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Efficient Computation of Gradients and Jacobians by Transparent Exploitation of Sparsity in Automatic Differentiation*

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Abstract. Automatic differentiation (AD) is a technique that augments computer codes with statements for the computation of derivatives. The computational workhorse of AD-generated codes for first-order derivatives is the linear combination of vectors. For many large-scale problems, the vectors involved in this operation are inherently sparse. If the underlying function is a partially separable one (e.g., if its Hessian is sparse), many of the intermediate gradient vectors computed by AD will also be sparse, even though the final gradient is likely to be dense. For large Jacobians computations, every intermediate derivative vector is usually at least as sparse as the least sparse row of the final Jacobian. In this paper, we show that transparent exploitation of the sparsity inherent in derivative computation can result in dramatic gains in runtime and memory savings. For a set of gradient problems exhibiting implicit sparsity, we report on the runtime and memory requirements of computing the gradients with the ADIFOR (Automatic Differentiation of FORtran) tool, both with and without employing the SparsLinC (Sparse Linear Combinations) library, and show that SparsLinC can reduce runtime and memory costs by orders of magnitude. We also compute sparse Jacobians using the SparsLinC-based approach—in the process, automatically detecting the sparsity structure of the Jacobian—and show that these Jacobian results compare favorably with those of previous techniques that require a priori knowledge of the sparsity structure of the Jacobian.

Key words. Automatic Differentiation, Sparsity, Partial Separability, Sparse Jacobians, Large-Scale Optimization, MINPACK-2, ADIFOR, SparsLinC.

1 Introduction

The numerical computation of gradients and Jacobians is an important step in the solution of many nonlinear problems, such as numerical optimization, mesh computations, nonlinear least squares, and systems of differential and algebraic equations. For such problems, the derivative computation often is a major contributor to the overall cost, in terms of both runtime and memory requirements.

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Automatic differentiation (AD) is a technique by which codes for the computation of functions are augmented to produce codes for the computation of desired derivatives of those functions. AD relies on the fact that every function is executed on a computer as a sequence of elementary operations, such as + and −, and intrinsics, such as sin and log. By successive applications of the chain rule to the composition of those elementary operations, derivatives can be computed exactly (up to machine roundoff) and in a completely mechanical fashion. For a detailed description of AD, see [17].

Previous studies have shown that AD is a powerful technique for computing derivatives. Accuracy and runtime efficiency of AD are superior to difference approximations, and the reliability, flexibility and development-time efficacy of AD surpass those of hand-coding approaches. For example, the AD tool ADIFOR (Automatic Differentiation of FORtran) [4, 6, 7] has been used to generate derivative codes for many applications in areas such as large-scale optimization [2, 13], computational fluid dynamics [8, 9, 14], weather modeling [22], and groundwater modeling [11]. Other examples of available AD tools are Odysseé [23] and GRESS [20] for Fortran, and ADOL-C [18] and ADIC [10] for C programs.

The forward and reverse modes are the two basic modes of AD, and are distinguished by the manner in which the chain rule is applied for propagating derivatives. This basic difference impacts the computational complexity and flexibility of each mode. The forward mode, despite its flexibility and the predictability of the memory requirements, has thus far been considered impractical for the computation of large-scale gradients, because of its runtime complexity. In this paper, we will show that in some cases this perceived limitation can be overcome by exploitation of sparsity.

Computationally, the most expensive kernel of derivative codes generated by primarily forward mode tools such as ADIFOR and ADIC is a vector linear combination (VLC):

\[ w = \sum_{i=1}^{k} \alpha_i v_i, \]  

where \( w \) and the \( v_i \) are gradient vectors, the \( \alpha_i \) are scalar multipliers, and \( k \) is the arity. If we assume that the gradient vectors are all of same size, say \( d \), then a computation composed primarily of invocations of (1.1), implemented as a dense vector loop, would have computational complexity linear in \( d \), in terms both of runtime and memory requirements (this type of implementation is used in the default or NONSPARSE mode of ADIFOR, examples of which we will show in Section 3). On the other hand, if the gradient vectors are sparse, then strategies that attempt to exploit that sparsity, in terms of both the storage of vectors and the implementation of the computation, can be expected to reduce the overall computational complexity. For any given problem, the extent of this reduction in complexity will depend not only on the efficiency of the sparse algorithm, but also on the sparsity characteristics of the underlying function.

For many large-scale problems there is inherent sparsity in the computation of gradients and Jacobians. In gradient computations, the scalar output function, \( f : \mathbb{R}^n \rightarrow \mathbb{R} \), is usually dependent on all of the inputs, resulting in a dense gradient vector \( g = \nabla f \in \mathbb{R}^n \). However, it has been shown that if the Hessian of \( f \) is sparse, then \( f \) is partially separable.
that is, \( f \) can be expressed as
\[
f(x) = \sum_{i=1}^{m} f_i(x),
\]
where \( m \) is the number of element functions \( f_i(x) \), and each \( f_i(x) \) is typically a function of just a few of the components of \( x \), implying that each \( g_i(x) = \frac{\partial f_i(x)}{\partial x} \) will be sparse, even though \( g(x) = \sum_{i=1}^{m} g_i(x) \) is dense.

Sparsity in the derivative computations is likewise a salient issue in many large-scale Jacobian computations where the final Jacobian associated with a function \( f: \mathbb{R}^n \rightarrow \mathbb{R}^m \) is sparse. As we will see in Section 2, the sparsity of the final derivative result implies equal or greater potential for sparsity in the intermediate computations.

In this paper we report on the runtime and memory performance measures of differentiating large-scale optimization codes with ADIFOR when interfaced with the SparsLinC (Sparse Linear Combinations) library \([5, 7]\), and contrast these results with \textsc{NonSparse} ADIFOR results (i.e., without SparsLinC). The codes are taken from the MINPACK-2 test set problems \([1]\) and the molecular distance geometry class of global minimization functions \([21]\). In Section 2 we look at sparsity in the context of the computation of VLCs in AD and show how this can be exploited with SparsLinC. Computational results for the gradient and Jacobian experiments are shown in Sections 3 and 4, respectively. Finally, we present our conclusions in Section 5.

2 Sparsity in First-Order Derivative Computation

For large-scale problems, gradient vectors appearing in VLCs in AD computations are often very sparse. In this section, we define sparsity in the context of AD computations, and identify characteristic problem types where sparsity in derivative computations occurs. We also give a brief overview of SparsLinC and show how it exploits sparsity.

2.1 Definition of Sparsity in Automatic Differentiation

To define sparsity, we first revisit some basic AD and ADIFOR definitions. All AD tools require the user to specify, from among the program variables, the \textit{independent and dependent variables}; the independents being the ones with respect to which the partial derivatives of the dependents are to be computed. In addition, ADIFOR performs an activity analysis pass, whereby all variables lying on the dependency path from the independent to the dependent variables are nominated as \textit{active variables} (by definition, the independents and dependents are themselves active).

A \textit{directional derivative}, defined as
\[
\lim_{h \to 0} \frac{f(x + hd_i) - f(x)}{h},
\]
is the partial derivative of an active variable along a direction vector \( d_i \). In ADIFOR, the user specifies the complete set of desired directions by means of the \textit{seed matrix}. In the simplest case, each unit direction is defined by one of the independent variables, which is equivalent to setting the seed matrix to the identity matrix. A \textit{directional gradient vector
is defined to be the set of directional derivatives of any scalar active variable with respect to all directions specified in the seed matrix (the term scalar active variable here refers to active variables declared as scalars in the program and also to the individual elements of active variables that are declared as arrays). In the context of AD, the vector operands in (1.1) are directional gradient vectors.

In the \textbf{non} \textbf{sparse} representation, a directional gradient vector \( V \) would be declared in the derivative code as an array variable of length \( d \), where \( d \) is the number of directions. We denote the number of nonzeros in \( V \) at a given point \( t \) during the execution by \( V_{t, \text{nonz}} \). The percentage of zero entries or \textit{sparsity} of \( V_t \) is defined as

\[
V_{t, \text{sparsity}} := (1 - \frac{V_{t, \text{nonz}}}{d}) \cdot 100\%.
\] (2.2)

A good measure for the \textit{overall sparsity} present in a derivative computation is the median of the sparsities of all directional gradient vectors computed during the execution of the derivative code. We also define the umbrella term \textit{sparsity characteristics} to refer to the sparseness of all directional gradient vectors during the execution of a derivative code.

