Construction of the adjoint MIT ocean general circulation model and application to Atlantic heat transport sensitivity

Jochem Marotzke,1 Ralf Giering,2 Kate Q. Zhang,3 Detlef Stammer,4 Chris Hill, and Tong Lee2

Center for Global Change Science, Department of Earth, Atmospheric, and Planetary Sciences, Massachusetts Institute of Technology, Cambridge

Abstract. We first describe the principles and practical considerations behind the computer generation of the adjoint to the Massachusetts Institute of Technology ocean general circulation model (GCM) using R. Giering’s software tool Tangent-Linear and Adjoint Model Compiler (TAMC). The TAMC’s recipe for (FORTRAN-) line-by-line generation of adjoint code is explained by interpreting an adjoint model strictly as the operator that gives the sensitivity of the output of a model to its input. Then, the sensitivity of 1993 annual mean heat transport across 29øN in the Atlantic, to the hydrography on January 1, 1993, is calculated from a global solution of the GCM. The “kinematic sensitivity” to initial temperature variations is isolated, showing how the latter would influence heat transport if they did not affect the density and hence the flow. Over 1 year the heat transport at 29øN is influenced kinematically from regions up to 20ø upstream in the western boundary current and up to 5ø upstream in the interior. In contrast, the dynamical influences of initial temperature (and salinity) perturbations spread from as far as the rim of the Labrador Sea to the 29øN section along the western boundary. The sensitivities calculated with the adjoint compare excellently to those from a perturbation calculation with the dynamical model. Perturbations in initial interior salinity influence meridional overturning and heat transport when they have propagated to the western boundary and can thus influence the integrated east-west density difference. Our results support the notion that boundary monitoring of meridional mass and heat transports is feasible.

1. Introduction

The impending need to synthesize, basin-wide and globally, ocean data such as the entire World Ocean Circulation Experiment (WOCE) data set including altimetry and the surface-forcing data obtained from weather centers and scatterometers makes imperative the use of sophisticated ocean general circulation models (GCMs) to (1) interpolate in space and time between the observations and (2) diagnose unobservable but important quantities such as vorticity and heat transports. Conversely (and, indeed, prior to all these interpretations), the data stream must be used to test and improve the GCMs (the stated goal 1 of International WOCE). A very powerful and general approach to synthesis is the use of optimization methods; we will concentrate on the particular flavor that has become known as the “adjoint approach” in meteorology and oceanography [e.g., Talagrand and Courtier, 1987; Thacker and Long, 1988]. (For the roots of adjoint methods in control theory and their relationship to sequential estimation, see Wunsch [1996].)

The basic idea is quite simple: A model is defined by an algorithm (often realized through a computer code) and its independent variables, for example, initial conditions, boundary conditions, or empirical parameters. A “performance index” or “cost function” measures how well a model realization matches observations; the cost function is mostly, but not necessarily, some weighted least squares measure. The optimization determines the independent (or control) variables such that the cost function is minimized. If the model is nonlinear and large (often with 10⁵–10⁷ independent variables), iterative searches for the optimal solution are among the few practical strategies, but they need directional information (which direction is downhill?). This gradient or sensitivity of the cost function with respect to the control variables is calculated by what has become known as the “adjoint model.”

Coding the “adjoint” to a complex numerical code is extremely tedious, time-consuming, and error-prone, so it is not surprising that until very recently, only few adjoint ocean GCMs existed. The first effort was performed by R. B. Long, S. M. Hwang, and W. C. Thacker (R. B. Long et al., The finite-difference equations defining the GFDL-GCM and its adjoint, unpublished report, Atlantic Oceanographic and Meteorological Laboratory, Miami, Florida, 1989), who constructed the adjoint to Cox’s [1984] version of the Geophysical Fluid Dynamics Laboratory (GFDL) GCM. It took several years to transform the prototype into a tool applicable to the inversion of hydrographic data (see the descriptions by Tziper-
man et al. [1992a, b], Marotzke [1992], Marotzke and Wunsch [1993], Bergamasco et al. [1993], and Yu and Malanotte-Rizzoli [1996]. The model and its adjoint have eventually been used to estimate the time mean and seasonally varying general circulations of the North Atlantic [Marotzke and Wunsch, 1993; Yu and Malanotte-Rizzoli, 1996, 1998], the Indian Ocean [Lee and Marotzke, 1997, 1998], and the global ocean [Sirkes et al., 1996].

The second adjoint GCM, based on the Hamburg Large-Scale Geostrophic (LSG) Ocean GCM [Maier-Reimer et al., 1993], was constructed by R. Giering around 1990 and was documented and applied to Pacific tropical wave dynamics by Giering [1996]. Schiller and Willebrand [1995] developed an approximate adjoint to the GFDL model (based only on heat and salt conservation), arguing that for optimization problems, high accuracy is often not required. This model was subsequently applied to the North Atlantic mean circulation [Schiller, 1995]. Moreover, an adjoint to the Laboratoire d'Océanographie Dynamique et de Climatologie (LODYC) GCM has been developed (P. Delecluse, personal communication, 1996).

The above list, while incomplete, indicates that adjoint GCMs have not found very widespread use in the oceanographic community despite their power (see, e.g., the comparison by Marotzke and Willebrand [1996] between general circulation and simpler-dynamics inverse models) and the popularity of the GCMs themselves. From our own experience it is clear that a large part of the problem lies in the unyielding nature of adjoint code. An adjoint tracks the sensitivity of output with respect to input (see section 2 below); every change in the GCM therefore has to be transferred to the adjoint to calculate correctly the change in sensitivity. It became clear that the construction of an adjoint ocean model ideally would be done with a software tool [e.g., Thacker, 1991].

A serious problem with most codes is that the sensitivity usually is obtained from numerical differentiation; this is not accurate enough for many applications. Moreover, this process is time-consuming. It is clear that a large part of the problem lies in the unyielding nature of adjoint code. An adjoint tracks the sensitivity of output with respect to input (see section 2 below); every change in the GCM therefore has to be transferred to the adjoint to calculate correctly the change in sensitivity. It became clear that the construction of an adjoint ocean model ideally would be done with a software tool [e.g., Thacker, 1991].

The following contains a heuristic account of how to compute an adjoint code; only the basic concepts will be explained. While there is overlap with the presentations by Talagrand [1991], Errico [1997], and Giering and Kaminski [1998], we differ from the first two in that we stress here the connection between the theoretical considerations and the actual (automatic) construction of adjoint code, a full account of which is given by Giering and Kaminski [1998].

2. Fundamentals of Adjoint Sensitivity Calculations and Adjoint Code Construction

The following contains a heuristic account of how to compute an adjoint code; only the basic concepts will be explained. While there is overlap with the presentations by Talagrand [1991], Errico [1997], and Giering and Kaminski [1998], we differ from the first two in that we stress here the connection between the theoretical considerations and the actual (automatic) construction of adjoint code, a full account of which is given by Giering and Kaminski [1998]. Computational differentiation, in general, has recently been reviewed in the conference proceedings edited by Berez et al. [1996].

