



**A new WRF-Chem
treatment for
studying regional
scale impacts**

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A new WRF-Chem treatment for studying regional scale impacts of cloud-aerosol interactions in parameterized cumuli

L. K. Berg, M. Shrivastava, R. C. Easter, J. D. Fast, E. G. Chapman, and Y. Liu

Pacific Northwest National Laboratory, Richland, WA, USA

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Correspondence to: L. K. Berg (larry.berg@pnnl.gov)

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Abstract

A new treatment of cloud-aerosol interactions within parameterized shallow and deep convection has been implemented in WRF-Chem that can be used to better understand the aerosol lifecycle over regional to synoptic scales. The modifications to the model to represent cloud-aerosol interactions include treatment of the cloud droplet number mixing ratio; key cloud microphysical and macrophysical parameters (including the updraft fractional area, updraft and downdraft mass fluxes, and entrainment) averaged over the population of shallow clouds, or a single deep convective cloud; and vertical transport, activation/resuspension, aqueous chemistry, and wet removal of aerosol and trace gases in warm clouds. These changes have been implemented in both the WRF-Chem chemistry packages as well as the Kain–Fritsch cumulus parameterization that has been modified to better represent shallow convective clouds. Preliminary testing of the modified WRF-Chem has been completed using observations from the Cumulus Humilis Aerosol Processing Study (CHAPS) as well as a high-resolution simulation that does not include parameterized convection. The simulation results are used to investigate the impact of cloud-aerosol interactions on regional scale transport of black carbon (BC), organic aerosol (OA), and sulfate aerosol. Based on the simulations presented here, changes in the column integrated BC can be as large as -50% when cloud-aerosol interactions are considered (due largely to wet removal), or as large as $+40\%$ for sulfate in non-precipitating conditions due to the sulfate production in the parameterized clouds. The modifications to WRF-Chem version 3.2.1 are found to account for changes in the cloud drop number concentration (CDNC) and changes in the chemical composition of cloud-drop residuals in a way that is consistent with observations collected during CHAPS. Efforts are currently underway to port the changes described here to WRF-Chem version 3.5, and it is anticipated that they will be included in a future public release of WRF-Chem.

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1 Introduction/motivation

There remains a significant amount of uncertainty related to both the aerosol direct forcing and aerosol indirect effects (e.g. Solomon et al., 2007; Boucher et al., 2013). Numerical models of the atmosphere are one of the common tools used to investigate these effects. High-resolution simulations using horizontal grid spacing less than 10 km, which can explicitly represent convective clouds and cloud-aerosol interactions, have been widely used for short-term studies of cloud-aerosol interactions (e.g. Qian et al., 2009; Wang et al., 2011; Fan et al., 2012). They have not, however, generally been used for long-term simulations because of the associated computational expense. For long-term simulations, coarser horizontal resolution is generally required that necessitates the use of a cumulus parameterization even if the cloud-aerosol interactions associated with sub-grid scale convective clouds are poorly represented (e.g. Zhao et al., 2011). Thus, improved treatments of cloud-aerosol interactions in cumulus parameterizations are needed for investigations of the impact of cloud aerosol interactions (Stevens and Feingold, 2009).

Shrivastava et al. (2013) compared changes in the aerosol chemical composition and cloud microphysical structure associated with cloud-aerosol interactions in fields of shallow cumuli to data collected during the Cumulus Humilis Aerosol Processing Study (CHAPS; Berg et al., 2009). The main goal of CHAPS was to document evidence of cloud-aerosol interactions in fields of shallow cumuli. The simulations presented by Shrivastava et al. (2013) were completed with sufficiently high resolution that a convective parameterization was not required allowing them to investigate cloud-aerosol interactions in relatively small shallow clouds that would be sub-grid scale at coarser resolutions. Among their findings were systematic changes in the chemistry of activated particles and cloud microphysics within shallow cumuli. They found that nitric acid vapor uptake by cloud droplets led to increased nitrate content in the cloud droplet residuals. They also reported changes in cloud microphysical properties, with increases in

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cloud droplet number concentration and decreases in droplet effective radius with an increase in pollutant loading.

The Weather Research and Forecasting model coupled with Chemistry (WRF-Chem) (Grell et al., 2005; Fast et al., 2006) is frequently used to simulate conditions over a range of spatial scales and has been used to study a wide range of atmospheric phenomena associated with atmospheric chemistry and aerosols (e.g. McKeen et al., 2007; Ntelekos et al., 2009; Grell et al., 2011; Pfister et al., 2011; Ahmadov et al., 2012; Matsui et al., 2013). To date, however, the treatment of cloud-aerosol interactions has largely been limited to grid-resolved clouds (Chapman et al., 2009). This is the case for WRF coupled with the Community Atmospheric Model version 5 (CAM5; Neale et al., 2012) physics parameterizations, although cloud-aerosol interactions in convectively detrained stratiform clouds are treated (Ma et al., 2013). One exception is the recent modification of the Grell cumulus parameterization (Grell, 1993; Grell and Dévényi, 2002) to include aerosol interactions in the conversion of cloud water to rainwater and the evaporation of rain (Grell and Freitas, 2013). Lim et al. (2013) added a treatment of aerosol activation to the Zhang–McFarlane parameterization (Zhang and McFarlane, 1995) while Zhao et al. (2013) modified the Kain Fritsch scheme to better account for transport and wet scavenging of dust, but each of their modifications do not include treatment of aqueous chemistry in the clouds nor have they been added to the publicly released version of WRF-Chem. To address this missing process, we have modified WRF-Chem to include treatment of a number of factors and processes important for accurately treating aerosol within sub-grid convective clouds, including: fractional coverage of active and passive clouds, vertical transport, activation and resuspension, wet removal, and aqueous chemistry for cloud-borne particles. It should be noted, however, that the modifications do not include feedbacks of aerosol on the amount of precipitation, impacts of the aerosol on the cumulus microphysics, or feedbacks between the cumulus microphysics and the radiation. These additions are topics for subsequent research.

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The work presented here describes the implementation of a treatment of cloud-aerosol interactions for sub-grid parameterized convective clouds in WRF-Chem that is compared to results presented by Shrivastava et al. (2013). Section 2 describes changes to both the standard cumulus parameterization and the treatment of processes affecting aerosols and trace gases in the sub-grid convective clouds. These changes include improved treatment of cloud fraction as well as treatment of cloud droplet number concentration, vertical transport, activation/resuspension, aqueous phase chemistry, and wet removal. Section 3 provides a description of the WRF-Chem configuration, simulation design, and emissions data used in the study. The data used from CHAPS are presented in Sect. 4. Analysis of the WRF-Chem simulations are presented in Sect. 5. Rather than focusing on the single geographic region, results are presented from three different locations that were selected to highlight the performance of the model in situations with shallow, deep, and a mixture of both types of sub-grid convective clouds; and to document the impact on the regional scale transport, cloud microphysics, and the chemical composition of cloud droplets. It should be noted that Shrivastava et al. (2013) presented an extensive evaluation of the performance of WRF-Chem during CHAPS; therefore, we also compare the results using the new convective cloud parameterization with their high-resolution results.

2 Modifications to WRF-Chem

The primary goal of this effort has been to improve the representation of cloud-aerosol interactions in parameterized sub-grid convective clouds within WRF-Chem. To address this goal, a number of modifications shown schematically in Fig. 1, have been made to WRF-Chem in order to account for cloud-aerosol interactions within these clouds. These modifications include changes to the Kain–Fritsch (KF) cumulus scheme (Kain and Fritsch, 1990; Kain, 2004) as well as changes designed to account for transport, transformation, and removal of aerosols and trace gases within sub-grid convective clouds.

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The WRF-Chem model architecture separates physical processes involving sub-grid cumulus, microphysics for grid-resolved clouds, boundary-layer turbulence, and radiation from processes involving aerosol and trace gases. We have followed this separation, so that code changes involve both a cumulus physics routine that determines the presence of sub-grid convective clouds, their properties, and their impacts on heat, moisture, and momentum, and a separate cumulus aerosol and trace gas routine that treats vertical transport, activation/resuspension, aqueous chemistry, and wet removal of aerosols and trace gases. Modifications to the cumulus physics routine are described in Sects. 2.1.1, 2.1.2, and 2.2.1. The cumulus aerosol and trace gas routine, which is new to WRF-Chem, is described in Sect. 2.2.2.

2.1 Modifications to the Kain–Fritsch cumulus parameterization

2.1.1 Trigger function for convection

Recently, the KF scheme has been modified to improve the treatment of shallow cumuli, which are defined by the KF scheme to be less than 2 to 4 km in height, depending on the temperature at the lifting condensation level. These changes were made primarily within the standard KF (Kain and Fritsch, 1990; Kain, 2004) convective parameterization and involved replacing the default ad-hoc trigger function used in the parameterization with one explicitly linked to the boundary-layer turbulence. This was accomplished using the Cumulus Potential (CuP) scheme (Berg and Stull, 2005) leading to the new KF-CuP parameterization (Berg et al., 2013). These changes were designed to better account for sub-grid variability by applying a range of temperature and moisture perturbations from the grid-box mean as the convective trigger, thus allowing a population of shallow clouds with different thermodynamic properties to coexist in a model grid column. In the case of deep convection, only the single most probable temperature and moisture perturbation that triggered clouds is applied to be consistent with the standard implementation of the KF scheme. The sub-grid distribution of temperature and humidity was parameterized using probability density functions (PDFs) of temperature and

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humidity that were based on the jump of potential temperature and moisture at the surface and at the boundary-layer top (Berg and Stull, 2004). These modifications, along with the cloud fraction changes (Sect. 2.1.2), were shown to significantly increase the frequency of occurrence of simulated shallow clouds over the Southern Great Plains, leading to improved forecasts of both cloud fraction and downwelling shortwave irradiance (Berg et al., 2013). It should also be noted that while the new trigger function is not scale aware, it could easily be modified to adjust the PDF based on the model grid spacing.