One implication of (2.2) is that \( d \) sets an upper bound on the overall sparsity in a problem. For example, if \( d = 10 \), overall sparsity is bounded above by 90\% (given that, from the definition of active variables, usually \( V_{t, \text{nonz}} \geq 1 \)). This underscores a necessary (but not sufficient) condition in terms of the practicality of a sparse solution: in order for SparsLinC—or any other strategy for exploiting sparsity in AD—to improve the runtime performance of derivative computation, the number of directions with respect to which we wish to compute derivatives should be “large.” Otherwise, the directional gradient vectors will be short, and runtime efficiencies gained by exploiting sparsity will be defeated by implementation overheads. The determination of what is considered a large sparse problem is to a great extent dependent upon the nature of the problem.

Another important fact relating to sparsity in derivative computations is that final sparsity of the least sparse directional gradient vector corresponding to the dependent variables is in all likelihood the lower bound on the sparsities of all intermediate directional gradient vectors during the execution. This means that the overall sparsity of the problem may be much higher than that of the final derivative result. This fact follows from the formation of structural merges in VLCs. The notational equation (2.3) is a depiction of a structural merge, that is, the additive propagation of nonzero structures of the right-hand-side vectors to the left-hand-side vector:

\[
\begin{align*}
\begin{pmatrix}
\diamond \\
\diamond \\
\diamond
\end{pmatrix}
\leftarrow
\begin{pmatrix}
\diamond \\
\end{pmatrix}
+ 
\begin{pmatrix}
\diamond \\
\end{pmatrix}
+ 
\begin{pmatrix}
\diamond \\
\diamond
\end{pmatrix}.
\end{align*}
\] (2.3)

Here, the symbol \( \diamond \) represents a nonzero entry, zero entries are not shown, \( d = 5 \), and the nonzero index set of the resulting left-hand-side vector is the union of index sets of the right-hand-side vectors.* Given that a left-hand-side vector in a VLC will usually

*This discussion precludes the possibility of the occurrence of (i) numerical zeroes resulting from exact
Figure 2.1: Sparsity of vectors on the left-hand side of vector linear combinations performed by SparsLinC during the execution of the gradient code for GL2, \( n = 2,500 \).

appear as a right-hand-side vector in a subsequent VLC, it is easy to see that the impact of the statement-level structural merge in (2.3) is that sparsity diminishes as the computation proceeds. It is the converse of this result that motivates our work, namely, if we suspect that the final derivative object we seek is sparse, then we can expect that all directional gradients vectors (i.e., operands of all VLCs in the derivative computation) will be as sparse—and, with great likelihood, more sparse.

Figure 2.1 depicts the sparsity characteristics of directional gradient vectors in the gradient computation of the GL2 problem, which will be introduced in Section 3. The computation involved over 30,000 VLCs. For every 250 VLCs, we gathered statistics on the number of nonzeros in the resultant left-hand-side vectors and plotted the minimum, maximum, median, and average on the graph. We note that the fact that the maximum drops from about 2,500 nonzeros to about 500 at around the 26,000-th VLC is not indicative of a decrease in the number of nonzeros of a particular vector, but rather that among the vectors appearing on the left-hand side of the 250 VLCs previous to the 26,000-th, the one with the maximum number of nonzeros had about 500. In fact, the final gradient vector is fully dense (with 2,500 nonzeros).

The main conclusion to be drawn from Figure 2.1 is that the median and average curves are always far below 2,500, the maximum vector size, thus implying that significant savings could be achieved in the computation of VLCs by exploiting sparsity. We note that the overall sparsity inherent in a derivative computation is largely a function of sparsity present in intermediate computations, and not so much the sparsity of the final result.

Cancellation (e.g., \( a + (-a) \)), and (ii) zero multipliers. In our experience, exact cancellation rarely occurs in derivative computation, and currently, SparsLinC does not check for it (i.e., numerically zero vector entries are treated like nonzero entries). SparsLinC does, however, check for zero multipliers, and vectors with zero multipliers are not referenced in VLC computations.
2.2 Characteristic Sparse Derivative Problems

Partially separable functions and their gradients arise in many computational contexts, in particular in large-scale optimization problems. In Section 1 we defined these functions (1.2) as a summation of element functions. Since each element function depends only on a very few components of the independent variable, each corresponding directional gradient vector is very sparse, and typically, the only dense vector present in the computation is the gradient of the partially separable function itself (the example in Figure 2.1 is one such case).

In terms of how to exploit the sparsity in these gradient computations, it is important to note that the mathematical notion of separability defined in (1.2) may not always be reflected in terms of how the code for a partially separable function is written (i.e., the separability may not be reflected in the code structure). An algorithmic approach for computing derivatives that transparently (i.e., without the need to rewrite code) exploits this “under-the-rug” sparsity in a code is the only approach to exploit this property without the need to rewrite the code. Our results in Section 3 will demonstrate the efficiency of computing gradients of partially separable functions using SparsLinC as a means of exploiting sparsity.

Large sparse Jacobian problems occur frequently in many contexts, and computing them without effective sparse strategies either has been prohibitively expensive or has depended on a priori knowledge of the closure of the sparsity patterns of the Jacobian, where closure denotes the union of the sparsity patterns for all values of the independent variables. In contrast to partially separable functions where some gradient vectors may be dense, the AD computation of sparse Jacobians, by definition, involves only sparse directional gradient vectors.

In Section 4 we briefly review the compressed Jacobian approach, which has previously been used [16, 2] as an effective strategy for computing sparse Jacobians in cases where the closure sparsity pattern is known. We then present our results of computing sparse Jacobians with known closures using the ADIFOR/SparsLinC approach, and contrast these results with those based on the compressed Jacobian method.

2.3 Exploiting Sparsity with SparsLinC

Our intent is to take advantage of inherent sparsity in directional gradient vectors to reduce both memory requirements and runtime of derivative computation. By devising a scheme that would preclude the storage of zero entries as well as extraneous “zero-sum” computations resulting from the zero entries, we can save on memory and runtime. The dynamic nature of the directional gradient vectors during the execution of AD-generated derivative codes, in terms of the number of nonzero entries in a vector at a given time, as well as the goal of obviating the need for a priori sparsity information requires that the vectors be represented by using dynamic data structures.

SparsLinC is a library of C routines that provide an implementation of VLCs employing dynamic data structures to represent the directional gradient vectors. When invoked in its SPARSE mode, ADIFOR generates code in which each VLC is implemented as a call to a SparsLinC routine, as opposed to the DO-loop implementation in the default (NONSPARSE)
mode. Whereas for every scalar active variable, \texttt{NONSPARSE} ADIFOR allocates a corresponding directional gradient vector of size $d$ containing the derivative values (many of which may possibly remain zero for the duration of the computation), \texttt{SPARSE} ADIFOR allocates an \texttt{INTEGER} variable that is interpreted as a pointer to SparsLinC's representation of the vector.

The design of SparsLinC is based on representing the nonzero information in each vector (i.e., the nonzero values and their corresponding vector indices) in one of three data structures—one to represent a vector with zero or one nonzero entries; another for a vector with a few nonzeroes, and a third for a vector with contiguous ranges of nonzeroes. The SparsLinC polyalgorithm heuristically switches between these representations. SparsLinC performs the necessary management of its internal vector representations, including the implementation of a "grow as you go" strategy for the recruitment of dynamically allocated memory for the storage of the sparse vectors.

The basic module of SparsLinC performs a VLC by constructing the left-hand-side vector as it traverses—in ascending order of indices—the vectors on the right-hand side. A special "plus-equals" module takes advantage of the existing data structure of the left-hand-side vector in cases in which the same vector also appears on the right-hand side, thus resulting in a more efficient computation.

Two features of SparsLinC are of particular relevance to the present discussion: the transparent exploitation of partial separability, and automatic detection of the sparsity pattern of the Jacobian. Previous attempts at exploiting partial separability in the context of AD have been based on manual modification of the code for the function in such a way as to transform the derivative computation from a dense gradient problem to a sparse Jacobian problem [12]. The transparent exploitation of partial separability in SparsLinC does not require code modification. Instead, it relies on the sparse representation and processing of the VLCs involving the element functions, and on the efficiency of the plus-equals module for the accumulation of the dense gradient vector.