Denote the "state" (i.e., all prognostic variables) of a numerical model at time step \( n \) by the vector \( \mathbf{x}_n \), \( 0 \leq n \leq N \). For simplicity, let us assume that the only independent or control variables are the initial conditions \( \mathbf{x}_0 \) of the model and
that the cost function $J$ depends only on the final state $X_N$; that is, one can write
\begin{equation}
J = f \circ X_N,
\end{equation}
where $f$ is a scalar function that maps the state vector $X_N$ onto the real axis, and the open circle stands for "operates on." The cost function is linked to the control variables by repeated application of the numerical model (once per time step),
\begin{equation}
J = f \circ \Psi_N \circ \Psi_{N-1} \circ \Psi_{N-2} \cdots \circ \Psi_2 \circ \Psi_1 \circ X_0
\end{equation}
\begin{equation}
= J[f(\Psi_N[\cdots(\Psi_1(X_0)\cdots)])].
\end{equation}

It is advantageous to consider the entire equation (2) as the model; that is, the definition of the cost function is part of the model. The sensitivity of $J$ to the control vector is given by the chain rule
\begin{equation}
\frac{\partial J}{\partial X_0} = 1 \circ f' \circ \Psi_N \circ \Psi_{N-1} \circ \Psi_{N-2} \cdots \circ \Psi_2 \circ \Psi_1 \circ 1,
\end{equation}
where the prime indicates derivative with respect to the argument, and 1 is the unit matrix. Every $\Psi_n$ is the Jacobian matrix of the model at time step $n$, i.e., the sensitivity of the state after time step $n$ to the state before time step $n$. Why the trivial factor $f' = dJ/df = 1$ has been retained will become clear immediately. Equation (3) is a $1 \times L$ matrix (i.e., a row vector), where $L$ is the size of the state vector. As it stands, (3) is evaluated from the right to the left; that is, $N$ matrix multiplications with $L \times L$ matrices must be performed. Notice that the Jacobians never need be computed explicitly if it is possible to differentiate the algorithm calculating $X_n$ from $X_{n-1}$. Every matrix multiplication corresponds to applying separately the so-called tangent linear model (TLM) to the $L$ columns of an $L \times L$ matrix (beginning with multiplying all $L$ unit vectors). The last operation is the multiplication with row vector $f'$. This is the forward mode of automatic differentiation, which has been implemented in the very general software tool ADIFOR [Bischof et al., 1992].

Alternatively, one can take the transpose of (3), which gives
\begin{equation}
\left(\frac{\partial J}{\partial X_0}\right)^T = (\Psi_1^T \circ \cdots \circ \Psi_N^T) \circ (f')^T \circ 1,
\end{equation}
\begin{equation}
\frac{\partial J}{\partial X_0} = \frac{\partial f}{\partial X_0}.
\end{equation}

Proceeding again from right to left, the number 1 is first operated upon by column vector $(f')^T$, followed by an application of the transpose Jacobian at time step $N$ and so on. Notice that (4) involves the application of an $L \times L$ matrix to a single-column vector; more generally, if $J$ were a vector valued function with $K$ components, $K$ applications would be involved. Equation (4) describes the reverse mode of automatic differentiation and is equivalent to computing the solution of the adjoint model since the adjoint operator is the transpose Jacobian. Notice that $f$ need not formally constitute an inner product, which would be required to use rigorously the term adjoint, in the same sense as in the theory of differential equations [see Talagrand, 1991]. In this regard we use a more general definition of the adjoint model, namely, as the transpose of the Jacobian matrix. Taking the point of view that the adjoint model evaluates the chain rule leads to its most parsimonious definition that we are aware of, and using the term "output function" is more appropriate here than cost function.

Notice, however, that this procedure leads to the same equations (model) as those derived for the Lagrange multipliers in constrained optimization [e.g., Thacker and Long, 1988]. For large $L$ the reverse mode is clearly more computationally efficient than the forward mode but at the expense that all the variables permitting the execution of the linearized model step must be available in reverse order of the original computation. Hence they must be stored throughout the integration of the nonlinear model or recomputed when needed. In contrast, the TLM can be applied by running along one integration of the nonlinear model. Storage requirements will be taken up again later.

The interpretation of the $\Psi_n$ as integrating the model over an entire time step is by no means required. Rather, it could denote any intermediate step between input of the controls and output of the cost function. In the extreme, $\Psi_n$ stands for the execution of a single line of computer code, which can be viewed as mapping the entire state vector plus derived variables (plus additional parameters irrelevant here) onto a single output variable. Constructing an adjoint thus means, according to (2)–(4), linearizing every code assignment and executing the transposes of all linearized assignments in reverse order, initializing with the number 1. This recipe is now demonstrated at the lowest level (we take the "atomistic" view).

Implicitly, all variables but the output variable are kept constant during an assignment, which can hence be written generally if not very efficiently as
\begin{equation}
\begin{pmatrix}
x^{in}_1 \\
x^{in}_2 \\
\vdots \\
x^{in}_n
\end{pmatrix}
= 
\begin{pmatrix}
x^{out}_1 \\
x^{out}_2 \\
\vdots \\
x^{out}_n
\end{pmatrix}
= 
\begin{pmatrix}
g(x^{in}, a) \\
\vdots \\
g(x^{in}, a)
\end{pmatrix},
\end{equation}

where $a$ is the vector of all input variables except the variable $x^{in}$. Since $x^{in}$ and $x^{out}$ allocate the same memory in the computer, $x^{in}$ is overwritten by $x^{out}$. The Jacobian of the assignment or mapping (5) is
\begin{equation}
\left(\frac{\partial x^{out}}{\partial x^{in}}\right) = 
\begin{pmatrix}
1 & 0 & 0 & \cdots & 0 \\
0 & 1 & 0 & \cdots & 0 \\
\vdots & \vdots & \vdots & \ddots & \vdots \\
0 & \cdots & 0 & 1 & 0 \\
0 & \cdots & 0 & 0 & 1
\end{pmatrix}.
\end{equation}

The adjoint of the assignment is defined by the transpose Jacobian, which links adjoint input and output variables (marked by an overbar) through
\begin{equation}
\left(\frac{\partial x^{out}}{\partial x^{in}}\right) = 
\begin{pmatrix}
1 & \frac{\partial g}{\partial x^{in}_1} & 0 & \cdots & 0 \\
0 & 1 & \cdots & 0 & 0 \\
\vdots & \vdots & \ddots & \ddots & \vdots \\
0 & \cdots & 0 & 1 & 0 \\
0 & \cdots & 0 & 0 & 1
\end{pmatrix}.
\end{equation}

Written out, this becomes
\begin{equation}
\tilde{a}^{out} = \tilde{a}^{in} + \frac{\partial g}{\partial a} x^{in},
\end{equation}
\begin{equation}
x_{r^{out}}^{new} = \frac{\partial g}{\partial x_{r^{in}}} x_{r^{in}}^{new}.
\end{equation}
In the assignment corresponding to (8b) the output again overwrites the input; hence the stated order of the assignments matters. Since every assignment usually involves only a small number of input variables, most components of (8) are, in practice, trivial: The adjoint variable remains unchanged since its physical variable has no impact on the output. These remarks illustrate how (2)–(4) have to be changed to include boundary conditions as control variables: Every mapping step \( \Psi_n \) then formally has as arguments not only the model state but also the forcing functions at all time steps. The mapping does not depend on the forcing except at the matching time step; before and after, the adjoint variable to the forcing is kept constant.

