

The Adjoint of CMAQ

AMIR HAKAMI ¹, DAVEN K. HENZE ¹, JOHN H. SEINFELD ^{1,*}, KUMARESH SINGH ²,
ADRIAN SANDU ², SOONTAE KIM ³, DAEWON BYUN ³, QINBIN LI ⁴

¹ Department of Chemical Engineering, California Institute of Technology, Pasadena, California.

5 ² Department of Computer Science, Virginia Polytechnic Institute and State University, Blacksburg,
Virginia.

³ Geosciences Department, University of Houston, Houston, Texas.

⁴ Tropospheric Modeling and Analysis, Jet Propulsion Laboratory, Pasadena, California.

ABSTRACT. An adjoint model for the internationally used Community Multiscale Air Quality
10 (CMAQ) modeling platform of the U.S. EPA is developed. The adjoint version for CMAQ (CMAQ-
ADJ) provides the user community with forward (direct decoupled method or DDM) and backward
(adjoint) sensitivity analysis capabilities. Current implementation is for gas-phase processes. Discrete
adjoints are implemented for all processes with the exception of horizontal advection, for which,
because of inherent discontinuities in the advection scheme, the continuous approach is superior. The
15 adjoint of chemistry is constructed by interfacing CMAQ with the Kinetic Pre-Processor (KPP), which
provides for increased flexibility in choice of chemical solver and facilitates implementation of new
chemical mechanisms. The adjoint implementation is evaluated both on a process-by-process basis and
for the full model. In general, adjoint results show good agreement with brute-force and DDM
sensitivities. As expected for a continuous adjoint implementation in a nonlinear scheme, the agreement
20 is not perfect for horizontal transport. Sensitivities of various air quality, public health, and
environmental metrics with respect to emissions are calculated using the adjoint method. In order to

* Corresponding author; phone: (626) 395-4635; fax: (626) 796-2591; email: seinfeld@caltech.edu.

show applicability to regional climate studies, as an example, sensitivities of these metrics with respect to local temperatures are calculated.

Introduction

25 Three-dimensional atmospheric chemical transport models (CTMs) are used to predict spatial and temporal distributions of airborne pollutants. In recent years, they have been increasingly modified to provide information, not only about concentrations, but also sensitivities of atmospheric levels with respect to various parameters. Owing in part to increased computational resources, new methods have been developed to more efficiently calculate sensitivity coefficients (derivatives) of model outputs with
30 respect to various inputs. Sensitivity information provided by atmospheric models can be used in various applications such as design of optimal pollution control strategies, inverse modeling and model parameter estimation, and air quality forecasting and data assimilation.

Local sensitivity analysis techniques can be divided into two general categories of forward and
35 backward methods. In the often-used forward method, sensitivities are propagated forward (along the model trajectories) from the perturbed source into various receptors/outputs (1-3). By nature, the methods in this category are efficient in simultaneously providing sensitivity information about all receptors with respect to a few specific parameters (4-7). In backward (adjoint) sensitivity analysis, the perturbation is made at the receptor end and is propagated backward in time and space through an
40 auxiliary set of equations. As a result, adjoint sensitivity analysis provides simultaneous sensitivity information about specific receptors with respect to all sources and parameters.

Adjoint sensitivity analysis in its current form can be traced back to the early stages of nuclear reactor physics in 1940s and 1950s (comprehensive views on the history of the adjoint method for sensitivity
45 analysis are given in 8, 9). Adjoint sensitivity analysis was later applied to various environmental problems (10-15). In particular, it has been used extensively in meteorology and oceanography for various applications such as sensitivity analysis (16, 17), variational data assimilation (18-20),

parameter estimation (21), etc. In the last decade, adjoint analysis has been extended to three-dimensional CTMs. Elbern *et al.* (22-25) developed an adjoint for the EURAD CTM and performed
 50 chemical data assimilation and inverse modeling of emissions. Vukicevic and Hess (26) implemented the adjoint method in the tracer model HANK and performed sensitivity analysis with respect to various parameters. More recently, adjoint versions for other global or regional CTMs have been developed for various applications such as inverse modeling and sensitivity analysis; these include regional models CHIMERE (27-29), STEM (30-32), Polair (33-34), CIT (35, 36), and DRAIS (37), and global models
 55 IMAGES (38, 39), TM4 (40), and GEOS-Chem (41). However, among the regional and air quality models for which an adjoint version is available, none is widely used in the modeling community. In this work, we develop an adjoint for the Community Multiscale Air Quality (CMAQ) model of U.S. EPA (42, 43). CMAQ is generally considered as the most widely used air quality model in the U.S. and across the world. Furthermore, we demonstrate how the adjoint model can be used for sensitivity
 60 analysis of various air quality and/or environmental metrics. The adjoint method, in tandem with DDM, provides the users of CMAQ with a powerful set of analysis tools that can tackle a wide spectrum of problems.

Development of the Adjoint

65 Atmospheric Chemical Transport Models (CTMs) are based on the atmospheric diffusion equation (44, 45),

$$\frac{\partial C_i}{\partial t} = -\mathbf{u} \cdot \nabla C_i + \frac{1}{\rho} \nabla \cdot (\rho \mathbf{K} \nabla C_i) + R_i + E_i \quad (1)$$

where C_i is the mixing ratio of species i (a function of three-dimensional space and time), \mathbf{u} is the
 70 vector wind field, \mathbf{K} is the diffusivity tensor, E_i represents elevated emissions, ρ is the air density, and R_i is the net chemical reaction rate for the species. Equation (1) is solved subject to specified initial and boundary conditions. In operator notation, the model operator matrix, \mathbf{M} , maps the input parameters vector $\boldsymbol{\alpha}$ (including initial conditions) into the output domain,

75

$$\mathbf{C} = \mathbf{M}\boldsymbol{\alpha} \quad (2)$$

where \mathbf{C} is the concentration vector at the end of the integration period (t_F). For most common applications, principal input parameters are initial and boundary concentrations and emission rates, but they may also include other parameters such as chemical reaction rate constants, dry deposition velocities, etc. Equation (1) and the operator \mathbf{M} can be referred to as the *forward model* and operator, respectively. Note that owing to nonlinearity in gas-phase chemistry, \mathbf{M} is a nonlinear operator. Transport processes (horizontal and vertical advection and diffusion) in Equation (1), on the other hand, are physically linear processes, although these terms may be integrated numerically by nonlinear schemes in the discretized form.

85 A perturbation in the input parameters (e.g., emissions δE_i) in Equation (1) results in a perturbation in each predicted concentration, δC_i , which is governed by,

$$\frac{\partial \delta C_i}{\partial t} = -\mathbf{u} \cdot \nabla (\delta C_i) + \frac{1}{\rho} \nabla \cdot (\rho \mathbf{K} \nabla \delta C_i) + \mathbf{F}_i \delta \mathbf{C} + \delta E_i \quad (3)$$

where \mathbf{F}_i is the i th row of the Jacobian of the chemical reaction rates ($F_{ij} = \partial R_i / \partial C_j$). Equation (3) can be represented in operator form as,

$$\delta \mathbf{C} = \mathbf{L} \delta \boldsymbol{\alpha} \quad (4)$$

where \mathbf{L} is the Jacobian of the model operator at the base case conditions. Equations (3) and (4) represent the linearized form of the forward model/operator, and are referred to as the Tangent Linear Model (TLM) and operators, respectively. Note that Equation (3) is also equivalent to the system of sensitivity equations solved in the direct decoupled method (DDM) (2, 3) for sensitivity analysis; TLM and DDM can be used interchangeably to refer to the forward method for calculation of first-order local (in the sense of a small perturbation) sensitivity coefficients, $\delta C_i / \delta E_j$.

100 If one defines a scalar cost function of the concentration field computed by the model (forms of the cost function for problems of interest in the application of CMAQ are discussed in Section 4) as,

$$J = \iint_{t,\omega} g(\mathbf{C}, t, \omega) d\omega dt \quad (5)$$

where ω is the generalized spatial coordinate, then an *adjoint model* to the TLM/DDM can be
 105 derived by applying Lagrange multipliers and integration by parts to Equation (3) and its associated initial and boundary conditions,

$$-\frac{\partial \lambda_i}{\partial t} = \nabla \cdot (\mathbf{u} \lambda_i) + \nabla \cdot (\mathbf{K} \nabla \lambda_i) + \mathbf{F}_i^T \boldsymbol{\lambda} + \varphi_i \quad (6)$$

where \mathbf{F}_i^T is the i th row of the transposed Jacobian (or the transpose of the i th column of the original
 110 Jacobian) of the chemical reaction rates, λ_i is the adjoint variable for species i such that at each time and location $\lambda_i = \partial J / \partial C_i$, and φ_i is the forcing term for the adjoint equations,

$$\varphi_i = \frac{\partial g}{\partial C_i} \quad (7)$$

Details of the derivation of adjoint equations and related initial and boundary conditions for a CTM
 115 can be found in Elbern et al., (2000), Sandu et al., (2005), Hakami et al. (2005), Martien et al. (2006), and Henze et al. (2007). Equation (6) is challenging to solve owing to the concentration-dependent cost function and the associated forcing term, φ_i . The negative sign preceding the transient term in Equation (6) indicates that the adjoint equations are integrated backwards in time. Calculation of the Jacobian of the reaction rates in Equations (3) and (6) requires values of the state vector (concentrations). For
 120 TLM/DDM calculations this is achieved easily, as the integrations for the TLM/DDM and forward models may be advanced together. For adjoint calculations, however, concentrations need to be stored (so-called *checkpointed*) in forward simulations and then used for backward integration. Checkpointing is also necessary for any other nonlinear process that is simulated in the backward mode.

125 The TLM/DDM equations are driven/forced by perturbations in the inputs/sources [e.g., δE in
Equation (3)]. These perturbations are then propagated forward in time to produce a field of sensitivity
coefficients with respect to the perturbed input/source parameter. On the other hand, the forcing terms
for the adjoint equations are perturbations in a scalar, receptor-based cost function [i.e., φ_i in Equation
(6)] that are propagated backward in time. Therefore, integration of the adjoint equations results in a
130 field of sensitivity coefficients of the cost function with respect to model inputs. TLM/DDM is a source-
based forward sensitivity method suitable for calculating sensitivities of a large number of outputs with
respect to a few inputs. Adjoint sensitivity analysis, on the other hand, is a receptor-based, backward
method that is most efficient in calculating sensitivities of a few outputs with respect to numerous
inputs. The duality in the range of efficiency and applicability for forward (TLM/DDM) and backward
135 (adjoint) sensitivity analysis methods makes them complementary approaches for addressing a wide
spectrum of problems.