The detection of the sparsity pattern of Jacobians is of interest in a number of computations (e.g., see [13]). As we shall see in Section 4, sparsity detection is a prerequisite for compressing Jacobians using coloring methods. In our Jacobian experiments we use prepackaged MINPACK-2 routines for detecting the sparsity pattern of the Jacobian for each problem. However, in the absence of such special routines (or in cases where the sparsity pattern varies for differing values of the independent variable), we can use SparsLinC for this purpose. The computation of the Jacobian using SparsLinC yields the sparsity pattern of the Jacobian as a natural byproduct of the work it does in computing the Jacobian.

A detailed description of the SparsLinC library will be reported elsewhere, and the full Fortran interface specification can be found in [5].

3 Computing Gradients of Partially Separable Functions

The MINPACK-2 Test Problem Collection [1] contains a number of unconstrained optimization problems from a variety of application areas. Table 3.1 shows the six MINPACK-2 problems we used in our gradient experiments. For each problem, we computed the gradient using both the \texttt{NONSPARSE} ADIFOR approach and the \texttt{SPARSE} ADIFOR/SparsLinC approach, and compared the memory requirements and runtimes of the two approaches (in what follows we will simply refer to the two approaches as \texttt{NONSPARSE} and \texttt{SPARSE}, respec-
tively). We performed our experiments on two workstation platforms: Sun SPARCstation IPX, and IBM RS6000-370.

The gradient values obtained for all problems using both the NONSPARSE and SPARSE approaches agreed to within approximately machine precision of the hand-coded derivatives, available in MINPACK-2.

3.1 Memory Requirements

In terms of feasibility, memory is a critical issue in the computation of large gradients. The memory required for a straightforward AD implementation of the gradient computation would be roughly equivalent to augmenting the memory required for the function computation by a factor of \( n \), where \( n \) is number of components of the independent variable, alternately referred to as the problem size. For large \( n \), this linear expansion can lead to excessive paging, or simply be prohibitive. For example, a NONSPARSE ADIFOR implementation of the gradient computation for the GL2 problem of size 160,000 would require more than 400 gigabytes of virtual memory.

One approach to breaking the linear memory dependence is stripmining of derivative computation. This approach involves dividing the gradient computation into strips, each strip consisting of the differentiation with respect to a few components of the independent variable, and each strip having reasonable memory requirements (as we will see in Section 4.3, stripmining can also be used in Jacobian computations). In the case of ADIFOR-generated code, stripmining can be done conveniently through the seed matrix mechanism. The runtime penalty for this approach is an extraneous function recomputation per strip, which can be effectively amortized for large strip sizes. Since each strip is computed independently, stripmining can also be used for easy parallelization of the gradient computation [9].

Table 3.2 contains a summary of the memory requirements* of our gradient experiments for the case of \( n = 160,000 \). We use the notation \( M[C] \) to denote the memory requirements of a computation \( C \), and we report only one set of numbers, since memory requirements on the two test-bed platforms are identical. The first column in Table 3.2 shows the memory

<table>
<thead>
<tr>
<th>Name</th>
<th>Description of the Problem</th>
</tr>
</thead>
<tbody>
<tr>
<td>EPT</td>
<td>Elastic-Plastic Torsion</td>
</tr>
<tr>
<td>GL2</td>
<td>Ginzburg-Landau (2-d) superconductivity</td>
</tr>
<tr>
<td>MSA</td>
<td>Minimal Surface Area</td>
</tr>
<tr>
<td>ODC</td>
<td>Optimal Design with Composite materials</td>
</tr>
<tr>
<td>PJB</td>
<td>Pressure distribution in a Journal Bearing</td>
</tr>
<tr>
<td>SSC</td>
<td>Steady State Combustion</td>
</tr>
</tbody>
</table>

*The Unix command `size executable-filename` reports the total amount of statically allocated memory (i.e., the memory requirements that can be assessed at link-time such as array sizes, etc.) needed to load and run the executable. In the case of SparsLinC, where memory is also allocated dynamically, we call a SparsLinC routine that reports the total amount of dynamically allocated memory, and add this to the statically allocated memory, in order to arrive at the total memory requirements.
required for computing the original function, \( F \). The next double column shows first the memory requirements, \( M\{G_{AD-16}\} \), of a stripmined \texttt{NONSPARSE} computation with a strip size \( p_s = 16 \), and then the ratio of this gradient memory requirement to that of the corresponding original function. We note that in each case except one, \( M\{G_{AD-16}\}/M\{F\} \) is slightly greater than \( p_s \), matching our expectation of linear augmentation in memory requirements. The exception is the MSA problem, where the \( M\{G_{AD-16}\}/M\{F\} \) ratio is smaller than \( p_s \). Here the computations involving a large work array in the original function are unrelated to derivative computation; hence ADIFOR does not generate a corresponding augmented derivative work array (for a full discussion of this issue, see “Variable Nomination” in [7]). We also note that the equivalence of the \( M\{F\} \)’s and \( M\{G_{AD-16}\} \)’s of the EPT, ODC, PJB and SSC problems is due to the similarity of the code structures of these problems.

The second double column in Table 3.2 shows the memory requirements of the \texttt{SPARSE} computation, \( M\{G_{Sparse}\} \), along with the ratio of this gradient memory requirement to that of the corresponding original function. Here we see the terseness of SparsLinC’s memory allocation scheme in that \( M\{G_{Sparse}\} \) is always within a factor 4.9–11.0 of \( M\{F\} \). The variation among these ratios is indicative of the variation in the sparsity characteristics of their corresponding gradient computations. Note that in all cases, \( M\{G_{Sparse}\} < M\{G_{AD-16}\} \), despite the fact that \( G_{Sparse} \) computes the whole gradient at once, whereas \( G_{AD-16} \) computes the gradient a strip at a time.

### 3.2 Runtime

Figure 3.1 is a summary of all our gradient runtime results. For each of the optimization problems in Table 3.1, we timed the \texttt{SPARSE} and \texttt{NONSPARSE} gradient computations for several problem sizes ranging from \( n = 2,500 \) to \( n = 160,000 \). Similar to the case of memory requirements, here we were interested in the augmentation factor of the runtime of the gradient computation with respect to the runtime of the function computation. Subsequently, the plots shown in the six panels of Figure 3.1 are all gradient-to-function runtime ratio plots.

Each panel shows the gradient-to-function ratios for two of the MINPACK-2 problems, for both \texttt{SPARSE} and \texttt{NONSPARSE} gradient computations on a given machine. Note also that

<table>
<thead>
<tr>
<th>Problem</th>
<th>( M{F} )</th>
<th>( M{G_{AD-16}} )</th>
<th>( M{G_{AD-16}}/M{F} )</th>
<th>( M{G_{Sparse}} )</th>
<th>( M{G_{Sparse}}/M{F} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>EPT</td>
<td>1.29</td>
<td>23.06</td>
<td>17.8</td>
<td>14.15</td>
<td>11.0</td>
</tr>
<tr>
<td>GL2</td>
<td>2.59</td>
<td>45.03</td>
<td>17.4</td>
<td>19.65</td>
<td>7.6</td>
</tr>
<tr>
<td>MSA</td>
<td>2.57</td>
<td>24.34</td>
<td>9.5</td>
<td>12.60</td>
<td>4.9</td>
</tr>
<tr>
<td>ODC</td>
<td>1.29</td>
<td>23.06</td>
<td>17.8</td>
<td>13.05</td>
<td>10.1</td>
</tr>
<tr>
<td>PJB</td>
<td>1.29</td>
<td>23.06</td>
<td>17.8</td>
<td>14.16</td>
<td>11.0</td>
</tr>
<tr>
<td>SSC</td>
<td>1.29</td>
<td>23.06</td>
<td>17.8</td>
<td>14.15</td>
<td>11.0</td>
</tr>
</tbody>
</table>
Figure 3.1: Log-Log plots for each platform of gradient-to-function runtime ratios of the SPARSE and NONSPARSE runs for all gradient problems.
Figure 3.2: Sparsity of vectors on the left-hand side of vector linear combinations performed by SparsLinC during the execution of the gradient code for MSA, \( n = 2,500 \).

for each panel both axes are logarithmic in scale (\( \log_{10} \)).

The most salient result evident in Figure 3.1 is the disparity between the \texttt{SPARSE} and \texttt{NONSPARSE} runtime ratios for each problem. In all cases the \texttt{NONSPARSE} gradient computation displays a purely linear behavior with respect to the function computation over the range of problem sizes. Note that in all cases the linear coefficient is very close to 1 (by virtue of the fact that all the \texttt{NONSPARSE} plots are close to the \( y = x \) diagonal line). As we mentioned in Section 1, this is due to the D0-loop implementation of VLCs in the derivative code generated by ADIFOR in the default or \texttt{NONSPARSE} mode. On the other hand, the \texttt{SPARSE} runtime ratios, though not independent of \( n \), show a markedly reduced dependency on \( n \).