A special case arises when in the assignment corresponding to (5) the input variable \( x_i \) does not influence the output \( x_o \); that is, \( x_i \) is assigned a new value, but the right-hand side does not contain \( x_i \). This means that \( \partial g / \partial x_i = 0 \) in (8) and hence \( x_i \) = 0: The adjoint variable traces the influence of the cost function backward; since \( x_i \) is overwritten, it has no effect on the later calculations, and its adjoint is set to zero.

The above exercise illustrates how the adjoint to an arbitrary assignment in program code has to be formulated. The instructions of (8) are the core of the software tool TAMC [Giering, 1996; Giering and Kaminski, 1998]. They are presented here to make plausible that indeed, computer-generated adjoints can be done using very general principles. To illustrate these ideas, Appendix A shows an example program that is simple in its calculations yet illustrates the power of the TAMC in dealing with nonlinear algorithms. In practice, the TAMC performs the following steps:

1. The code is parsed to identify variables, syntax, etc. This step is required by every compiler. For further processing an internal abstract representation of the code is generated.

2. Data flow is analyzed. This step superficially resembles what optimizing compilers do but is performed on the entire code (globally) rather than locally (e.g., a single loop). It is needed to avoid a myriad of trivial adjoint statements that propagate zero adjoint values for constants and variables that do not influence the cost function. The data flow analysis identifies “active variables,” both in the sense of what influences the cost function and of what is influenced by the control variables.

3. The adjoint code is constructed line by line on the basis of (8). Most program codes contain higher-level structures beyond assignments; their treatment is described in detail by Giering and Kaminski [1998]. A loop can be considered as a formalized sequence of individual assignments. If any pass through the loop depends on the outcome of any other pass, the adjoint loop is performed in reverse order. Otherwise, the order is not important, and the adjoint loop is executed in the same order as in the physical model. Conditional statements (e.g., if statically unstable, perform corrective adjustment) are not differentiable at the branch point; away from it, however, linearization is achieved by providing the adjoint with the outcome of the condition during the physical model integration.

4. Once a piece of adjoint code is constructed, the parts of the solution of the physical model that are required are determined. An additional reverse data flow analysis is then applied to construct the sufficient but minimal recomputation of these (“program slicing”). Furthermore, the data flow analysis is needed to identify when the Jacobian depends on the solution of the physical model, the results of which then have to be stored or recomputed.

### 3. Preparation of the MIT GCM Code

The overarching design principle in constructing the GCM adjoint has been that there must be no human intervention necessary in the adjoint code because whatever change is manually inserted into the adjoint would be lost again upon a later application of the TAMC. This means, in particular, that if a particular data flow structure in the model leads to a conflict and errors in the adjoint, we always modify the physical model and never directly the adjoint, although the latter sometimes might be far simpler. Occasionally, errors in the adjoint suggested where the TAMC should be modified or extended.

A second principle has been to keep the optimization portion of the code, needed when the model is fitted to data, separate from the model and the adjoint. From the optimization routine’s perspective the model is merely a device that returns a single number, the cost function, if given some input vector. The adjoint delivers the gradient of the cost function with respect to the input. Hence the optimization is oblivious to the inner workings of the model, which makes modularity of the code desirable, so that different optimization procedures and, in principle, different physical models could be readily inserted. A basic logical structure emerges in which, at the top level, an optimization is performed with respect to some dimensionless independent (or control) variables, which usually are deviations from some prior estimates, scaled by the uncertainty of the prior (see Wunsch [1996] for a general discussion). When the model is called to perform an evaluation of the cost function, the vector of control variables is mapped onto the physical variables of the model in an interface subroutine. Its adjoint subroutine maps the solution of the adjoint model onto a dimensionless gradient vector.

The greatest obstacle to applying the adjoint technique to large-scale problems comes from the dependence of the transpose Jacobian on the physical model solution if the model is nonlinear (or, in other words, the dependence of adjoint code on the model solution). Two extreme strategies exist: The first is recomputation of every model statement up to the point where the solution is needed, leading to a computational load proportional to the square of the “complexity” (number of operations) of the model. The second strategy stores every intermediate quantity ever computed by the code, leading to a storage requirement roughly proportional to the code’s complexity. Apparently, only the latter strategy, which is infeasible for large-scale problems, was considered by the group that developed the “forward mode” automatic differentiation tool ADIFOR [Bischof et al., 1992].

In contrast, the TAMC by default performs “total” recomputation, at subroutine level, with the remaining storage requirements handled automatically. It is plausible, however, that by compromising between the recomputation and storage requirements a manageable computational load can be attained. Indeed, the example in Appendix A shows that by introducing storage directives, recalculation can be avoided. In completely general code the efficient introduction of storage directives is not straightforward. However, most large oceanographic and meteorological computing tasks involve the time integration of GCMs, thereby introducing a very natural guiding principle: The model state (comprised of all prognostic variables) is stored once per time step, and all diagnostic quantities are recomputed. While by no means guaranteed to be optimal, this principle provides a very simple and useful default strategy, which can subsequently be refined.
Storing the model state at every time step still would make memory requirements prohibitive for even a moderate-sized GCM, but the TAMC supports an elegant strategy called checkpointing, which was proposed by Griewank [1992] and later (manually) implemented into a quasi-geostrophic model and its adjoint by Restrepo et al. [1995]. A good summary of the method was provided by Hersbach [1998], who applied the predecessor of the TAMC to a surface wave model. The integration period of \( N \) time steps is subdivided into \( M \) segments each of length \( N/M \); the result of the forward model is stored at the end of each segment but not at other time steps. The forward model is rerun over the last segment, and its history is stored at every time step; then the adjoint is run backward over the last segment. This is repeated for all segments, moving backward in time (from the last to the first segment). The result is the exact adjoint, at the cost of one extra forward integration.

For a given number \( N \) of time steps and assuming that the "cost" of every type of storage is the same the optimum number \( M \) of segments is calculated from minimizing the total storage, which is proportional to

\[
R = \frac{N}{M} + M. \tag{9}
\]

At any given time an entire segment of length \( N/M \) and the number \( M \) of segments must be held in memory or on disk. Minimal total storage is achieved for \( M_{\text{opt}} = N^{1/2} \) and is proportional to \( R_{\text{min}} = 2N^{1/2} \), which for \( N \sim 10^4 \) (i.e., roughly 1 model year at a time step of 1 hour) is a reduction by a factor of 50. Many refinements to this simplest checkpointing strategy are possible; for example, early segments can be made longer than later segments, and further levels of recomputation can be added. Griewank [1992] proved that storage can be made to grow no faster than logarithmically, at the cost of logarithmically increasing run time. Here we use only one or two levels of recomputation to limit not only storage but also run time cost.

