Development of the adjoint of GEOS-Chem

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Abstract

We present the adjoint of the global chemical transport model GEOS-Chem, focusing on the chemical and thermodynamic relationships between sulfate – ammonium – nitrate aerosols and their gas-phase precursors. The adjoint model is constructed from a combination of manually and automatically derived discrete adjoint algorithms and numerical solutions to continuous adjoint equations. Explicit inclusion of the processes that govern secondary formation of inorganic aerosol is shown to afford efficient calculation of model sensitivities such as the dependence of sulfate and nitrate aerosol concentrations on emissions of SO\(_x\), NO\(_x\), and NH\(_3\). The adjoint model is extensively validated by comparing adjoint to finite difference sensitivities, which are shown to agree within acceptable tolerances; most sets of comparisons have a nearly 1:1 correlation and \(R^2>0.9\). We explore the robustness of these results, noting how insufficient observations or nonlinearities in the advection routine can degrade the adjoint model performance. The potential for inverse modeling using the adjoint of GEOS-Chem is assessed in a data assimilation framework through a series of tests using simulated observations, demonstrating the feasibility of exploiting gas- and aerosol-phase measurements for optimizing emission inventories of aerosol precursors.

1 Introduction

Chemical transport models (CTMs) enhance our ability to understand the chemical state of the atmosphere and allow detailed analysis of issues ranging from intercontinental pollution transport to the coupling of anthropogenic processes, regional pollution and climate change. Of particular interest in these realms is explicit consideration of the role of aerosols, the importance of which is well documented. Given the substantial uncertainty that remains in many aspects of detailed aerosol simulations, it is critical to further examine how the numerous parameters in such models steer their predictions, especially estimates of emissions inventories for aerosols and their precursors. The
complexity of the thermodynamic and photochemical processes that govern secondary formation of aerosols precludes simple assessment of the dependence of model predictions on such parameters. Working to arrive at CTMs that more reliably reproduce observations, adjoint modeling is often employed as a method for determining the sensitivity of model predictions to input parameters and for optimizing these parameters to enforce agreement between the model predictions and an observational data set (data assimilation).

Several inverse modeling studies have analyzed sources of aerosols and aerosol precursors on regional scales. As of yet, most studies have been fairly coarse, limited to optimization of a few scaling factors for emissions inventories spanning large domains. Park et al. (2003) used multiple linear regression to estimate annual mean sources of seven types of primary carbonaceous aerosol over the United States. A Kalman filter approach was used to estimate improved monthly emissions scaling factors for NH$_3$ emissions over the United States using observations of ammonium wet deposition in works by Gilliland and Abbitt (2001) and Gilliland et al. (2003, 2006). Mendoza-Dominguez and Russell (2000, 2001) optimized domain-wide emissions scaling factors for eight species over the eastern Unites States using observations of gas-phase inorganic and organic species and specified fine particles. Source apportionment models have also been refined using inverse modeling (Knipping et al., 2006; Schichtel et al., 2006).

Data from satellite observations offer tremendous potential for inverse modeling of aerosols (Collins et al., 2001; Kahn et al., 2004). In order to best exploit these, and other, large data sets, it is desired to extend inverse analysis of aerosol models to global scales and to finer decomposition of the emissions domains. Such goals require consideration of inverse modeling methods designed for large sets of variable parameters. The adjoint method is known to be an efficient means of calculating model sensitivities that afford examination of numerous parameters, where these values can subsequently be used in tandem with an observational data set for data assimilation. First appearing in the field of atmospheric science in the early 1970s (Marchuk, 1974; Lamb et al., 1975), the method later came to be applied extensively in meteorology, e.g., Talagrand and Courtier (1987); Erice and Vukicevic (1992). In the last decade, the adjoint approach has expanded to include ever more detailed CTMs, beginning with the abbreviated Lagrangian stratospheric model of Fisher and Lary (1995) and the Lagrangian tropospheric model of Elbern et al. (1997). Vukicevic and Hess (2000) used the adjoint method to perform a sensitivity study of an inert gas-phase tracer over the Pacific, while Elbern and Schmidt (1999) presented the first adjoint of a 3-D Eulerian CTM to include chemistry. These initial works have been followed more recently by similar development and application of adjoint models of several CTMs: CHIMERE (Vautard et al., 2000; Menut et al., 2000; Schmidt and Martin, 2003), IMAGES (Muller and Stavrakou, 2005; Stavrakou and Muller, 2006), Polair (Mallet and Sportisse, 2004, 2006), TM4 (Meirink et al., 2006), the California Institute of Technology urban-scale model (Martien et al., 2006; Martien and Harley, 2006), and DRAIS (Nester and Panitz, 2006). The adjoint of the regional model STEM also has been developed (Sandu et al., 2005a) and deployed (Hakami et al., 2005, 2006; Chai et al., 2006).

Of all the previous 3-D adjoint modeling studies, none includes detailed treatment of aerosols, likely owing to the difficult prospect of deriving the adjoint of the model routines dealing with aerosol thermodynamics. The study of Hakami et al. (2005) deals only with inert carbonaceous aerosols, and the work of Dubovik et al. (2004), though global in scale, does not include full chemistry or aerosol thermodynamics. Detailed adjoint modeling of aerosols began with the theoretical investigations of Henze et al. (2004) and Sandu et al. (2005b). However, these are preliminary studies performed on idealized box model systems. In the current work we present the first adjoint of a global CTM that includes dynamics, full tropospheric chemistry, heterogeneous chemistry, and aerosol thermodynamics. We demonstrate the potential value of this tool for quantifying and constraining the many factors that govern global secondary inorganic aerosol formation. In addition, we note the general usefulness of the adjoint model of GEOS-Chem for a wide variety of applications, such as constraining CO emissions...
2 Forward and inverse models

The GEOS-Chem model is used to simulate global aerosol distributions (version 6.02.05 with a horizontal resolution of 4° × 5° and 30 layers up to 0.01 hPa, GEOS-3 meteorological fields). This version of the model includes detailed gas-phase chemistry coupled with heterogeneous reactions, inorganic aerosol thermodynamics, and oxidative aging of carbonaceous aerosols (Park et al., 2004). A few of the specific equations for various model processes are given in Sect. 3.3, along with their corresponding adjoints. We note here that gaseous SO\textsubscript{2} and primary sulfate are co-emitted in GEOS-Chem using a single emissions inventory, referred to as SO\textsubscript{x}, which is partitioned between the two species on a regional basis, with sulfate comprising 5% of SO\textsubscript{x} emissions in Europe, 1.7% in North America, and 3% elsewhere (Chin et al., 2000).

The standard model has been modified to facilitate the specific inverse modeling goals of the present study. We neglect stratospheric chemistry, which over the course of the short simulations considered here should not have a substantial impact. The standard GEOS-Chem tropospheric chemical mechanism comprises 87 species and 307 reactions integrated using the SMVGearII solver of Jacobson (1995). We retain this standard chemical mechanism; however, we implement a different numerical solver. The details of this are given in Appendix A. To summarize, we implement a 3rd order Rosenbrock solver that not only facilitates construction of the adjoint model, but also improves forward model efficiency. We also consider employing a lower order advection solver and using offline concentrations of sulfate aerosol for calculation of photolysis rates and heterogeneous reaction probabilities, see Sect. 3.5.1.


2.1 Inverse modeling

An adjoint model is used to calculate the gradient of a cost function, J, with respect to a set of model parameters, p, ∇\textsubscript{p}J. For data assimilation applications, the cost function is defined to be

\[
J = \frac{1}{2} \sum_{\mathbf{c} \in \Omega} (\mathbf{c} - \mathbf{c}_{\text{obs}})^T \mathbf{S}_{\text{obs}}^{-1} (\mathbf{c} - \mathbf{c}_{\text{obs}}) + \frac{1}{2} \mathbf{r}^T \mathbf{S}_p^{-1} \mathbf{r}
\]

where \( \mathbf{c} \) is the vector of species concentrations, \( \mathbf{c}_{\text{obs}} \) is the vector of species observations mapped to the model domain space, \( \mathbf{S}_{\text{obs}} \) is the observation error covariance matrix, \( \mathbf{p} \) is a vector of active model parameters throughout the model domain, \( \mathbf{p}_a \) is the initial estimate of these parameters, \( \mathbf{S}_p \) is the error covariance estimate of these parameters, \( \gamma \) is a regularization parameter, and \( \Omega \) is the domain (in time and space) over which observations and model predictions are available. We will sometimes use the notation \( c \) and \( p \) to represent single elements of the vectors \( \mathbf{c} \) and \( \mathbf{p} \). Using the variational approach, the gradient ∇\textsubscript{p}J is supplied to an optimization routine and the minimum of the cost function is sought iteratively. At each iteration, improved estimates of the model parameters are implemented and the forward model solution is recalculated. In this study, the magnitude of each variable parameter is actually adjusted using a scaling factor, \( \sigma \), such that \( \mathbf{p} = \sigma \mathbf{p}_a \). We use the L-BFGS-B optimization routine (Byrd et al., 1995; Zhu et al., 1994), which affords bounded minimization, ensuring positive values for the scaling factors.