Note that in the case of four of the problems (EPT, MSA, ODC, and SSC), gradient computation is clearly feasible with SparsLinC, whereas the \texttt{NONSPARSE} computation is so costly that it would be considered impractical. For example, the runtime of computing the MSA and ODC gradients of size 160,000 on the SPARC IPX is less than 100 times the function runtime, which is a reasonable cost for some applications, given that SparsLinC requires no rewriting of code. We also note that this cost can be further reduced by efficient hand-coding of the gradient or, in the case of partially separable functions, by reformulating the gradient computation as a sparse Jacobian computation [3].

The variance between the degree of disparity between \texttt{NONSPARSE} and \texttt{SPARSE} results is due to the differing sparsity characteristics of the MNPACK-2 problems. Figure 3.2 depicts the sparsity characteristics of the MSA problem, for the case of \( n = 2,500 \). In contrast to Figure 2.1, where for parts of the GL2 problem we observe medians of approximately 1000 nonzeros per vector, in the case of the MSA computation the median is always approximately 1. Correspondingly, we note in Figure 3.1 the markedly smaller gradient-to-function ratios for the \texttt{SPARSE} MSA computation as compared with the \texttt{SPARSE} GL2
Figure 3.3: Logarithmic plot for each platform of \texttt{SPARSE} divided by \texttt{NONSPARSE} runtimes for all gradient problems.
computation.

One aspect of the results in Figure 3.1 is that despite the qualitative similarity of the corresponding curves on the SPARC IPX and the RS6000, the quantitative results differ in certain characteristic ways. We note that in general, the disparity between the SPARSE and NONSPARSE results are somewhat larger on the SPARC IPX. As is clear from the plots, this is caused both by the SPARSE ratios being smaller on the SPARC IPX (compare for example the SPARSE MSA results on the two platforms) and by the NONSPARSE ratios being larger.

These performance differences are a reflection of architectural features of the two platforms. The SPARC IPX essentially has a scalar processor and a flat memory hierarchy. Hence, vector operations execute only marginally faster, and memory locality (i.e., the reuse of data and the accessing of adjacent memory locations) is not much of an issue. In contrast, the RS6000 architecture employs a superscalar chip and a cache-based memory architecture. Hence, this machine performs better if executing short vector operations, since these operations can fill the short pipes and take advantage of memory locality. This is advantageous in the NONSPARSE computation where the VLCs are implemented via DO-loops. On the other hand, indirect addressing, used extensively in the SparsLinC algorithm, while fairly inconsequential on the SPARC, may lead to a performance degradation on the RS6000, as memory locality suffers.

Figure 3.3 is an alternative way of presenting the results in Figure 3.1. Here we have plotted for each machine and each problem the log_{10} of the ratio of NONSPARSE to SPARSE runtimes in order to demonstrate the orders of magnitude runtime improvements achieved by SparsLinC. Note that the line definition box also includes for each problem the true ratio of NONSPARSE to SPARSE runtimes for the case of n = 160,000.

For the largest problem size we note that the improvements in the gradient computation due to SparsLinC range from a factor of nearly 10 in the case of the PJB problem running on the RS6000, to a factor of greater than 1,000 for the ODC problem on the SPARC IPX. We also note that for large problem sizes the runtime efficiencies achieved by SparsLinC near a plateau. We suspect that this is because the dense gradient computation increasingly dominates the overall computational task.

4 Computing Sparse Jacobians

For our sparse Jacobian experiments we selected five problems from the MINPACK-2 problem set of systems of nonlinear equations and one problem from the molecular distance geometry (MDG) class of global minimization functions. The MINPACK-2 problems are identified in Table 4.1. These are a standard set of test problems; the reader is referred to [1] for detailed descriptions.

The MDG problem is described in Moré and Wu [21] as follows: Given bond lengths \( \delta_{i,j} \) between a subset \( S \) of the atom pairs, determine whether there is a molecule that satisfies these bond length constraints. Moré and Wu formulated this problem in terms of finding the global minimum of the function

\[
f(x) = \sum_{i,j \in S} w_{i,j} \left( \| x_i - x_j \|^2 - \delta_{i,j}^2 \right)^2,
\]

where \( w_{i,j} \) are positive weights, and \( x_i, x_j \in \mathbb{R}^3 \) are the positions of the \( i \)-th and \( j \)-th
Table 4.1: MINPACK-2 Nonlinear Equations

<table>
<thead>
<tr>
<th>Name</th>
<th>Description of the Problem</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDC</td>
<td>Flow in a Driven Cavity</td>
</tr>
<tr>
<td>FIC</td>
<td>Flow in a Channel</td>
</tr>
<tr>
<td>IER</td>
<td>Incompressible Elastic Rod</td>
</tr>
<tr>
<td>SFD</td>
<td>Swirling Flow between Disks</td>
</tr>
<tr>
<td>SFI</td>
<td>Solid Fuel Ignition</td>
</tr>
</tbody>
</table>

atoms in the molecule. Here, we reformulate the MDG problem in terms of finding the global minimum of the nonlinear least squares problem

$$\min_{x \in \mathbb{R}^{n}} \| F(x) \|_2,$$

(4.2)

where $F(x) \in \mathbb{R}^{m}$, $n = 3 \times$ the number of atoms (3 coordinates per atom), $m =$ the number of $\delta_{ij}$ terms that are specified in order for the problem to be deterministic ($m \geq n$), and for each $\delta_{ij}$, the corresponding ($k$-th) component of $F(x)$ is defined by

$$F_k(x) = w_{ij} \left( \| x_i - x_j \|^2 - \delta_{ij}^2 \right)^2,$$

$k \equiv (i, j), \ k \in S.$

This transformation, while not altering the nature of the underlying computation or solution, poses the derivative problem as a sparse Jacobian computation.

We compared the ADIFOR/SparsLinC approach with an approach that uses a graph-coloring algorithm to compute the compressed Jacobian. In this section, we describe the compressed Jacobian approach. Then, we present the memory requirements and runtime results of both approaches for the MINPACK-2 and the MDG test problems.

4.1 The Compressed Jacobian Approach

The compressed Jacobian approach has been used as an effective strategy for computing sparse Jacobians in conjunction with both finite differencing [16] and automatic differentiation methods [2]. The prerequisite for applying the compressed Jacobian approach to a given sparse Jacobian computation is a priori knowledge of the sparsity pattern of the Jacobian.

The basic idea underlying this approach is that all the relevant (i.e., nonzero) information in the full Jacobian $f'(x)$ of size $m \times n$ can be represented in a compressed Jacobian $C(x)$ of size $m \times p$, where $C(x) = f'(x) S$, for some seed matrix $S$, and usually $p \ll n$. Essentially, $S$ maps each group of *structurally orthogonal* columns of $f'(x)$ (i.e., columns that do not have a nonzero in the same row position) to a column of $C(x)$. Because of the structural orthogonality property we can uniquely extract all entries of the original Jacobian matrix from the compressed Jacobian [2].

If the sparsity pattern of the Jacobian can be determined, graph-coloring techniques can be used to arrive at $S$ and $p$. These algorithms produce a partitioning of the columns of the Jacobian into $p$ structurally orthogonal groups by coloring the column-intersection graph associated with the Jacobian. In our experiments we employ the graph-coloring software
described in [15] to obtain $S$, and then compute $C(x)$ by initializing the ADIFOR-generated derivative code of $f(x)$ with the seed matrix set to $S$.

From the point of view of computational complexity, the consequence of the compressed Jacobian approach is clear. Specifically, the memory and runtime requirements of the Jacobian computation will approximately augment function requirements by a factor $p$ rather than $n$. For many sparsity patterns, in particular regular grid problems, $p$ is small and independent of $n$. Hence, the compressed Jacobian can be computed in a constant times the function computation time, regardless of $n$.

Naturally, the compressed Jacobian approach does introduce additional computational requirements, namely, (i) computing the sparsity pattern; (ii) performing the graph coloring; and (iii) subsequent to the computation of the compressed Jacobian, unraveling to arrive at the full Jacobian. However, often Jacobian computation is a subproblem within an optimization problem, and in an optimization algorithm we invariably need to compute a sequence $\{f'(x_k)\}$ of Jacobians for some sequence $\{x_k\}$ of iterates. In most cases we need to perform (i) and (ii) only once, since we can specify the closure of the sparsity patterns (i.e., a sparsity pattern that, for every iterate $x_k$, contains the sparsity pattern of $\{f'(x_k)\}$). If we are not able to specify the closure, the compressed Jacobian approach requires a call to the graph-coloring software at each iteration. Finally, the computational requirements of (iii) are relatively small.