2.1.2 Cloud fractional area

In their modifications to the standard KF scheme, Berg et al. (2013) included a simple treatment of the cloud fraction associated with sub-grid scale convective clouds. Their method was based on representative time scales associated with cumulus, which the method defined to be a function of the cloud depth, turbulence intensity, and the moisture in the cloudy layer. In the work presented here, an additional treatment was added to determine the cloud fractional area for instances with deep convection. Rather than develop a new representation of the total cloud fraction for deep convection the empirical treatment used in CAM5 is applied. In this parameterization the cloud fraction associated with deep convection is a function of the convective mass flux (Neale et al., 2012), and is represented as:

$$\sigma_{dp,cu} = k_{1,dp} \ln(1 + k_2 M_{dp,cu}), \quad (1)$$

where $\sigma_{dp,cu}$ is the cloud fraction associated with deep-convective clouds, $k_{1,dp}$ is an adjustable parameter set to 0.1, k_2 is assumed to be 675, and $M_{dp,cu}$ is the updraft mass flux of the convective clouds (in $\text{kg m}^{-2} \text{s}^{-1}$). The values of both $k_{1,dp}$ and k_2 were selected to be the same as the values used in CAM5, and are identical to those used by Ma et al. (2013) in their implementation of the CAM5 physics in WRF and made publically available in version 3.5. It could be argued that a parameterization of cloud

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fraction developed for a relatively coarse resolution model like CAM5 is not appropriate for a regional scale model like WRF, which can be run at a wide range of resolutions. When run at high horizontal resolution, however, the cumulus parameterization is generally turned off so that the parameterization of sub-grid convective cloud fraction is not utilized. Given the constants define above, Eq. (1) predicts the maximum cloud fraction in the grid cell associated with deep convection to be approximately 45 %. Similar to the methodology applied by Berg et al. (2013) for cases of shallow cumuli, the deep-cloud fraction computed using Eq. (1) is applied in the radiation parameterization but has no impact on either the convective tendencies for heat, moisture, momentum or on the cumulus transport of aerosols and trace gases. It is, however, used in the computations related to aqueous chemistry described in Sect. 2.2.

The cloud fraction associated with both shallow and deep sub-grid convective clouds is broken further into two sub-types: active and passive clouds (e.g. Stull, 1985). Active clouds are those that have vigorous updrafts and contribute to the upward cloud mass fluxes. The fractional area of active cumulus for shallow clouds is defined as the fraction of the PDF of temperature and humidity applied in the convective scheme that forms clouds, while for deep clouds it is the standard KF updraft fraction area. Passive clouds consist primarily of decaying clouds without a well-organized updraft. The fractional area of the passive clouds is determined as the difference between the total cloud fraction (computed following Berg et al., 2013, for shallow clouds, and Eq. 1 for deep clouds) and the active cloud fraction that is determined within the KF-CuP scheme. Passive clouds are treated as quiescent and are assigned zero vertical velocity, so that there is no vertical mass flux. They are assumed to have the same total cloud water and ice content as the active clouds, but be non-precipitating, so there is no wet removal associated with passive clouds. In addition, when convective cloud is triggered in a model grid column, the cloud population is assumed to be in steady state over the cloud lifetime defined in the cumulus parameterization (30 min for shallow clouds and 30 to 60 min for deep clouds).

2.2 Modifications to account for cloud-aerosol interactions

Chapman et al. (2009) described a treatment of cloud-aerosol interactions for grid resolved clouds within WRF-Chem. For cloudy grid cells, the standard version of WRF-Chem treats both cloud borne (activated) and interstitial (nonactivated) particles as separate transported species. A number of modifications to the standard WRF-Chem version 3.2.1 have been implemented in this study to specifically address cloud-aerosol interactions in sub-grid convective clouds. These modifications include calculations for:

- cloud droplet number mixing ratio
- cloud microphysical (conversion rates, and cloud water and cloud ice mixing ratios) and cloud macrophysical properties (updraft fractional area, updraft and downdraft mass fluxes, and entrainment) averaged over the population of shallow convective clouds, or for the single deep convective cloud value, and
- vertical transport, activation/resuspension, aqueous chemistry, and wet removal of aerosols and trace gases.

WRF-Chem has several different aerosol and trace gas representations, which are referred to as chemistry packages. Our changes for sub-grid convective clouds were implemented with the Model for Simulating Aerosol Interactions and Chemistry (MO-SAIC; Zaveri et al., 2008) sectional aerosol model and the SAPRC-99 photochemical mechanism (Carter, 2010). Extension to other WRF-Chem chemistry packages would be relatively straightforward, especially those packages for which aqueous chemistry and aerosol activation modules (or interfaces) already exist.

2.2.1 Aerosol effects on the sub-grid cumulus

Within the default KF scheme, as well as other cumulus parameterizations applied in WRF, a highly simplified treatment of cloud microphysics is used. Cloud water is produced in updrafts and converted to precipitation based on a prescribed e-folding height,

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and additional assumptions are made involving frozen condensate and precipitation and detrainment to downdrafts (e.g. Kain and Fritsch, 1990). While such a simplified treatment has been successful for mesoscale weather forecasting, it is not sufficient for studying cloud-aerosol interactions that are intimately linked to the cloud microphysics.

5 Thus, the activation of cloud drops in convective drafts must be considered. The activation is largely a function of the cloud updraft speed. In the modified version of WRF-Chem described here, the updraft velocities associated with the buoyancy excess are computed using the temperature and humidity perturbations for the range of parcels identified by the KF-CuP parameterization that form clouds. Further, the droplet acti-
10 vation for each perturbation is computed by applying an entraining parcel conceptual model using the Abdul-Razzak and Ghan (2000) parameterization modified to account for entrainment following Barahona and Nenes (2007). Once the droplet number concentrations are computed for each perturbation value of temperature and humidity in the PDF, they are averaged together to provide a single value of cloud droplet num-
15 ber concentration for each grid cell. Above cloud base, the number of cloud droplets is further reduced by entrainment, where the entrainment rates are determined using the KF scheme (averaged over all of the parcel perturbations to yield a single entrainment rate). At present, secondary activation is not considered for the sub-grid convective clouds, nor does the activation feedback on the cumulus clouds via changes in the
20 conversion of cloud water to rain (as treated by Grell and Freitas, 2013).

This calculation of droplet number allows for the treatment of the aerosol first indirect effect in sub-grid convective clouds. Currently, this droplet number information is not used by the radiation routines, but this addition is planned as part of our ongoing research on aerosol-cloud interactions.

25 2.2.2 Effects of sub-grid cumulus on aerosol and trace gases

A new module was introduced to WRF-Chem to calculate the effects of sub-grid convective clouds on aerosols and trace gases, including vertical transport, activation/resuspension, aqueous chemistry in cloud droplets, and wet removal. The new

module has separate sections that treat the active clouds (as well as vertical transport in the subsiding environment surrounding the active clouds) and passive clouds (for which the only processes are activation/resuspension and aqueous chemistry).

In models of the cloud (and precipitation) effects on aerosols and trace gases, one must consider the attachment state (Ghan and Easter, 2006) of (aerosol) particles and gases. For example, interstitial aerosol particles (i.e., particles suspended in air) may become attached to, dissolved in, or suspended in various hydrometeors (cloud and rain drops, ice crystals, snow and graupel particles). When the aerosol representation involves several size bins (8 in our study) and multiple chemical species within each bin (14 in our study), the computational expense of explicitly treating all possible attachment states is considerable, and simplifying assumptions are often used. For example, in Chapman et al. (2009) the treatment of cloud-aerosol interactions focused on grid-resolved warm clouds. For all attachment states, aerosol material (sulfate, nitrate, etc.) associated with cloud droplets of grid-resolved clouds was treated explicitly as transported prognostic species, while moderately soluble gases dissolved in cloud droplets were assumed to be in equilibrium with the gas-phase and were treated diagnostically, and aerosol material and gases that became associated with precipitation particles (rain, snow, graupel) and also ice crystals were assumed to be quickly removed from the atmosphere and were not treated explicitly. A similar but somewhat simpler approach is used in our treatment of sub-grid cumulus effects. For all attachment states, the aerosol species associated with cloud droplets in the sub-grid convective clouds are treated explicitly, but only within the convective cloud routines. This approximation is reasonable because of the relatively short life-time of the parameterized convective clouds (30–60 min) and the fact that the parameterization is intended for use with model horizontal grid spacings of approximately 10 km or more. Thus within the convective cloud routines, the treatment of attachment state is similar to that used for grid-resolved clouds in Chapman et al. (2009).