Alternatively, for sensitivity analysis, the cost function can be defined as simply a set of model predictions,

\[
J = \sum_{\mathbf{c} \in \Omega} g(\mathbf{c})
\]

where \( \Omega \) is the set of times at which the cost function is evaluated. The desired gradient values are the sensitivities of this set of model predictions to the model parameters.
2.2 Adjoint modeling

Equations for calculating the desired gradient using the adjoint method can be derived from the equations governing the forward model or from the forward model code. The prior approach leads to the continuous adjoint, while the latter leads to the discrete adjoint (Giles and Pierce, 2000). The continuous adjoint equations for CTMs have been derived previously, using methods based upon the Lagrange duality condition (Vukicevic and Hess, 2000; Pudykiewicz, 1998; Schmidt and Martin, 2003) or Lagrange multipliers (Elbern et al., 1997). Consideration of the continuous adjoint equations is often useful for interpreting the significance of the adjoint values. Many previous studies have also described the derivation of discrete adjoints of such systems (Sandu et al., 2005a; Muller and Stavrakou, 2005). An advantage of the discrete adjoint model is that the resulting gradients of the numerical cost function are exact, even for nonlinear or iterative algorithms. Furthermore, portions of the discrete adjoint code can often be generated directly from the forward code with the aid of automatic differentiation tools. Continuous adjoint gradients may differ from the actual numerical gradients of $J$, and continuous adjoint equations (and requisite boundary/initial conditions) for some systems are not always readily derivable. Here we present a brief description of the discrete adjoint method for the sake of defining a self-consistent set of notation for this particular paper; we refer the reader to the cited works for further derivations and discussions of continuous and discrete adjoints.

The GEOS-Chem model can be viewed as a numerical operator, $F$, acting on a state vector, $c$:

$$c^{n+1} = F(c^n)$$

where $c$ is the vector of all $K$ tracer concentrations, $c^n = [c_1^n, c_2^n, \ldots, c_K^n]^T$ at step $n$. In practice, $F$ comprises many individual operators representing various physical processes. For the moment we will simply say that $F$ represents a portion of the discrete forward model which advances the model state vector from step $n$ to step $n+1$.

For simplicity, consider a cost function evaluated only at the final time step $N$ with no penalty term. We wish to calculate the gradient of the cost function $J$ with respect to the model state vector $c$ at any step in the model,

$$\nabla_c J = \frac{\partial J(c^N)}{\partial c^n}$$

We define the local Jacobian around any given step as

$$\frac{\partial c^{n+1}}{\partial c^n} = \frac{\partial F(c^n)}{\partial c^n} = F_c^n$$

Using the chain rule, we can expand the right hand side of the above equation to explicitly show the calculation of $c^{n+1}$ from $c^n$,

$$\nabla_c J = (F_c^n)^T (F_c^{n+1})^T \cdots (F_c^{N-1})^T \frac{\partial J(c^N)}{\partial c^N}$$

Evaluating the above equation from left to right corresponds to a forward sensitivity calculation, while evaluating from right to left corresponds to an adjoint calculation. When $K$ is larger than the dimension of $J$, which in this case is a scalar, the adjoint calculation is much more efficient (Giering and Kaminski, 1998).

For the adjoint calculation, we define the adjoint state variable $\lambda_c^n$,

$$\lambda_c^n = \frac{\partial J(c^N)}{\partial c^n}$$

This can also be expanded,

$$\lambda_c^n = \left[ \frac{\partial c^{n+1}}{\partial c^n} \right]^T \frac{\partial J(c^N)}{\partial c^{n+1}}$$

$$= (F_c^n)^T \frac{\partial J(c^N)}{\partial c^{n+1}}.$$
The equation above suggests how to solve for the adjoint variable iteratively. Initializing the adjoint variable at the final time step

$$\lambda_{c}^{N} = \frac{\partial J(c^{N})}{\partial c}$$  

we solve the following equation iteratively from \(n=N, \ldots, 1,\)

$$\lambda_{c}^{n-1} = (F_{c}^{n})^{T} \lambda_{c}^{n}$$  

The value of \(\lambda_{c}^{0}\) is then the sensitivity of the cost function with respect to the model initial conditions,

$$\lambda_{c}^{0} = \nabla_{c}J$$  

The scheme above shows why calculating the adjoint variable is often referred to as “reverse integration” of the forward model, as we step from the final time to the initial time. This should not be confused with simply integrating the forward model equations backwards in time.

In order to calculate the sensitivity of \(J\) with respect to other model parameters such as emissions, a similar analysis (Sandu et al., 2003) shows that the gradient of the cost function with respect to these parameters,

$$\lambda_{p}^{0} = \nabla_{p}J$$  

can be found by iteratively solving the following equation,

$$\lambda_{p}^{n-1} = (F_{p}^{n})^{T} \lambda_{c}^{n} + \lambda_{p}^{n}$$  

where the subscripts \(c\) and \(p\) indicate sensitivity with respect to \(c\) and \(p\), respectively, and

$$F_{p}^{n} = \frac{\partial F^{n}}{\partial p}$$

When a penalty term is included in the cost function, the gradient becomes

$$\nabla_{p}J = \lambda_{p}^{0} + \gamma S_{p}^{-1}(p - p_{a})$$

3 Constructing and validating the adjoint of GEOS-Chem

Here we present the derivation of the adjoint of GEOS-Chem. Whenever feasible, the adjoint model is based upon the discrete formulation, using automatic differentiation tools for assistance. We use the freeware multipurpose program TAMC (Giering and Kaminski, 1998), and KPP, a public domain numerical library for constructing the adjoint of chemical mechanisms (Sandu et al., 2003; Daescu et al., 2003). Always some, if not significant, manual manipulation of the code is required to use such tools. We often combine automatically generated adjoint code with manually derived discrete adjoint code to improve efficiency and transparency of the adjoint model.

Validation of the adjoint model is an important part of introducing an adjoint model of this size and complexity. As a test of a newly constructed adjoint model, many studies present the results of a twin experiment, wherein pseudo-observations are generated using the forward model itself, allowing for formulation of an inverse problem for which the correct solution is known (Fisher and Lary, 1995; Elbern et al., 1997; Errera and Fonteyn, 2001). Such tests are useful for exploring the performance of the adjoint model in a data assimilation setting. Further verification is necessary to ensure that the gradients calculated using the adjoint model are accurate in and of themselves. It has been our experience that even when serious errors exist in the adjoint model, the optimization routine can often still converge reasonably well in the twin experiment framework, indicating that such performance, while necessary, is not sufficient evidence of proper adjoint model construction. For this reason, we spend a great deal of effort checking the implementation of our adjoint model by comparing the derivatives of each subprocess in the model, separately and collectively, calculated using the adjoint method to derivatives calculated using the one sided finite difference approximation,

$$\Lambda = \frac{J(\sigma + \delta \sigma) - J(\sigma)}{\delta \sigma}$$

We use \(\delta \sigma = 0.1–0.01\) for most tests, and have checked that values of \(\Lambda\) do not differ
significantly from the more accurate, yet more computationally intensive, centerd finite difference gradients. For these validation tests, it suffices to use a simplified cost function that does not depend upon any observational data set, as in Eq. (3), defining $g$ to be a predicted tracer mass, either gas- or aerosol-phase, in a single grid cell, or the total mass burden over a larger spatial domain. For simplicity, most results are given in units of kg/grid cell or kg.