The adjoint of a linear operator L can also be defined using the following duality principle,

$$140 \quad \langle \mathbf{u}, \mathbf{L}\mathbf{v} \rangle_n = \langle \mathbf{L}^* \mathbf{u}, \mathbf{v} \rangle_n \quad (8)$$

where L^* is the adjoint operator, and $\langle \cdot, \cdot \rangle_n$ denotes the inner product in R^n . It can be seen that $L^* = L^T$.

Applying Equation (8) to the TLM/DDM model:

$$\langle \delta \mathbf{C}, \mathbf{L} \delta \boldsymbol{\alpha} \rangle = \langle \mathbf{L}^* \delta \mathbf{C}, \delta \boldsymbol{\alpha} \rangle \quad (9)$$

145 Or:

$$(\mathbf{L} \delta \boldsymbol{\alpha})^T (\mathbf{L} \delta \boldsymbol{\alpha}) = \delta \boldsymbol{\alpha}^T \mathbf{L}^* (\mathbf{L} \delta \boldsymbol{\alpha}) \quad (10)$$

which can be used for verification of the adjoint operator from a validated TLM/DDM model (46).

150 Two approaches exist for integration of the adjoint equations. In the continuous approach, the adjoint
 Equations (6) are discretized and solved numerically. Alternatively, discrete adjoint models may be
 developed directly from the discretized DDM model. The two approaches in general produce different
 results, as adjoint and discretization operations are not commutable (47). Note that the same distinction
 can be made between continuous and discrete approaches for development of TLM/DDM models.

155

In Equations (2) and (4) it is assumed that evolution of concentrations or sensitivities is described by a
 single operator. In practice, CTMs use an operator splitting scheme, and they also integrate the
 governing equations over multiple time steps. The overall operator can be considered as the
 composition of all internal steps. In other words, for a model of N time-steps and m processes at each
 160 step,

$$M = \left(M_{p_1}^1 \circ M_{p_2}^1 \circ \dots \circ M_{p_m}^1 \right) \circ \dots \circ \left(M_{p_1}^N \circ M_{p_2}^N \circ \dots \circ M_{p_m}^N \right) \quad (11a)$$

$$L = \left(L_{p_1}^1 \circ L_{p_2}^1 \circ \dots \circ L_{p_m}^1 \right) \circ \dots \circ \left(L_{p_1}^N \circ L_{p_2}^N \circ \dots \circ L_{p_m}^N \right) \quad (11b)$$

$$L^* = \left(L_{p_m}^N \right)^T \circ \left(L_{p_{m-1}}^N \right)^T \circ \dots \circ \left(L_{p_1}^N \right)^T \circ \dots \circ \left(L_{p_m}^1 \right)^T \circ \left(L_{p_{m-1}}^1 \right)^T \circ \dots \circ \left(L_{p_1}^1 \right)^T \quad (11c)$$

165 **Implementation of the Adjoint (CMAQ-ADJ)**

In this work adjoint (and DDM) methods are implemented in CMAQ version 4.5.1 for gas-phase
 processes. (For identification purposes, this code is referred to as CMAQ-ADJ). As there are no
 significant changes in these processes in the recently released version 4.6, the adjoint model
 implementation can be equally applied to CMAQ 4.6. We use a hybrid approach in development of the
 170 adjoint (and DDM) model(s), wherein discrete adjoints for chemistry, diffusion, and vertical advection,
 and continuous adjoints for horizontal advection are employed. As explained later, these choices are
 made based on accuracy of the adjoints, as well as their physical significance.

A variation of DDM (4, 7) has been previously implemented in CMAQ (48). Implementation of
 175 DDM is briefly addressed to complement the discussion of adjoint implementation. The previous

implementation of DDM in CMAQ employed a continuous approach. In the present DDM implementation a discrete approach for chemistry integration is used to enhance accuracy (see discussion on the chemistry adjoint below). Complete validation of the DDM results is not shown here, but in general they are in good agreement with brute-force finite difference calculations (Figure 1). Note that minor disagreements between DDM and brute-force results in Figure 1 are primarily a result of the use of a continuous approach for calculating DDM sensitivities of horizontal advection. Using a discrete approach for horizontal advection will enhance agreement between BF and DDM, but may produce physically inconsistent results (see discussion on horizontal advection below). The same problem will occur in vertical advection for earlier versions of CMAQ where a nonlinear scheme was employed.

185

Validating DDM results with brute-force (BF) sensitivities is a rather straightforward task as they both provide forward sensitivity fields. Validation of adjoint results can not be achieved as easily. For each pair of forward (DDM or BF) and backward (adjoint) sensitivity simulations, there is only one point-of-comparison available for validation purposes. Depending on the definition of the forward sensitivity parameter (perturbed source) and adjoint cost function (perturbed receptor), the point-of-comparison can be the sensitivity of a single output or an integrated concentration metric to changes in a single input or collective change to a set of inputs. Regardless of the source and receptor metrics involved, available points-of-comparison between each pair of forward and backward sensitivity fields is reduced to a single scalar. Therefore, a complete validation of adjoint variables for all locations, times, and species is pragmatically infeasible. In the following sections we describe the methods used for implementation of adjoint analysis in CMAQ, and then for validation purposes we will introduce reduced models where points-of-comparison between forward and backward sensitivity fields are increased. For example, for chemistry validation, forward and backward simulations are carried out only for chemistry (transport processes turned off). Therefore, the 3-D model can be considered as an ensemble of numerous box models, each of which provides a point-of-comparison. The reduced

200

chemistry-only model will then provide a three-dimensional (in space only) field of points-of-comparison between forward and backward sensitivity fields.

Chemistry Adjoint. In order to implement forward and backward sensitivity analysis capabilities, the Kinetic PreProcessor (KPP) version 2.2 (49, 50) has been integrated into CMAQ. KPP can be efficiently used to generate required subroutines for any chemical mechanism. Recent versions of KPP have been extended to include DDM and adjoint sensitivity analysis capabilities (51-53). KPP significantly enhances the flexibility of CMAQ for using new or modified chemical mechanisms. Chemical solvers offered in KPP are generic and independent of mechanism, and therefore, migration to a new chemistry can be achieved seamlessly. The latest version of KPP offers a choice from multiple Rosenbrock (54, 55) and Runge-Kutta solvers (including families of fully-implicit three-stage, and singly diagonally-implicit Runge-Kutta methods), where within each family various solvers differ in accuracy and stability properties. KPP generates all required subroutines for continuous and discrete DDM and adjoint sensitivity calculations. In general, KPP solvers compare favorably with CMAQ's SMVGEAR (56), ROS3, and Euler backward iterative (EBI) (57) solvers. At the default tolerance values, cpu time requirement for KPP's Rosenbrock solvers are comparable with EBI and lower than CMAQ's SMVGEAR and ROS3 (Table 1). Work-precision diagrams (55) for model simulations with original CMAQ solvers, as well as those of KPP are shown in Figure 2. These diagrams suggest that KPP's Rosenbrock solvers provide fairly accurate solutions at relatively low computational cost and outperform original CMAQ solvers.

State vectors (concentrations) are checkpointed at synchronization (chemistry) time steps into netCDF files. Currently, checkpointing is available only for fixed time steps in each simulation day. Therefore, predetermined synchronization steps are set for each day. Checkpoints are written during the forward simulations. During the backward simulations, checkpointed concentrations are read at the beginning of each synchronization time step. The required checkpoints at the internal chemistry time steps are

recalculated by forward integration of the concentrations for the synchronization time step (i.e., two-level checkpointing, see 30, 58). Calculated chemistry-only adjoint sensitivities are in close agreement with BF values (Figure 3).

230

Horizontal advection adjoint. CMAQ version 4.5 uses the Piecewise Parabolic Method (59) to integrate the flux form of the one-dimensional horizontal advection equation,

$$\frac{\partial(\eta)}{\partial t} = -\frac{\partial(u\eta)}{\partial x} \quad (12)$$

235 where η is the mass-based concentration vector ($\eta = \rho C$, where ρ is air density). If total mass continuity holds for the advection process, then Equation (12) is equivalent to:

$$\frac{\partial C}{\partial t} = -u \frac{\partial C}{\partial x} \quad (13)$$

240 Applying a mass-based adjoint variable (for conservation) to Equation (13) results in the following one-dimensional adjoint advection equation,

$$-\frac{\partial(\lambda_m)}{\partial t} = \frac{\partial(u\lambda_m)}{\partial x} \quad (14)$$

where λ and λ_m are mixing ratio and mass-based adjoint variables ($\lambda_m = \lambda/\rho$). Therefore, the one-dimensional adjoint advection equations solved in the continuous adjoint implementation are,

245

$$-\frac{\partial(\lambda/\rho)}{\partial t} = \frac{\partial(u\lambda/\rho)}{\partial x} \quad (15)$$

250 where all Dirichlet boundary conditions for the adjoints are set to zero. In forward simulations mixing ratios are “coupled” with (multiplied by) densities (i.e., they are converted to mass-based concentrations) for advection processes. Subsequently, after the completion of each advection step, mass-based concentrations are “decoupled” from (divided by) densities and converted back to mixing ratios. Unlike the forward calculations, in the backward simulation, conversion of mixing ratio to mass-

based adjoints is accomplished by division by densities (see Table 2 for an operational scheme of the forward and backward simulations).

255 Figure 4 shows comparisons between adjoint and BF horizontal advection sensitivities. The simulations for Figure 4 include only horizontal advection in the x -direction, where the figure shows an x -cross section (for 23 vertical layers) of sensitivities of ozone at the final time step in the 20th column of the domain with respect to initial ozone concentrations in the 20th column. By considering advection in the x -direction as the sole process, a two-dimensional field of point-of-comparison between forward and backward sensitivity fields is available for visualization. BF sensitivities are calculated by various changes to the initial ozone at the 20th column. As can be seen, the general features of the advected fields are similar, but there are noticeable differences between the forward and backward fields. The adjoint field is smoother while negative sensitivities in the BF field are physically meaningless; negative sensitivities represent only numerical noise in the BF fields. The behavior seen in Figure 4 is consistent with previous studies of the adjoints of nonlinear advection schemes (41, 60, 61). Also note that BF fields are inconsistent among themselves (for various perturbations), and deviate further from the backward field for smaller perturbations. This indicates that differences between BF and adjoint fields are not a result of nonlinearities in the advection scheme, as they increase with decreasing perturbations. These differences instead result from discontinuity and non-differentiability in the forward scheme that causes irrecoverable inconsistencies between BF simulations (62). If a discrete adjoint approach is employed in these simulations, the resulting adjoint field would resemble a noisy BF sensitivity field. As a result of these inconsistencies, we conclude that for the current nonlinear advection scheme in CMAQ, continuous adjoint implementation of the horizontal advection scheme is superior to discrete implementation.