The interesting difference between the MINPACK-2 and the MDG Jacobians is that for each MINPACK-2 problem the chromatic number is independent of $n$, whereas the chromatic number for the MDG Jacobians grows as a function of $n$. This difference is due to the “regularity” of the sparsity structures of the MINPACK-2 Jacobians as we change $n$, and the lack of the same in the case of the MDG Jacobians.

Figure 4.1 shows the sparsity patterns for some SFD and MDG Jacobians (here each dot represents the occurrence of a nonzero in the final Jacobian). In the case of each problem, we have chosen two small problem sizes in order to visually compare their respective sparsity structures. For the MDG problem of size $n = 81$, we have additionally plotted the upper left-hand corner of the Jacobian, since it is difficult to distinguish the location of the nonzeros in the full Jacobian.

In comparing the sparsity patterns of the two SFD problems, we note that an identically shaped nonzero block structure appears repeatedly along the main diagonal as $n$ is increased. The implication of this regularity of structure for the graph-coloring algorithm is that a fixed number of colors are sufficient for the representation of the compressed Jacobian, that is, $p$ remains constant for all $n$. Though the algorithms used in graph coloring are based on partitioning methods, one can visualize the results as each block in the full Jacobian sliding left to occupy the corresponding set of rows in the $p$ columns of the compressed Jacobian. Though distinct in shape, the sparsity patterns of the remaining MINPACK-2 Jacobians also have this regularly repeating feature, since they are all regular grid problems.

In contrast to the MINPACK-2 problems, for the two MDG Jacobians we note that as $n$ grows, the shape of the structures in the Jacobian sparsity pattern changes. This is clearly apparent when comparing the sparsity pattern of the MDG Jacobian for $n = 24$ with the corresponding slice of the MDG Jacobian for $n = 81$. The reason for this variation has to do with the problem specification itself. The formulation of the MDG problem is such that commensurate with increasing $n$, the number of atoms in the molecule is increased. In
Figure 4.1: SFD Jacobian sparsity patterns for $n = 42$, and $n = 98$, and MDG Jacobian sparsity patterns for $n = 24$, and $n = 81$. 
order for the problem to remain deterministic, corresponding to the increase in \(n, m\) (i.e.,
the number of distances between the atoms \(\delta_{i,j}\)'s) which must be specified) also increases.
The net effect is the structural change evident in the Jacobian sparsity, which in turn leads
to a different chromatic number being computed by the coloring algorithm for each \(n\).

In Sections 4.2 and 4.3 we will compare the performance of the ADIFOR/SparsLinC
approach with that of the ADIFOR/compressed Jacobian approach when the chromatic num-
ber, \(p\), is independent of \(n\) and when it is not, respectively. We refer to the
ADIFOR/SparsLinC approach as the **Sparse** approach in the sense that the full (sparse)
Jacobian is computed, and we will refer to the compressed Jacobian/ADIFOR approach as
the **NonSparse** approach, since the compressed Jacobian is a nonsparse remapping of the
sparse Jacobian.

For our purposes of comparing the results of these two functionally equivalent
approaches, we find it useful to think of the **NonSparse** approach as the **Static** approach to
exploiting sparsity in AD, since it statically allocates arrays of length \(p\) for the storage of
the directional gradient vectors, and since it requires a priori knowledge of the sparsity pat-
tern of the Jacobian. Conversely, the **Sparse** approach can be thought of as the **Dynamic**
approach to exploiting sparsity in AD, since it uses dynamic memory allocation for the
storage of the directional gradient vectors, and since the sparsity pattern is computed as a
by-product during the computation of the Jacobian.

When \(p\) is independent of \(n\) and is relatively small, as in the case of the MINPACK-2
problems, two issues influence whether the static or dynamic approach is preferable. The
first is the issue of convenience. The dynamic approach does not require knowledge of the
closure of the sparsity patterns; hence, from the viewpoint of code development, it is the
more convenient approach. Note that if it is not possible to compute closure, it may be the
only approach.

The second issue concerns the sparsity of the compressed Jacobian, which turns out to
be highly correlated with the relative performance of the **Sparse** computation as compared
with the **NonSparse** one. Consider the case where the compressed Jacobian is fully dense.
Then the static approach is ideal, since it contains no extraneous zero-sum computations
nor does it allocate extraneous storage. Though by design the dynamic approach also avoids
extraneous computations and storage, it is burdened by overheads associated with dynamic
memory allocation and the maintenance of dynamic data structures. On the other hand,
if the compressed Jacobian is sparse, the dynamic approach becomes preferable, since its

<table>
<thead>
<tr>
<th>Problem</th>
<th>(\text{nnz}(f'(x)))</th>
<th>(p)</th>
<th>density of (C(x))</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDC</td>
<td>13 (m)</td>
<td>19</td>
<td>68%</td>
</tr>
<tr>
<td>FIC</td>
<td>8 (m)</td>
<td>9</td>
<td>89%</td>
</tr>
<tr>
<td>IER</td>
<td>11 (m)</td>
<td>17</td>
<td>65%</td>
</tr>
<tr>
<td>SFD</td>
<td>12 (m)</td>
<td>14</td>
<td>86%</td>
</tr>
<tr>
<td>SFI</td>
<td>5 (m)</td>
<td>7</td>
<td>71%</td>
</tr>
</tbody>
</table>
Table 4.3: Structural Information about the Molecular Distance Geometry Jacobians

<table>
<thead>
<tr>
<th>$n$</th>
<th>$m$</th>
<th>$p$</th>
<th>density of $C(x)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>375</td>
<td>2801</td>
<td>78</td>
<td>8%</td>
</tr>
<tr>
<td>525</td>
<td>4051</td>
<td>78</td>
<td>8%</td>
</tr>
<tr>
<td>648</td>
<td>7111</td>
<td>111</td>
<td>5%</td>
</tr>
<tr>
<td>1029</td>
<td>15583</td>
<td>150</td>
<td>4%</td>
</tr>
<tr>
<td>1536</td>
<td>30689</td>
<td>195</td>
<td>3%</td>
</tr>
<tr>
<td>2187</td>
<td>55729</td>
<td>246</td>
<td>2%</td>
</tr>
<tr>
<td>3000</td>
<td>94951</td>
<td>303</td>
<td>2%</td>
</tr>
<tr>
<td>3993</td>
<td>153671</td>
<td>366</td>
<td>2%</td>
</tr>
<tr>
<td>5184</td>
<td>238393</td>
<td>435</td>
<td>1%</td>
</tr>
<tr>
<td>6591</td>
<td>356929</td>
<td>510</td>
<td>1%</td>
</tr>
<tr>
<td>8232</td>
<td>518519</td>
<td>591</td>
<td>1%</td>
</tr>
<tr>
<td>10125</td>
<td>733951</td>
<td>678</td>
<td>1%</td>
</tr>
</tbody>
</table>

overheads are offset by the many extraneous computations and unused memory in the static approach.

Tables 4.2 and 4.3 contain information relating to the structure of the MINPACK-2 and the MDG Jacobian problems, respectively. In the case of the MINPACK-2 problems, $m = n$ and $p$ remains constant. Hence, in Table 4.2 we have shown for each problem the number of nonzeros in the Jacobian, $\text{nnz}(f'(x))$, as well as the chromatic number, $p$, computed by the graph-coloring software. As previously discussed, $p$ is independent of $n$ for every MINPACK-2 problem because the sparsity structure of the corresponding Jacobian is also independent of $n$. On the other hand, for the MDG problem the Jacobian structure is distinct for each problem size $n$. Hence, in Table 4.3 we have shown the corresponding $n$, $m$, and $p$ values. For all $n$, the number of nonzeros in the MDG Jacobian is given by $\text{nnz}(f'(x)) = 6m$.

The last column in both Tables 4.2 and 4.3 show the density of $C(x)$, computed as $\text{nnz}(f'(x))/mp$. In Sections 4.2.1 and 4.2.2, we will show the correlation of these densities with the memory requirements and runtime performances of the two approaches.