For large CTMs, it is not feasible to compare adjoint and finite difference gradients for every control parameter as the finite difference calculation requires an additional forward model evaluation per parameter; hence, usually only a representative sample of parameters is selected for validation. Many studies mention similar validation (Menut, 2003; Schmidt and Martin, 2003; Nester and Panitz, 2006), though only a few show actual comparisons (Mallet and Sportisse, 2004), while even fewer consider more sophisticated validations, such as comparing adjoint sensitivities to direct decoupled (DDM) sensitivities (Martien et al., 2006). Other possible methods for validating the adjoint code include checking the Lagrange condition (Giles and Pierce, 2000) and the complex variable method (Reuther et al., 1999), though these methods are used mostly in other fields.

3.1 Aerosol thermodynamics

The equilibrium thermodynamic model MARS-A (Binkowski and Roselle, 2003) is used to calculate the partitioning of total ammonia and nitric acid between the aerosol and gas phases. While it is a relatively simple model compared to others such as SCAPE (Kim et al., 1993) or ISORROPIA (Nenes et al., 1998), the MARS-A model is still fairly complex. It uses an iterative algorithm to find equilibrium concentrations, considering two primary regimes defined by the ionic ratio of ammonium to sulfate and several sub-regimes defined by conditions such as relative humidity.

Several factors have historically prevented rigorous treatment of aerosol thermodynamics from inclusion in adjoint modeling studies of CTMs, or even adjoint studies of aerosol dynamics (Henze et al., 2004; Sandu et al., 2005b). Division of the possible thermodynamic states into distinct regimes causes many discontinuities in the derivatives, precluding easy derivation of continuous adjoint equations and raising doubts to the value of such sensitivities. Furthermore, several coding tactics often employed in these types of models render them intractable for direct treatment using automatic differentiation tools.

We develop the adjoint of MARS-A in pieces, separating the model into several subprograms, the adjoints of which are then created using TAMC. Tracking variables are added to the forward model routine to indicate which of these subroutines to call during the adjoint calculation. Initial unequilibrated concentrations at the beginning of each external time step are saved in checkpoint files during the forward calculation. Intermediate values are recalculated from these during the adjoint integration. This type of two-level checkpointing strategy has been shown to optimally balance storage, memory and CPU requirements (Griewank and Walther, 2000; Sandu et al., 2005a).

The accuracy of the resulting adjoint code is tested by comparing adjoint gradients to finite difference gradients. These comparisons can be made directly throughout the entire model domain by turning off all transport processes. Figure 1 shows comparisons for the sensitivity of surface level nitrate aerosol mass with respect to scaling factors for emissions of surface level anthropogenic SO$_x$ and NH$_3$ after a week-long simulation. The gradients agree quite well, confirming the accuracy of the thermodynamic adjoint code. Discussion of the values of model sensitivities is given in Sect. 4.

3.2 Chemistry

KPP (v2.2) (Sandu et al., 2003; Daescu et al., 2003) is used to automatically generate code for the adjoint of the tropospheric chemistry solver, which calculates derivatives with respect to the initial species concentrations. We are also interested in the gradient with respect to the emission rates for those species whose emissions are incorporated into the chemical mechanism itself, such as NO$_x$. (as opposed to those that are simply injected into the model grid cells at intermediate times, such as SO$_x$). Though not a standard feature, KPP does provide the necessary subroutines for this task. The equa-
to demonstrate the accuracy of the adjoints of the chemistry routine, we calculate the derivative of the species concentrations at the end of a single chemistry time step (1 h) with respect to the emissions of NO\textsubscript{x} (emitted as NO) in a box model test. For this test, the chemical environment is that of a polluted, urban grid cell in the afternoon, the concentrations are in units of molecule cm\textsuperscript{-3}, and the adjoint sensitivities are with respect to the emissions themselves (molecule cm\textsuperscript{-3} s\textsuperscript{-1}); hence, \(\lambda\) has units of s\textsuperscript{-1}. Table 1 gives the initial \((c(\tau=0))\) and final \((c(\tau=1\text{ h}))\) species concentrations, the value of the adjoint derivatives \((\lambda_{\text{ENO}})\), the value of the same derivatives calculated using the finite difference method \((\lambda_{\text{ENO},0})\) and the ratio \(\lambda_{\text{ENO}}/\lambda_{\text{ENO,0}}\). A perturbation \(\delta_{\text{ENO}}=0.1\) was used for these tests. All adjoint derivatives are in good agreement with the finite difference derivatives.

The above test was reassuring, yet limited in scope for a global CTM. To test our adjoint model over a wide variety of chemical conditions, we also compare the accuracy of the adjoint derivatives of the chemical mechanism in global simulations over much longer time scales. We turn off all transport related processes in the model and calculate the adjoint and finite difference derivatives of surface level tracer masses with respect to NO\textsubscript{x} emissions in each location after a week-long simulation. Many chemical changes associated with aerosols are treated separately from the main tropospheric chemistry mechanism in GEOS-Chem, such as aqueous reactions, dry deposition, chemical aging, and emission of SO\textsubscript{2} and NH\textsubscript{3} (Park et al., 2004). The adjoints of these processes are constructed separately (manually and with TAMC) and included in the following tests.

Figure 2 shows the adjoint and finite difference sensitivities of several species with respect to surface level, anthropogenic emissions of NO\textsubscript{x}. We choose to show sensitivities of species such as acetone and methacrolein to NO\textsubscript{x} emissions to highlight the potential value of the adjoint model for analysis of non-aerosol species. We see from these, and similar tests for other active species (not shown), that the sensitivities cal-

culated using the adjoint model consistently agree with those using the finite difference method over a wide range of conditions.

The code generated by KPP allows computation of either the continuous or discrete adjoints of the chemical mechanism. The continuous adjoint equation can be solved faster than the discrete adjoint equation at a given tolerance level, as calculation of the latter requires recalculation of intermediate values from the forward integration and computation of the Hessian during the adjoint integration, see Appendix B. At tight tolerance levels (i.e. very small internal time steps), the results of these methods should converge. However, for tolerance levels appropriate for global modeling, the continuous adjoint is only approximate, as \(\lambda+\delta\lambda\), where \(||\delta\lambda||\ll C-Tol\). Given that the computational expense of the Rosenbrock solver increases substantially for tighter tolerance levels (see Appendix A), it is more efficient to use the discrete adjoint, even though this requires an additional forward integration. This is in contrast to the approach of Errera and Fonteyn (2001), who chose to approximate the necessary intermediate values by linearly interpolating from values stored at each external time step, an approach likely more appropriate for their stratospheric chemistry application.

GEOS-Chem accounts for the effect of aerosol concentrations on the radiation available for photolysis reactions and on the available surface area for the heterogeneous reactions included in the main chemical mechanism. The influence of the concentration of sulfate-ammonium-nitrate aerosols on such rates is not currently accounted for in the adjoint model. We assume such an effect is less than 5\% (Liao et al., 1999; Martin et al., 2003), especially as the absorbing aerosols (black carbon, mineral dust) are not active variables during these tests. The general agreement between \(\lambda\) and \(\Lambda\), only the latter of which accounts for this effect, indicates this assumption is adequate, at least for simulations of this length. Further tests indicate that this assumption is valid for most, though not all, cases, see Sect. 3.5.1.
3.3 Convection, turbulent mixing, and wet removal

Wet removal of tracers in GEOS-Chem is generally treated as a first-order process, leading to discrete forward model equations of the form,

\[ c_k^{n+1} = c_k^n e^{-r_{w,k} \Delta t} \]  

(17)

Since the loss rate \( r_{w,k} \) for most species does not depend upon any active variables (Jacob et al., 2000), the corresponding adjoint is simply

\[ \lambda_k^n = \lambda_k^{n+1} e^{-r_{w,k} \Delta t} \]  

(18)

The adjoints of these routines are generated using hand-created code, retaining efficiency and legibility. However, the in-cloud formation and cycling of sulfate aerosol from SO\(_2\) is decidedly nonlinear, as the soluble fraction of SO\(_2\) is limited by availability of H\(_2\)O, and a fraction of the SO\(_2\) is reintroduced into the gas phase as sulfate when droplets evaporate (Park et al., 2004). Such nonlinearities that span multiple program modules are treated both manually and with the help of TAMC, requiring additional recalculation and checkpointing of intermediate values.