275

Vertical advection adjoint. CMAQ uses an upwind first order finite difference scheme for solving the vertical advection equation,

$$\frac{\partial(\eta)}{\partial t} = -\frac{\partial(w\eta)}{\partial z} \quad (16)$$

280 In CMAQ version 4.5, vertical advection is used as a mass conservation step. During horizontal advection, air densities are also advected alongside concentrations. In each ensuing vertical advection step, the vertical wind profile is calculated such that transported air density for that time step at each level matches meteorological densities from MM5. The calculated vertical wind profile is then applied to all species. It is necessary to use a similar vertical wind profile during backward calculations to that
 285 used in the forward simulations. Therefore, either transported air densities after each horizontal advection step or the calculated vertical wind profile at each vertical advection step during forward simulations need to be checkpointed.

Similar to horizontal advection, the corresponding adjoint equation for vertical advection is,

$$290 \quad -\frac{\partial(\lambda/\rho)}{\partial t} = \frac{\partial(w\lambda/\rho)}{\partial z} \quad (17)$$

The same numerical subroutine may be used with the reverse vertical wind profile. In a discrete context, the concentrations evolve by applying the following forward operator in vertical advection,

$$\begin{bmatrix} C_1 \\ C_2 \\ \vdots \\ C_{l-1} \\ C_l \end{bmatrix}^{n+1} = \begin{bmatrix} 1+d_1 & u_1 & 0 & 0 & 0 \\ l_2 & 1+d_2 & u_2 & 0 & 0 \\ 0 & \ddots & \ddots & \ddots & 0 \\ 0 & 0 & l_{l-1} & 1+d_{l-1} & u_{l-1} \\ 0 & 0 & 0 & l_l & 1+d_l \end{bmatrix} \begin{bmatrix} C_1 \\ C_2 \\ \vdots \\ C_{l-1} \\ C_l \end{bmatrix}^n \quad (18)$$

295 where the elements of the operator depend on the magnitude and direction of the recalculated vertical wind profile. Applying the adjoint (transpose) of the linear operator above (discrete adjoint) yields different results than using the forward operator with the reverse wind profile (continuous adjoint). Note that in previous versions of CMAQ (before v4.5) where a nonlinear scheme is used for vertical advection, a continuous approach would be preferable. The discrete adjoint provides better accuracy and
 300 is therefore used for vertical advection. Good agreement between the adjoint and BF sensitivity fields

can be seen in Figure 5 where sensitivities of surface ozone to initial surface ozone concentrations are shown for a set of forward and adjoint simulations that include vertical advection only.

Vertical diffusion adjoint. CMAQ provides an option for species emissions to be processed during chemistry or vertical diffusion integrations. In the present adjoint implementation, emissions are injected during vertical diffusion for which the following equation is solved,

$$\frac{\partial C}{\partial t} = \frac{\partial}{\partial z} \left(K \frac{\partial C}{\partial z} \right) \quad (19)$$

with the corresponding adjoint equation as:

$$-\frac{\partial \lambda}{\partial t} = \frac{\partial}{\partial z} \left(K \frac{\partial \lambda}{\partial z} \right) \quad (20)$$

Deposition velocities are included in the first layer as part of the boundary conditions. The same numerical scheme can be used for adjoint integrations by excluding only the emissions. Equation (20) is evolved to the next time step by applying the following operator,

$$C^{n+1} = (LHS)^{-1} [(RHS)C^n + E^n] \quad (21)$$

where *LHS* and *RHS* are tri-diagonal left hand side and right hand side matrices in a Crank-Nicholson discretization, respectively. Therefore, the discrete adjoint operator is,

$$\lambda^n = (RHS)^T (LHS)^{-T} \lambda^{n+1} \quad (22)$$

For vertical diffusion, discrete and continuous adjoints produce nearly identical results. We use the continuous approach in the present implementation. The gradients of the cost function with respect to emissions can be calculated from the adjoint variables during backward integrations:

$$\frac{\partial J}{\partial E^n} = \left(\frac{\partial C^{n+1}}{\partial E^n} \right)^T \left(\frac{\partial J}{\partial C^{n+1}} \right) = (LHS)^{-1} \lambda^{n+1} \quad (23)$$

Note that the required factorization in Equation (23) is already carried out for continuous adjoint integration. Therefore, calculation of the gradient of the cost function with respect to emissions of each species is accomplished by the minimal cost of one additional back-substitution in the modified tri-diagonal solver used in the adjoint integration.

Figure 6 compares BF and adjoint sensitivity fields for sensitivities of ozone at the surface with respect to surface NO emissions when simulations include only chemistry and vertical diffusion. As the accuracy of chemistry adjoints has been previously verified, the agreement shown in Figure 6 serves as validation of both the adjoint of vertical diffusion and calculation of emission gradients.

Horizontal diffusion adjoint. In CMAQ species undergo horizontal diffusion according to the following equation,

$$\frac{\partial C}{\partial t} = \frac{1}{\rho} \frac{\partial}{\partial x} \left(\rho K \frac{\partial C}{\partial x} \right) \quad (24)$$

the continuous adjoint of which is,

$$-\frac{\partial \lambda}{\partial t} = \frac{\partial}{\partial x} \left(\rho K \frac{\partial (\lambda/\rho)}{\partial x} \right) \quad (25)$$

The horizontal diffusion operator is symmetrical, and therefore, self-adjoint. As a result: (a) the same numerical scheme can be used to horizontally diffuse concentrations and adjoints in forward and backward simulations; and (b) continuous and discrete horizontal diffusion adjoints are identical. We use the same subroutine for horizontal diffusion of adjoints with an internal “decoupling” from (division by) densities (see Table 2). Adjoint and BF sensitivity fields show good agreement in Figure 7 where only horizontal diffusion in one direction is included in the simulations and sensitivities of final ozone at the 21st x -cross section with respect to initial ozone at the 20th column are shown.

Overall implementation. In the present implementation, forward and backward models are run as separate, independent models. For executing the adjoint model, one needs to carry out the forward simulation only once and store the generated checkpoint files. However, for subsequent adjoint simulations, execution of the forward model is not required, as the same checkpoint files can be re-used. This leads to significant computational savings when multiple backward simulations are performed. In general, the order in which processes are called during the backward simulation is reverse of that in the forward simulation (Table 2). As only chemistry requires knowledge of concentrations in the current implementation, checkpoints are written and read before each chemistry call. However, transported air densities are written to checkpoint files after each horizontal advection scheme in the forward mode, and read before each vertical advection scheme in the backward simulation.

In summary, validation of full adjoint results (including all processes) with forward sensitivity fields (BF or DDM) is possible only for a few sensitivity coefficients. Validation of the adjoint of isolated processes through comparison with forward sensitivity fields can be carried out for a larger number of points-of-comparison (Figures 3-7), and provides for a more robust verification procedure. Full model results are also in good agreement with BF sensitivities as shown in Figure 8 for few select points and types of sensitivity coefficients.

370

Computational efficiency. The original CMAQ provides 3 options for chemical solvers in Euler Backward Iterative (EBI), a vectorized Gear solver (SMVGEAR), and ROS3 from the family of Rosenbrock solvers. With the implementation of KPP, users of CMAQ-ADJ have access to 5 Rosenbrock and 4 Runge-Kutta solvers. Unlike the original implementation of ROS3 in CMAQ, KPP implementation does not employ cell blocking for improved performance on vectorized machines. Table 1 shows comparison of computational times for forward, DDM, and adjoint models with various solvers. All tests are carried out for a 1-day simulation for a $45 \times 46 \times 18$ computational domain with 36 km horizontal resolution. In these tests, the CB-IV chemical mechanism is used. Also note that these

simulations are carried out on non-vectorized machines. Work-precision diagrams for simulations with
 380 all solvers are shown in Figure 2. Also shown in Table 1 are relative costs of forward and backward
 sensitivity calculations. Backward calculations require at least twice the computational time as forward
 calculations (significantly more for Runge-Kutta solvers), as in the two-level checkpointing scheme, a
 second-level forward integration is performed for retrieval of concentrations at the internal time steps in
 between checkpoint intervals. Finally, note that the relative costs in Table 1 are for a single
 385 receptor/cost function (in adjoint mode) or source/sensitivity parameter (in DDM mode). When
 sensitivities for multiple receptors or sources are calculated simultaneously, the overall cost per
 receptor/source for both DDM and adjoint simulation is significantly reduced. This is a result of the
 shared computational cost in matrix factorization among various receptors/sources. The potential saving
 is more substantial for large chemical mechanisms where chemistry integration is a larger contributor to
 390 the overall computational cost.

Applications in receptor-based sensitivity analysis

The adjoint method has been used widely for variational data assimilations and inverse modeling. In
 these applications, the cost function can be defined generally as,

395

$$\begin{aligned}
 J &= J_{Observations} + J_{Background} \\
 &= \frac{1}{2}(\mathbf{C} - \mathbf{C}_{obs})^T \mathbf{O}^{-1}(\mathbf{C} - \mathbf{C}_{obs}) + \frac{1}{2}(\mathbf{E} - \mathbf{E}^b)^T \mathbf{B}^{-1}(\mathbf{E} - \mathbf{E}^b)
 \end{aligned}
 \tag{26}$$

where \mathbf{C}_{obs} is a set of observed concentrations, \mathbf{E}^b is an *a priori* (background or initial guess) estimate
 of inputs, and where \mathbf{O} and \mathbf{B} are observation and background error covariance matrices. The cost
 400 function consists of two parts: the first part is a measure of model prediction errors, and the second part
 is a penalty for deviation from a priori estimates of model inputs. In typical applications the adjoint
 method is used to calculate the gradients of the cost function with respect to initial concentrations (for
 data assimilation applications) or model parameters such as emissions (for inverse modeling
 applications). These gradients are then used in an iterative optimization algorithm in order to minimize

405 the cost function, which reduces the mismatch of model predictions and observations is reduced by
adjusting the inputs (e.g. emissions) within a reasonable range. Variational methods provide an
important approach for constraining emissions of various species on a spatially resolved basis (36).

