Some highlights on Source-to-Source Adjoint AD

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Abstract

Algorithmic Differentiation (AD) provides the analytic derivatives of functions given as programs. Adjoint AD, which computes gradients, is similar to Back Propagation for Machine Learning. AD researchers study strategies to overcome the difficulties of adjoint AD, to get closer to its theoretical efficiency. To promote fruitful exchanges between Back Propagation and adjoint AD, we present three of these strategies and give our view of their interest and current status.

1 Adjoint AD models

Algorithmic Differentiation (AD) \footnote{9,14} is one way to obtain the analytical derivatives of a vector function \( F : X \in \mathbb{R}^n \mapsto Y \in \mathbb{R}^m \), when this function is provided as an algorithm. There are other ways, such as differentiating the mathematical equations that the algorithm implements, if they exist. For AD the algorithm is the function: its run-time sequence of instructions is a composition of elementary functions that define the function. By applying the chain rule of calculus, AD builds a new sequence of instructions and eventually a new algorithm that computes the desired derivatives.

Using AD, one can obtain first-order derivatives as well as higher-order or Taylor developments. Being based on the chain rule, these derivatives are analytic, meaning exact up to machine precision and free of the approximation errors that would result from other approaches, such as divided differences \( \dot{Y} \simeq \frac{(F(X + h \ast \dot{X}) - F(X))/h} \). Yet their algorithmic cost remains similar to divided differences, and even much cheaper in the case of gradients.

Focusing on first-order derivatives from now on, associativity of matrix multiplication implies that the chain rule can be applied in various orders. Among those are two extremes:

- Derivative matrices can be multiplied forward, as the original program goes. This is called the tangent mode of AD. One provides one (or several) initial direction vector \( \dot{X} \) in the input space of the program, and the tangent differentiated code progressively computes the derivatives of every intermediate variable with respect to the inputs, along direction \( \dot{X} \). Upon completion, the tangent code has computed into \( \dot{Y} \) the directional first-order derivative of the result \( Y \) with respect to \( X \), at the given point \( X \) and along the given direction \( \dot{X} \). Notice that the divided differences approach is by essence tightly bound to tangent AD, as it also goes from one given \( \dot{X} \) to \( \dot{Y} \). Complexity-wise and like divided differences, the run-time of the tangent code is only a small multiple (1 to 3) of the original code. Computing all first-order derivatives of this vector function \( F : \mathbb{R}^n \rightarrow \mathbb{R}^m \) requires \( n \) runs of the tangent code along the \( n \) vectors of the Cartesian basis of the input space.

- Derivative matrices can also be multiplied backwards. This is called the adjoint (or reverse) mode of AD. Given one (or several) weighting \( \bar{Y} \) in the output space, the adjoint differentiated code progressively computes the derivatives of the scalar \( (\bar{Y}|Y) \) with respect to every intermediate variable, in the reverse of the original program execution order. Upon completion, the adjoint code has computed into \( \bar{X} \) the gradient with respect to \( X \) of the function \( (\bar{Y}|F(X)) \). Divided differences are unable to emulate adjoint mode. Complexity-wise,
the run-time of the adjoint code is also a small multiple (slightly higher than for tangent
code, 5 to 10 in practice) of the original code. Computing all first-order derivatives of
\( F : \mathbb{R}^n \to \mathbb{R}^m \) requires \( m \) runs of the adjoint code along the \( m \) vectors of the Cartesian
basis of the output space. Still, computing derivatives in reverse order poses additional
technical problems that we will discuss later.

It turns out that many applications actually require a gradient \( (m = 1) \) or more generally \( m \ll n \). In
that case adjoint mode is remarkably more efficient than tangent mode or than divided differences
approximation. Primarily in view of applications such as gradient-based optimization or parameter
estimation, AD researchers have spent considerable effort in building efficient adjoint codes. In
Machine Learning, gradients are crucial for training neural networks, and “Back Propagation”,
a close cousin of adjoint AD, has been recognized as a key technology. To investigate fruitful
exchanges between Back Propagation and adjoint AD, we’d like to give here a few highlights on
adjoint AD technology, and more precisely about AD tools that provide the adjoint mode through
Source-to-Source Transformation (ST-AD).