Checkpointing is implemented in the following way. The time-stepping loop is split into two nested loops, contained in two subroutines, outer and inner. Subroutine outer runs over \( M \) segments, defined above. Subroutine inner performs \( N/M \) time steps each time it is called. By placing specific directives before and at the very beginning of these loops, the TAMC generates the checkpointing adjoint code. Two different storage devices ("tapes") are used; in inner the prognostic and some diagnostic variables are stored in auxiliary common blocks, while in outer the prognostic variables are stored in a direct access file on disk. This technique provides the most efficient adjoint code on a vector computer; on a parallel computer with large core memory the outer tape can also be stored in memory, providing higher performance since input/output to disk is rather slow.

An extra saving is achieved by recognizing that in minimization problems the adjoint is used only to provide the downhill direction, which is useful even if only approximate [Schiller and Willebrand, 1995; Marotzke and Wunsch, 1993] implemented the very efficient shortcut of storing the model history only at certain intervals, so the adjoint operator is approximated; the permissible length of storage interval is problem-dependent. Through explicitly specified storage locations the TAMC generates code to reuse storage; this trick can be combined with the checkpointing strategy and implemented into the adjoint code without human intervention. Even under rapidly varying circumstances, additional storage savings by a factor of 5 have been possible. Storing with reduced precision further decreases the memory requirements.


The TAMC-generated adjoint GCM has already been used for very complex tasks, which justifies the considerable investment in development. Stammer et al. [1997] performed a global assimilation with a 2°-resolution model for the year 1993 using TOPEX/Poseidon altimetry and 10 day averages of synoptic surface fluxes. Although global assimilation experiments with adjoint GCMs had been performed earlier [e.g., Sirkes et al., 1996], Stammer et al.'s [1997] application appears to be the first with synoptic forcing and data. Zhang and Marotzke [1999] estimated the Indian Ocean general circulation from climatological and a basin-scale ocean model; for the first time with an ocean GCM they estimated the parameters of open boundary conditions. Both these developments were enormously simplified through the flexible construction and modification of the adjoint model.

We now describe in some depth the application of the adjoint model to a sensitivity analysis of our ocean GCM. Although straightforward in principle and extensively done with atmospheric models [see Hall et al., 1982; Errico, 1997], such a sensitivity calculation appears never to have been done with an ocean GCM. Previously, the solution of an adjoint ocean GCM has mainly been used as directional information in a minimization procedure but otherwise discarded. Here we calculate the sensitivity of an important function of the model solution, the heat transport across 29°N in the Atlantic, to the initial conditions in temperature and salinity. More specifically, the heat transport is the annual mean for 1993 and taken from the solution of Stammer et al. [1997], while the initial conditions are the temperatures and salinities estimated for January 1, 1993. The latitude of 29°N is chosen because it is close to the maximum northward heat transport. Hence the output function is

\[
Q = P^{-1} c_p \rho_0 \int \int \int v T dx \, dz \, dt, \tag{10}
\]

where \( P \) is the averaging period, \( c_p \) is specific heat, \( \rho_0 \) is a reference density, \( v \) is meridional velocity, and \( T \) is potential temperature.

Plate 1a shows the sensitivity of the annual mean heat transport at 29°N, \( Q \), to the sea surface temperature of January 1, 1993. The large positive contribution near the western boundary and south of 29°N is readily understood from the surface velocity field (Figure 1b). The considerable northward boundary current contributes a larger temperature flux if its temperature is higher; downstream of the section, there is no contribution from this effect. With a northward velocity component of 10 cm s⁻¹ a temperature anomaly can be transported in the boundary current over 3000 km in 1 year, roughly consistent with the southward extent of the region of positive sensitivity in Plate 1a. In contrast, simple passive advection of temperature anomalies cannot explain the negative sensitivity away from the western boundary and right to the north of the section because the surface flow across the section is weakly northward over the entire year. Neither can passive advection account for the large negative sensitivities along the western boundary to the north of the section or the large positive sensitivity off Africa.
To gain further insight, we have to consider the effect of a temperature perturbation more broadly. A change in temperature has a dynamically active component, through its effect on density and hence thermal wind shear, and a dynamically passive component (temperature change on an isopycnal, contributing to the combination of temperature and salinity that is locally orthogonal to the density change and was called “veronicity” by Munk [1981, p. 282]). For small amplitudes, the “active” component of a temperature perturbation has the same consequences as a salinity perturbation, scaled by \(-\alpha/\beta\), where \(\alpha\) and \(\beta\) are the thermal and haline expansion coefficients, respectively:

\[
\alpha = -\frac{1}{\rho} \frac{\partial \rho}{\partial T}_s, \quad \beta = \frac{1}{\rho} \frac{\partial \rho}{\partial S}_T.
\]

Hence we can identify the sensitivity of heat transport to the dynamically active part of the initial temperature by considering the sensitivity of heat transport across 29\(^\circ\)N to initial salinity perturbations, which is shown in Plate 1b and given by

\[
\left( \frac{\partial Q}{\partial T} \right)_s = \left( \frac{\partial Q}{\partial \rho} \right)_T \left( \frac{\partial \rho}{\partial T} \right)_s + \left( \frac{\partial Q}{\partial T} \right)_\rho = -\alpha \rho \left( \frac{\partial Q}{\partial \rho} \right)_T, \quad (13)
\]

using (11), where the last term on the right-hand side is the kinematic sensitivity. The sensitivity of \(Q\) to initial salinity variations is shown in Plate 1b and given by

\[
\left( \frac{\partial Q}{\partial S} \right)_T = \left( \frac{\partial Q}{\partial \rho} \right)_T \left( \frac{\partial \rho}{\partial S} \right)_T = \beta \rho \left( \frac{\partial Q}{\partial \rho} \right)_T. \quad (14)
\]

We can now isolate the kinematic sensitivity of heat transport to initial temperature variations by rearranging (13) and using (14) to obtain

\[
\left( \frac{\partial Q}{\partial T} \right)_\rho + \alpha \rho \left( \frac{\partial Q}{\partial \rho} \right)_T = \left( \frac{\partial Q}{\partial T} \right)_s + \alpha \beta \left( \frac{\partial Q}{\partial S} \right)_T. \quad (15)
\]

A worked example of (15) is given in Appendix B, where the result is demonstrated for a box model that is “run” over one time step. Plate 1c shows the “sum” of Plates 1a and 1b weighted according to (15). The large positive contributions in the western boundary current south of 29\(^\circ\)N confirm the simple picture, outlined above, of warm perturbations leading to increased heat transport by the northward mean flow. In contrast, the kinematic sensitivity to initial SST is negative over the ocean interior and shows considerable symmetry about the section. This latter feature reflects the model’s finite difference scheme (heat transport is calculated at a velocity gridpoint, so the two neighboring temperatures to the north and south must be averaged to form the product of velocity and temperature). The minima in Plate 1c across 29\(^\circ\)N reflect the banded structure of generally southward velocity at level 2 (Figure 1b), except west of 70\(^\circ\)W. Temperatures in the top two layers are strongly coupled in winter (the mixed layer is \(\sim 100\) m deep; not shown), so in the interior, where the meridional surface velocities are small, the flow at level 2 dominates the advection of a temperature anomaly. Overall, the kinematic sensitivity displayed in Plate 1c reflects the classical picture of a subtropical gyre, with a strong poleward western boundary current and broad equatorward return flow. It is the ability to isolate the kinematic sensitivity that allows us to state, loosely, that we have identified a passive tracer from temperature and salinity without having to run a passive-tracer equation or an ensemble of Lagrangian drifters.