The cumulus physics routine determines if sub-grid convective cloud is present within a model grid column and the physical properties of the cumulus clouds (shallow or

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deep; life-time; updraft and downdraft mass fluxes, entrainment, and vertical velocity; mixing ratios of cloud water, ice, and precipitation; and microphysical conversion of cloud water to cloud ice and precipitation) that are used in the cumulus effects routine. Within the KF-CuP scheme, when deep convection is diagnosed within a grid column, the deep clouds are assumed identical, and there is a single vertical profile for updraft and downdraft mass fluxes and each microphysical parameter. When shallow convection is diagnosed, there is a population of shallow clouds with different profiles, and downdrafts are not treated. In the cumulus-effects-on-aerosols routine, calculations are made using the properties of an average (over the population) shallow cloud, rather than doing calculations for each shallow cloud in the population.

Active cloud calculations are performed first, followed by passive cloud calculations. The treatment of active sub-grid cumulus effects on aerosols and gases is very similar to the unified treatment described in the Supplement of Wang et al. (2013). The active-cloud updrafts and downdrafts are treated as steady-state entraining plumes. The updraft and downdraft mass fluxes obey

$$\frac{\partial M_Y}{\partial z} = \frac{(E_Y - D_Y)}{\Delta z} \quad (2)$$

where the Y subscript is either U for updraft or D for downdraft, M_Y is the mass flux ($\text{kg m}^{-2} \text{s}^{-1}$) defined at vertical layer boundaries, and E_Y and D_Y are the entrainment and detrainment in a layer, and Δz is the layer thickness. The compensating mass flux in the environment, M_E , is equal to $-(M_U + M_D)$. The active-cloud calculations involve integrating conservation equations for grid-cell mean mixing ratios of aerosol and trace gas species over the lifetime of the cumulus cloud. A time sub-step is used such that the transport of air out of a layer (by M_E , E_U , and E_D) during the sub-step does not exceed the layer's air mass $\rho \Delta z$, where ρ is the air density.

For each time sub-step, steady-state vertical profiles of aerosol and trace gas species in the updraft and downdraft are first calculated. This is done by integrating steady-state continuity equations upwards (for updrafts) or downwards (for downdrafts).

For aerosol species in the updraft, the continuity equation is

$$\frac{\partial(M_U q_{X,U})}{\partial z} = \frac{(E_U q_{X,E} - D_U q_{X,U})}{\Delta z} + \rho A_U [(\dot{q}_{X,U})_{\text{ACTI}} + (\dot{q}_{X,U})_{\text{WETR}} + (\dot{q}_{X,U})_{\text{AQCH}}] \quad (3)$$

Here $q_{X,E}$ and $q_{X,U}$ are aerosol mixing ratios in the environment (E) and updraft (U), respectively, the X subscript is either AI for interstitial aerosol species or ACC for convective-cloud-borne (activated) aerosol species. The environment mixing ratios for interstitial aerosol are assumed equal to the grid-cell mean values, and are zero for convective-cloud-borne aerosol. A_U is the updraft fractional area and is equal to $(M_U/\rho w_U)$, where w_U is the updraft vertical velocity. The last three terms on the right hand side are the rates of change due to activation (ACTI), in-cloud wet removal (WETR), and aqueous-phase chemistry within cloud droplets (AQCH). For interstitial aerosol, only the activation term is non-zero.

Aerosol activation is calculated as described in Sect. 2.2.1, but for shallow convective clouds, the average (over different clouds) vertical velocity is used. The Abdul-Razzak and Ghan (2000) parameterization provides activation fractions (f_{ACT}) for aerosol number and mass species in each size bin. The activation rate in Eq. (3) is then

$$(\dot{q}_{\text{ACC,U}})_{\text{ACTI}} = -(\dot{q}_{\text{AI,U}})_{\text{ACTI}} = (f_{\text{ACT}} q_{\text{AI,U}})/\Delta t_U \quad (4)$$

where $\Delta t_U = \Delta z/w_U$ is the time for updraft air to move across a layer.

The wet removal rate for cloud-borne aerosol in Eq. (3) is given by

$$(\dot{q}_{\text{ACC,U}})_{\text{WETR}} = -(f_{\text{WETR}} q_{\text{ACC,U}})/\Delta t_U \quad (5)$$

where f_{WETR} is the fractional removal of cloud-borne aerosols in the updraft as they move across a layer. This fractional removal is currently equal to the fractional conversion of cloud-water to precipitation across the layer, which is provided by the cumulus physics routine. Cloud water can also be converted to cloud ice, but currently this is not included as part of the aerosol wet removal, as the fate of cloud ice (conversion to

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precipitation or detrainment near cloud top) can vary. In the future, ice processes could be incorporated in the cumulus effects routine by treating cloud-ice-borne aerosol in addition to cloud-droplet-borne aerosol.

The aqueous-phase chemistry rate in Eq. (3) is obtained by calling the WRF-Chem cloud-chemistry routine for grid-resolved clouds (Chapman et al., 2009). This routine calculates mixing ratio changes from gas uptake and aqueous-phase reactions in an air parcel (or layer) over a specified time step, and it is applied to updraft air moving across a layer in time Δt_U .

For trace gases in the updraft, the continuity equation is

$$\frac{\partial(M_U q_{G,U})}{\partial z} = \frac{(E_U q_{G,E} - D_U q_{G,U})}{\Delta z} + \rho A_U [(\dot{q}_{G,U})_{WETR} + (\dot{q}_{G,U})_{AQCH}] \quad (6)$$

where $q_{G,E}$ and $q_{G,U}$ are gas mixing ratios in the environment and updraft, respectively. The environment gas mixing ratios are assumed equal to the grid-cell mean values. The $q_{G,U}$ includes both gas-phase and dissolved in convective cloud-water species (e.g., gaseous SO_2 plus S(IV) in cloud water). The WRF-Chem cloud-chemistry routine gives the aqueous-phase chemistry rate in Eq. (6), as well as the fraction of the gas that is dissolved in convective cloud-water ($f_{G,CCW}$). The wet removal rate for gases only considers the removal of gases dissolved in cloud droplets; uptake of gases by rain is currently neglected. The wet removal rate in Eq. (6) is then

$$(\dot{q}_{G,U})_{WETR} = -(f_{WETR} f_{G,CCW} q_{G,U}) / \Delta t_U \quad (7)$$

Downdrafts are assumed to be sub-saturated and contain no cloud droplets or convective-cloud-borne aerosol. Thus activation, wet removal, and aqueous-phase chemistry are not treated in downdrafts. The downdraft continuity equations are then

$$\frac{\partial(M_D q_{X,D})}{\partial z} = \frac{(E_D q_{X,E} - D_D q_{X,D})}{\Delta z} \quad (8)$$

where X is either AI for interstitial aerosol species or G for gases.

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Once the aerosol and gas profiles in the updraft and downdraft have been calculated, conservation equations for grid-cell mean mixing ratios of aerosol and trace gas species are integrated for the time sub-step. These conservation equations have the form

$$\rho \frac{\partial \bar{q}_X}{\partial t} = -\frac{\partial}{\partial z} [M_U q_{X,U} + M_D q_{X,D} + M_E q_{X,E}] + \rho A_U [(\dot{q}_{X,U})_{ACTI} + (\dot{q}_{X,U})_{WETR} + (\dot{q}_{X,U})_{AQCH}] \quad (9)$$

where the X subscript is either AI, ACC, or G, and the updraft rate of change terms come from the updraft calculations described above. The integration is explicit in time and uses simple upstream finite differencing for the vertical transport terms. After the integration sub-step, the grid-cell mean mixing ratio of convective-cloud-borne aerosol (\bar{q}_{ACC}) may be non-zero at or near levels where the updraft detrains. This convective-cloud-borne aerosol is partially transferred to grid-resolved cloud-borne aerosol (fraction transferred equal to grid-resolved cloud fraction) and partially resuspended to interstitial aerosol. At the end of all the active-cloud integration sub-steps, the new grid-cell mean aerosol and gas mixing ratios reflect the effect of the active cumulus cloud over the cloud lifetime.

The passive cumulus effects calculations are performed next. These calculations are relatively simple in comparison, as there is no vertical transport or wet removal of aerosol. The cumulus physics routine provides the passive cumulus cloud fraction and cloud water mixing ratio at each vertical level. Initial mixing ratios of interstitial aerosol and trace gases are set equal to the grid-cell mean mixing ratios at the end of the active cumulus effects calculation. Some of the interstitial aerosol is then activated, in order to provide an initial chemical composition of the cloud water. Because vertical velocity is assumed zero in the passive clouds, the Abdul-Razzak and Ghan (2002) parameterization cannot be used. Instead, we assume that the activated fraction for each aerosol chemical component (and size bin) is the same as the activated fraction in the steady-state updraft of the active cumulus. Aqueous-phase chemistry calculations

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are then made for this passive cloud fraction, again over the lifetime of the cumulus. Finally, the passive cumulus fraction of the grid cell is mixed with the remainder of the grid cell, and convective-cloud-borne aerosol is partially transferred to grid-resolved cloud-borne and partially resuspended to interstitial.