Turbulent mixing in the boundary layer in the forward model is calculated according to a mass-weighted mixing algorithm applied every dynamic time step (30 min for our case),

\[ \mu_{k,j}^{n+1} = \sum_{l=1}^{L} \frac{m_j \mu_{k,l}^n}{m_T} \]  

(19)

where \( \mu_{k,j} \) is the mixing ratio \((c/\rho, \rho \text{ is the density of air})\) of tracer \( k \) in layer \( j \), \( m_j \) is the air mass in a single layer \( l \), \( m_T \) is the total air mass in the boundary layer column, and \( L \) is the number of layers in the boundary layer. Rewritten in matrix form, this equation reads,

\[
\begin{bmatrix}
\mu_{k,1} \\
\vdots \\
\mu_{k,L}
\end{bmatrix}^{n+1} = 
\begin{bmatrix}
m_1 & \cdots & m_L \\
\vdots & \ddots & \vdots \\
m_1 & \cdots & m_L
\end{bmatrix}^{-1} 
\begin{bmatrix}
\mu_{k,1} \\
\vdots \\
\mu_{k,L}
\end{bmatrix}^n
\]

(20)

Direct application of Eq. (11) yields the corresponding adjoint equation,

\[
\begin{bmatrix}
\lambda_{\mu_{k,1}} \\
\vdots \\
\lambda_{\mu_{k,L}}
\end{bmatrix}^n = 
\begin{bmatrix}
m_1 & \cdots & m_L \\
\vdots & \ddots & \vdots \\
m_1 & \cdots & m_L
\end{bmatrix}^{-1} 
\begin{bmatrix}
\lambda_{\mu_{k,1}} \\
\vdots \\
\lambda_{\mu_{k,L}}
\end{bmatrix}^{n+1}
\]

(21)

which can be simply written as,

\[ \lambda_{\mu_{k,j}}^n = \frac{m_j \sum_{l=1}^{L} \lambda_{\mu_{k,l}}^{n+1}}{m_T} \]  

(22)

Deep convection is calculated in the forward model using cumulus cloud fluxes and an RAS type algorithm, see Appendix A of Allen et al. (1996). We calculate the discrete adjoint of this scheme using TAMC, noting that TAMC initially generates code that is accurate, yet several orders of magnitude slower than necessary due to several superfluous loops that have to be removed manually. The numerical scheme for the forward calculation iteratively solves a set of essentially linear equations, with an internal time step of five minutes. If we neglect a single conditional statement that checks only for rare floating point exceptions, then storage or recalculation of the intermediate values is not required for the adjoint calculation.

The adjoint model performance for a simulation including convection, turbulent mixing, and wet deposition is tested by comparison of finite difference derivatives to the adjoint derivatives of concentrations of a soluble tracer with respect to its initial concentrations in a location exhibiting strong convection, deposition, and mixing. Horizontal
transport, chemistry, and aerosol thermodynamics are turned off for these tests. We use a perturbation of one percent for the finite difference calculation. The ratio $\lambda_c/\Lambda_c$ for simulations that are 6 h, 1 d and 3 d in length are 0.9998, 1.0002 and 1.0003, from which we see consistent satisfactory agreement between the two methods. Performance is similar in other tested locations.

3.4 Advection

For reasons of practicality, we implement an adjoint of the continuous advection equations, rather than the discrete advection algorithm, as the latter is not directly amenable for use with automatic differentiation tools, nor is the derivation of discrete adjoints of advection routines the focus of this work. Below we explain the derivation of the continuous adjoint equation and afterwards address some of the issues that are wedded to this approach.

We consider the 1-D example of the advection equation for a tracer in mass concentration units,

$$\frac{\partial c}{\partial t} = -\frac{\partial uc}{\partial x}$$ (23)

where $u$ is the wind velocity in the $x$-direction. The forward numerical model actually solves the flux form of Eq. (23) in terms of the mixing ratio (Lin and Rood, 1996),

$$\frac{\partial (\rho \mu)}{\partial t} = -\frac{\partial (\rho \mu u)}{\partial x}$$ (24)

Assuming that the continuity equation for $\rho$ is satisfied, this can be rewritten in the advection form,

$$\frac{\partial \mu}{\partial t} = -u \frac{\partial \mu}{\partial x}$$ (25)

The continuous adjoint of Eq. (25) is

$$-\frac{\partial (\lambda \mu)}{\partial t} = \frac{\partial (\lambda \mu u)}{\partial x}$$ (26)

where $\lambda$ is the adjoint of the mixing ratio. Note that we have assumed that the winds (or any other met fields) are not active variables; taking the adjoint with respect to the meteorology is another task in itself (Giering et al., 2004). Applying the simple transform $\tilde{\lambda}_\mu = \lambda \mu / \rho$, and substituting this into Eq. (26), we arrive at the following adjoint equation,

$$-\frac{\partial (\rho \tilde{\lambda}_\mu)}{\partial t} = \frac{\partial (\rho \tilde{\lambda}_\mu u)}{\partial x}$$ (27)

which is similar in form to Eq. (24).

If we assume that $\rho$ is relatively constant over a single dynamic time step and that the advection is linear, then we can simply solve Eq. (27) using the same numerical code that was used to solve Eq. (24) in the forward model, scaling the adjoint by $1/\rho$ before and re-scaling by $\rho$ afterwards, which is equivalent to solving Eq. (26).

In practice, nonlinearities are often introduced into numerical advection schemes to provide stability, positive definiteness, and to maintain monotonic behavior, leading to considerable differences between the discrete and continuous adjoints. The discrete adjoint of such schemes can lead to numerical modes in the adjoint solution (Sirkes and Tziperman, 1997). Also, for flux correcting, and even linear, advection schemes, the difference between the continuous and the discrete adjoints can be appreciable under certain conditions (Liu and Sandu, 2006$^2$). GEOS-Chem nominally employs a monotonic piecewise parabolic (PPM) advection routine (Lin and Rood, 1996), the discrete adjoint of which could possibly be subject to such numerical artifacts. On the other hand, the continuous adjoint will generate sensitivities that are consistent with the forward model equations, but not the numerical forward model itself. Several other adjoint modeling studies have investigated the trade-offs between various approaches (Vukicevic and Hess, 2000; Schmidt and Martin, 2003). We implement the continuous adjoint, and we assess how these issues affect our results in the following section.

3.5 Combined performance

3.5.1 Accuracy

Again we compare the gradients calculated using the adjoint model to those calculated using the finite difference method, this time including all model processes. We calculate the sensitivity of global aerosol distributions of sulfate, ammonium, and nitrate to surface emissions of anthropogenic SO$_2$, NO$_x$ and NH$_3$ in select locations. As noted previously, such comparisons are quite time consuming to perform on a global scale owing to the expense of the finite difference calculations. Attempting to cover a wide range of conditions, while keeping the number of required calculations within reason, we choose to analyze ten locations for each set of emissions considered, see Fig. 3. The simulations are one day in length, and the cost function (Eq. 2) is evaluated only once at the end of the day. We use a perturbation of $\delta \sigma = 0.1$ for the finite-difference calculations.

Figure 4 shows the adjoint gradients compared to the finite difference gradients for each of nine relationships. From visual inspection of the scatter plots, it is clear that the agreement is generally within reason given the fact that using a continuous adjoint for advection is expected to cause some amount of discrepancy. Regression lines, slopes, and $R^2$ values are given for each set of comparisons. The absolute difference between the two methods is often more substantial for the larger values. As the gradients in a given set usually span several orders of magnitude, many of the slopes are biased by a few such larger values and are not representative of the overall fit. However, accounting for such heteroscedasticity by re-scaling the gradients by $1/p$ or performing weighted regressions that place less emphasis on the larger values still leads to the same general results. Picking twice as many test cells, different test cells, or a different value of $\delta \sigma$ also was not found to substantially alter the overall comparisons.

Initial comparison (not shown) of gradients for five of the 90 tests showed underestimation of adjoint sensitivities by more than an order of magnitude. Four of these tests were for the sensitivity of sulfate with respect to NH$_3$ emissions while one was for the sensitivity of nitrate with respect to SO$_x$ emissions. Using offline concentrations for calculation of the contribution of sulfate aerosol to photolysis rates and heterogeneous reaction probabilities in the main tropospheric chemical mechanism for these tests alleviated the discrepancy, demonstrating that while this feedback is generally negligible, it is occasionally quite strong. Future work will extend the adjoint model to account for this feedback.