Since Plate 1c has allowed us to identify the kinematic contributions to heat transport sensitivity, we can now return to Plates 1a and 1b to complete our interpretation. The contributions from temperature and salinity in the western boundary
MAROTZKE ET AL.: ADJOINT ATLANTIC HEAT TRANSPORT SENSITIVITY 29,35

Plate 1. (a) Sensitivity of mean 1993 Atlantic heat transport across 29°N (dotted line) to surface temperature on January 1, 1993. The contour interval is $0.02 \times 10^{12}$ W K$^{-1}$, and the zero contour is drawn in red. (b) Sensitivity of mean 1993 Atlantic heat transport across 29°N (dotted line) to surface salinity on January 1, 1993. The contour interval is $0.04 \times 10^{12}$ W per practical salinity unit (psu), and the zero contour is drawn in red. (c) As in Plate 1a but for the kinematic sensitivity.

Current to the north of the section and off Africa cancel nearly perfectly in Plate 1c and are hence purely dynamical; that is, they influence the heat transport only through their effects on density and velocity. Near the western boundary an increase in surface density leads to a decrease in sea level and hence a stronger baroclinic shear in the Gulf Stream. This perturbation would propagate southward, as a mix of Kelvin and topographic waves [e.g., Döscher et al., 1994], and influence heat transport across 29°N from a significant distance. The nearly antisymmetric contributions off Florida and off Africa, just north and south of the section, respectively (Plates 1a and 1b), lead to increased northward thermal wind shear and hence surface flow at both boundaries. The location right at the boundary is crucial to effect zonally integrated northward surface flow; in contrast, an isolated mid-ocean anomaly would induce flow around itself without additional zonally integrated transport.

A tentative explanation for the negative sensitivity to mid-ocean initial salinity (Plate 1b) can be given as follows. A positive salinity anomaly in the center of the subtropical gyre causes a positive density and hence negative sea level anomaly. If this anomaly can spread westward as a barotropic Rossby wave (without, however, reaching the western boundary), the net result is a weakening of the subtropical gyre and its associated heat transport. A positive temperature anomaly has the opposite effect; notice that only a narrow latitudinal range is involved. The near-total compensation between dynamical and kinematic temperature sensitivities south of the section is conspicuous but appears accidental, judging from an analogous sensitivity calculation of heat transport across 33°N (not shown).

The sensitivity of heat transport across 29°N to initial conditions deeper down differs significantly from those at the surface (Plate 2). Somewhat trivially, a unit perturbation in deep temperature or salinity creates a larger heat or salt content change because of the larger layer thickness at greater depth. In addition, there is a considerably larger dynamical contribution of initial temperature changes to the heat transport; correspondingly, there is a larger contribution from initial salinity (Plate 2b). As a consequence, the kinematic sensitivity to a deep initial temperature change (again, reflecting the horizontal velocity field at the same depth; not shown) plays a negligible role; the cancellation between the dynamical sensitivities is almost complete. At 1160 m the sensitivity to temperature shows a pattern with negative contributions north of the section, concentrated near the western boundary, while south of the section, the contributions are positive and arise along the entire section. Additionally, positive contributions arise along the eastern boundary, from as far south of the section as 10°N. Most remarkable is perhaps that both shallow and deep temperature and salinity anomalies at the rim of the Labrador Sea can influence, within 1 year, the heat transport.
Plate 1. (continued)
Plate 2. (a) Sensitivity of mean 1993 Atlantic heat transport across 29°N (dotted line) to temperature at 1160 m on January 1, 1993. The contour interval is $0.5 \times 10^{12} \text{ W K}^{-1}$, and the zero contour is drawn in red. 
(b) Sensitivity of mean 1993 Atlantic heat transport across 29°N (dotted line) to salinity at 1160 m on January 1, 1993. The contour interval is $2 \times 10^{12} \text{ W (psu)}^{-1}$, and the zero contour is drawn in red.
Plate 3. Perturbation in Atlantic meridional overturning stream function (in Sverdrups), at 60 day intervals, beginning with day 30, following a salinity perturbation of amplitude 0.01 at 1160 m between latitudes 28° and 38°N and between longitudes 68° and 58°W. Positive values indicate clockwise anomalous transport.
5. Dynamical Considerations

5.1. Sensitivity of Heat Transport to Salinity Perturbations

We now change our perspective and consider how a deep salinity anomaly influences meridional heat transport over a considerable distance. Apart from the dynamical information this procedure yields, it provides an independent test of the adjoint sensitivity calculation. A salinity perturbation is added to the initial conditions of the optimized run of Stammer et al. [1997] in that salinity at 1160 m depth is perturbed by 0.01 between latitudes 28° and 38°N and between longitudes 68° and 58°W (a total of 25 grid cells). This location for the initial anomaly is chosen to encompass the sensitivity maximum shown in Plate 2b, but its center is slightly to the east so that its westward migration can also be studied.

Figure 2 shows the resulting perturbations in horizontal velocity at 160 m depth, at 60 day intervals, beginning with day 30. The salinity anomaly at 1160 m depth creates a response at all depths (counterclockwise vortex above, clockwise vortex below), which migrates westward with the speed of first-mode Rossby waves. The perimeter of the vortex reaches the western boundary after ~90 days. The subsequent evolution south of the anomaly resembles that of a shoaling Kelvin wave, with a northward current emerging along the western boundary; when the wave reaches the equator, it “draws” water from the east. The behavior in Figure 2 is entirely consistent with the Kelvin wave patterns described by Kawase [1987]. Notice that within 1 year, there is only a weak response south of the equator.

Plate 3 shows the perturbation in meridional overturning stream function at the same times as the velocity plots of Figure 2. A coherent positive pattern (indicating clockwise anomalous transport) is established roughly at the latitudes of the perturbation within 90 days, which is when the anomaly vortex has reached the western boundary. This pattern intensifies until day 270, when its southern edge begins to extend southward across the equator. In contrast, the core of the stream function anomaly migrates southward more slowly (from ~30° to 20°N, between days 90 and 330). Below ~3000 m depth the southern edge migrates southward more slowly than above.