After the passive cloud calculations, the grid-cell mean mixing ratios of aerosols and trace gases reflect the effects of active and passive cumulus over the cloud lifetime. These mixing ratios are returned to the host code as the updated mixing ratios. In our simulations, a primary time step (for dynamics) of 15 s was used, and a chemistry time step (for most processes involving trace gases and aerosols) of 5 min was used. The sub-grid cumulus lifetimes, as defined within the cumulus parameterization, ranged between 30 and 60 min, and the cumulus effects on aerosols/gases are calculated once only when a cumulus is triggered in a grid column. On subsequent chemistry time steps, no more cumulus effect calculations are performed until a new cumulus is triggered in a column. An alternate approach would be to save the cumulus effects tendencies for aerosols and gases, then apply them gradually over the cumulus lifetime, analogous to the approach used in the cumulus physics for temperature, moisture, and momentum. We chose this one-time update approach for aerosols and gases for simplicity and to reduce memory costs associated with storing the cumulus effects tendencies for the many aerosol and gas species. The net changes to the aerosol would be the same in either case, but the changes are applied somewhat sooner in the once-only approach (when a cloud triggers rather than over its lifetime), producing small differences in a simulation that could grow over time.

3 WRF-Chem configuration

3.1 Experiment setup

WRF-Chem version 3.2.1 was configured in a way similar to that described by Shrivastava et al. (2013). A single domain, 2240 km on a side, over the central United States

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was used with 10 km horizontal grid spacing. WRF-Chem was also configured to use 64 vertical levels, with approximately 25 levels in the lowest 1 km of the atmosphere. The various parameterizations utilized in the simulations, not including the modifications described in Sect. 2, are listed in Table 1. Multi-day WRF-Chem simulations for the period of 1 June through 30 June 2007 were completed in individual 36 h blocks. The first 12 h of each block were discarded and the final 24 h saved for analysis. Meteorological initial and boundary conditions for each block were taken from the Global Forecast System (GFS). Boundary conditions of trace gases and aerosols were derived from the MOZART global simulation (Emmons et al., 2010b). Initial conditions for trace gases and aerosol were taken from the end of the previous simulation block.

Three sets of simulations are used to investigate the regional impacts of cloud-aerosol interactions associated with both shallow and deep convection (Table 2). In all three simulations, the shallow and deep cumulus physics are enabled. However, the cumulus effects on aerosols and trace gases are selectively enabled in the different simulations. The first simulation includes aerosol processing associated with both shallow and deep clouds, and is referred to as DeepShallow. This simulation can be used to estimate the regional impact on aerosol properties due to cloud processing associated with all clouds in the domain (including both grid resolved and parameterized clouds). The second simulation has aerosol processing by shallow convection turned on and by deep convection turned off, and is referred to as ShallowOnly. The difference between DeepShallow and ShallowOnly is used to document the impact of aerosol processing by deep convection alone and is identified as the Deep-Effect in this work. The third simulation is conducted with all aerosol processing by sub-grid convective clouds turned off (Control) and is the default treatment in WRF-Chem. The difference between ShallowOnly and Control simulations show the impact of shallow clouds and will be identified as the Shallow-Effect in the rest of the manuscript. An additional simulation was completed for a subset of the study period to document the impact of aqueous phase cloud chemistry on aerosol composition. This was accomplished by repeating the DeepShallow simulation for 25 June 2007 with the convective

cloud aqueous chemistry turned off. This run was initialized using the aerosol from the end of the previous DeepShallow simulation block.

3.2 Emissions

Hourly emissions used in this study are the same as those used by Shrivastava et al. (2013). In brief, hourly emissions of aerosol and trace gases are derived for the desired 2007 period by assuming a linear variation in the US Environmental Protection Agency's National Emissions Inventory (NEI; e.g., <http://www.epa.gov/ttn/chief/net/2005inventory.html>) for 2005 and 2008, supplemented with biomass burning gas and aerosol emissions taken from the 2007 Fire Inventory produced by NCAR (FINN07) (Wiedinmyer et al., 2011). The NEI contains two sizes of particulate matter emissions: particles with diameters less than or equal to $2.5\ \mu\text{m}$ ($\text{PM}_{2.5}$) and those less than or equal to $10\ \mu\text{m}$ (PM_{10}). NEI $\text{PM}_{2.5}$ emissions are divided into categories of sulfate, nitrate, organic aerosol, elemental carbon, and unspiciated primary $\text{PM}_{2.5}$, following Hsu et al. (2006). As in Shrivastava et al. (2013), all unspiciated $\text{PM}_{2.5}$ is lumped into the MOSAIC other inorganic material (OIN) category. For the simulations presented here, OIN accounts for approximately 77% of the $\text{PM}_{2.5}$ mass emissions. The MOZART model (Emmons et al., 2010a) was used to provide the inflow of dust through the boundaries of the WRF-Chem domain with these values assumed to be OIN. $\text{PM}_{2.5}$ and PM_{10} emissions are mapped to eight size bins for the sectional size distribution representation following Fast et al. (2006). Particles in each size bin are assumed to be internally mixed and the same size distribution is assumed for all species. VOC emissions were speciated using the SAPRC-99 mechanism and biogenic VOC emissions are estimated using the Model of Emissions of Gases and Aerosols from Nature (MEGAN <http://bai.acd.ucar.edu>) (Guenther et al., 2006). The 138 biogenic species in MEGAN are grouped into three classes for use with WRF-Chem. Primary emissions are further modified to account for semi-volatile and intermediate volatility organic compounds (S/IVOC) that are large potential anthropogenic SOA precursors and are co-emitted with primary organic aerosols (POA) (Shrivastava et al., 2008). In this study, emissions

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of SVOC are assumed to be twice that of POA for anthropogenic sources, while IVOC emissions are estimated to be 1.5 times the sum of SVOC and POA emissions, for a total S/IVOC emissions equal to 6.5 times POA (Hodzic et al., 2010; Tsimpidi et al., 2010; Shrivastava et al., 2011). A two-species VBS mechanism is used here, with both POA and secondary organic aerosols (SOA) assumed to have a very low volatility (Shrivastava et al., 2011). In previous work, Shrivastava et al. (2013) showed that this 2-species VBS mechanism resulted in reasonable predictions of organic aerosols compared to measurements made during CHAPS, as described in the next section.

4 Data

In this study, a subset of model results are compared to data collected during CHAPS, which was conducted during June 2007 and included the deployment of the Department of Energy's Gulfstream-1 (G-1) aircraft. During CHAPS the G-1 was configured for in situ sampling of aerosol chemical and optical properties (Berg et al., 2009). The flight path was specifically designed to measure conditions below, within, and above a population of shallow cumuli near Oklahoma City, Oklahoma. The size distribution of aerosol and cloud drops was measured using a Droplet Measurement Technology (DMT) Passive Cavity Aerosol Spectrometer Probe (PCASP) and DMT Cloud Aerosol Spectrometer (CAS). The G-1 was equipped with two aerosol inlets: an isokinetic inlet for sampling aerosol in clear air and interstitial aerosol within clouds, and a Counter Flow Virtual Impactor (CVI) to sample only cloud droplets. An Aerodyne Aerosol Mass Spectrometer (AMS) was used to analyze the composition of non-refractory aerosol sampled via both inlets. In their work, Shrivastava et al. (2013) evaluated the performance of WRF-Chem for the same period and found reasonable agreement with the observations when the model was run with relatively fine spatial resolution that explicitly represented convection. They reported some discrepancies between the simulated and observed aerosol optical properties, but these were attributed to assumptions related

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were selected because they contain primarily shallow convection (MSN), deep convection (AUS), or a mixture of both (OKC) (see Fig. 3) and allow us to better understand the behavior of the model and its parameterizations over a range of conditions. The MSN box has a very high frequency of shallow clouds distributed over the box with the nearest up-stream deep convection occurring over central Illinois. The AUS box has a very small frequency of simulated sub-grid shallow clouds and a much larger frequency of simulated sub-grid deep convective clouds. In contrast to the other two boxes, the OKC box includes a mixture of both shallow and deep convection.

While MOSAIC represents multiple aerosol constituents, only BC, OA, and sulfate have been selected for analysis within the three boxes. These particular constituents were selected because of their climatic relevance, and their representative behavior. BC is, to a first approximation, only impacted by transport, activation/resuspension, dry deposition, and wet removal – and in the case of non-precipitating convection acts essentially as a passive tracer. Although freshly emitted BC is hydrophobic, the internal mixing assumption applied in the model causes it to quickly reside in hygroscopic particles. Interpretation of cloud-aerosol interactions and vertical sulfate transport is more complicated than for BC because sulfate can be produced within cloud droplets via aqueous-phase oxidation of dissolved sulfur dioxide gas as well as removed via precipitation (e.g. Koch et al., 2003). While the majority of OA in the atmosphere is secondary and is somewhat hygroscopic, its behavior within convective clouds is similar to that of BC aerosol because the aqueous chemistry related to OA production is not fully understood and currently is not included in the model.