The adjoint method is a powerful tool for receptor-oriented, sensitivity analysis. As a receptor-based
410 method, adjoint sensitivity analysis is particularly suitable for addressing policy problems. Hakami et al.
(63) used the adjoint of a CTM for ozone nonattainment sensitivity analysis over the continental United
States. They demonstrated that the adjoint method is a powerful framework for formal analysis of
interstate, trans-boundary, and intercontinental transport of pollution. Here, we briefly describe a few
other potential applications for adjoint sensitivity analysis at regional scales using the adjoint of CMAQ.
415 These applications differ only in the definition of the cost function.

Local nonattainment analysis. In this application the cost function measures the amount of deviation
from an air quality standard (63),

$$420 \quad J = \sum_{x,y,t,+} [C_i(x,y,0,t) - \gamma]^2 \quad (27)$$

where γ represents an air quality standard or threshold, and the plus sign in summation indicates that
only points with concentrations above the threshold are included in the cost function. Adjoint analysis
provides the sensitivity of nonattainment at a particular location with respect to various parameters, for
example emissions at all grid cells. In other words, the analysis will identify those sources contributing
425 most to a certain nonattainment episode. If the cost function is defined simply as an average or peak
concentration, then adjoint analysis performs, to a first-order approximation, as a source apportionment
method. In Figure 9, such analysis is carried out for ozone in Houston-Galveston area and for a
threshold of 80 ppbv. The gradients of the overall nonattainment (as defined in equation 27) with
respect to NO_x emissions provide first-order estimates of the contribution of various sources to the cost
430 function (nonattainment).

Population exposure analysis. If the cost function is defined as a population exposure metric, then the resulting sensitivities identify the most influential parameters (emissions) affecting population exposure,

435

$$J = \sum_{x,y,t} [P \times (C_i - \gamma)] \quad (28)$$

γ represents an exposure threshold, and P is the population at each location. Note that Equation (28) can be extended to include dose-response relationships for one or multiple pollutants to represent public health risks. As a target-based method, adjoint analysis can identify the most influential emission sources that contribute to the overall exposure/risk metric (e.g., Figure 10). As expected, this definition of the cost function emphasizes on areas with large population densities.

440

Environmental exposure. A cost function similar to that in Equation (28) can be used to quantify environmental stress resulting from increased pollution levels. For example, an environmental exposure metric based on the W-126 function can be defined to quantify the impact of increased ozone on crops and vegetation (64, 65),

445

$$J = \sum_{x,y,t} \left[\frac{C_i}{1 + 4403 \exp(-216C_i)} \right] \quad (29)$$

450

Using the above as the cost function in adjoint analysis, corresponding (emission) gradients can be calculated (Figure 11). Areas of significant gradients (right panel) indicate sources where emission control can result in largest reductions in the cost function (left panel). The adjoint method affords users the flexibility to combine environmental and public health metrics in a single cost function for integrated analysis of air pollution effects.

455

Effect of temperature variation on air pollution levels. In a manner similar to the calculation of emission gradients, the sensitivity of the cost function with respect to changes in temperature can be calculated,

460
$$\frac{\partial J}{\partial T^n} = \left(\frac{\partial C^{n+1}}{\partial T} \right)^T \left(\frac{\partial J}{\partial C^{n+1}} \right) = \left(\frac{\partial k^n}{\partial T} \right)^T \left(\frac{\partial C^{n+1}}{\partial k^n} \right)^T \lambda^{n+1} \quad (30)$$

As written, Equation (30) describes only the effect of temperature on the cost function via chemistry. Changes in temperatures will also affect biogenic (and anthropogenic) emissions. The results of applying Equation (30) in the adjoint analysis for population and environmental exposures above [Equations (28) and (29)] are shown in Figure 12. It is interesting that temperature gradients show
465 significant spatial distributions, something that is possible to capture only in a receptor-oriented method, i.e., adjoint sensitivity analysis. Spatial distributions such as those in Figure 12 can be used in conjunction with results from regional climate studies to formally quantify the impact of future climate conditions on regional air quality. These gradients can also be used to quantify contributions to the uncertainties in future (or current) air quality from uncertainty/variability in regional temperatures.

470

Acknowledgements

This work was supported by National Science Foundation award NSF ITR AP&IM 0205198 and by funding from the Jet Propulsion Laboratory. The work of A. Sandu and K. Singh was also supported by
475 the Houston Advanced Research Council through the award H59/2005.

References

- 1- Dickinson, R. P.; Gelinas, R. J. Sensitivity analysis of ordinary differential equation systems - Direct method. *J Comput Phys* **1976**, 21, 123-143.
- 480 2- Dunker, A. M. Efficient calculation of sensitivity coefficients for complex atmospheric models. *Atmos Environ* **1981**, 15, 1155-1161.

- 3- Dunker, A. M. The Decoupled direct method for calculating sensitivity coefficients in chemical kinetics. *J Chem Phys* **1984**, 81, 2385-2393.
- 485 4- Yang, Y. J.; Wilkinson, J. G.; Russell, A. G. Fast, direct sensitivity analysis of multidimensional photochemical models. *Environ Sci Technol* **1997**, 31, 2859-2868.
- 5- He, S.; Carmichael, G. R.; Sandu, A.; Hotchkiss, B.; Damian-Iordache, V. Application of ADIFOR for air pollution model sensitivity studies. *Environ Modell Softw* **2000**, 15, 549-557.
- 6- Dunker, A. M.; Yarwood, G.; Ortmann, J. P.; Wilson, G. M. The decoupled direct method for sensitivity analysis in a three-dimensional air quality model - Implementation, accuracy, and efficiency. 490 *Environ Sci Technol* **2002**, 36, 2965-2976.
- 7- Hakami, A.; Odman, M. T.; Russell, A. G. High-order, direct sensitivity analysis of multidimensional air quality models. *Environ Sci Technol* **2003**, 37, 2442-2452.
- 8- Cacuci, D. G.; Schlesinger, M. E. On the application of the adjoint method of sensitivity analysis to problems in the atmospheric sciences. *Atmosfera* **1994**, 7, 47-59.
- 495 9- Marchuk, G. I.; Shutyaev, V.; Bocharov, G. Adjoint equations and analysis of complex systems: Application to virus infection modelling. *J Comput Appl Math* **2005**, 184, 177-204.
- 10- Marchuk, G. I. *Numerical Solution of the Problems of the Dynamics of the Atmosphere and the Ocean* (in Russian); Gidrometeoizdat: St. Petersburg, 1974.
- 11- Cacuci, D. G. Sensitivity theory for nonlinear systems .1. Nonlinear functional analysis approach. *J* 500 *Math Phys* **1981**, 22, 2794-2802.
- 12- Cacuci, D. G. Sensitivity theory for Nonlinear systems .2. Extensions to additional classes of responses. *J Math Phys* **1981**, 22, 2803-2812.
- 13- Marchuk, G. I. *Mathematical models in environmental problems*; North-Holland ; sole distributors for the U.S.A and Canada, Elsevier Science Pub. Co.: Amsterdam ; New York, New 505 York, 1986.
- 14- Cacuci, D. G.; Ionescu-Bujor, M. Deterministic local sensitivity analysis of augmented systems - I: Theory. *Nucl Sci Eng* **2005**, 151, 55-66.

- 15- Cacuci, D. G.; Ionescu-Bujor, M.; Navon, I. M. *Sensitivity and Uncertainty Analysis*; Chapman & Hall/CRC Press: Boca Raton, 2003.
- 510 16- Hall, M. C. G.; Cacuci, D. G.; Schlesinger, M. E. Sensitivity analysis of a radiative-convective model by the adjoint method. *J Atmos Sci* **1982**, 39, 2038-2050.
- 17- Hall, M. C. G.; Cacuci, D. G. Physical interpretation of the adjoint functions for sensitivity analysis of atmospheric models. *J Atmos Sci* **1983**, 40, 2537-2546.
- 18- Ledimet, F. X.; Talagrand, O. Variational algorithms for analysis and assimilation of meteorological observations - Theoretical aspects. *Tellus A* **1986**, 38, 97-110.
- 515 19- Talagrand, O.; Courtier, P. Variational assimilation of meteorological observations with the adjoint vorticity equation .1. Theory. *Q J Roy Meteor Soc* **1987**, 113, 1311-1328.
- 20- Courtier, P.; Talagrand, O. Variational assimilation of meteorological observations with the adjoint vorticity equation .2. Numerical results. *Q J Roy Meteor Soc* **1987**, 113, 1329-1347.
- 520 21- Navon, I. M. Practical and theoretical aspects of adjoint parameter estimation and identifiability in meteorology and oceanography. *Dynam Atmos Oceans* **1998**, 27, 55-79.
- 22- Elbern, H.; Schmidt, H.; Ebel, A. Variational data assimilation for tropospheric chemistry modeling. *J Geophys Res-Atmos* **1997**, 102, 15967-15985.
- 23- Elbern, H.; Schmidt, H. A four-dimensional variational chemistry data assimilation scheme for Eulerian chemistry transport modeling. *J Geophys Res-Atmos* **1999**, 104, 18583-18598.
- 525 24- Elbern, H.; Schmidt, H.; Talagrand, O.; Ebel, A. 4D-variational data assimilation with an adjoint air quality model for emission analysis. *Environ Modell Softw* **2000**, 15, 539-548.
- 25- Elbern, H.; Schmidt, H. Ozone episode analysis by four-dimensional variational chemistry data assimilation. *J Geophys Res-Atmos* **2001**, 106, 3569-3590.
- 530 26- Vukicevic, T.; Hess, P. Analysis of tropospheric transport in the Pacific Basin using the adjoint technique. *J Geophys Res-Atmos* **2000**, 105, 7213-7230.
- 27- Menut, L.; Vautard, R.; Beekmann, M.; Honore, C. Sensitivity of photochemical pollution using the adjoint of a simplified chemistry-transport model. *J Geophys Res-Atmos* **2000**, 105, 15379-15402.