Figure 3 shows the time series of the resulting heat transport anomalies at a number of latitudes. To the north of the perturbation, there is virtually no response. At 35°N the initial response of 1.5 Terawatts (TW, 1 TW = 10^{12} W) decays over 100 days but regains strength to end up at ~2 TW. At both 24° and 29°N the heat transport anomaly starts very small but grows steadily to a maximum of ~4 and 3.5 TW, respectively, between days 200 and 250, roughly coincident with the maximum in meridional overturning. Finally, the perturbation heat transport anomaly at 9°N begins to increase considerably only after day 100. The evolution of the meridional overturning anomaly well explains that of the heat transport anomaly: There is no northward migration at all, while to the south of the perturbation the heat transport evolves in lockstep with the meridional overturning; notice that positive stream function values mean northward flow near the surface and southward flow at depth, causing positive heat transport anomalies.

The annual mean of the heat transport anomaly at 29°N is 2.3 TW. This compares excellently with that implied by the adjoint sensitivity calculation,

\[
\delta Q = N_{\delta S} \left( \frac{aQ}{aS} \right)_T \delta S \\
= 25 \times 1.0 \times 10^{13} \frac{W}{psu} \times 0.01 \text{ psu} = 2.5 \text{ TW},
\]

where \(N_{\delta S}\) is the number of perturbed grid points, \(\delta S\) is the magnitude of the perturbation, and \((aQ/aS)_T\) is the patch average of the adjoint sensitivities of Plate 2b. We have confirmed the linearity of our perturbation run by scaling the salinity perturbation down by an order of magnitude; the response is nearly identical except for the same order-of-magnitude reduction in amplitude.

We have experimented with other deep-salinity perturbations (not shown), and the picture that emerges explains why the pattern of positive influence is wider at lower latitudes than at higher latitudes (Plate 2b). When the anomaly is placed right at the boundary, it sooner reaches latitudes farther south; the Kelvin wave is set off immediately, rather than following the westward migration. Over 1 year hence a salinity perturbation...
from near 50°N can reach the "output latitude" of 29°N only if the perturbation starts right at the boundary. Anomalies farther south, in contrast, have enough time to migrate westward prior to setting off the Kelvin wave. Conversely, in the adjoint the "anti-Kelvin wave" induces eastward propagation, which has more time to migrate at lower latitudes (e.g., 40°N), because it is excited earlier. However, notice that the wave picture is complicated considerably by bottom topography; the time evolution of the adjoint solution (not shown) clearly shows that the maximum near 38°N in Plate 2b is due to trapping by topography and local recirculation.

5.2. Monitoring the Meridional Overturning

The above considerations have demonstrated the importance of density anomalies near the boundaries for meridional overturning and heat transport. Specifically, the sensitivity plots (Plates 1 and 2) show that dynamical sensitivity of heat transport is generally greatest to density perturbations near zonal boundaries. Moreover, the model runs of the previous subsection show that density anomalies influence the meridional transports once the anomalies reach the boundaries. Thus a suggestion arises for the monitoring of meridional overturning and heat transport by measuring density changes near the ocean margins as part of a climate-observing system. We now place this connection into a more general dynamical context. If meridional flow is everywhere in thermal wind balance,

\[ \partial_z v = -\frac{g}{f\rho_0} \partial_z \rho, \quad \text{(17)} \]

zonal integration across an ocean basin yields

\[ L_x \overline{\partial_z v} = -\frac{g}{f\rho_0} \left( \rho_e - \rho_w \right), \quad \text{(18)} \]

where \( L_x \) is the zonal extent of the basin, \( \rho_e \) and \( \rho_w \) are the densities at the eastern and western walls, respectively, and the overbar marks a zonal average. Other notation is standard; Cartesian coordinates are used for simplicity. Equations (17) and (18) reflect the well-known fact that the geostrophic transport between any two points is proportional to the pressure difference between these points, irrespective of their distance (provided that topography is nowhere intersected).

Assuming vertical sidewalls (i.e., \( L_x \) independent of depth) and the existence of a meridional stream function \( \psi \) (guaranteed if no mass enters or leaves through the zonal boundaries) leads to

\[ \frac{f\rho_0}{g} \partial_z \psi = -\frac{f\rho_0}{g} \partial_z (L_x \overline{v}) = \rho_e - \rho_w, \quad \text{(19)} \]

meaning that under the simplest of all circumstances the curvature of the meridional overturning stream function, with respect to the vertical, is proportional to the east-west density difference. This latter quantity took center stage in the theory of the purely buoyancy-driven meridional overturning circulation developed by Marotzke [1997] and Marotzke and Klinger [1999], but it does not appear to have found widespread use elsewhere. Notable exceptions are the effort of Lynch-Stieglitz et al. [1999], who used the oxygen isotope composition of benthic foraminifera to estimate the density drop and geostrophic transport across the Florida Straits, and the closely related "transport index" used by R. G. Curry and M. S. McCartney (manuscript in preparation, 1999) to characterize horizontal circulation changes inferred from hydrography.

The situation is considerably more complex than suggested by (19) if wind forcing and irregular bottom topography are admitted, which leads to contributions to the meridional overturning stream function that are not in thermal wind balance. Specifically, the external mode (vertical average) projects onto the meridional overturning in the presence of bottom topography [e.g., Robbins and Toole, 1997; Lee and Marotzke, 1998]. Moreover, meridional overturning variability on timescales of seasonal and shorter is dominated by the varying Ekman transport and its depth-independent return flow [Lee and Marotzke, 1998; Jayne, 1999]. None of these contributions to the meridional overturning has a straightforward relation to the density at the zonal boundaries, but as we show now, (19) can be a good approximation, even under nonidealized circumstances.

Plate 4 shows the left-hand and right-hand sides of (19) evaluated from the model solution obtained by Stammer et al. [1997] for the year 1993. For simplicity we only show regions of the North Atlantic where in the analysis we could easily avoid cutting through bottom topography. Between 500 and 1700 m depth the correspondence between the curvature of the meridional overturning stream function and the east-west density difference is quite good; the difference plot between the two shows considerably smaller contributions (Plate 4c). Largely, density is lower at the eastern boundary than at the western boundary, above 1700 m depth, meaning positive meridional transport shear, in turn consistent with northward mass transport in the thermocline. In contrast, the fit is poor above 500 m, presumably because of the reasons outlined above. A deeper investigation would go beyond the scope of this paper and will be left for future study.

6. Concluding Remarks

We have presented the foundations and a novel application of a powerful recent development in ocean modeling. A software tool has been used to computer generate the adjoint of a full-fledged ocean GCM, without human manipulation of the adjoint code itself to make it correct or efficient. Possible applications of the adjoint model are: (1) oceanographic inverse or optimization problems (synthesizing observations; the model described here has been applied in this mode by Stammer et al. [1997] and Zhang and Marotzke [1999]); (2) the calculation of fastest growing structures in dynamical instabilities [e.g., Farrell and Moore, 1992; Eckert, 1998; Eckert et al., submitted manuscript, 1999] (we have not yet applied the adjoint MIT model to a problem of this class); and (3) the sensitivity of parts of the model solution to independent parameters (only this application has been the focus of this paper).