5.1 Local impacts on aerosol vertical distribution

One important impact of convective clouds is the vertical redistribution of aerosol due to the impact of convective updrafts, downdrafts, entrainment mixing, enhanced subsidence, and wet removal associated with sub-grid clouds. Figure 4 shows examples of vertical north-south cross sections (through the center of the analysis boxes) of the amount of BC (including both interstitial and activated aerosol in the cloudy grid

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cells) for the DeepShallow case and the fractional change in BC loading between the DeepShallow and control simulations (indicated by the colors) as well as the cloud fraction (indicated by the gray shading) within the AUS and MSN boxes valid at 20:00 UTC on 25 July 2007. Within both the AUS and MSN boxes the largest BC mass loadings are found near the surface. There are also large amounts of BC 4–6 km above the surface in the AUS cross section that is apparent in both the DeepShallow (Fig. 4) and Control simulations (not shown). This elevated layer is not associated with convection but rather with long-range transport, most likely from a fire located central New Mexico (not shown) and a coal-fired power plant in Colorado.

At first glance it might be surprising that there are not columns of enhanced aerosol loading within the AUS clouds due to enhanced upward transport from the sub-cloud layer shown in Fig. 4. Their absence is primarily due to the wet removal of aerosol within the lowest levels of the clouds, as well as the cloud fraction (which ranges from 20 to 60% within the deep convective clouds shown in the figure), which reduces the relative impact of the aerosol in the updraft, within any given model grid cell. In the AUS cross section, the large fractional increase in BC between the DeepShallow and Control simulations for altitudes ranging from 3 and 5 km and the decrease above 5 km can be attributed to vertical transport by updrafts, downdrafts, and convection induced subsidence. At these altitudes (which are below the detrainment level), this transport replaces some of the air (and aerosol) in a grid cell with air from higher levels that has smaller BC concentrations.

Within the AUS cross section, the clouds extend from an altitude of approximately 0.5 km to nearly 15 km. The clouds in the MSN box are much shallower, extending from approximately 1 to 2 km as is more typical for boundary-layer cumuli (e.g. Berg and Kassianov, 2008). The decrease in amount of BC loading in the sub-cloud layer is caused by the venting of aerosol out of that layer by the convective clouds. In contrast to the AUS box that includes deep sub-grid convective clouds, the vertical extent of the transport of BC is more limited within the MSN box (Fig. 4b). This result is consistent with the much smaller vertical extent of the clouds in this box. Within the cloud layer, the

shallow cumuli still have an important impact on the vertical extent of the BC (Fig. 4d). The fractional difference in the BC between the DeepShallow and Control simulations approaches 50 % as the convective clouds transport BC from below the cloud into the cloud layer. The net effect of the non-precipitating cumuli is to mix BC over the sub-
5 cloud and cloud layers.

Similar to the case for BC, there is an elevated plume of sulfate aerosol near an altitude of 5 km in the AUS cross-section that is associated with long-range transport (Fig. 5a). In both the AUS and MSN cross-sections there is a large concentration of sulfate within the boundary layer that is associated with surface emissions. As with
10 BC, fractional differences between the DeepShallow and Control runs are much larger than 50 %. Within the AUS box there is a large fractional change in the amount of sulfate aloft that can be attributed to vertical transport by updrafts, downdrafts, and convection induced subsidence that are represented in the DeepShallow simulations (Fig. 5c). The situation is different in the MSN box, where all of the clouds are shallow
15 non-precipitating cumuli (Fig. 5b and d). In this case, the vertical transport is limited to the cloud layer (altitudes lower than approximately 2 km), where there is significant increase in the sulfate loading in the cloud layer (Fig. 5b). In contrast to the BC within the MSN box, the sulfate is enhanced in the ShallowOnly simulations both below and within the cloud layer. This is due to sulfate production within clouds, the detrainment
20 of cloudy air with enhanced sulfate, and subsequent downward transport of air back into the subcloud layer. There is no evidence of lofted sulfate in the levels above the shallow cumuli (Fig. 5b).

While the cross sections of aerosol loading are instructive and highlight important processes associated with both clouds and aerosol, they are not necessarily representative of conditions occurring over the entire area of the AUS, OKC and MSN boxes. The average vertical profiles of aerosol loading, and the associated fractional changes of BC, OA, and sulfate between the various simulations have been computed using
25 WRF-Chem output valid at 20:00 UTC on 25 July for each box.

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Horizontal averaged aerosol loading in the AUS box is shown in Fig. 6. Consistent with the results shown in Figs. 4 and 5, there is a systematic decrease of each species in the sub-cloud layer (altitudes below 0.5 km) in the DeepShallow simulations that is caused by the venting of aerosol-rich air by the deep convective clouds, and the profile of aerosol loading between 1 and 5 km is similar in shape to the Control simulations but is systematically shifted downward (which is most obvious for sulfate; Fig. 6c). The effects of deep and shallow convection shown in Fig. 6 are calculated as $[100 \cdot (\text{DeepShallow} - \text{ShallowOnly}) / \text{ShallowOnly}]$ and $[100 \cdot (\text{ShallowOnly} - \text{Control}) / \text{Control}]$, respectively. The importance of deep convection (black lines in panels b, d, and f of Fig. 6) is clearly much larger than the effect of shallow convection for the AUS box. Within the AUS box, there is a 15 to 40 % decrease of BC, OA, and sulfate within the sub-cloud layer associated with venting of air from the boundary layer by the convection in the simulations associated with the effect of the deep clouds (Fig. 6b, d, and f).

The “S” shaped portion of the profile showing the effects of deep clouds on the vertical profile of BC, OA, and sulfate between heights of 0.5 and 1.5 km in Fig. 6 is related to how the numerator (DeepShallow minus ShallowOnly) and the denominator (ShallowOnly) are both decreasing but at different rates. Near the top of the boundary layer (near an altitude of 0.5 km), the fractional change is enhanced in magnitude due to the downward transport of air with smaller aerosol loading leading to changes of approximately -30% for the BC and OA and -20% for sulfate. Just above the boundary layer (for altitudes ranging between 0.5 to 1.5 km), the fractional difference is reduced (as shown by the near overlap of all three simulations near a height of 0.75 km; Fig. 6a, c and e) consistent with the relatively small differences in the mass loading with height for the different simulations at these altitudes. Similarly, convection-induced transport, mixing and subsidence is likely responsible for the net decrease in aerosol mass between 1 and 2 km and the net increase in mass between 2 and 5 km as the profile of BC, OA, and sulfate has a similar shape to the Control simulations, only shifted downward in height.

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In contrast to the large amount of deep convection within the AUS box, the MSN box is dominated by shallow convection and the impact on the vertical profile of BC, OA, and sulfate is limited to heights below approximately 2 km due to the limited vertical extent of the clouds (Fig. 7a, c, and e). From the surface to approximately 0.75 km, shallow clouds reduce both BC and OA by about 10 %. In contrast, the effect of shallow clouds on the amount of sulfate in the sub-cloud layer is to increase the loading by more than 20 % compared to the control simulations. Within the cloud layer the BC and OA increase by 50 % and 40 %, respectively due to the lofting of aerosol by the clouds. The increase in sulfate is much larger than either BC or OA and is as large as 70 %. This increase is due to the production of sulfate within cloud droplets, followed by evaporation of detrained droplets.

The MSN box is dominated by shallow sub-grid convective clouds, but deep convection has a small but noticeable impact on the simulated aerosol mass loading within the box, as shown by the effects of both deep and shallow convection (Fig. 7b, d and f). The effect of deep convective clouds is to somewhat reduce the aerosol mass loading in the subcloud layer. There is, however, a larger decrease in the BC aerosol, OA, and sulfate aerosol within and near the top of the cloud layer (occurring at a height slightly higher than the largest changes associated with shallow convection) that indicate the role of regional transport processes and/or removal of aerosol mass upstream of the MSN box.

While the AUS and MSN boxes are dominated by deep or shallow convection, respectively, both cloud types are present in the OKC box, and cloud effects on aerosols within that box are expected to lie somewhere between those found in the AUS and MSN boxes. This is generally the case, with large fractional changes associated with both shallow and deep convection (Fig. 8b, d and f). It is interesting to note that the impact of shallow convection extends to a greater height (nearly 6 km) in this box than in the MSN box. Given that shallow cumuli are limited to a depth of 4 km in the KF-CuP scheme (as described in Sect. 2.2.1) the changes in aerosol loading between altitudes

of 4 and 6 km are attributable to a combination of vertical transport by the shallow cumuli and regional scale lifting that could occur upwind of the analysis box.

For the OKC box, the results from the simulations with parameterized convection can also be compared to the high-resolution simulations presented by Shrivastava et al. (2013) valid at 20:00 UTC on 25 June 2007 for the OKC box (Fig. 8). Some differences between the low resolution and high-resolution simulations are likely due to the averaging of the emissions over larger grid cells that produce smaller horizontal gradients in emissions that could lead to systematic differences in the aerosol loading. There are also differences in the simulated cloud field. For example, the grid-resolved simulations were free of deep convection (i.e., grid resolved clouds that one would interpret as deep convection) within the OKC analysis box while the low-resolution simulations presented here predicted a large amount of deep convection in the same box (not shown). Another factor is the model configuration used by Shrivastava et al. (2013) for their outer domain. Their simulations used only the standard KF scheme on the outer domain, which does not include a treatment of cloud-aerosol interactions in convective clouds. This could lead to differences in the aerosol loading upwind of the high-resolution domain. All of these differences could lead to inconsistencies in the predicted aerosol loading aloft in the grid-resolved and DeepShallow simulations as well as differences in the wet removal of aerosol associated with precipitating deep convection. The control simulation (which ignores cloud-aerosol interactions within sub-grid cumuli) has the largest aerosol loading within the boundary layer (below 0.5 km) regardless of the constituent of interest. In contrast, the DeepShallow simulations have the smallest aerosol mass loading within the boundary layer, and, in the case of BC and OA, slightly larger values above a height of 2 km. Similar to the results for the AUS box there is a decrease in the mass loading of sulfate both within the boundary layer and to an altitude of approximately 1.5 km. In contrast to the results from the MSN box, there is no evidence of sulfate enhancement by the shallow cumuli within the sub-cloud layer, as shown by the negative effect of the shallow clouds below an altitude of 1 km (Fig. 8d).