4.2 Computing Sparse Jacobians When $p$ Is Independent of $n$

4.2.1 Memory Requirements

Table 4.4 presents the total memory requirements of the two approaches for computing the Jacobian of the MINPACK-2 problems for the case of $n = 160,000$. In addition to the notation introduced in Table 3.2, we use $M\{J\}$ to denote the memory requirements of the \textsc{NonSparse} approach, which includes the memory needed for the graph-coloring computation, and $M\{J_{\text{Sparse}}\}$ to denote the memory requirements of the \textsc{Sparse} approach. For each problem, the ratio $M\{J\}/M\{F\}$ remains constant for all $n$. Hence, Table 4.4 is a sufficient
Table 4.4: Memory Requirements for MINPACK-2 Jacobians (in Mbytes; \( n = 160,000 \))

<table>
<thead>
<tr>
<th>Problem</th>
<th>( M{F} )</th>
<th>( M{J} )</th>
<th>( M{J}/M{F} )</th>
<th>( M{J_{\text{sparse}}} )</th>
<th>( M{J_{\text{sparse}}}/M{F} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDC</td>
<td>2.57</td>
<td>74.90</td>
<td>29.1</td>
<td>54.35</td>
<td>21.1</td>
</tr>
<tr>
<td>FIC</td>
<td>2.58</td>
<td>43.05</td>
<td>16.7</td>
<td>32.17</td>
<td>12.5</td>
</tr>
<tr>
<td>IER</td>
<td>2.57</td>
<td>67.22</td>
<td>26.1</td>
<td>50.65</td>
<td>19.7</td>
</tr>
<tr>
<td>SDF</td>
<td>2.58</td>
<td>60.84</td>
<td>23.6</td>
<td>41.77</td>
<td>16.2</td>
</tr>
<tr>
<td>SFI</td>
<td>2.57</td>
<td>33.94</td>
<td>13.2</td>
<td>28.35</td>
<td>11.1</td>
</tr>
</tbody>
</table>

summary of all the memory results.

As mentioned in Section 3.1, the \texttt{NONSparse} mode of ADIFOR tends to augment memory requirements of the function computation linearly. In the case of the compressed Jacobian computations the augmentation factor is the chromatic number, \( p \). However, since the graph-coloring algorithm also introduces additional memory requirements proportional to \( \text{nnz}(f'(x)) \), the \( M\{J\}/M\{F\} \) ratios in Table 4.4 are larger than the corresponding \( p \) values in Table 4.2, and we have

\[
M\{J\} \leq \kappa \ p \ M\{F\}
\]

with \( 1.5 \leq \kappa \leq 1.9 \). In general, \( \kappa \) is largest for the problems with denser compressed Jacobians. For example, for the FIC problem, the compressed Jacobian is 89% dense and \( \kappa = 1.9 \), whereas for the IER problem, the compressed Jacobian is 65% dense and \( \kappa = 1.5 \).

Given that the memory requirements of the \texttt{Sparse} approach are based on the need to represent the nonzero information in the derivative computation, and that this information is close in volume to the size of the compressed Jacobian (for the MINPACK-2 problems, within 65% – 89%), we expect the values of \( M\{J_{\text{sparse}}\}/M\{F\} \) to be somewhat correlated with the corresponding \( p \) for each problem. A comparison of the last column of Table 4.4 and the corresponding values of \( p \) shows this to be the case, where we have

\[
M\{J_{\text{sparse}}\} \leq \sigma \ p \ M\{F\}
\]

with \( 1.1 \leq \sigma \leq 1.6 \).

We note that the memory requirements of the \texttt{NONSparse} approach are 19% – 46% greater than the memory requirements of the \texttt{Sparse} approach. This is somewhat surprising, since in all cases the compressed Jacobians are fairly dense, and one might suspect that a static approach would be more parsimonious in terms of memory usage. The disparity is mainly attributable to the need for the additional memory required by the graph-coloring algorithm in the \texttt{NONSparse} approach, but also to the terseness of SparsLinC’s memory allocation scheme, and the fact that SparsLinC exploits intermediate sparsity in derivative computations.

Inequality (4.4) is a useful metric in terms of comparing the memory requirements of the \texttt{Sparse} with the \texttt{NONSparse} approach. However, SparsLinC has no knowledge of \( p \), which can be thought of as a global measure of sparsity. SparsLinC’s representation of directional gradient vectors is based on very localized information, namely, the number of nonzeroses
in each gradient vector. The most striking example demonstrating the difference between these global and local approaches toward storage would be the computation of a sparse Jacobian with one fully dense row and a dense diagonal. In this case \( p = n \); hence, the compressed Jacobian approach would fail entirely. On the other hand, SparsLinC would need to maintain only one vector of length \( n \), and \( n - 1 \) vectors of length 1, resulting in much less memory-intensive (as well as faster) code.

Hence, it would be more descriptive of the localized vector representations in SparsLinC to express the augmentation factor of the memory requirements of the \textit{SPARSE} approach with respect to those of the function, as proportional to the average number of nonzeros per row of the Jacobian,

\[
M \{J_{\text{Sparse}} \} \leq \sigma' \frac{\text{nnz}(f'(x))}{m} M \{F\}, \tag{4.5}
\]

where for the MINPACK-2 problems, \( 1.4 \leq \sigma' \leq 2.2 \). As we shall see in Section 4.3.1, inequality (4.5) is a more reliable generalization of the expected memory requirements of the \textit{SPARSE} approach.

### 4.2.2 Runtime

Figures 4.2 and 4.3 summarize the runtime results of the Jacobian problems on the SPARC IPX and RS6000, respectively. As in the case of the gradient experiments, the problem sizes range from \( n = 2,500 \) to \( n = 160,000 \). Each panel in these figures contains a plot of the \textit{SPARSE} Jacobian-to-function runtime ratios. Additionally, the results from the \textit{NONSPARSE} computation are shown with two plots in each panel, one showing the coloring-to-function runtime ratios, and the other showing the compressed Jacobian-to-function run-
time ratios. As we explained in Section 4.1, for most optimization algorithms, the coloring computation is performed only once, whereas iterates of the compressed Jacobian are computed repeatedly, hence the separation of the two computations in our plots. In addition to plotting these runtime ratios, we report their values for the case of $n = 160,000$ in Tables 4.5 and 4.6.

A main feature of these results is that for both approaches the runtime of the Jacobian computation is independent of $n$, as shown by the constant ratio plots. In the case of the NONSPARSE approach, this behavior is expected, since we know the Jacobian runtime will be approximately equal to the linear augmentation of the function runtime by a factor $p$, and as we have stated, $p$ is independent of $n$. Based on the NONSPARSE ratios in Tables 4.5 and 4.6, we can state that

$$T\{J_{Compressed}\} \leq \lambda \ p \ T\{F\},$$  \hspace{1cm} (4.6)$$

where $T\{\cdot\}$ denotes the runtime of each computation and where $0.6 \leq \lambda \leq 2.4$ on the SPARC IPX and $0.5 \leq \lambda \leq 1.1$ on the RS6000. The variation in $\lambda$ values for the different problems is due to the peculiarities of each code, in particular the extent to which ADIFOR can minimize the computations performed in the derivative code through activity analysis. The variation in $\lambda$ values between the two platforms is due to architectural differences reviewed in Section 3.2, though it is surprising to see that the RS6000 ratios are slightly greater for the FIC and SFD problems. We note that in all cases $\lambda$ is a small constant, which matches our expectation.

For the SPARSE Jacobian computation, the constant behavior with respect to function runtime is due to the sparsity present in all the intermediate directional gradient vectors. Unlike the SPARSE gradient computation, where at least one directional gradient vector (i.e., the final gradient vector) grows to be fully dense, in the Jacobian computation all intermediate vectors will be as sparse or more sparse than the least sparse row of the Jacobian. From the results of the coloring algorithm, we know that this row can have no more than $p$ nonzeros (otherwise the compressed Jacobian would not “fit” into $p$ columns). In terms of an augmentation over function time, we can state that

$$T\{J_{Sparse}\} \leq \omega \ p \ T\{F\},$$  \hspace{1cm} (4.7)$$

where $1.4 \leq \omega \leq 5.9$ on the SPARC IPX and $5.0 \leq \omega \leq 17.9$ on the RS6000. Similar to the case of the memory requirements of SPARSE approach, we can express the augmentation of the SPARSE runtime with respect to the function runtime as a factor of the average number of nonzeros per row of the Jacobian:

$$T\{J_{Sparse}\} \leq \omega' \ \frac{\text{nnz}(f'(x))}{m} \ T\{F\},$$  \hspace{1cm} (4.8)$$

where $2.1 \leq \omega' \leq 8.3$ on the SPARC IPX and $7.8 \leq \omega' \leq 20.1$ on the RS6000. We will revisit inequality 4.8 in Section 4.3.2.