Our results must in part be considered preliminary since the model solution obtained by Stammer et al. [1997] is only the first application of a global state estimation system. A subsequent experiment comprising a longer integration time and model improvements is being carried out; this run will be used for a more in-depth application of the ideas about sensitivity studies presented here. Moreover, in order to draw a conclusion concerning the importance of exact initialization at various locations one must multiply the sensitivities shown here by typical variabilities; this would reduce the apparent influence of the deep temperatures and salinities. On the other hand, the isolation of kinematic and dynamical influences of temperature anomalies is independent of the particular model realization; it shows a way of disentangling individual processes from a complex superposition of effects.

Our sensitivity analysis lays out a strategy for designing field
would be detrimental to the ability to link dynamically tem-
perature perturbations. Finally, the highly localized regions of
large dynamical influence of temperature changes on heat trans-
port, visible in Plate 1a at both ends of the heat transport line,
indicate that “boundary monitoring” of the meridional over-}
turning may be a feasible observational strategy. This last point
warrants further study because the thermal wind relation is not
guaranteed to hold in the vicinity of the boundary and the presence
of small-scale variability. It remains to be shown whether spatial
averaging would alleviate these concerns.

Appendix A: A Nontrivial Example of Automatic
Adjoint Code Construction

To illustrate the ideas outlined in section 2, an example
program is shown that is simple in its calculations yet illustrates
the power of the TAMC in dealing with nonlinear algorithms.
The FORTRAN code (Figure A1) and its TMC-generated
adjoint (Figure A2) are shown with minimal editing, essentially
taking out the declarative statements to make the codes comp-
act enough for illustration. Moreover, the numerical results
are inserted step by step as comments. Before the subroutine
under consideration is one of many and would
have as input arguments adfc and the adjoints to the control
variables that resulted from previous adjoint subroutines.)
The adjoint statements are executed in reverse order, as are the do
loops, where necessary. Repeatedly, the solution of the model
is restored from an array or file and then used in the adjoints
to (linearized) nonlinear statements. Most notable is the ad-
joint to the convective adjustment. The “static stability” of
the physical model solution is checked, and the convective adjust-
ment performed on the adjoint variables. The outcome of this
check is predetermined once the physical model has been run,
which is tantamount to the statement that we have performed
a linearization of the branching statement (this is, of course,
not possible right at the branch point). After the $i = 2$ step,
$\text{adx}(2)$ and $\text{adx}(3)$ are completely mixed, while after the $i = 1$
step, $\text{adx}(1) = \text{adx}(2)$, both in direct analogy to the convective
adjustment in the model. The physical interpretation is that
since the properties of two adjacent “boxes” are mixed, each
box contributes equally much to the value after mixing; hence
the adjoint variables are equal, too.

To provide an independent test of the outcome of the ad-
joint model, we now perturb the control variables by “tracers”
such that

$$
\mathbf{x}_0 = (1 + e_1, 3 + e_2, 3 + e_3)
$$

This leads to the intermediate results of model (see program
code for definitions; perturbations are carried to first order
only),

$$
\begin{align*}
\mathbf{x}_1 &= (2 + 4e_1, 10 + 2e_2, 10 + 2e_1 + 6e_3) \\
\mathbf{x}_2 &= (6 + 3e_1 + 3e_2, 6 + 3e_1 + 3e_2, 10 + 2e_1 + 6e_3),
\end{align*}
$$

and after convective adjustment is complete, to

$$
\begin{align*}
\mathbf{x}_3 &= (6 + 3e_1 + 3e_2, 8 + 2.5e_1 + 1.5e_2 + 3e_3, \\
&\quad 8 + 2.5e_1 + 1.5e_2 + 3e_3).
\end{align*}
$$

Hence

$$
f_c = 40.25 + 15.5e_1 + 10.5e_2 + 15e_3,
$$

so that in accord with the adjoint result,

$$
\frac{\partial}{\partial \mathbf{x}_0} f_c(\mathbf{x}_0 = (1, 3, 3)) = \frac{\partial}{\partial e} f_c(e = 0) = (15.5, 10.5, 15).
$$

\[A6\]
Appendix B: Derivation of Kinematic Sensitivity in a Simple Example

As the simplest example of a nonlinear ocean model, with the flow dependent on temperature and salinity and a spatial "extent," consider the two-box model of the thermohaline circulation originally devised by Stommel [1961] and later used extensively for studying multiple equilibria and atmospheric interactions of the thermohaline circulation [e.g., Marotzke, 1996]. The model consists of two ocean boxes, which are well mixed and have constant depth; box 1 represents the high-latitude ocean, and box 2 represents the low-latitude ocean (Figure B1). H₁ and H₂ are ocean heat gain through the surface, and Hₛ is a virtual surface salinity flux, which can be simply related to a surface freshwater flux. All surface fluxes are assumed constant. The conservation equations for the ocean are

\[ \begin{align*}
\dot{T}_1 &= H_1 + |q|(T_2 - T_1), \\
T_2 &= H_2 - |q|(T_1 - T_2), \\
\dot{S}_1 &= -H_S + |q|(S_2 - S_1), \\
\dot{S}_2 &= H_S - |q|(S_1 - S_2),
\end{align*} \]

The flow strength q is related to the meridional density gradient by a linear law,

\[ q = k[\alpha(T_2 - T_1) - \beta(S_2 - S_1)], \tag{B5} \]

where a linear equation of state has been assumed and \( \alpha \) and \( \beta \) are the thermal and haline expansion coefficients, respectively. Positive q means poleward surface flow, while negative q means equatorward surface flow. Here we can assume that q is positive without loss of generality and hence can omit the absolute magnitude signs in the equations. We denote initial temperatures by \( \theta_1 \) and \( \theta_2 \) and initial salinities by \( \Gamma_1 \) and \( \Gamma_2 \).

The initial flow field is

\[ q_0 = k[\alpha(\theta_2 - \theta_1) - \beta(\Gamma_2 - \Gamma_1)]. \tag{B6} \]

The model is stepped forward by one time step according to

\[ T_{1/2} = \theta_{1/2} + \Delta t H_{1/2} \pm \Delta t q_0(\theta_2 - \theta_1), \tag{B7} \]

and done similarly for salinities. The flow after one time step is

\[ q = k[\alpha(T_2 - T_1) - \beta(S_2 - S_1)]. \tag{B8} \]

Notice that in contrast to (B5), the variables in (B8) are discretized in time. Analogous to the sensitivity analysis in the main text, the output function is chosen to be the heat transport,

\[ J = q(T_2 - T_1). \tag{B9} \]

Despite the simplicity of the model, analyzing the sensitivity of J with respect to the initial conditions is fairly involved because of the multiple dependencies of the variables. The complete algebra is rather tedious, and we will not present it.
Plate 4. Diagnostics from the global solution of Stammer et al. [1997] over the North Atlantic sector: (a) left-hand side of (19), $f\rho_0/g\partial_z\psi$; (b) right-hand side of (19), $\rho_E - \rho_W$; and (c) residual of (19), $(\rho_E - \rho_W) - f\rho_0/g\partial_z\psi$. The contour interval is 0.1 kg m$^{-3}$, and the zero contour is drawn in red.
subroutine admodel( nc, x0, fc, adx0, adfc )

C***********************************************************************
C** This routine was generated by the                              **
C** Tangent linear and Adjoint Model Compiler, TAMC 5.0.10        **
C***********************************************************************

...  