There are many possible architectures to implement the backward computation of adjoint AD [5]. We
call them adjoint AD models. An AD tool will implement (at least) one of these models. We like to
illustrate the model implemented by our tool Tapenade [11] and also by OpenAD [16] by figure 1: since the deriva-
tive propagation goes backwards, the adjoint code must first run the original code
to memorize the control flow that will be
used backwards. We show this “forward sweep” as a thick arrow to the right. Only then can derivative
computation start, retrieving control flow information on the way. We show this “backward sweep”
as a thick arrow to the left. Think of an original instruction and its derivative counterpart as aligned
vertically. The derivative of an instruction may in general need the values that are used by the original
instruction. If these needed values are overwritten by the sequel of the forward sweep, we must devise
a mechanism, symbolized by the few red arrows, to make them available again during the backward
sweep. In our model, this is basically done by storage. Other models exist that use recomputation [7],
that we will not discuss here for the sake of brevity. In figure 2 arrow thickness also symbolizes
storage and retrieval of these intermediate values. This is the bottleneck of adjoint AD, as this storage
grows linearly with the program run time. In the next sections we will discuss techniques that cope
with this issue: static Data-Flow analysis can reduce significantly the amount of storage needed, but
still growing linearly with run time. Checkpointing is a memory/recomputation tradeoff that can
make the storage grow only logarithmically with run time. In special cases such as Fixed-Point loops,
specialized techniques can also reduce this storage drastically.

Before that, we close this section with a contrast with AD models based on Operator-Overloading
(OO-AD). In OO-AD tools such as AdolC [17], dco [13], CoDiPack [1]…, the forward sweep
additionally records every executed instruction into a “tape”, in a format resembling three-address
code. The backward sweep doesn’t exist as a source any more. Rather, the tape is interpreted
backwards by a special program that computes the derivatives. The advantage is that the tape format
is easier to generate than it is to build the source of a reverse sweep, so that OO-AD can address more
sophisticated application languages such as C++. It takes extra run-time to write this tape and the
tape itself cannot be exposed to the compiler for optimization, however OO-AD tools are making
impressive progress to cope with that. The main drawback is that the tape is much larger than the
data recorded by ST-AD. Consequently the techniques in the next sections can be of value also for
OO-AD.

In the small world of AD, ST-AD and OO-AD are too often seen as opposed. This is bound to change.
To begin with an abstract remark, ST-AD can be viewed as OO-AD followed by Partial Evaluation.
Partial Evaluation applied to the backward tape interpreter, with parts of the tape considered as known
because they come from fixed parts of the source (think of Basic Blocks), ideally results in a code
that takes as input only the control flow decisions and the intermediate values, i.e. a backward sweep
of ST-AD. More pragmatically, sophisticated language constructs do not lend themselves to a purely
static transformation, and ST-AD must store more and more dynamic information such as pointer
base addresses, or corresponding ends of point-to-point message communication. Finally, ST-AD
and OO-AD can collaborate: ST-AD can preprocess a source in view of OO-AD, and OO-AD can
call ST-AD generated code for critical, computation-intensive sections.
2 Data-Flow Analysis

ST-AD tools use several data-flow analyses to produce simplified derivative code, that are indeed classical forward propagation and backward slicing customized to the structure of these codes. The most important are, in order of appearance during AD:

1. **Activity**: detects the variables that deserve a derivative at a given point. An active variable must be *Varied*, i.e. its derivative is not structurally zero. This part of the analysis is therefore a forward propagation of zero derivatives. An active variable must at the same time be *Useful*, i.e. its derivative is used in the sequel. This part of the analysis is therefore a backward slicing of unused derivatives. As Varied analysis is a forward propagation, some OO-AD tools perform it too, in that case as a run-time Data-Flow Analysis.

2. **Diff-Liveness**: detects primal variables that are not needed for derivative computation, although they may be used for the original, primal computation. This analysis is therefore a backward slicing of unused primals in the forward sweep.

3. **TBR**: detects the primal variables that are never used in any derivative (e.g. they only appear in linear expressions [10]). Their value need not be stored by the forward sweep. This analysis can be seen as a backward slicing of store/restore of primals, in both sweeps.

These analyses face the same issues as any Data-Flow Analysis. Being undecidable, they must often use conservative over-approximations. Arrays are always considered atomic, which is another source of over-approximation. In particular, the deprecated coding style known as the “big work array” considerably reduces the benefit of these analyses. On the other hand, to reduce approximation, ST-AD tools make these analyses interprocedural, context-sensitive, and flow-sensitive. The price to pay is that there is no “separate AD” in the fashion of separate compilation.

In theory, one could spare these analyses, and resort instead to aggressive optimisation by the compiler [15]. However, these analyses exploit knowledge of the structure of the adjoint code, that the compiler is not aware of. In particular, these analyses exploit the correspondence between a forward, primal instruction and its derivative in the backward sweep. These two locations can be arbitrarily far apart in the adjoint code, making peep hole or window-based optimizations fail.