The last column in Tables 4.5 and 4.6 show the ratios of SPARSE runtime to NONSPARSE runtime for each problem on each platform. These ratios—all of which are above 1—represent the penalty paid for the convenience of the dynamic approach. As expected, there is a strong correlation between these ratios and the sparsity of the corresponding compressed Jacobian. For example, on both platforms the highest ratios occur for the FIC problem, which has the
Figure 4.2: Runtime ratios of MINPACK-2 Jacobian problems on the SPARC IPX. For each problem, NONSPARSE computation-to-function, SPARSE computation-to-function and coloring-to-function runtime ratios are shown for problem sizes $n = 2,500$ to $n = 160,000.$
Figure 4.3: Runtime ratios of MINPACK-2 Jacobian problems on the RS6000. For each problem, NONSPARSE computation-to-function, SPARSE computation-to-function and coloring-to-function runtime ratios are shown for problem sizes $n = 2,500$ to $n = 160,000$. 

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least sparse compressed Jacobian. Conversely, the ratios for the IER problem, which has the most sparse compressed Jacobian, are among the smallest. In general, the sparser the compressed Jacobian, the better the runtime performance of the \textsc{Sparse} approach relative to that of the \textsc{NonSparse} approach.

The coloring algorithm also demonstrates constant runtime behavior with respect to function runtime, with the constant factor usually being somewhere between the corresponding factors of the \textsc{NonSparse} and \textsc{Sparse} computations. The closure of the sparsity patterns of each MINPACK-2 Jacobian problem is known; hence, the impact of the one-time graph-coloring computation on the overall optimization computation would be minimal. Had the closure been unknown, the coloring computation would need to be repeated at each iteration, with the possible consequence of the \textsc{Sparse} approach having a faster runtime. For example, in Table 4.5 note that for the IER and SFI problems on the SPARC IPX, the sum of the runtime ratios of the coloring and \textsc{NonSparse} compressed Jacobian computations exceeds the corresponding ratio from the \textsc{Sparse} computations.

4.3 Computing Sparse Jacobians When $p$ Grows with $n$

4.3.1 Memory Requirements

Table 4.7 presents the total memory requirements of the two approaches for computing the Jacobians of the MDG problem, for various problem sizes (we use the same notation introduced in Table 4.4). In contrast to Table 4.4 where the memory requirements were shown only for the largest-sized problem ($n = 160,000$), Table 4.7 shows the memory requirements for all problem sizes, ranging from $n = 375$ to $n = 10,125$. This is done for two reasons. First, since now $p$ grows with $n$, so too does the ratio $M\{J\}/M\{F\}$, in contrast to the constant $M\{J\}/M\{F\}$ in the case of the MINPACK-2 Jacobians where $p$ was constant.

Second, largely because of the growing value of $p$, the executable for the largest-sized problems became too large to fit in the virtual memory available on either platform. Hence, we resorted to the stripmining approach (see Section 3.1) for the \textsc{NonSparse} computation, which dramatically reduced memory requirements, while slightly increasing the runtime. We used strip size $p_s = 16$ for the problems of dimension $n = 3,000$ to $n = 8,232$. However, this choice of $p_s$ for $n = 10,125$ led to performance degradation as a result of paging. Hence, we used $p_s = 8$ for the largest problem size (we discuss our choice of strip sizes further in Section 4.3.2). In Table 4.7, the memory requirements of the stripmined \textsc{NonSparse} computations are marked with * for $p_s = 16$ and ** for $p_s = 8$.

The results in Table 4.7 show that for the non-stripmined \textsc{NonSparse} computation of the MDG Jacobians, the inequality (4.3) holds as it did in the case of the MINPACK-2 Jacobians, with the difference that here $p$ is not constant. Nonetheless, we can compute $\kappa$ in (4.3) for the MDG problem and ascertain that, as expected, its value varies only slightly ($0.35 \leq \kappa \leq 0.41$) for the problem sizes $n = 375$ to $n = 2,187$. We note that here $\kappa$ is a factor 3–5 smaller than in the MINPACK-2 cases. This is mainly due to the presence of several large work arrays of dimension $m$ in the MDG function code, which are unrelated to any derivative computation; hence, ADIFOR does not generate corresponding augmented derivative work arrays.

For the stripmined \textsc{NonSparse} computation of the MDG Jacobians, we need a new in-
Table 4.7: Memory Requirements for the Distance Geometry Jacobians (in Mbytes)

<table>
<thead>
<tr>
<th></th>
<th>$M{F}$</th>
<th>$M{J}$</th>
<th>$M{J}/M{F}$</th>
<th>$M{J_{\text{Sparse}}}$</th>
<th>$M{J_{\text{Sparse}}}/M{F}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>No Stripmining</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>375</td>
<td>0.07</td>
<td>2.22</td>
<td>31.7</td>
<td>0.70</td>
<td>10.0</td>
</tr>
<tr>
<td>525</td>
<td>0.10</td>
<td>3.19</td>
<td>31.9</td>
<td>0.98</td>
<td>9.8</td>
</tr>
<tr>
<td>648</td>
<td>0.18</td>
<td>7.46</td>
<td>41.4</td>
<td>1.65</td>
<td>9.2</td>
</tr>
<tr>
<td>1029</td>
<td>0.38</td>
<td>21.16</td>
<td>55.7</td>
<td>3.49</td>
<td>9.2</td>
</tr>
<tr>
<td>1536</td>
<td>0.75</td>
<td>52.66</td>
<td>70.2</td>
<td>6.77</td>
<td>9.0</td>
</tr>
<tr>
<td>2187</td>
<td>1.36</td>
<td>118.29</td>
<td>87.0</td>
<td>12.22</td>
<td>9.0</td>
</tr>
<tr>
<td>Stripmined ($p_s = 16$)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3000</td>
<td>2.30</td>
<td>19.88*</td>
<td>8.6</td>
<td>20.71</td>
<td>9.0</td>
</tr>
<tr>
<td>3993</td>
<td>3.72</td>
<td>32.00*</td>
<td>8.6</td>
<td>33.47</td>
<td>9.0</td>
</tr>
<tr>
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<td>5.76</td>
<td>49.47*</td>
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<td>51.85</td>
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</tr>
<tr>
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<td>8.62</td>
<td>73.88*</td>
<td>8.6</td>
<td>77.48</td>
<td>9.0</td>
</tr>
<tr>
<td>8232</td>
<td>12.51</td>
<td>107.11*</td>
<td>8.6</td>
<td>112.16</td>
<td>9.0</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10125</td>
<td>17.70</td>
<td>103.76**</td>
<td>5.9</td>
<td>158.47</td>
<td>9.0</td>
</tr>
</tbody>
</table>

equality, since now the linear augmentation is dependent on the strip size and not the chromatic number:

$$M\{J_{\text{Stripmined}}\} \leq \kappa' p_s M\{F\}$$  \hspace{1cm} (4.9)

with $0.54 \leq \kappa' \leq 0.74$. Note that the lower range of $\kappa'$ values corresponds to $p_s = 16$, implying that the memory requirement penalty of the stripmining method decreases with respect to the function memory requirements as the strip size increases. By virtue of the same reasoning, we can explain why $\kappa < \kappa'$, since essentially the non-stripmined computation can be thought of as a stripmined computation with one strip of size $p_s = p$.

We also observe from Table 4.7 that the $M\{J_{\text{Sparse}}\}/M\{F\}$ ratio for the SPARSE approach varies only slightly for different problem sizes. As noted in Section 4.1, $\text{nnz}(f'(x)) = 6m$ for the MDG problem; hence the average number of nonzeros per row of the Jacobian is always 6, for all $n$. Consequently, inequality (4.5) holds, and the corresponding range of $\sigma'$ values for the MDG problem is given by, $1.5 \leq \sigma' \leq 1.7$.