Napelenok et al. (2006) performed a complementary analysis, calculating the sensitivities of local aerosol distributions with respect to domain-wide precursor emissions over the United States with a forward sensitivity method (DDM-3D), using finite-difference calculations to check their results. While they found similarly good agreement for the more direct relationships (such as sensitivity of sulfate with respect to SO$_2$ emissions, or ammonium with respect to NH$_3$ emissions), they had difficulty verifying the variability in the sensitivities of some of the more indirect relationships (such as the sensitivity of sulfate to NH$_3$ emissions or nitrate to SO$_2$ emissions). Granted, they used the more complex and rigorous thermodynamic model ISORROPIA; they suggested that such discrepancies were due to numerical diffusion, with spatial oscillations of the sensitivities indicative of errors due to transport.

In our tests, transport does not drastically degrade the consistency of the correlation between the two approaches; all of the $R^2$ are near unity. There is, however, some amount of bias in the comparisons, as indicated by slopes ranging from 0.8 to 1.3, and this does appear to be a result of transport. Figure 5 contains scatter plots of the sensitivities of sulfate with respect to NO$_x$ emissions for several additional tests. Panel (a) shows the base case simulation described previously. Turning off advection in the model leads to improved agreement, $m=1.03$, see panel (b). Including advection, but evaluating the cost function only in a single location, rather than globally, leads to a very unsmooth adjoint field and triggers many nonlinear aspects of the numerical scheme in a manner inconsistent with advection of the relatively smooth concentration field in the forward model; hence, agreement between adjoint and finite difference gradients under these conditions is worse, see panel (c). For data assimilation, this type
of discrepancy could potentially be an impediment if only sparse or infrequent measurements are available. Panel (d) shows the results using global concentrations, but using a first order, upstream, linear transport scheme, for which it might be expected that discrepancies between the continuous adjoints and the numerical (finite difference) gradients would be diminished; however, the diffusive nature of such schemes actually increases the bias, \( m = 1.27 \).

All of the tests so far have been based on a single evaluation of the cost function at the end of a day long simulation. Alternatively, in accordance with the goals of a particular sensitivity study or the availability of observational data in an inverse modeling application, the cost function can be evaluated several times throughout the simulation. We will refer to the time between consecutive evaluations of the cost function as the assimilation window (though observations aren’t assimilated for sensitivity analysis). The effects of changing the assimilation window and the total simulation length are shown in Fig. 6. The base case is shown in panel (a), which is again the sensitivity of sulfate to NO\(_x\) emissions. Next we consider evaluating the cost function once every 6 h, rather than only once at the end of the day. Not surprisingly, decreasing the assimilation window to 6 h for a day-long simulation improves agreement, panel (b). Doubling both the simulation length and the assimilation window to two days degrades the agreement considerably, panel (c), while doubling only the simulation length but maintaining a one-day assimilation window improves the agreement, panel (d). Inaccuracies in the numerical gradients of the cost function as calculated with the adjoint model depend more upon the assimilation window than the total simulation length.

Overall, we find the accuracy of the adjoint gradients to be satisfactory. The adjoint model clearly captures the dependence of inorganic aerosol burdens on the chemical and thermodynamic interactions that lead to their formation. We have identified possible sources of bias in the adjoint gradients that may degrade their value as model sensitivities. This could be more critical for analysis of longer lived species whose distributions are dictated largely by transport. Such biases are less of a hindrance for data assimilation as the adjoint gradients still accurately capture the largest aspects of the variances, though a larger number of iterations may be required to minimize the cost function.

### 3.5.2 Computational efficiency

Here we report computational resource requirements for running the adjoint model of GEOS-Chem on a Linux workstation with dual Intel Itanium 1.5 GHz processors and 4 GB of RAM. The adjoint model utilizes multiple processors on shared memory architectures as efficiently as the forward model. It requires 16 KB of checkpoint storage space per simulated day per grid cell; this amounts to 11 GB of storage space per week with the current model configuration. This is comparable to the storage requirements of other adjoint models of CTMs such as STEM, 40 KB per day per cell (Sandu et al., 2005a), or the CIT model, 100 KB per day per cell (Martien et al., 2006), taking into account that the time step is 30 min in GEOS-Chem (for this study), and was 15 min for STEM, and 3 min for the CIT model. The computational cost of the adjoint model (backward only) of GEOS-Chem is 1.5 times that of the forward model, requiring 2.5 h for a week long iteration (forward and backward). Adjoint models of other CTMs report this ratio as: STEM: 1.5, CHIMERE: 3–4, IMAGES: 4, Polair: 4.5–7, CIT: 11.75. We see that the adjoint of GEOS-Chem is quite efficient; in general, adjoint codes that are derived by hand or use specialized tools such as KPP are most efficient. Such efficiency is the trade-off for the labor involved in manually constructing an adjoint model of this size and complexity.

### 4 Sensitivity analysis

In this section we briefly demonstrate how the adjoint model can be used as an efficient method of investigating the sensitivity of modeled aerosol concentrations to their precursor emissions. Many of the results shown previously in Sect. 3 contain interesting sensitivity information. For example, the box model tests of the chemistry adjoint...
were performed in a polluted urban grid cell. The high concentrations of pollutants leads to an inverse relationship between NO and other oxidants such as O$_3$, H$_2$O$_2$, and OH, hence, the value of $\lambda_{\text{NO}_x}$ is negative for these species (Table 1). One may have also noticed that the signs of the sensitivities of nitrate aerosol to emissions of SO$_x$ and NH$_3$ appear correlated in the tests of the thermodynamic adjoint, Fig. 1. The physical distribution of these two regimes is not itself significant as the artificial nature of a “thermodynamics only” simulation likely leads to nonphysical conditions. However, it is interesting to note such definitive shifts in the dependence of nitrate aerosol on emissions of SO$_x$ and NH$_3$. These sharp discontinuities arise from discretization of the sets of allowable species in the MARS-A algorithm, delineated by conditions such as the ammonium to sulfate ratio and relative humidity. The sensitivities of nitrate aerosol to both NH$_3$ and SO$_x$ appears to be largely negative in regions where nitrate aerosol dominates and positive in regions where nitric acid dominates (not shown). Hence, the regions of positive and negative sensitivities roughly correspond to those that are primarily sulfate-rich and sulfate-poor, respectively.

Sensitivity calculations for the full model are performed for a week-long simulation. Figure 7 shows the sensitivity of global burdens of sulfate, nitrate and ammonium aerosol to surface level emissions of anthropogenic SO$_x$, NO$_x$ and NH$_3$. The cost function is evaluated once daily. Other results retrieved from the same calculations (not shown) are sensitivities of these species with respect to the following emissions: stack SO$_x$, stack NO$_x$, biofuel SO$_x$, biomass burning SO$_x$, ship SO$_x$, biofuel NH$_3$, biomass burning NH$_3$, and natural NH$_3$.

The sensitivities in Fig. 7 encompass a wide range of relationships between aerosols and their primary precursors. Some of these relationships are practically intuitive, such as the sensitivities of sulfate to SO$_x$ emissions or of nitrate to NO$_x$ emissions, both of which are generally large and positive. The sensitivity of ammonium to emission of NH$_3$ is also positive, and the sensitivities of ammonium to SO$_x$ and NO$_x$ emissions are always positive, owing to uptake of NH$_3$ on inorganic aerosol by sulfate and nitrate.

Some of the relationships in Fig. 7 are less obvious, such as the negative sensi-

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tivity of sulfate to emissions of NH$_3$. This effect is smaller in magnitude than some of the others, because the relationship between NH$_3$ emissions and sulfate aerosol concentrations is less direct. As total sulfate is conserved in the MARS-A aerosol equili-

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brium model, this effect is not due to thermodynamic interactions between ammonium and sulfate. The only species directly affected by NH$_3$ or ammonium concentrations are nitrate and nitric acid, via thermodynamic interactions. Therefore, the relationship between NH$_3$ and sulfate is dictated by the interactions between sulfate and nitrate, and, hence, NO$_x$. The sensitivity of nitrate to SO$_x$ is largely negative, owing to ther-

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modynamic competition between nitrate and sulfate for ammonium. The sensitivity of nitrate to NH$_3$ is entirely positive, due to the necessary presence of excess NH$_3$ for HNO$_3$ to condense. The combination of these two effects explains the overall negative relationship between sulfate and emissions of NH$_3$.