- 28- Vautard, R.; Beekmann, M.; Menut, L. Applications of adjoint modelling in atmospheric chemistry: sensitivity and inverse modelling. *Environ Modell Softw* **2000**, 15, 703-709.
- 535
- 29- Menut, L. Adjoint modeling for atmospheric pollution process sensitivity at regional scale. *J Geophys Res-Atmos* **2003**, 108 (D17), 8562, doi:10.1029/2002JD002549.
- 30- Sandu, A.; Daescu, D. N.; Carmichael, G. R.; Chai, T. F. Adjoint sensitivity analysis of regional air quality models. *J Comput Phys* **2005**, 204, 222-252.
- 540
- 31- Hakami, A.; Henze, D. K.; Seinfeld, J. H.; Chai, T.; Tang, Y.; Carmichael, G. R.; Sandu, A. Adjoint inverse modeling of black carbon during the Asian Pacific Regional Aerosol Characterization Experiment. *J Geophys Res-Atmos* **2005**, 110, D14301, doi:10.1029/2004JD005671.
- 32- Chai, T. F.; Carmichael, G. R.; Sandu, A.; Tang, Y. H.; Daescu, D. N. Chemical data assimilation of Transport and Chemical Evolution over the Pacific (TRACE-P) aircraft measurements. *J Geophys Res-Atmos* **2006**, 111, D02301, doi:10.1029/2005JD005883.
- 545
- 33- Mallet, V.; Sportisse, B. A comprehensive study of ozone sensitivity with respect to emissions over Europe with a chemistry-transport model. *J Geophys Res-Atmos* **2005**, 110, D22302, doi:10.1029/2005JD006234.
- 34- Quelo, D.; Mallet, V.; Sportisse, B. Inverse modeling of NO_x emissions at regional scale over northern France: Preliminary investigation of the second-order sensitivity. *J Geophys Res-Atmos* **2005**, 110, D24310, doi:10.1029/2005JD006151.
- 550
- 35- Martien, P. T.; Harley, R. A. Adjoint sensitivity analysis for a three-dimensional photochemical model: Implementation and method comparison. *Environ Sci Technol* **2006**, 40, 2663-2670.
- 36- Martien, P. T.; Harley, R. A. Adjoint sensitivity analysis for a three-dimensional photochemical model: Application to Southern California. *Environ Sci Technol* **2006**, 40, 4200-4210.
- 555
- 37- Nester, K.; Panitz, H. J. Sensitivity analysis by the adjoint chemistry transport model DRAIS for an episode in the Berlin Ozone (BERLIOZ) experiment. *Atmos Chem Phys* **2006**, 6, 2091-2106.
- 38- Muller, J. F.; Stavrou, T. Inversion of CO and NO_x emissions using the adjoint of the IMAGES model. *Atmos Chem Phys* **2005**, 5, 1157-1186.

- 560 39- Stavrakou, T.; Muller, J. F. Grid-based versus big region approach for inverting CO emissions using Measurement of Pollution in the Troposphere (MOPITT) data. *J Geophys Res-Atmos* **2006**, 111, D15304, doi:10.1029/2005JD006896.
- 40- Meirink, J. F.; Eskes, H. J.; Goede, A. P. H. Sensitivity analysis of methane emissions derived from SCIAMACHY observations through inverse modelling. *Atmos Chem Phys* **2006**, 6, 1275-1292.
- 565 41- Henze, D. K.; Hakami, A.; Seinfeld, J. H. Development of the adjoint of GEOS-Chem. *Atmos Chem Phys*, in press, 2007.
- 42- Byun, D. W.; Ching, J. K. S. Science algorithms of the EPA Models-3 Community Multiscale Air Quality (CMAQ) Modeling System, U. S. Environmental Protection Agency, USEPA/600/R-99/030; Research Triangle Park, NC, 1999.
- 570 43- Byun, D. W.; Schere, K. L. Review of the governing equations, computational algorithms, and other components of the Models-3 Community Multiscale Air Quality (CMAQ) modeling system. *Applied Mechanics Reviews* **2006**, 59, 51-77.
- 44- Seinfeld, J. H.; Pandis, S. N. *Atmospheric Chemistry and Physics : From Air Pollution to Climate Change*; 2nd ed.; J. Wiley: Hoboken, N.J., 2006.
- 575 45- Jacobson, M. Z. *Fundamentals of Atmospheric Modeling*; 2nd ed.; Cambridge University Press: New York, 2005.
- 46- Navon, I. M.; Zou, X.; Derber, J.; Sela, J. Variational data assimilation with an adiabatic version of the NMC spectral model. *Mon Weather Rev* **1992**, 120, 1433-1446.
- 47- Sirkes, Z.; Tziperman, E. Finite difference of adjoint or adjoint of finite difference? *Mon Weather*
580 *Rev* **1997**, 125, 3373-3378.
- 48- Cohan, D. S.; Hu, Y.; Hakami, A.; Odman, M. T.; Russell, A. G. Implementation of a direct sensitivity method into CMAQ, Models-3 Annual Conference, Chapel Hill, NC, 2002.
- 49- Damian, V.; Sandu, A.; Damian, M.; Potra, F.; Carmichael, G. R. The kinetic preprocessor KPP - a software environment for solving chemical kinetics. *Comput Chem Eng* **2002**, 26, 1567-1579.
- 585 50- Daescu, D.; Carmichael, G. R.; Sandu, A. Adjoint implementation of Rosenbrock methods applied to variational data assimilation problems. *J Comput Phys* **2000**, 165, 496-510.

- 51- Sandu, A.; Sander, R. Technical note: Simulating chemical systems in Fortran90 and Matlab with the Kinetic PreProcessor KPP-2.1. *Atmos Chem Phys* **2006**, 6, 187-195.
- 52- Sandu, A.; Daescu, D. N.; Carmichael, G. R. Direct and adjoint sensitivity analysis of chemical
590 kinetic systems with KPP: Part I - theory and software tools. *Atmos Environ* **2003**, 37, 5083-5096.
- 53- Daescu, D. N.; Sandu, A.; Carmichael, G. R. Direct and adjoint sensitivity analysis of chemical
kinetic systems with KPP: II - Numerical validation and applications. *Atmos Environ* **2003**, 37, 5097-
5114.
- 54- Hairer, E.; Wanner G. *Solving Ordinary Differential Equations II. Stiff and Differential-Algebraic*
595 *Problems*; Springer: Berlin, 1991.
- 55- Sandu, A.; Verwer, J. G.; Blom, J. G.; Spee, E. J.; Carmichael, G. R.; Potra, F. A. Benchmarking
stiff ODE solvers for atmospheric chemistry problems .2. Rosenbrock solvers. *Atmos Environ* **1997**, 31,
3459-3472.
- 56- Jacobson, M. Z.; Turco, R. P. Smvgear - a Sparse-Matrix, Vectorized Gear code for atmospheric
600 models. *Atmos Environ* **1994**, 28, 273-284.
- 57- Hertel, O.; Berkowicz, R.; Christensen, J.; Hov, O. Test of two numerical schemes for use in
atmospheric transport chemistry models. *Atmos Environ* **1993**, 27, 2591-2611.
- 58- Griewank, A.; Walther, A. Algorithm 799: Revolve: An implementation of checkpointing for the
reverse or adjoint mode of computational differentiation. *Acm T Math Software* **2000**, 26, 19-45.
- 605 59- Colella, P.; Woodward, P. R. The Piecewise Parabolic Method (PPM) for gas dynamical
simulations. *J Comput Phys* **1984**, 54, 174-201.
- 60- Vukicevic, T.; Steyskal, M.; Hecht, M. Properties of advection algorithms in the context of
variational data assimilation. *Mon Weather Rev* **2001**, 129, 1221-1231.
- 61- Liu, Z.; Sandu, A. Analysis of discrete adjoints for upwind numerical schemes. *Lect Notes Comput*
610 *Sc* **2005**, 3515, 829-836.
- 62- Thuburn, J.; Haine, T. W. N. Adjoint of nonoscillatory advection schemes. *J Comput Phys* **2001**,
171, 616-631.

63- Hakami, A.; Seinfeld, J. H.; Chai, T. F.; Tang, Y. H.; Carmichael, G. R.; Sandu, A. Adjoint sensitivity analysis of ozone nonattainment over the continental United States. *Environ Sci Technol* 615 **2006**, 40, 3855-3864.

64- Lefohn, A. S.; Runeckles, V. C. Establishing standards to protect vegetation - Ozone exposure dose considerations. *Atmos Environ* **1987**, 21, 561-568.

65- U. S. EPA. Air quality criteria for ozone and related photochemical oxidants, U. S. Environmental Protection Agency, EPA/600/P-93/004bF, Research Triangle Park, NC, 1996.

620

FIGURE CAPTIONS

Figure 1: Comparison between BF (left) and DDM (right) sensitivities of ozone with respect to initial ozone concentration after 6 hours of simulation.

625 Figure 2: Work-precision diagrams for day-long simulations with various CMAQ (black) and KPP solvers (Rosenbrock in red, and Runge-Kutta in blue). Significant digit accuracy (SDA) for each species is defined as $SDA_i = -\log(RMSRE_i)$, where RMSRE is the root mean square of relative error in comparison with a reference solution. Diagrams are shown for the overall SDA (minimum across all species, top) and ozone (bottom). Simulations with default/recommended settings are indicated by
630 enlarged, green markers. For this application, most Rosenbrock solvers fall in the desirable performance region.

Figure 3: Comparison of Adjoint and BF sensitivities of final ozone concentration with respect to initial NO concentration; chemistry-only simulations are carried out for 6 hours.

Figure 4: Comparison of various BF with adjoint sensitivity of ozone at the 20th column with respect to
635 initial ozone at the 20th column. Only horizontal advection in the x direction included in the simulations. DDM sensitivity (bottom) shows reasonable agreement with adjoint.

Figure 5: Simulations with only vertical advection for calculation of the sensitivities of surface ozone with respect to initial surface ozone.

Figure 6: Simulations with only chemistry and vertical diffusion, normalized sensitivities of surface
640 ozone with respect to surface emissions of NO.

Figure 7: Simulations with only horizontal diffusion in x-direction for calculation of ozone sensitivities at the 21st column with respect to initial ozone at 20th column.

Figure 8: Full model simulations, sensitivities of surface ozone with respect to surface NO initial concentrations (left) and emissions (right) for selected locations.

645 Figure 9: Spatial distribution of the cost function for nonattainment in Houston-Galveston Area (left) and associated gradients with respect to NO_x emissions (right). Values are normalized to the total cost function and presented in percent.

Figure 10: Spatial distribution of the cost function for population exposure (left) and associated gradients with respect to NO_x emissions (right). Values are normalized to the total cost function and presented in percent. Exposure threshold is 60 ppb.
650

Figure 11: Spatial distribution of the cost function for W-126 metric of vegetation exposure from ozone (left) and associated gradients with respect to NO_x emissions (right). Values are normalized to the total cost function and presented in percent.

Figure 12: Temperature gradients of population exposure (left) and W-126 (right) metrics as calculated during backward integration.
655

Table1. Comparison of computational efficiencies for the forward, DDM and adjoint models with various chemical solvers at the default/recommended settings.