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This is likely due to differences in the fractional cover of shallow sub-grid convective clouds between the two boxes (Fig. 9).

5.2 Regional scale impacts

The results presented in Sect. 5.1 highlight that the parameterization is performing reasonably and can be used to investigate the regional impacts of cloud-aerosol interactions within the areas defined by the analysis boxes. The primary advantage of using a parameterization to represent convective clouds is the ability to run simulations over a large domain, which enables the evaluation of regional scale impacts of cloud-aerosol interactions that is not possible using high-resolution simulations. Differences in the column-integrated mass loading are one method that can be used to investigate changes in mass loading of atmospheric aerosol over large areas. BC represents particles that are essentially passive tracers (ignoring wet and dry removal) that do not undergo aqueous phase chemistry in simulated clouds. Overall, there is a significant reduction in the column integrated BC and OA across the model domain (Fig. 10). The primary removal mechanism added in the DeepShallow simulations (compared to the Control simulations) is the wet removal associated with the parameterized precipitation. This leads to systematic decreases of as much as -50% in the amount of BC. It is interesting to note that there is a net decrease of BC within the MSN box in which there is no convection and very little grid resolved precipitation, indicative of wet removal upwind of the box during the simulation and pointing to regional scale impacts of cloud-aerosol interactions. There are also small areas in which the column integrated BC loading is larger in the DeepShallow than control simulations. These features are also present in the ShallowOnly case (not shown). The increase in the column integrated BC in the AUS box is the result of slight differences in the path of the aerosol plume coming from the Houston, Texas area. Different aerosol loadings in the simulations produce different feedbacks on meteorology (i.e aerosol indirect effects in grid-resolved clouds and aerosol direct effects), leading to small differences in winds. In the DeepShallow

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simulations the main part of the plume is shifted a small distance to the north, giving rise to the apparent increase in the BC loading just downwind of Houston.

The OA follows a pattern similar to what is seen for BC, but the fractional change is smaller in magnitude. Currently in WRF-Chem the OA are unaffected by aqueous chemistry within the clouds, but can be affected by changes in the amount of precursor gases. Vertical transport of SOA precursor gases (which are not wet-removed in our parameterization) to higher and colder altitudes can result in more partitioning to the particle phase. These changes lead to areas, such as the central swath through the OKC box, and over parts of the southeastern United States, where there is an increase in the column integrated OA. Based on these simulations the change in OA can be significant, approaching a column integrated increase of 10 to 15 % for some areas.

In contrast to BC, the wet removal of sulfate can be counteracted by its production in cloud. In the AUS box, sulfate wet removal is larger than production, leading to a small net decrease in sulfate when cloud-aerosol interactions (including aqueous chemistry) associated with deep and shallow clouds are considered. Within the MSN box (and over much of the upper-midwest), there is no convective and very little grid resolved precipitation so that the production of sulfate aerosol by aqueous chemistry dominates and there is a significant increase in the column burden of sulfate when non-precipitating clouds are present (Fig. 10). The additional sulfate is limited to the cloud layer and below, but as shown in Figs. 5 and 7 this enhanced sulfate can spread over a deeper layer of the atmosphere. Relative to the control case where the impacts of cumulus are ignored, our results indicate that cumulus can increase the column sulfate burden by as much as 40 %. While the simulations shown here were rather short, longer integration times could lead to significant differences downwind of the area of sulfate production due to enhanced vertical mixing and regional scale transport.

5.3 Impact on cloud microphysics

Using data collected during CHAPS, Berg et al. (2011) measured differences in cloud microphysical properties as a result of differences in the amount of aerosol within

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individual clouds and the cloud draft velocity. They used perturbation of CO (CO' ; defined as the difference between the instantaneous measured CO and the average CO observed during a flight leg) as an indicator of increased aerosol. They found systematic increases in the cloud droplet number concentration (CDNC) associated with both increases in CO' and the cloud updraft strength, which highlighted the importance of considering both the aerosol loading and the cloud dynamics. In their analysis of high-resolution WRF-Chem simulations, Shrivastava et al. (2013) found results consistent with those reported by Berg et al. (2011). A similar analysis has been completed here using results from the DeepShallow simulations in the OKC analysis box, but limited to only grid columns with shallow convection. The cloud microphysical properties were computed for only the cloudy updrafts, as this is the part of the parameterized clouds where the sub-cloud particle loading can influence the cloud microphysical properties via drop activation. A probability density function (PDF) of simulated CO' and perturbation vertical velocity (w' , defined in a way analogous to CO') is shown in Fig. 11. In this case the parameterized updraft speeds were found to range from 1.0 to 3.5 ms^{-1} which are consistent with the updraft speeds in Fig. 1 of Berg et al. (2011).

For the parameterized sub-grid convective clouds the CDNC is found to increase with increasing values of CO, showing an increase from about 500 to 800 cm^{-3} (an increase by about a factor of 1.6) as the CO' ranges from clean (-35 ppbv) to dirty ($+35$ ppbv) for model grid cells where the updraft ranges from 2.0 to 2.5 ms^{-1} (Fig. 11). The results are fairly noisy with relatively large standard deviations highlighting the wide range of additional factors that can impact the CDNC. The slope of the CDNC vs. CO' regression line for w' equal to 2.0–2.5 ms^{-1} is computed to be 4.2 cm^{-3} ppbv $^{-1}$, which is smaller than the 7.2 cm^{-3} ppbv $^{-1}$ reported by Berg et al. (2011), but is close to the value of 4.5 cm^{-3} ppbv $^{-1}$ derived from the results of Shrivastava et al. (2013). The different slopes seen in the observations, those reported by Shrivastava et al. (2013) and this study could be related to the smoothing of emissions, which has been documented in the literature in regards to both simulated cloud characteristics (Gustafson et al., 2007) and aerosol loadings (Gustafson et al., 2011). The results shown by the

different studies should be considered with care, however, because of the different vertical velocity ranges used in each case. While not ideal, the different w ranges were applied because of differences in the spatial and temporal scales associated with the observations and high- and low-resolution simulations.

5.4 Chemical composition of cloud drops

Changes to chemical properties of the particles associated with passage through clouds are an important aspect of cloud aerosol interactions. One of the goals of the CHAPS study was to document changes in the chemical composition of particles that served as CCN (activated) or remained inactive (interstitial). During CHAPS, measurements showed both the activated and interstitial aerosol were dominated by organics and sulfate (Fig. 12). In their analysis, Berg et al. (2009) also reported enhanced nitrate in the dried cloud drop residuals that were sampled via a counter flow virtual impactor (CVI). They attributed this to the uptake of gas-phase nitric acid by cloud drops. In their analysis of high-resolution WRF-Chem simulations, Shrivastava et al. (2013) also found enhanced nitrate when aqueous phase chemistry, which includes trace gas-liquid phase equilibria, was turned on. When aqueous phase chemistry was turned off, however, the particle nitrate in cloud drop residuals and interstitial particles was nearly the same, indicating the importance of the uptake and dissociation of gas-phase nitric acid within cloud drops.

A similar analysis has been completed for the OKC box using results from ShallowOnly simulations. The mass loading of the interstitial aerosol within the shallow clouds is generally smaller in this study than the loading reported by Shrivastava et al. (2013) for either the observations (Fig. 12) or high-resolution simulations (Fig. 7 of Shrivastava et al., 2013). This behavior may, in part, be attributed to the averaging of the emissions over the larger model grid cell in the vicinity of Oklahoma City and the location of the simulated shallow clouds in the two studies. In contrast to the interstitial particles, the simulated mass loading of the activated aerosol is larger in all three simulations (grid-resolved, ShallowOnly with cloud chemistry on, and ShallowOnly with

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cloud chemistry off) than the loading that was observed during CHAPS. The over-estimation of simulated aerosol mass may, in part, be due to the cut size used by the CVI operated on the aircraft that would exclude small cloud drops. In contrast to the aerosol mass loading, the observed and simulated aerosol volume fractions are in good agreement. Thus, even if the mass loading is incorrect, the consistent volume fractions indicates that the chemical processing within the model clouds is behaving in a way that is consistent with the observations. Similar to the observations and high-resolution simulations, there is an increase in the volume fraction of nitrate in activated (cloud-borne) aerosol compared to interstitial aerosol.