Within the global trends noted above, there is also much discernible local variability. While explication of every feature is not within the scope of this work, there are some aspects worth addressing. For example, there are a few locations where the sensitivity of sulfate to NH$_3$ emissions changes abruptly from predominantly negative to locally positive. Some of these actually correspond to similarly abrupt shifts between areas that are sulfate-poor to areas that are sulfate-rich, such as the tip of South America and immediately west of the Iberian Peninsula. In-cloud formation of sulfate under certain conditions reinforces negative sensitivities of sulfate on NO$_x$ emissions. In other conditions or times of the day, emission of NO$_x$ can actually lead to a decrease in nitric acid, and, hence, nitrate. Though tempting to attribute these types of seemingly small anomalies to numerical artifacts of the adjoint model, these sudden shifts in sign are, at least, correctly representative of the discrete forward model. Further study is required to determine the physical significance of these results.

While the adjoint model accounts for nonlinearities in the relationships between emissions and aerosols, the results of the adjoint calculation are still merely tangent linear derivatives (gradients) which are likely to be valid over only a limited range of values for the parameters (emissions). We explore the robustness of the aerosol sensitivity calcu-
lations with respect to the magnitude of the emissions. Figure 8 shows the sensitivity of
nitrate with respect to NO\textsubscript{x} emissions calculated when the emissions are multiplied by
uniform scaling factors of 0.75 and 1.25; the relative differences between these values
and base case sensitivities shown in Fig. 7. The sensitivities can differ substantially on
a point to point basis (>50\%), particularly near boundaries between the positive and
negative sensitivities or in areas where the sensitivities are very small. The differences
are generally much less (<20\%) in areas with the largest sensitivities such as Europe,
Eastern Asia and the Eastern United States. Despite these relative differences, the
sensitivity field, viewed on the global (log) scale, remains nearly identical to the base
case values. While individual sensitivities may be valid only over a limited range, the
sensitivity field as a whole appears fairly robust.

Overall, the adjoint model is a promising tool for examining the dependence of aerosol
concentrations on emissions, taking into account effects of gas-phase chemistry,
heterogeneous reactions, and equilibrium thermodynamics. We note that the time
required to calculate all of these sensitivities was less than 10 times the cost of a single
forward model evaluation, while obtaining these results using the finite difference
method would have required >5000 times the cost of a forward run.

5 Inverse modeling tests

Several inverse modeling tests are performed to assess the capabilities of the adjoint
model in a data assimilation application. Using the twin experiment framework, pseudo
observations, \(c_{\text{obs}}\), are generated with the forward model using a base set of emissions
parameters, \(p=p_0\). An active subset of the parameters used to generate these
observations is then perturbed using scaling factors, \(p=\alpha p_0\). The inverse model uses
the pseudo-observations to recover the original unperturbed values of these active pa-
rameters.

We begin by generating a week-long set of observational data using the forward
model with all scaling factors set equal to unity. For these initial tests, we perturb one
set of emissions by re-scaling the emissions in every cell by a factor of two, and we use
observations in every grid cell once every 24 h to force the data assimilation. As there
is no error in these observations, equal weight is ascribed to each (\(S_{\text{obs}}^{-1}\) is the identity
matrix), and the error covariance of our initial (perturbed) estimate of the emissions
scaling factors is infinite (\(S_{\text{p}}^{-1}\) is zero). Such conditions are unrealistic and serve only
to test the adjoint model under the most ideal conditions possible.

In the first set of tests (DA1), we perturb the emission inventories of (a) surface
level anthropogenic SO\textsubscript{2}, (b) biomass burning SO\textsubscript{2} and (c) biofuel SO\textsubscript{2}. We assimilate
observations of sulfate for the week of 1–7 July 2001. Figure 9 shows the progression
of the normalized (divided by the initial value) cost function at iteration \(i\) during the
optimization procedure, \(J_i/J_1\). The cost function quickly reduces by at least five orders
of magnitude in each case. The correct emissions inventories are essentially entirely
recovered.

In the next test (DA2), we perturb the emission inventory of NH\textsubscript{3} from anthropogenic
sources, and assimilate observations of aerosol ammonium. This is a slightly more
difficult inversion as ammonium measurements alone do not fully constrain NH\textsubscript{3} emis-
sions (Gilliland et al., 2006). As demonstrated in Sect. 3.5.1, ammonium is indirectly,
yet appreciably, coupled to gas-phase oxidants. Utilizing observations of O\textsubscript{3}, O\textsubscript{2}, NO\textsubscript{2}
and NO\textsubscript{3} in conjunction with ammonium observations noticeably increases the con-
vergence rate over using either type of observations alone, see Fig. 10. This demonstra-
tes, albeit in a highly idealized fashion, the potential for exploiting multi-phase mea-
surements as constraints for aerosol modeling.

The final test (DA3) attempts to mimic a slightly more realistic scenario than the pre-
tvious tests: improving estimates of global anthropogenic SO\textsubscript{2} and NO\textsubscript{2} emission in-
tories using surface measurements of sulfate, nitrate, and ammonium aerosol. In this
case, the emissions inventories are perturbed regionally by 5–30\% with an additional
random factor of order 5\%. For example, the anthropogenic SO\textsubscript{2} and NO\textsubscript{2} emissions in
North America are perturbed by factors of 0.8\(+r\) and 0.85\(+r\), respectively, while emis-
sions in Asia are perturbed by factors of 1.2\(+r\) and 1.3\(+r\), where \(r\) is a random number
uniformly distributed between 0 and 0.05. The error covariance matrix \( S_p \) is calculated using an ascribed error of 100% and is assumed to be diagonal. Observations are used once per day in only half of the land-based surface grid cells. The reduction of the cost function after 15 iterations is shown in Fig. 11. The difference between the true emission inventories for \( \text{SO}_x \) and \( \text{NO}_x \) and the estimated inventory at the first and final iterations are shown in Fig. 12. While the cost function has reduced by nearly two orders of magnitude, the optimization procedure has clearly yet to reach a minimum. In applications of this type, the procedure is often halted according to an appropriate convergence criteria. While there are substantial improvements in the \( \text{SO}_x \) emissions and the \( \text{NO}_x \) emissions in Europe and Asia, the \( \text{NO}_x \) emissions in North America have yet to converge. Further iterations might be justified; however, this would risk over-optimization in other areas, where we begin to see oscillatory behavior in the solution (deviations switching from positive to negative from iteration to iteration).

6 Summary and conclusions

The derivation of the adjoint model of GEOS-Chem has been presented in a piecewise fashion. We have implemented the first adjoint of an aerosol equilibrium thermodynamic model (MARS-A, Binkowski and Roselle, 2003), derived using the automatic differentiation tool TAMC (Giering and Kaminski, 1998), which required significant manual pre- and post-processing owing to the structure and complexity of the code. To facilitate construction of the adjoint of the GEOS-Chem gas-phase chemical mechanism, we implemented a Rosenbrock solver using the KPP numerical library (Sandu et al., 2003). This has allowed for automatic generation of the adjoint of the chemical mechanism and also improved forward model performance (see Appendix A). The adjoints of wet removal, deep convection, and turbulent mixing were derived manually and with the aid of TAMC. We have used the continuous adjoint method to treat advection, wherein the same numerical algorithm is used to solve the continuous adjoint advection equation as was used for tracer advection in the forward model.

All aspects of the adjoint model have been tested both separately and together by comparing the adjoint gradients to finite difference gradients. Each individual discrete adjoint routine showed satisfactory performance over a wide range of conditions. The adjoint gradients of the cost function evaluated using the full model are well correlated with the numerical gradients, as measured using finite difference calculations, with most \( R^2 > 0.95 \). However, depending upon the nature and frequency of the analysis points, they can exhibit appreciable bias (10–30%) owing to the fact that the continuous adjoint of the advection scheme is not an exact representation of the numerical model sensitivities (Sirkes and Tziperman, 1997; Liu and Sandu, 2006). Overall, the hybrid approach adopted here is not ideal, but appears not to overly degrade the usefulness of the adjoint model. Nevertheless, this treatment necessitated additional inspection of the inverse model performance, wherein the effect of the frequency and distribution of measurements on the accuracy of the model sensitivities calculated using the adjoint model were shown to depend primarily on the smoothness of the observation field and the length of the assimilation window.

The adjoint model clearly demonstrates the importance and relative strengths of many complex nonlinear relationships connecting concentrations of aerosol species and their precursor emissions. Though indirect, relationships such as the dependence of sulfate aerosol concentrations on emission of \( \text{NH}_3 \) or \( \text{NO}_x \) are captured by the adjoint model, and can be determined globally in an efficient manner. The sign and magnitude of many of these sensitivities exhibit a rich array of features owing to the influence of environmental factors such as the sulfate to ammonium ratio, cloud processing of \( \text{SO}_2 \), and variability in the \( \text{NO}_x \) and \( \text{O}_3 \) levels.