C RESET LOCAL ADJOINT VARIABLES
  do ipl = 1, 3
    adx(ipl) = 0.
  end do
ady = 0. ! Nonlocal adjoint variables: adfc=1 adx0=(0,0,0)

C OPEN FILES OF TAPE: storefile
  open(60,file='storefile_1_model_x',ACCESS='DIRECT',RECL=1*3*8)

C ADJOINT COMPUTATIONS
  read(60,REC=1) x ! Restore from file: x = x3 = (6,8,8)
  adx(3) = adx(3)+3*adfc ! adx(3)=3
  adx(2) = adx(2)+2*adfc ! adx(2)=2
  adx(1) = adx(1)+2*adfc*(x(1)-5.5) ! adx(1)=1
  adfc = 0.

C Adjoint of convective adjustment.
  do i = 2, 1, -1
    do ipl = 1, 3 ! Restore from array xh in COMMON block:
      x(ipl) = xh(ipl,1)
    end do ! i=2- x=x2=(6,6,10)
    if (x(i) .lt. x(i+l)) then ! i=2-6<10? Yes.
      adx(i) = adx(i)+adx(i+l) ! i=2, adx(2)=5
      adx(i+l) = 0. ! i=2, adx(3)=0 ! i=1-adx(2)=2.5
      adx(i+l) = adx(i+l)+0.5*adx(i) ! i=2, adx(3)=2.5 ! i=1-adx(2)=1.75
      adx(i) = 0.5*adx(i) ! i=2, adx(2)=2.5 ! i=1-adx(1)=1.75
    endif
  end do
  doi=l, 3
  x(i) = x0(i)
  adx0(i) = adx0(i)+adx(i)
  adx(i) = 0.
end do

C CLOSE FILES OF TAPE: storefile
  close(60)
end

! adx = (1.75, 1.75, 2.5)
! i=1: ady=1.75 !i=2: ady= 3.5 !i=3: ady = 6
! adx(i) = 2*adx(i)*x(i)
! ady = 6 adx=(3.5,10.5,15)
! adx(1) = adx(1)+2*ady*x(1) ! adx(1) = 15.5
ady = 0.
do i = 1, 3
  adx0(i) = adx0(i)+adx(i)
adx(i) = 0.
end do
! adx0 = (15.5,10.5,15)

Figure A2. Tangent-Linear and Adjoint Model Compiler (TAMC)-generated adjoint of the code displayed in Figure A1. Included as comments are the values of the variables after the statements have been executed.
This is indeed the sensitivity of the transport of a purely passive tracer, after one time step, to the initial condition, as can be seen by inserting the time-stepping equation (B7) into the transport definition (B9) and assuming that \( q \) and \( q_0 \) are known a priori.

**Appendix C: The MIT General Circulation Model**

The MIT GCM has been documented extensively by Marshall et al. [1997a, b], so a brief description suffices here. The model is used in its hydrostatic version, which is based on the primitive equations on a sphere under the Boussinesq approximation. Spatial coordinates are longitude, latitude, and height. Conservation of horizontal and vertical momentum, volume, heat, and salt and an equation of state are written in the form

\[
\begin{align*}
\partial \mathbf{u} &= -\nabla p/\rho_0 + \mathbf{G}_u, \\
\partial \rho &= -g \rho \\
\nabla \mathbf{v} + \partial \mathbf{w} &= 0 \\
\partial T &= G_T \\
\partial S &= G_S \\
\rho &= \rho(S, T, p),
\end{align*}
\]

where \( \mathbf{u} \) and \( \nabla \mathbf{v} \) are the horizontal components of velocity and the gradient operator, respectively, \( T \) is the potential temperature, and \( p \) is the deviation of the pressure from that of a resting ocean of constant reference density \( \rho_0 \). The \( G \) and \( \mathbf{G} \) stand for all contributions to the tendency terms except the pressure gradient forces in (C1), i.e., for advective, Coriolis, metric, forcing, dissipation, and mixing terms including a convective adjustment. The nonhydrostatic version of the model will not be considered here, and the quasi-hydrostatic approximation [Marshall et al., 1997b] would produce only minor modifications in (C2). The model equations are solved on a staggered grid ("C" grid) [Arakawa and Lamb, 1977] using the usual boundary conditions (insulating side walls and bottom, surface fluxes of momentum, heat, and (equivalent) salinity, and either no-slip or free-slip conditions tangential to side walls or the bottom). Horizontal velocity, temperature, and salinity are marched forward in time using (C1), (C4), and (C5) in an Adams-Bashforth discretization. Vertical velocity, density, and pressure are diagnostic variables; the first two of them are readily calculated from (C3) and (C6), respectively.

The computation of pressure is less straightforward and hence roughly sketched here, in particular, since this is important for the construction of the adjoint. The deviation of pressure from the reference state is separated into "surface" and "hydrostatic" contributions according to

\[
p(\lambda, \phi, z) = p_S(\lambda, \phi) + p_{HY}(\lambda, \phi, z),
\]

where \( p_S \) is the pressure at \( z = 0 \) (the surface) and \( p_{HY} \) is calculated from the hydrostatic relationship (C2) using the integration constant \( p_{HY}(z = 0) = 0 \). (The calculation of \( p_S \) and \( p_{HY} \) does not change even in the nonhydrostatic case [Marshall et al., 1997a, b]). An equation for the surface pressure is obtained by integrating the continuity equation (C3) vertically over the entire depth. As shown in Appendix 2 of Marshall et al. [1997a], the end result is an elliptic equation for the surface pressure, which is solved through an iterative con-
jugate gradient algorithm. The discrete elliptic operator is a symmetric (and hence self-adjoint) matrix, meaning that the adjoint of the procedure calculating the surface pressure is identical to the procedure itself. This leads to considerable simplification; constructing the adjoint to the iterative procedure would lead to either excessive recomputation or storage because nonlinear operations (e.g., scalar products and ratios [Press et al., 1992]) are involved. However, since the equation for surface pressure and hence its solver are self-adjoint the subroutine itself is called from the adjoint, with input and output variables appropriately interchanged by the TAMC.

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MAROTZKE ET AL.: ADJOINT ATLANTIC HEAT TRANSPORT SENSITIVITY


R. Giering and T. Lee, Jet Propulsion Laboratory, Pasadena, CA 91109.

C. Hill, Center for Global Change Science, Department of Earth, Atmospheric, and Planetary Sciences, Massachusetts Institute of Technology, Cambridge, MA 02139.

J. Marotzke, School of Ocean and Earth Science, University of Southampton, Southampton Oceanography Centre, Southampton SO14 3ZH, England, U. K. (Jochem.Marotzke@soc.soton.ac.uk)

D. Stammer, Department of Physical Oceanography, Scripps Institution of Oceanography, Mail Code A-024, La Jolla, CA 92039.

K. Q. Zhang, Department of Geophysics and Planetary Sciences, California Institute of Technology, Pasadena, CA 91125.

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