6 Summary and conclusions

A new treatment of cloud-aerosol interactions within parameterized shallow and deep convection has been implemented in WRF-Chem with the goal of improving regional scale simulations of the aerosol lifecycle and cloud-aerosol interactions. The modifications designed to represent cloud-aerosol interactions include treatment of the cloud droplet number mixing ratio; key cloud microphysical and macrophysical parameters (including the updraft fractional area, updraft and downdraft mass fluxes, and entrainment) averaged over the population of shallow clouds, or a single deep convective cloud; and vertical transport, activation/resuspension, aqueous chemistry, and wet removal of aerosol and trace gases in warm clouds. These changes have been implemented in the WRF-Chem chemistry package as well as the Kain–Fritsch cumulus parameterization (Kain and Fritsch, 1990; Kain, 2004), which has been modified to better represent shallow convective clouds (Berg et al., 2013). Preliminary testing of the new version of WRF-Chem has been completed, with results compared with data from the CHAPS field experiment (Berg et al., 2009, 2011) as well as high-resolution simulations (Shrivastava et al., 2013).

The preliminary results are encouraging and demonstrate the advantages of the modifications that have been made to WRF-Chem. It is shown that both deep and

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shallow convective clouds have an important impact on the horizontal and vertical distribution of aerosol loading. Three different domain sub-regions were selected for detailed analysis, including locations near Madison, Wisconsin (MSN), Austin, Texas (AUS), and Oklahoma City, Oklahoma (OKC), the latter corresponding to the site of CHAPS and the domain used in previous high-resolution simulations. These regions were selected to represent instances dominated by shallow (MSN), deep (AUS), or a mix of both (OKC) types of convective clouds. In each case the WRF-Chem simulations behaved in a manner consistent with expectations and consistent with both the CHAPS data and the results of high-resolution simulations. In the case of shallow clouds, enhanced mixing leads to a deepening of the layer containing BC and decreased amounts of BC near the surface. Results are similar for OA, but the net impact was found to be smaller. In contrast to BC, sulfate aerosol was enhanced throughout the layer due to sulfate production within clouds. In the vicinity of AUS, the impact of shallow convective clouds is minimal. There was a decrease in BC, OA, and sulfate in the sub-cloud layer due to vertical transport associated with deep convective clouds. There were also significant changes in the aerosol loading aloft that were the result of the impacts of updrafts, downdrafts, entrainment mixing, enhanced subsidence, and wet removal associated with the sub-grid clouds. In the area near OKC, both the deep and shallow sub-grid convective clouds had a significant impact on the simulated aerosol loading. The shallow sub-grid clouds led to a decrease of aerosol in the sub-cloud layer and an increase of aerosol aloft. The parameterized deep-convective clouds led to decreases in the BC and OA over the lowest 2 km and sulfate over the lowest 3 km of the atmosphere. There were differences between the simulations completed for this study and the high-resolution simulations presented by Shrivastava et al. (2013), but these differences could be attributed to treatment of the emissions between the high and low resolution grids, differences in the simulated cloud fields, and differences in the upwind conditions associated with the two sets of simulations.

One of the motivations for the development of the improved parameterization is to allow the investigation of regional and synoptic scale aerosol transport. In our case-study

period, there is a significant reduction in the BC and OA over much of the central United States. The primary removal mechanism added in the new treatment is the wet removal associated with the parameterized precipitation. Thus, the differences in the aerosol loading highlight the importance of wet removal on the aerosol lifecycle at the regional scale. In contrast to BC and OA, there are large regions in which there are increases in the column-integrated sulfate due to the production of sulfate and absence of wet removal in nonprecipitating clouds.

The behavior of the modified version of WRF-Chem in regards to the cloud microphysical properties and chemical composition of aerosol is also investigated. The results show that the modified version of WRF-Chem is able to reproduce the first aerosol indirect effect in a way that is consistent with both high-resolution simulations and observations from CHAPS. The CDNC associated with the parameterized clouds was found to be less sensitive to pollutant loading than was observed (Berg et al., 2011) but was similar to that reported by Shrivastava et al. (2013) in their high resolution simulations. It should be noted that the current modifications do not include the treatment of other indirect effects, which will be included at a later date. The chemical composition of the simulated cloud-drop residuals is compared to the composition measured with an AMS operated behind a CVI inlet during CHAPS. While there were differences in the simulated and observed mass loadings, the simulated and observed mass fractions were consistent, including the presence of enhanced amounts of nitrate in the cloud drop residuals. WRF-Chem is also able to accurately represent the increase in nitrate found in the observed cloud-drop residuals. Overall, these findings provide evidence that the modified version of WRF-Chem is able to represent key features of the cloud-aerosol interactions in a realistic way. While the results presented here utilized WRF-Chem version 3.2.1, the code is being ported to WRF-Chem version 3.5. We anticipate including these changes in a future public release of WRF-Chem.

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Supplementary material related to this article is available online at
[http://www.geosci-model-dev-discuss.net/7/2651/2014/
gmdd-7-2651-2014-supplement.zip](http://www.geosci-model-dev-discuss.net/7/2651/2014/gmdd-7-2651-2014-supplement.zip).

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Table 2. Definitions of simulations completed as part of the study. The parameterized cumulus dynamics are applied in all simulations.

Simulation	Aerosol Processing by Shallow and Deep Cu
DeepShallow	Aerosol Processing Shallow Cu: On Aerosol Processing Deep Cu: On
ShallowOnly	Aerosol Processing Shallow Cu: On Aerosol Processing Deep Cu: Off
Control	Aerosol Processing Shallow Cu: Off Aerosol Processing Deep Cu: Off

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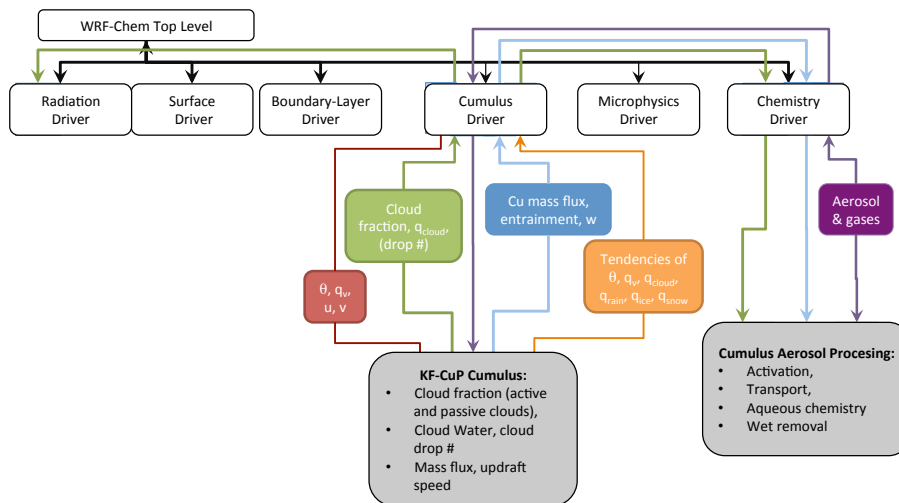


Fig. 1. Summary of modifications to the standard implementation of WRF-Chem. Colored boxes indicate information passed between subroutines related to the thermodynamics (red), cloud microphysical and macrophysical properties (green), cloud dynamics (blue), thermodynamic tendencies (orange) and aerosol and trace gases (purple), while gray boxes indicate the new or modified parameterizations applied in WRF-Chem. Arrows indicate information flow within the model. Note that the droplet number generated in the KF-CuP parameterization is not currently used in the Radiation Driver.

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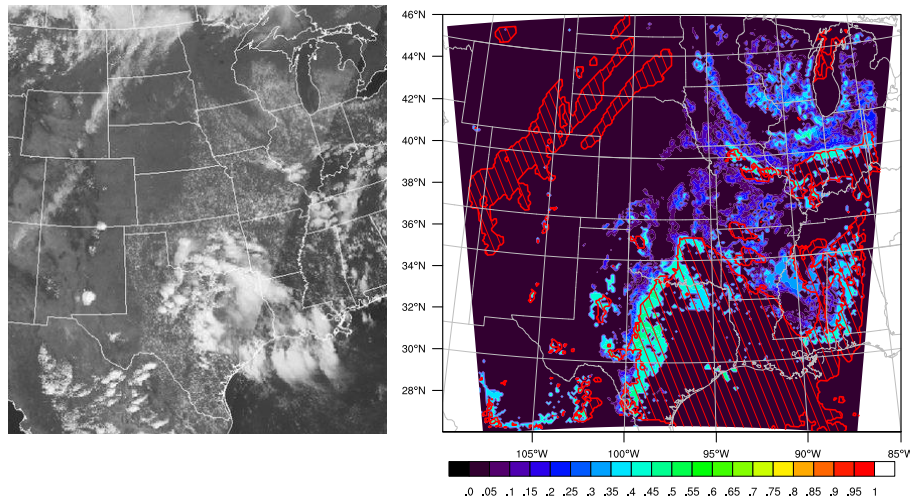


Fig. 2. GOES visible satellite image valid at 20:15 UTC, 25 June 2007 (left), and simulated cloud fraction associated with the KF-CuP parameterization (colors), and areas with grid resolved clouds (hashed; right).