Solver	Relative computational efficiency ¹		
	Forward Model ²	DDM ^{3,4}	Adjoint ³
CMAQ-EBI	1.00	-	-
CMAQ-ROS3	2.10	-	-
CMAQ-SMVGear	3.69	-	-
KPP-ROS2	1.59	1.88	2.02
KPP-ROS3	1.08	1.96	2.02
KPP-ROS4	1.18	2.11	2.11
KPP-RODAS3	0.96	2.12	2.09
KPP-RODAS4	1.18	2.39	2.18
KPP-RADAU-2A	2.08	7.81	7.87
KPP-LOBATTO	2.66	7.93	7.25
KPP-GAUSS	2.66	8.13	5.41
KPP-RADAU-1A	1.99	7.60	7.96

1- All simulations are carried out sequentially for 24-hours, on 64-bit, 2.0 GHz dual-core Optron processors. 2- Values are normalized to forward simulation with EBI solver. 3- Values are normalized to the forward simulation with the same solver. 4- Values include the time required for concentration integrations.

Table 2. Forward and backward simulation schemes in CMAQ.

Forward Model	Adjoint Model
INIT (t=0)	INIT (t=tF)
DO (Synchronization steps)	DO (Synchronization steps)
DO (Advection steps)	DO (Advection steps)
V-DIFF	FORCE-ADJ
COUPLE	NEXTIME (-TSTEP)
H-ADV	READ CHECKPOINT (CONC)
WRITE CHECKPOINT (Density)	CHEM-ADJ
V-ADV	H-DIFF-ADJ
H-DIFF	READ CHECKPOINT (Density)
DECOUPLE	DECOUPLE
WRITE CHECKPOINT (CONC)	V-ADV-ADJ
CHEM	H-ADV-ADJ
NEXTIME (TSTEP)	COUPLE
END DO	V-DIFF-ADJ
WRITE CONC	END DO
END DO	WRITE ADJ
	END DO

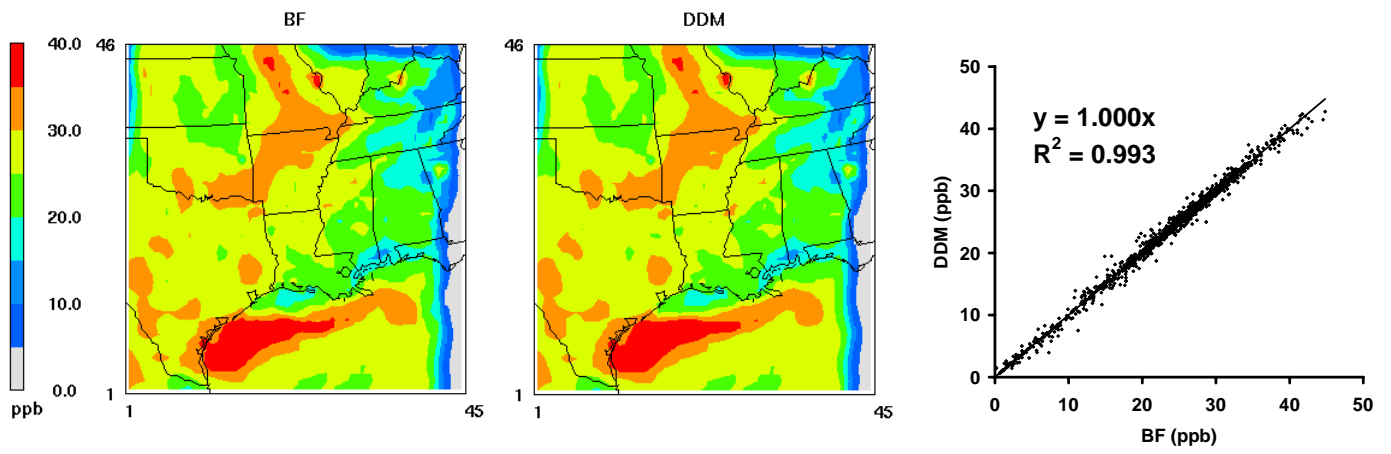


Figure 1: Comparison between BF (left) and DDM (right) sensitivities of ozone with respect to initial ozone concentration after 6 hours of simulation.

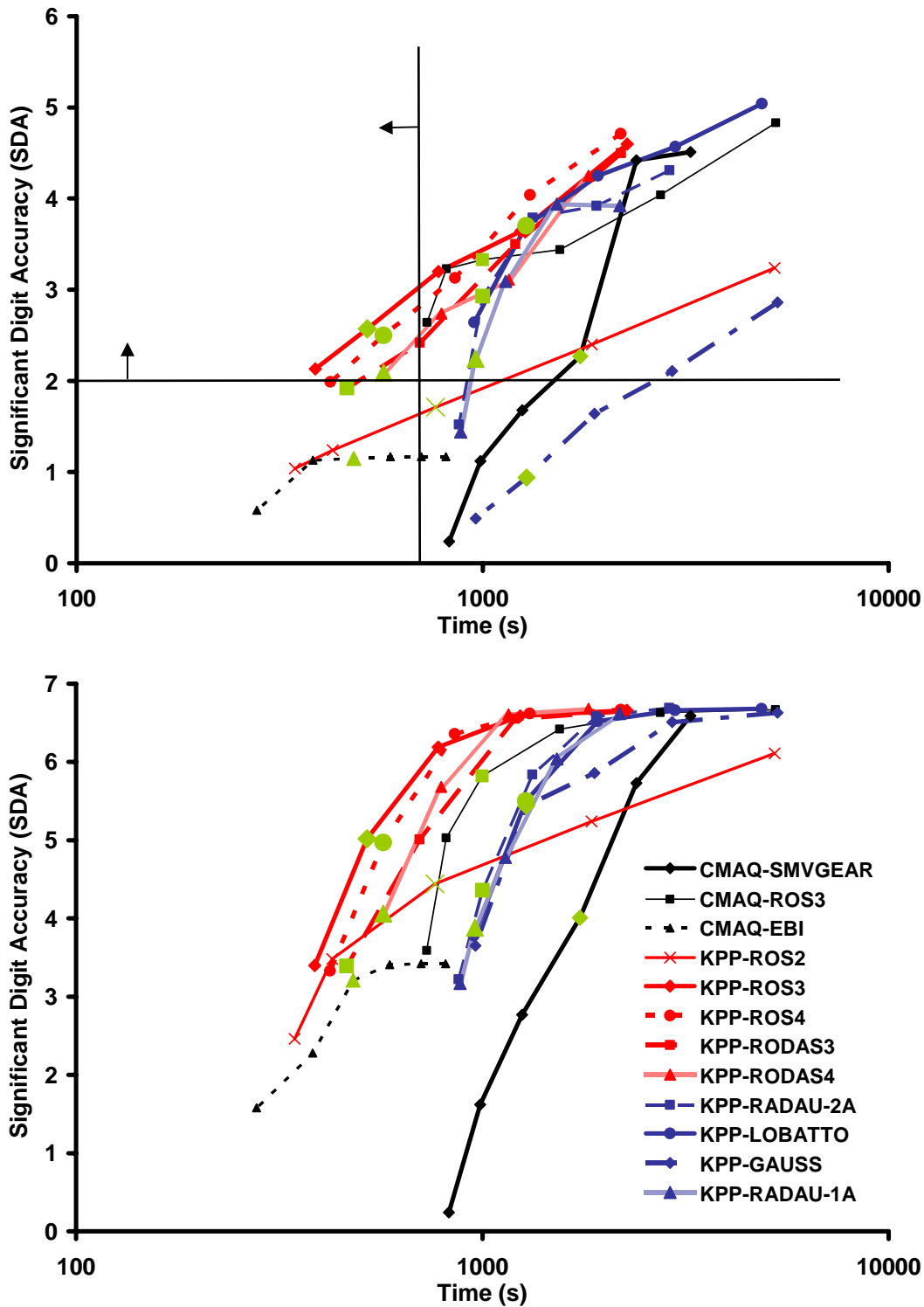


Figure 2: Work-precision diagrams for day-long simulations with various CMAQ (black) and KPP solvers (Rosenbrock in red, and Runge-Kutta in blue). Significant digit accuracy (SDA) for each species is defined as $SDA_i = -\log(RMSRE_i)$, where RMSRE is the root mean square of relative error in comparison with a reference solution. Diagrams are shown for the overall SDA (minimum across all species, top) and ozone (bottom). Simulations with default/recommended settings are indicated by enlarged, green markers. For this application, most Rosenbrock solvers fall in the desirable performance region.

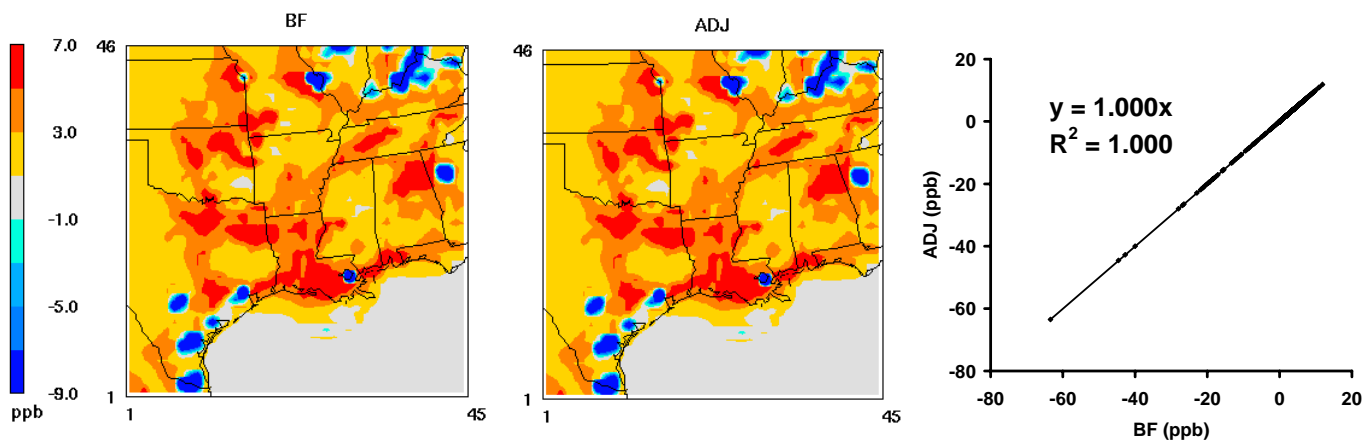


Figure 3: Comparison of adjoint and BF sensitivities of final ozone concentrations with respect to initial NO concentrations; chemistry-only simulations are carried out for 6 hours.

