparse) rows. As a result, we can compute \( \nabla f \) at a cost that is potentially much less than \( n \) times the evaluation of \( f \).

7 Conclusions

This report demonstrated how to properly use ADIFOR-generated derivative codes. One of the strengths of ADIFOR is that it does not assume a particular calling sequence of the function to be differentiated. We gave examples that showed how to properly use ADIFOR-generated codes for various styles of codes. We also showed how to exploit a known sparsity structure of the derivative matrix in the initialization of ADIFOR code. By properly initializing the derivative objects corresponding to independent variables, we can merge structurally orthogonal columns and hence compute derivatives at greatly reduced cost. We also mentioned partially separable functions, where this technique can also be applied advantageously to the computation of dense gradient objects.

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References