We have also demonstrated the capabilities of the adjoint model in mock data assimilation applications. An adjoint model of this type allows for the possibility of exploiting multi-phase observations to constrain emissions of aerosol precursors. Here we have focused on regional variability of the emissions inventories, though the emissions can also be adjusted on a temporal basis. For real data assimilation projects, many application specific issues inherent in this type of inverse modeling have yet to be resolved,
such as specification of the error covariance matrices $S_{obs}$ and $S_p$. The dependance of adjoint model performance is known to depend strongly on such factors (Chai et al., 2006), proper formulation of which is necessary to ensure scaling of the inventories that are physically realistic (Stavrakou and Muller, 2006). Real world application will also likely require conditioning of the cost function to improve convergence rate (Meirink et al., 2006) and tuning of the regularization parameter (Hakami et al., 2005). Subsequent studies will focus on expanding the adjoint model to capture feedbacks such as the effect of sulfate aerosol concentrations on photolysis rates and heterogeneous reaction probabilities, seen here to occasionally be quite important. Work on the adjoint of the aerosol equilibrium model ISORROPIA (Nenes et al., 1998) is also in progress. Further application of the GEOS-Chem model will focus also on the exploitation of multi-phase measurements from sources such as surface stations, aircraft, and satellites as model constraints. The adjoint of GEOS-Chem has already been used to constrain emissions of carbon monoxide from Asia using satellite (MOPITT) measurements (Kopacz et al., 2006), demonstrating the potential for addressing a wide range of scientific questions with this type of inverse model.

Appendix A

Implementation of a Rosenbrock solver and comparison to SMVGEARII

Solving large systems of chemical rate equations in CTMs requires the use of special numerical tools, or solvers, that are specifically designed for this purpose. Taking the adjoint of such solvers manually, or using generic automatic differentiation tools, can be an onerous task. We desire to create the adjoint of the full chemical mechanism in GEOS-Chem using the KPP software library (Sandu et al., 2003), which is a set of tools specifically built for automatic differentiation of chemical mechanisms and the numerical algorithms used to solve these systems. In order to make use of these tools, we must first implement the KPP generated numerical integration routines in the forward model. We investigate the feasibility and ramifications of replacing the current solver in GEOS-Chem, SMVGEARII (Jacobson, 1995), with a KPP generated Rosenbrock solver. We consider the amount of work required to make such a switch, the efficiency of the Rosenbrock solver compared to the SMVGEARII solver, and the overall effect that such a switch has on the model predictions after a week-long simulation.

After manually transposing the SMVGEARII mechanism input files to KPP input files, the KPP tools easily generate a set of Fortran code that solves the given system for a variety of supported Rosenbrock type integrators in a box model setting. Minimal manual adjustment to this code was required to interface with the 3-D GEOS-Chem model and to allow support for OpenMP parallelization. Some amount of modifications to the KPP code itself will be required to fully automate this process.

Next we consider the efficiency of the Rosenbrock solver and the SMVGEARII solver in a global simulation with only chemistry. For each species, in every cell, we compare the concentrations from benchmark solutions at the end of a day-long simulation to concentrations from a reference solution for each solver. The benchmark calculations span a set of tolerance levels $\{10^{-1} \leq RTOL \leq 10^{-5}, 10^3$ molecules cm$^{-3} \geq ATOL \geq 10^{-2}$ molecules cm$^{-3}\}$ while the reference solutions were computed using tight tolerances ($RTOL=10^{-8}, ATOL=10^2$ molecules cm$^{-3}$). Looser tolerance levels result in repeated failure to converge in numerous grid cells.

To assess the accuracy of the two methods, following Sandu et al. (1997) we define the significant digits of accuracy (SDA) as

$$\text{SDA} = - \log_{10}(\max_k ER_k)$$

where $ER_k$ is a spatially modified root mean square norm of the relative error of the benchmark solution ($\hat{c}_{k,j}$) with respect to a reference solution ($c_{k,j}$) for species $k$ in
grid cell \( j \),

\[
ER_k = \frac{1}{|\theta_k|} \sum_{j \in \theta_k} \left| \frac{c_{k,j} - \hat{c}_{k,j}}{c_{k,j}} \right|^2
\]

For \( \Theta \) total grid cells, \( \theta_k \) is the set of all locations of significant concentrations of species \( k \), \( \{0 \leq \theta \leq \Theta : c_{k,j} \geq a\} \). A threshold value of \( a = 10^6 \) molecules cm\(^{-3}\) is chosen to avoid inclusion of errors from locations where concentrations of a given species are less than chemically meaningful values.

We present the results in the form of a work – precision diagram, wherein the value of SDA for each test is plotted versus the average computational expense for the solver to integrate the chemical mechanism for one hour. When calculating this average, we do not consider the time required during the initial six hours of the simulation, as each solver requires a bit of “spin up” time in order to adjust internal time steps to values more appropriate than the default starting step size according to the stiffness of the local system. Such spin up time is negligible with respect to the total computational cost of any simulation longer than a few days.

Figure A1 shows the work-precision diagram for the global benchmark simulations. The Rosenbrock solver is nearly twice as efficient as the SMVGEARII solver during these tests. Based on this analysis, we choose to run the Rosenbrock solver at tolerance levels that yield an SDA of \( \sim 1.0 \) as the standard setting for this work.

For practical applications, we are interested in the difference in the total model predictions, including all model processes, incurred by switching to the Rosenbrock solver. We compare the daily average concentrations after a week-long simulation, including all model processes, calculated using the new standard Rosenbrock settings versus the standard SMVGEARII settings. Figure A2 shows the values of \( ER_k \) for each species \( k \) using the Rosenbrock solver to generate the test solution and SMVGEARII for the reference solution. This figure shows that after switching to this Rosenbrock solver, the solution is changed by less than 10% for most species. The difference is larger, between 10 and 15%, for HNO\(_2\), HNO\(_4\), IAP, INO\(_2\), ISOP, N\(_2\)O\(_5\), NO, NO\(_2\), PP, and RIP.

Determining whether or not this is an actual improvement in the accuracy of the forward model itself would require further comparison to observations. At the very least, the switch results an improvement in the numerical solution of the forward model equations for slightly less computational cost.

Overall, while a more detailed analysis (requiring optimization of specific species tolerance levels and the parameters that control internal step size expansion and contraction) is necessary to determine unequivocally which method is more efficient, in our experience, not only is the Rosenbrock method desirable because of its differentiability, but it also appears to improve forward model performance by providing more accurate solutions to the model’s chemical mechanism than the SMVGEARII solver for less computational expense. We have reported only the results using the Rodas-3 set of Rosenbrock coefficients; however, additional tests were performed using the other available sets (Ros-2,Ros-3,Ros-4,Rodas-4), and the trends were similar. It must also be emphasized that these comparisons should not be generalized to other platforms or CTMs; the SMVGEARII algorithm is designed to perform most efficiently on vector platforms by re-ordering the grid cells every external chemistry time step, an operation which serves only to increase the cost of this method by \( \sim 5\% \) on non-vector machines such as those used in this study, and most other GEOS-Chem studies.

Appendix B

Discrete adjoint derivatives with respect to reaction rate constants

We desire to calculate the gradient of the cost function with respect to NO\(_x\) emissions. In GEOS-Chem, the emission and dry deposition of many species, such as NO\(_x\), are incorporated into the tropospheric chemical mechanism as reactions such as,

\[
\frac{dc_{NO}}{dt} = ENO_x
\]
where \( \text{ENO}_x \) is the \( NO_x \) emission rate, emitted as NO. The strong influence of \( NO_x \) on the overall chemistry precludes using the continuous adjoint equation of the above equation,

\[
\lambda_E = \int \lambda_c \, dt \tag{B2}
\]

Hence, we must calculate the sensitivity of the discrete chemical solver itself with respect to the reaction rate coefficients. We present a derivation of these equations here, as they have not yet been presented elsewhere, and they are necessary for accurate calculation of the desired adjoint sensitivities.