In summary, the memory requirements of the NONSPARSE approach are greater than those of the SPARSE approach by a large factor, ranging from 3.2 to 9.7 for the non-stripmined problems. This result is due to the compressed Jacobians of the MDG problems being very sparse, and more and more so as $n$ grows. Stripmining can be a very effective method of reducing the memory requirements of the NONSPARSE approach—and, as we shall see, at the cost of a modest runtime penalty. Finally, we note that though we did not need to stripmine the SPARSE computation, this method is equally possible and straightforward.
4.3.2 Runtime

Figures 4.4 and 4.5 show the Jacobian-to-function runtime ratios for the MDG problems on the SPARC IPX and RS6000, respectively, with sizes \( n = 375 \) to \( n = 10125 \). Each figure includes three plots showing the ratios of the NONSPARSE (dotted), SPARSE (solid), and graph-coloring (dashed) runtimes to the function runtimes. We also present the detailed Jacobian-to-function and coloring-to-function runtime ratios on the SPARC IPX and the RS6000 in Tables 4.8 and 4.9, respectively. Note that, as before, the runtime ratios of a stripmined NONSPARSE computation are marked with * for \( p_s = 16 \) and ** for \( p_s = 8 \).

The main conclusion that can be drawn from the plots in Figures 4.4 and 4.5 is that the the Jacobian-to-function runtime ratios appear to grow linearly with \( n \) for the NONSPARSE approach, whereas these ratios are independent of \( n \) for the SPARSE approach. Hence, the SPARSE approach is clearly the method of choice for problems where the chromatic number grows as a function of the problem size and yet the sparsity of the corresponding Jacobians remains constant (i.e., \( \text{nnz}(f'(x)) \) proportional to \( m \)). The efficient runtime performance of the SPARSE approach relative to the NONSPARSE approach is attributed to the fact that the already very sparse compressed Jacobians get even sparser as we increase the value of \( n \) (see the last column of Table 4.3).

In contrasting the numbers in Tables 4.8 and 4.9, we note that the previously discussed architectural effects (see Section 4.2.2) once again favor the SPARSE approach on the SPARC IPX and the NONSPARSE approach on the RS6000. In particular, the crossover point at which the SPARSE computation runs faster than the NONSPARSE computation occurs at \( n = 525 \) on the SPARC IPX and at \( n = 2,187 \) on the RS6000. Also, for the largest problem, the speedup of the SPARSE versus the NONSPARSE computation is 10 on the SPARC IPX and 3.5 on the RS6000.

We observe that for both the straightforward and the stripmined NONSPARSE computations, the inequality (4.6) holds; however, in contrast to the MINPACK-2 problems, \( p \) is not a constant. For the MDG problem, we have \( 1.4 \leq \lambda \leq 1.8 \) on the SPARC IPX and \( 1.3 \leq \lambda \leq 1.5 \) on the RS6000. Likewise, for the SPARSE computation inequality (4.8) holds with \( 18 \leq \omega^' \leq 22 \) on the SPARC IPX and \( 43 \leq \omega^' \leq 49 \) on the RS6000.

The strip sizes were chosen such that they would result in reasonable memory values. As was previously explained, stripmining is needed in order for the NONSPARSE computation to fit in memory, and comes at the cost of an additional (extraneous) function evaluation per strip. In terms of the effect of stripmining on the runtime of the NONSPARSE computation, we note that the NONSPARSE plots in Figures 4.4 and 4.5 appear to have three distinct slopes corresponding to the three computational flavors: (i) straightforward or non-stripmined, (ii) stripmined with \( p_s = 16 \), and (iii) stripmined with \( p_s = 8 \). As expected, the slope of the line segment corresponding to (iii) is greater than that of (ii), because halving the strip size, doubles the overhead function computations. By virtue of the same overhead effects, one might expect the slope of the line segment corresponding to (ii) to be greater than that of (i). However, the contrary is true. We suspect that this fact is due to the degradation in runtime performance of the larger non-stripmined problems brought about by excessive paging.

Finally, we note in Figures 4.4 and 4.5 that the coloring-to-function ratio results—in like manner to their the MINPACK-2 counterparts—are constant, with the constant factor
Table 4.8: Runtime Ratios of the MDG Problem on the SPARC IPX

<table>
<thead>
<tr>
<th>n</th>
<th>Coloring</th>
<th>NONSPARSE</th>
<th>SPARSE</th>
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<tbody>
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<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>No Stripmining</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>375</td>
<td>31.6</td>
<td>113</td>
<td>124</td>
<td>1.1</td>
</tr>
<tr>
<td>525</td>
<td>31.3</td>
<td>121</td>
<td>128</td>
<td>1.1</td>
</tr>
<tr>
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<td>187</td>
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<td>0.7</td>
</tr>
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<td>126</td>
<td>0.6</td>
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<tr>
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<td>332</td>
<td>127</td>
<td>0.4</td>
</tr>
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<td>420</td>
<td>123</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>3000</td>
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<td>44.7</td>
<td>557*</td>
<td>117</td>
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<td>645*</td>
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<td>108</td>
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<td>49.6</td>
<td>857*</td>
<td>122</td>
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<td></td>
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</tr>
<tr>
<td>10125</td>
<td>54.5</td>
<td>1233**</td>
<td>123</td>
<td>0.1</td>
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</table>

Table 4.9: Runtime Ratios of the MDG Problem on the RS6000

<table>
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<th>SPARSE</th>
<th>SPARSE/NONSPARSE</th>
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<tr>
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<td>111</td>
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<td>2.6</td>
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<td>213</td>
<td>285</td>
<td>1.3</td>
</tr>
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<td>288</td>
<td>1.0</td>
</tr>
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<td>378</td>
<td>293</td>
<td>0.8</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
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<td>84.7</td>
<td>396*</td>
<td>289</td>
<td>0.7</td>
</tr>
<tr>
<td>3993</td>
<td>92.3</td>
<td>472*</td>
<td>289</td>
<td>0.6</td>
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<tr>
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<td>579*</td>
<td>285</td>
<td>0.5</td>
</tr>
<tr>
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<td>662*</td>
<td>291</td>
<td>0.4</td>
</tr>
<tr>
<td>8232</td>
<td>122.6</td>
<td>777*</td>
<td>294</td>
<td>0.4</td>
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<tr>
<td>Stripmined ($p_b = 8$)</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10125</td>
<td>123.5</td>
<td>1022**</td>
<td>286</td>
<td>0.3</td>
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Figure 4.4: Jacobian-to-function runtime ratios on the SPARC IPX. **NONSPARSE** (dotted), **SPARSE** (solid), and graph-coloring (dashed).

Figure 4.5: Jacobian-to-function runtime ratios on the RS6000. **NONSPARSE** (dotted), **SPARSE** (solid), and graph-coloring (dashed).
being smaller than the corresponding factor for the SPARSE computation.

5 Conclusions

We have shown the effectiveness of the SparsLinC library in conjunction with the ADIFOR tool for exploiting sparsity in automatic differentiation. The fact that SparsLinC does not require a priori knowledge about the structure of sparsity in a particular problem, coupled with the fact that sparsity is exploited transparently, without the need for rewriting code, makes the ADIFOR/SparsLinC combination a valuable and convenient tool set. It can be used either for quick prototyping of the sparsity characteristics of a problem or as a means of getting improved efficiency in sparse derivative computations at very little cost in terms of program development time.

The ADIFOR/SparsLinC approach reduces the runtime cost of computing gradients of partially separable functions by as many as three orders of magnitude over the NONSPARSE ADIFOR approach, while imposing modest memory requirements. SparsLinC does not require a priori knowledge about where and how the partial separability is coded; if it exists, SparsLinC will exploit the consequent sparsity.

The relative efficiency of SparsLinC’s runtime results in computing sparse Jacobians as compared with the performance of the compressed Jacobians method is dependent on whether or not the chromatic number, \( p \), derived from graph-coloring of the Jacobian is dependent on the problem size, \( n \). (This comparison presupposes the feasibility of a priori computation of the closure sparsity pattern of the Jacobian needed for the compressed Jacobian approach.) If \( p \) is independent of \( n \), both the ADIFOR/SparsLinC approach and the ADIFOR/compressed Jacobians approach will run at a constant factor times the function runtime, with the compressed Jacobian approach typically being the faster of the two.

If \( p \) grows with \( n \), the compressed Jacobians approach will have a linear dependence on \( n \), which implies much higher runtimes and memory requirements, making ADIFOR/SparsLinC the clear method of choice in such cases.

Acknowledgments

We thank Jorge Moré for his assistance with the MINPACK-2 problem set, Paul Hovland and Aaron Ross for their contributions to building early prototypes of SparsLinC, Andrew Mauer and Tim Knauff for their assistance in debugging SparsLinC, Vitaly Shmatikov for his assistance with the sparsity characteristics plots, and Zhijun Wu for his assistance with the MDG problem.

References