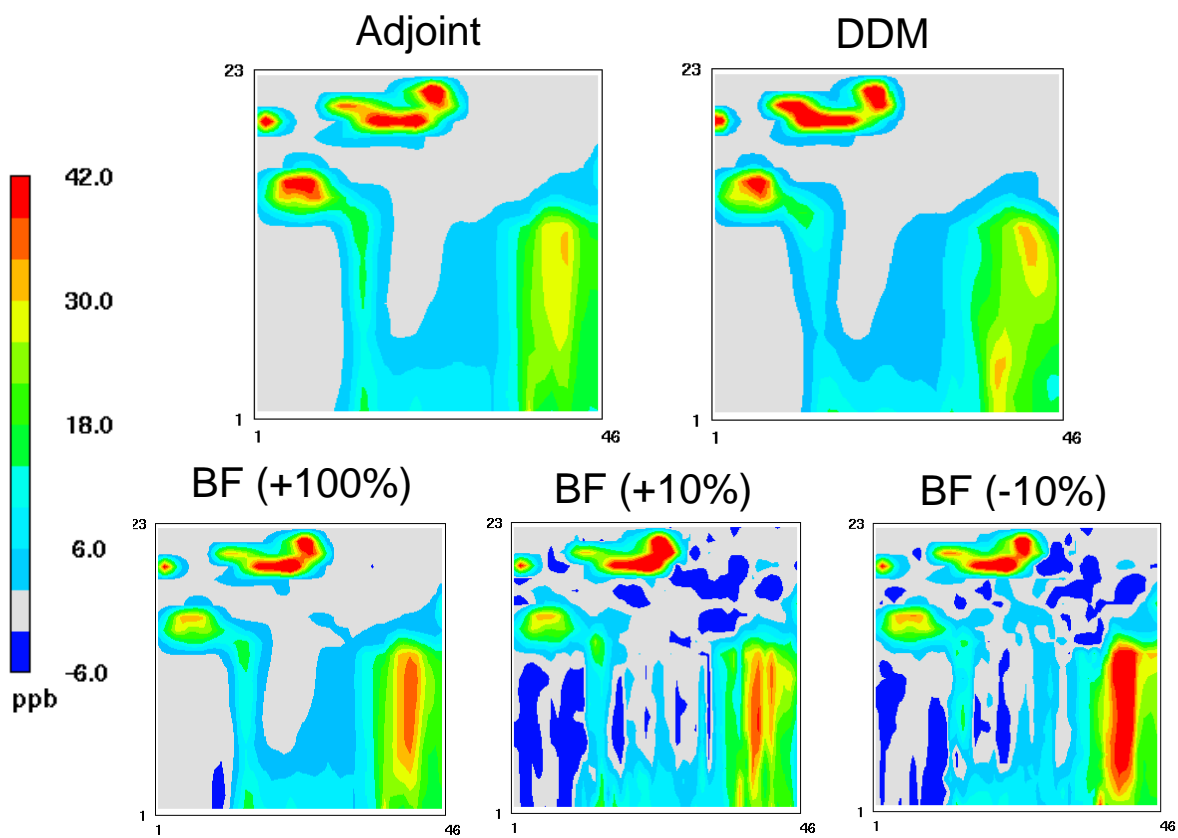


Figure 4: Comparison of BF and adjoint sensitivities of ozone at the 20th column with respect to initial ozone at the 20th column. Only horizontal advection in the x-direction is included in the simulations. DDM sensitivity (bottom) shows reasonable agreement with the adjoint.

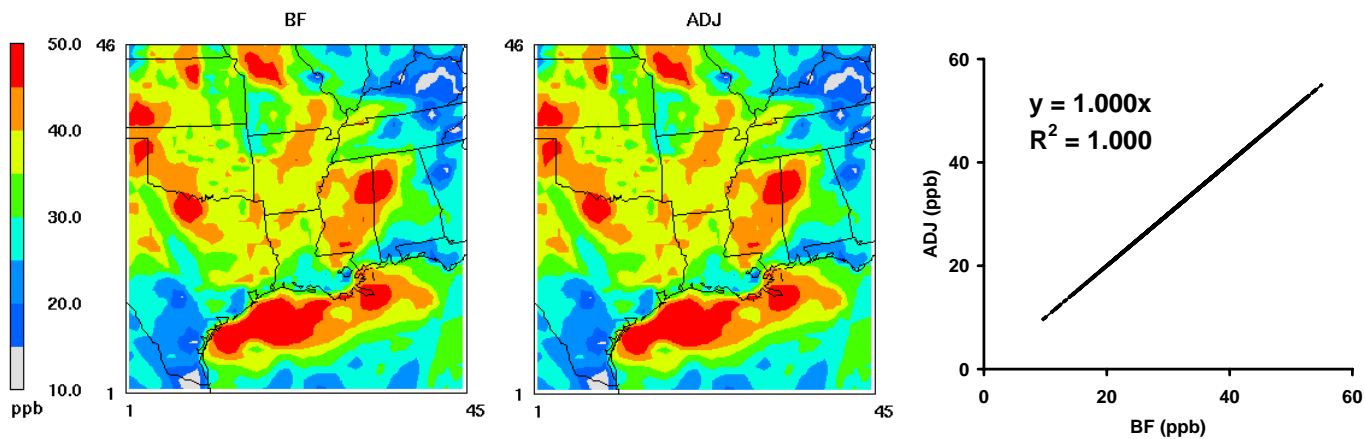


Figure 5: Simulations with only vertical advection for calculation of the normalized sensitivities of surface ozone with respect to initial surface ozone.

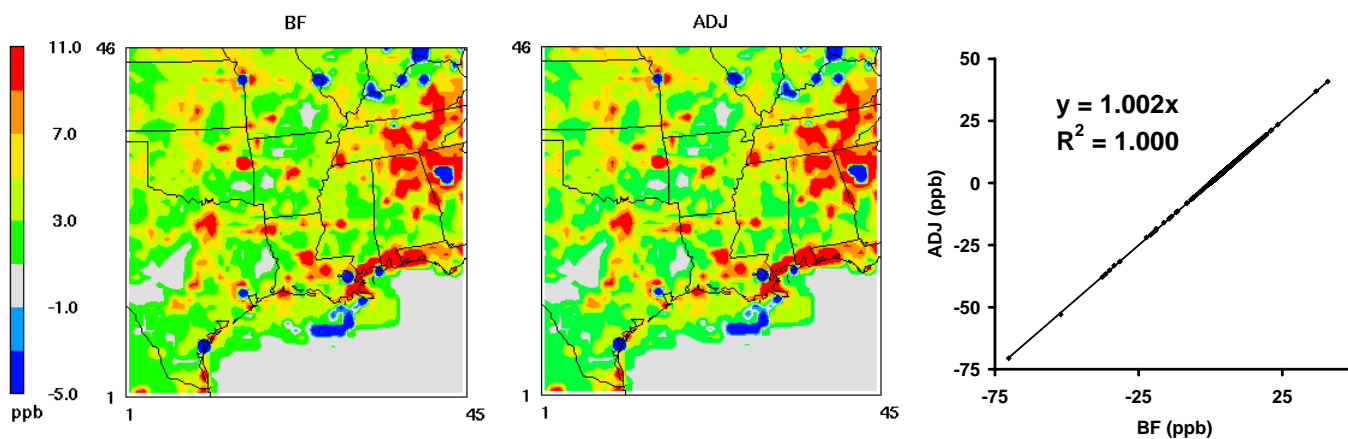


Figure 6: Simulations with only chemistry and vertical diffusion, normalized sensitivities of surface ozone with respect to surface emissions of NO.

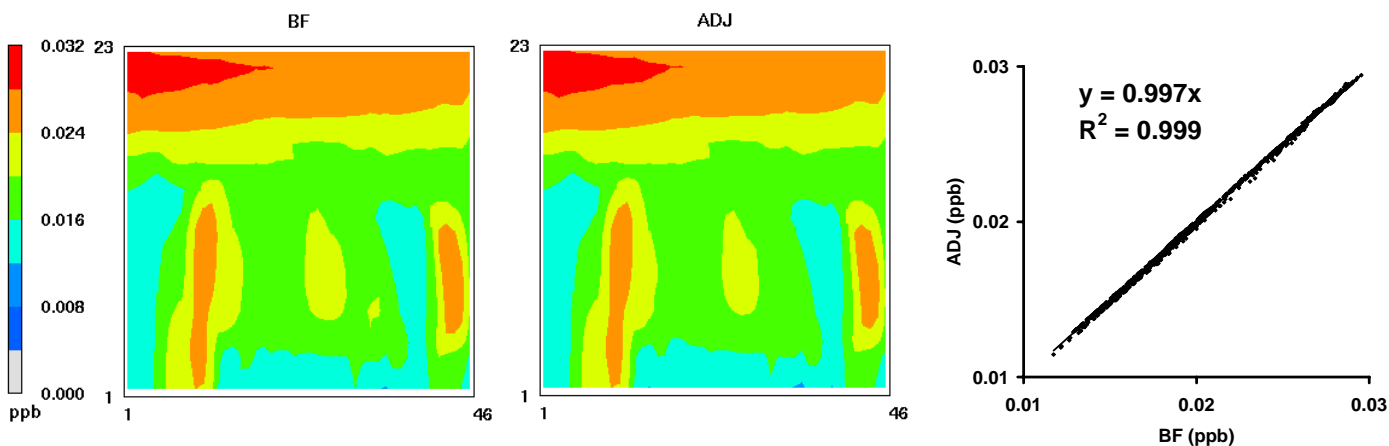


Figure 7: Simulations with only horizontal diffusion in x-direction for calculation of normalized ozone sensitivities at the 21st column with respect to initial ozone at 20th column.