Measurements with Tapenade on a dozen large test cases show that these analyses can improve significantly runtime and storage size of adjoint codes, in a proportion that varies largely. Run-time is reduced by 20 to 50%, with one outlier at 70%. Storage is reduced by 40 to 70%, with outliers at 0% and 92%. These improvements remain proportional: they are of limited help against the linear growth of storage of intermediate values. To answer that, one must resort to checkpointing.

3 Checkpointing

Checkpointing is a memory/recomputation tradeoff illustrated by figure 2 on a given section C of the code, the forward sweep stops storing data for the reverse sweep. In other words, what is run then is the original code, represented by a thin arrow to the right. The adjoint code can then resume until the backward sweep reaches the checkpointed section. Then C is executed forward again, with storage (thick arrow), and the backward sweep can then resume. The peak storage space used for intermediates is thus reduced, at the cost of duplicated execution. Duplicate execution also requires some data (a “snapshot”) to be stored immediately before the checkpointed section (black bullet) and restored before its duplicate execution (white bullet). For well-chosen C, the size of the snapshot is much smaller than that of the intermediate values that would be stored without checkpointing. Data-Flow analysis helps us keep the snapshot small: only what is needed for the forward-then-backward sweeps C of the checkpointed section C must be stored. Also what is modified by C followed by D must be stored. All in all the snapshot need not be bigger than \(\text{use}(C) \cap (\text{out}(C) \cup \text{out}(D))\). Notice

![Figure 2: Checkpointing: trading duplicate execution for storage space](image)
also that \( \text{out}(D) \) is smaller or equal to \( \text{out}(D) \) thanks to restoration of intermediate values. Similarly \( \text{use}(C) \subseteq \text{use}(C) \). We pushed this analysis slightly further in [4].

Checkpoints reach their full power when they are nested. One can check that the pattern of figure 2 can be repeated recursively inside each forward-then-backward sequence. Ideally an optimal checkpointing nesting will allow both peak storage size and number of repeated executions to grow like the logarithm of the primal program run time. Even if the true optimal checkpointing scheme has been demonstrated only on one particular code type [8], see figure 3, sub-optimal checkpointing schemes can be found for most programs, that are still logarithmic. Obvious constraints apply: both ends of a checkpointed section must be in the same procedure, and in the same control-flow context. The section must also be “reentrant”, i.e. can be executed twice with no harm, which is not granted if the section allocates memory without deallocating, or sends a message without receiving it [12]...

In practice, logarithmic growth is not exactly what is desired. The available memory only allows for a fixed maximum number of snapshots \( q \). It can be shown that the optimal checkpointing scheme then requires a number of repeated executions that grows like the \( q - \text{th} \) root of the primal program run time, which is still acceptable in general.

In Tapenade, the default checkpointing scheme is to checkpoint each procedure call, except those non-reentrant. The user has freedom to deactivate checkpointing of desired calls, and to define additional checkpointed sections in the code.

4 Fixed-Point Iterations

Most code transformations can in general take advantage of additional semantic knowledge on parts of the code. This is true for adjoint AD in several situations such as parallel loops (gather-scatter), linear solvers… We take here the example of Fixed-Point iterations. A Fixed-Point loop takes as input a state \( z \) and parameters \( x \). At each iteration, \( z \) is recomputed as \( z = \phi(z,x) \) until \( z \) converges up to some tolerance. The resulting, final \( z \) depends only on the parameters \( x \). The input \( z \) is called the initial guess and only influences the iteration number till convergence. Adjoint AD must exploit the fact that the initial iterations operate on an arbitrary, almost meaningless \( z \), so that corresponding intermediate values should not be stored. In fact, in the “Two-Phases” adjoint [3], see figure 4 only the intermediate values of the last, converged iteration need be stored. On the other hand these intermediate values will be used several times during the backward sweep. This requires a special stack mechanism for repeated access, which can be arranged easily.

Let’s conclude with an open question. When the Fixed-Point iteration is itself included in an iterative control, it is often profitable to reuse the previous Fixed-Point as the initial guess of the next Fixed-Point iteration. This is called a warm start. Since the backward sweep of the adjoint code reproduces the same structure of nested iterative constructs, it should be possible to benefit also from a warm-start effect. However, strict application of the algorithm of figure 4 does not exhibit this effect. Only after a manual modification, poorly justified and unproved, could we retrieve a warm-start effect in the adjoint. This deserves further study.
References