For completeness, we first present the equations for the Rosenbrock method, which advances the forward model solution \( \{c^n\} \) from one step to the next using the following formulas,

\[
c^{n+1} = c^n + \sum_{i=1}^{s} m_i k_i, \quad \text{Err}^{n+1} = \sum_{i=1}^{s} e_i k_i \tag{B3}
\]

\[
T_i = t^n + \alpha_i h, \quad C_i = c^n + \sum_{j=1}^{i} a_{i,j} k_j \tag{B4}
\]

\[
A = \left[ \frac{1}{h} - J^T(t^n, c^n) \right] \tag{B5}
\]

\[
A \cdot k_i = f(T_i, C_i) + \sum_{j=1}^{i-1} b_{i,j} k_j + t^n c^n \tag{B6}
\]

where \( s \) is the number of stages, \( a_i = \sum_j a_{i,j}, \quad Y_j = \sum_j Y_{j,i} m_i, \quad a_{i,j}, \quad b_{i,j}, \quad Y_{i,j}, \) and \( e_i \) are method coefficients, \( f(\cdot, \cdot) \) is the ODE derivative function: \( c' = f(t, c) \), \( f(\cdot, \cdot) \) is the partial time derivative: \( f_i(t, c) = \partial f(t, c) / \partial t, \quad J(\cdot, \cdot) \) is the Jacobian: \( J(t, c) = \partial f(t, c) / \partial c \), \( J_i(\cdot, \cdot) \) is the partial time derivative of the Jacobian: \( J_i(t, c) = \partial J(t, c) / \partial t, \) and \( H(\cdot, \cdot) \) is the Hessian: \( H(t, c) = \partial^2 f(t, c) / \partial c^2 \). \( A \) is the system matrix, and \( C_i, T_i, k_i \) are internal stage quantities defined by the method. The \( J \) and \( y \) used here are not likely to be confused with the use of these notations in the cost function definitions, and allow us to maintain consistent notation with the KPP documentation, which should be consulted for further explanations and values of the method coefficients.

The equation for the adjoint of the concentrations, \( \lambda_c \), is obtained by differentiating the method with respect to \( c_p \), see Eq. (11).

\[
A \cdot u_i = m_i \lambda^{n+1}_c + \sum_{j=i+1}^{s} \left( a_{i,j} v_j + b_{i,j} u_j \right) \tag{B7}
\]

\[
v_i = J^T(T_i, Y_i) \cdot u_i, \quad i = s, s-1, \ldots, 1 \tag{B8}
\]

\[
\lambda^n_c = \lambda^{n+1}_c + \sum_{i=1}^{s} (H(t^n, c^n) \cdot k_i)^T \cdot u_i + h J_{t}^T(t^n, c^n) \cdot \sum_{i=1}^{s} v_i u_i + \sum_{i=1}^{s} v_i \tag{B9}
\]

where \( v_i \) and \( u_i \) are internal stage vectors defined by the method. For GEOS-Chem, the reaction rates are constant over the internal time steps, hence we use the reduced form of this equation for autonomous systems,

\[
\lambda^n_c = \lambda^{n+1}_c + \sum_{i=1}^{s} (H(t^n, c^n) \cdot k_i)^T \cdot u_i + \sum_{i=1}^{s} J^T(T_i, C_i) \cdot u_i \tag{B10}
\]

Taking the derivative of the Rosenbrock method with respect to the reaction rate parameters, and applying Eq. (14), gives the following equation, again for autonomous systems,

\[
\lambda^n_p = \lambda^{n+1}_p + \sum_{i=1}^{s} (J_f(t^n, c^n) \cdot k_i)^T \cdot u_i + \sum_{i=1}^{s} J^T(T_i, C_i) \cdot u_i \tag{B11}
\]

Though Eq. (B11) is not implemented in the KPP generated adjoint code, KPP does generate the necessary routines for calculation of \( f_p \) \( (dFun_dRcoeff) \) and \( (J_f(t^n, c^n) \cdot k_i)^T \) \( (dJac_dRcoeff) \). For emissions, the function derivative is simply the
identity matrix, and the Jacobian derivative is zero as the emission ODE is independent of any other species concentrations, so the discrete adjoint of the emission rates is

$$\lambda_E^n = \lambda_E^{n+1} + \sum_{i=1}^{s} I \cdot u_i.$$  \hspace{1cm} (B12)

Acknowledgements. This work was supported by U.S. Environmental Protection Agency, grant R832158, and the National Science Foundation, grant NSF ITR AP&IM 0205198, which provided access to the TeraGrid resources at the National Center for Supercomputing Applications. We wish to thank M. Kopacz and A. Sandu for valuable insight and technical support.

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Fig. 1. Thermodynamic adjoint validation. Comparison of sensitivities of nitrate aerosol mass at the surface to emissions of anthropogenic NH$_3$ and SO$_x$ calculated using the adjoint method vs. the finite difference method. Transport processes are turned off for these tests to afford direct model-wide comparison.

Fig. 2. Chemistry adjoint validation. Full chemistry (only) simulation for 1 week. In the left column are the gradients of the masses of acetone (ACET), sulfate aerosol (SO$_4$), and methacrolein (MACR) at the surface with respect to anthropogenic emissions of NO$_x$. In the right column are the same quantities calculated using the finite difference method.
Fig. 3. Select points for accuracy tests. Black locations used for anthropogenic emissions of $\text{SO}_x$ and $\text{NO}_x$, grey points for $\text{NH}_3$, with one overlapping pair in Europe.

Fig. 4. Overall accuracy. Comparison of sensitivities of global aerosol burdens to anthropogenic precursor emissions calculated using the adjoint method vs. the finite difference method. A few of the plots contain insets with magnified views of a cluster of points.
**Fig. 5.** Effects of advection. Comparison of sensitivities of sulfate burdens to NO\textsubscript{x} emissions calculated using the adjoint method vs. the finite difference method. Case (a) is the base case which employs the standard PPM advection scheme and a global analysis domain. The other cases differ from the base case in the following manner: (b) advection is turned off; (c) the analysis domain is only a single location; (d) advection with a first order, linear scheme.

**Fig. 6.** Effects of simulation length and assimilation window. Comparison of sensitivities of sulfate burdens to NO\textsubscript{x} emissions calculated using the adjoint method vs. the finite difference method. Case (a) is the base case with a 24 h simulation length and a 24 h window. In the remaining cases, these parameters are adjusted to explore the effects of (b) decreasing the window; (c) increasing both the window and the length; (d) increasing the length.
Fig. 7. Sensitivities of global burdens of sulfate, nitrate and ammonium aerosol to emissions of anthropogenic SO\textsubscript{x}, NO\textsubscript{x} and NH\textsubscript{3} calculated using the adjoint model for a week-long simulation.

Fig. 8. Sensitivities of nitrate aerosol to emissions of anthropogenic NO\textsubscript{x} when the emission inventories are scaled by factors of 0.75 and 1.25, and the percent difference between these sensitivities and those calculated with the base case ($\sigma_{\text{ENO}x}=1.0$), shown in Fig. 7.
**Fig. 9.** Cost function reduction for tests DA1. A uniform perturbation is applied to emission inventories of (a) SO$_x$ (b) biomass burning SO$_2$ (c) biofuel SO$_2$. Complete daily measurements of sulfate aerosol are utilized for the data assimilation during a week-long simulation.

**Fig. 10.** Cost function reduction for tests DA2. A uniform perturbation is applied to emission inventories of anthropogenic NH$_3$. Complete daily measurements of (red-crosses) ammonium aerosol and (blue-diamonds) ammonium aerosol and gas-phase O$_x$ are utilized for the data assimilation during a week-long simulation.
Fig. 11. Cost function reduction for tests DA3. Emissions inventories of anthropogenic SO$_x$ and NO$_x$ emissions are perturbed regionally and optimized simultaneously utilizing sparse daily measurements of aerosol sulfate, ammonium, and nitrate during a week-long simulation.

Fig. 12. Emissions inventory estimates for test DA3. Difference between the estimated emission inventory at iteration $i$ and the “true” inventory, which was used to generate the pseudo-observations. Results are shown for the initial estimate (left column) and after 15 iterations (right column).
Fig. A1. Work-precision diagram for the Rosenbrock (blue circles), and SMVGEARII (red crosses) chemical solvers. Each solver is implemented in the 3-D model and run for one day using a 1 h external chemical time step. Plot shows the average time taken per external chemical time step versus the significant digits of accuracy (SDA) achieved. Tests performed using dual 1.5 GHz Itanium processors.

Fig. A2. Difference between the new standard GEOS-Chem simulation using the Rosenbrock solver with respect to the original GEOS-Chem solution using SMVGEARII after a week-long run. The effect of switching solvers is a ∼5–10% change in species concentrations.