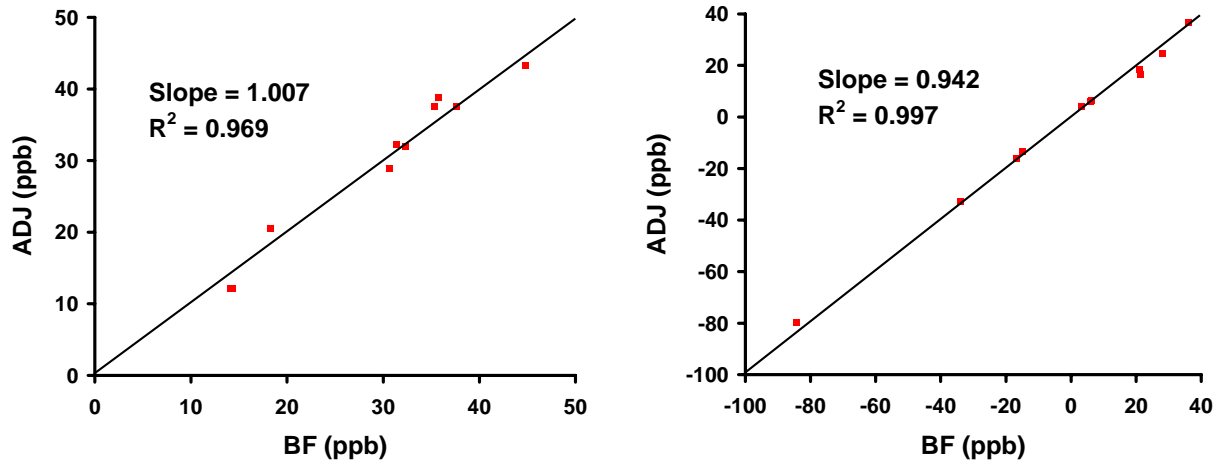


Figure 8: Full model simulations, sensitivities of surface ozone with respect to surface ozone initial concentrations (left) and surface NO emissions (right) for selected locations. Values are shown in comparison to the one-on-one lines.

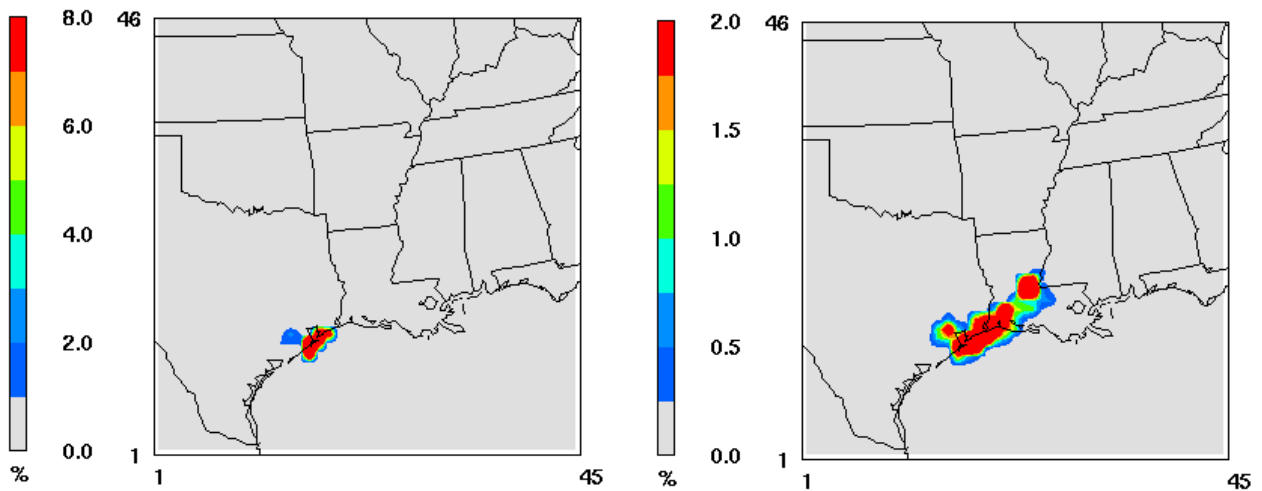


Figure 9: Spatial distribution of the cost function for nonattainment in Houston-Galveston Area (left) and associated gradients with respect to NO_x emissions (right). Values are normalized to the total cost function and presented in percent.

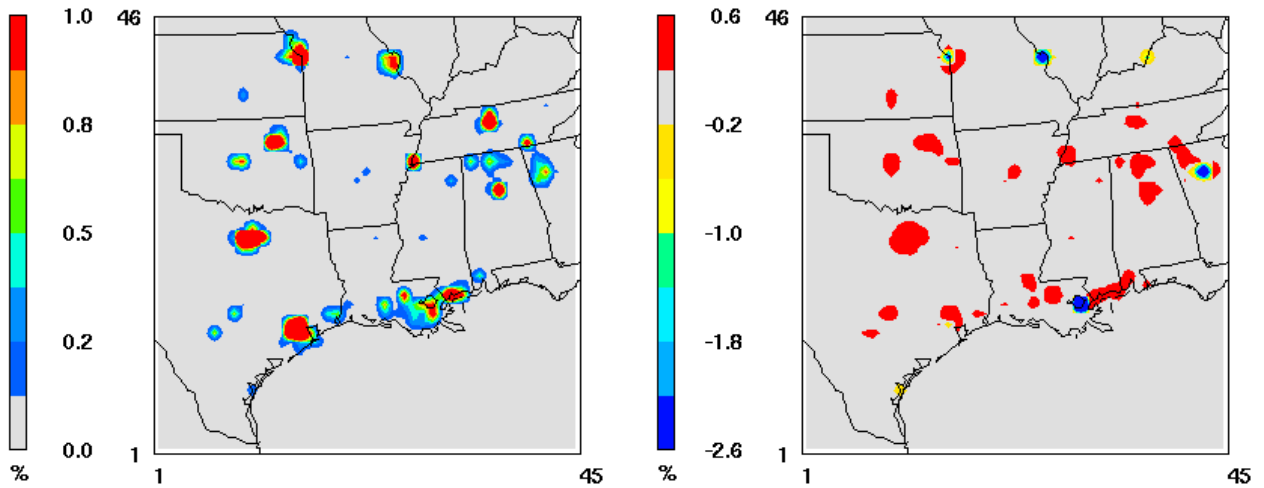


Figure 10: Spatial distribution of the cost function for population exposure (left) and associated gradients with respect to NO_x emissions (right). Values are normalized to the total cost function and presented in percent. Exposure threshold is 60 ppb.

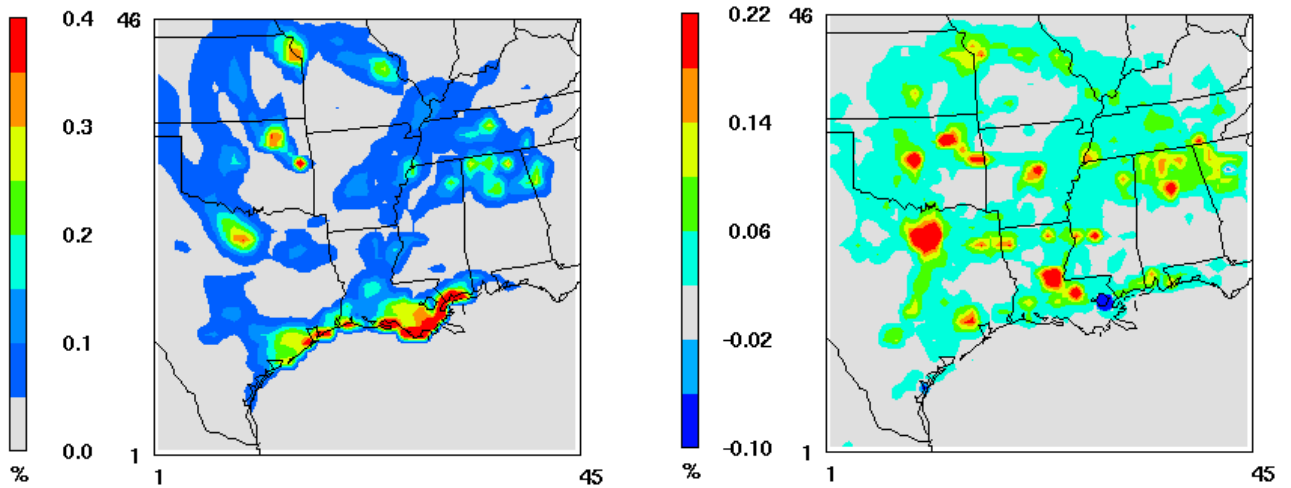


Figure 11: Spatial distribution of the cost function for W-126 metric of vegetation exposure to ozone (left) and associated gradients with respect to NO_x emissions (right). Values are normalized to the total cost function and presented in percent.

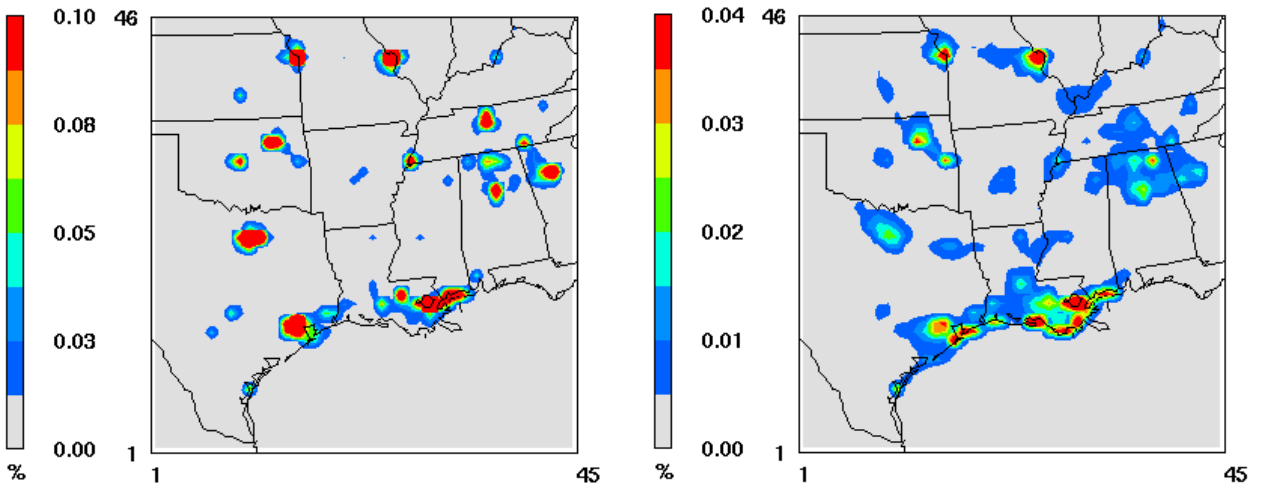


Figure 12: Spatial distribution of the cost function for W-126 metric of vegetation exposure to ozone (left) and associated gradients with respect to NO_x emissions (right). Values are normalized to the total cost function and presented in percent.