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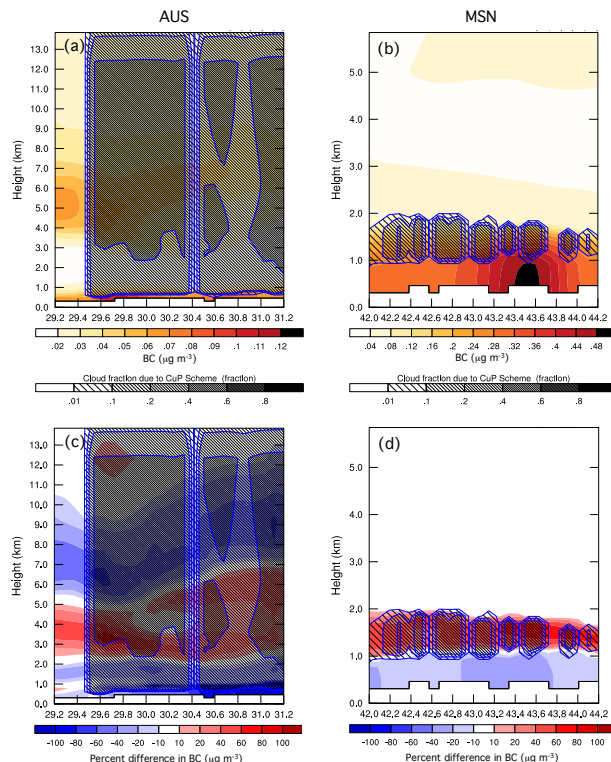


Fig. 4. Vertical north-south cross sections of BC in size bins 1 through 4 (colors top; $\mu\text{g kg}^{-1}$), including both interstitial and activated aerosol in the cloudy grid cells, and difference in BC mass loading between DeepShallow and control simulations (colors bottom; percentage) for conditions dominated by deep convective clouds (AUS; left) and shallow convective clouds (MSN; right) boxes at 20:00 UTC on 25 June 2007. Hatching indicates cloud fraction associated with sub-grid convective clouds. The horizontal axis is labeled in degrees of latitude and heights are height above mean sea level.

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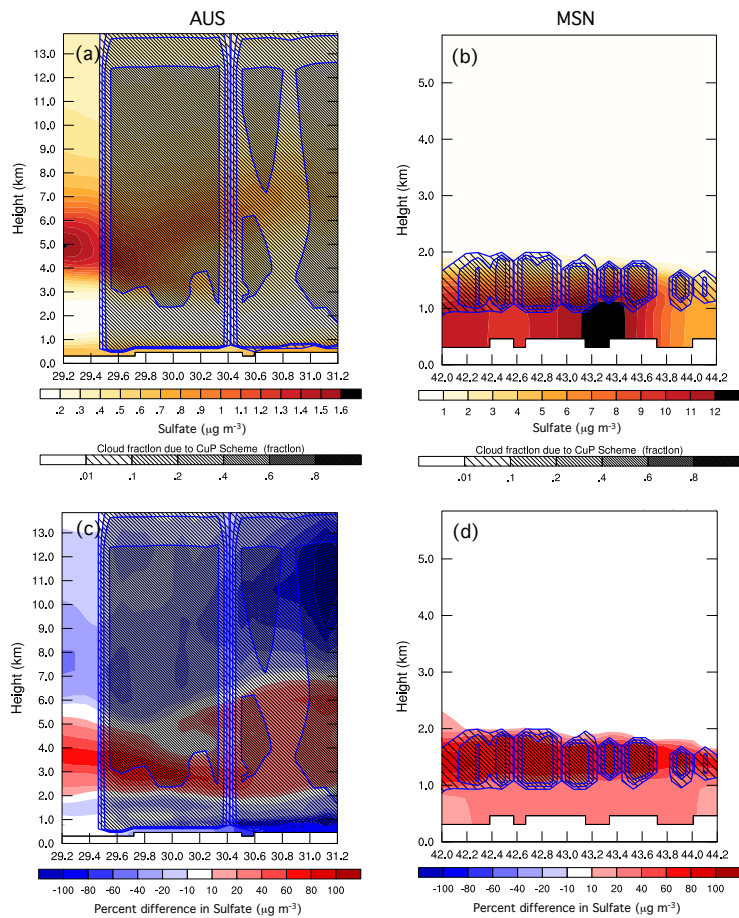


Fig. 5. Same as Fig. 4, but for sulfate.

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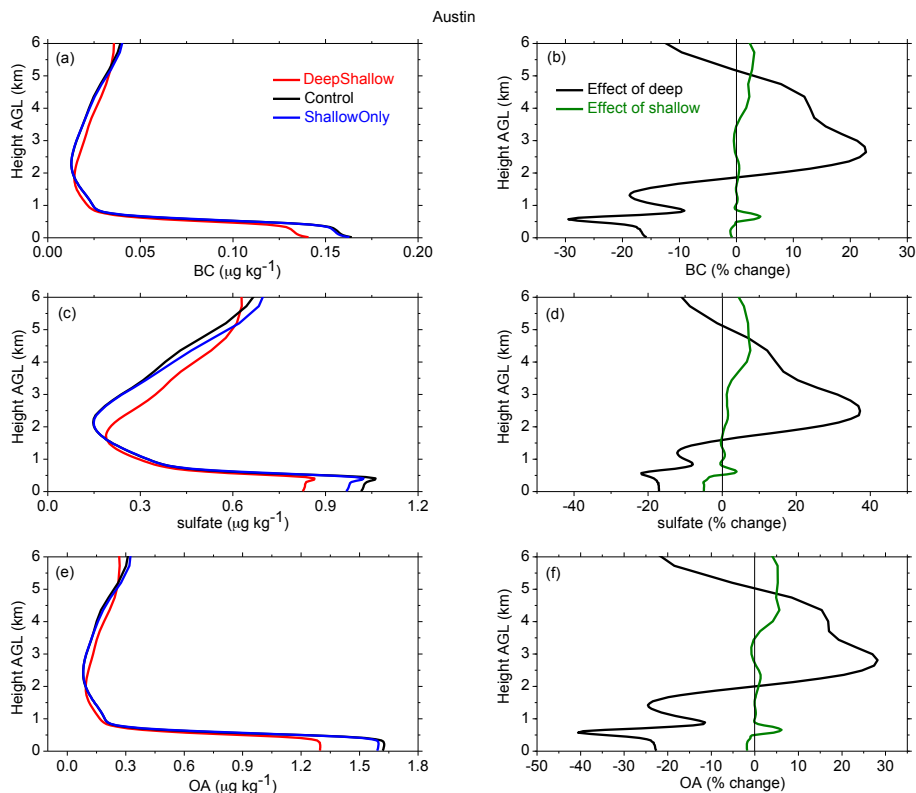
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Fig. 6. Simulated mass loading for size bins 1 through 4 for the effect of deep and shallow clouds on BC (a and b), sulfate (c and d), and OA (e and f) simulated for the cases DeepShallow (red), and ShallowOnly (blue), and Control (black) for the AUS box valid at 20:00 UTC on 25 June 2007. The effects of deep and shallow convection are calculated as $[100 \cdot (\text{DeepShallow} - \text{ShallowOnly}) / \text{ShallowOnly}]$ and $[100 \cdot (\text{ShallowOnly} - \text{Control}) / \text{Control}]$, respectively. Note different horizontal scales in the figure.

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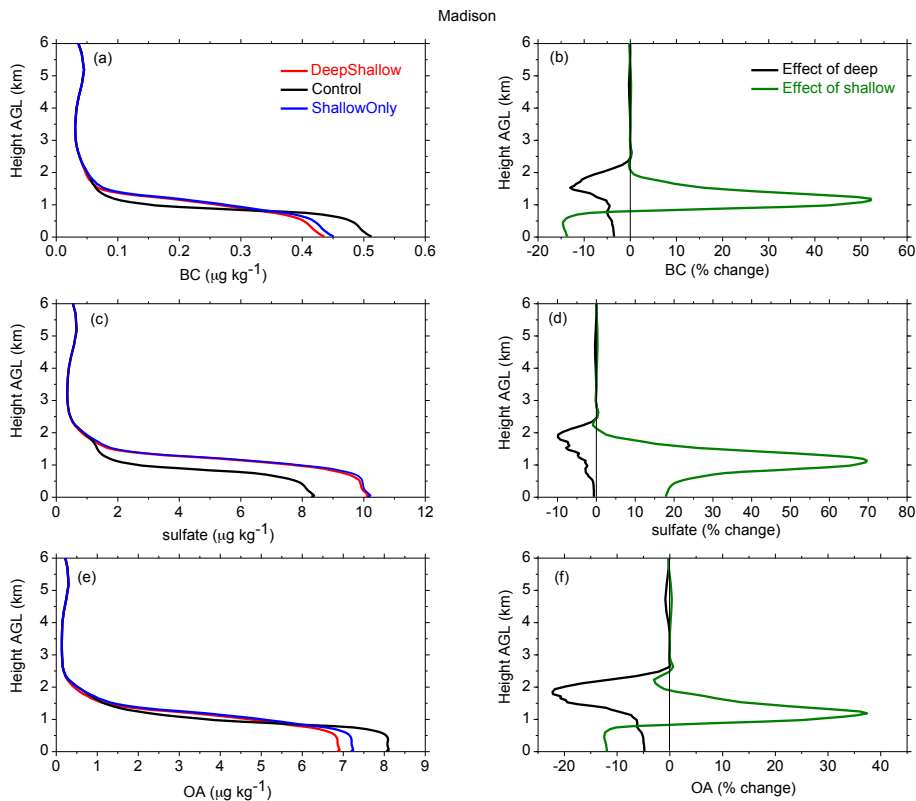


Fig. 7. As in Fig. 6 but for MSN box.

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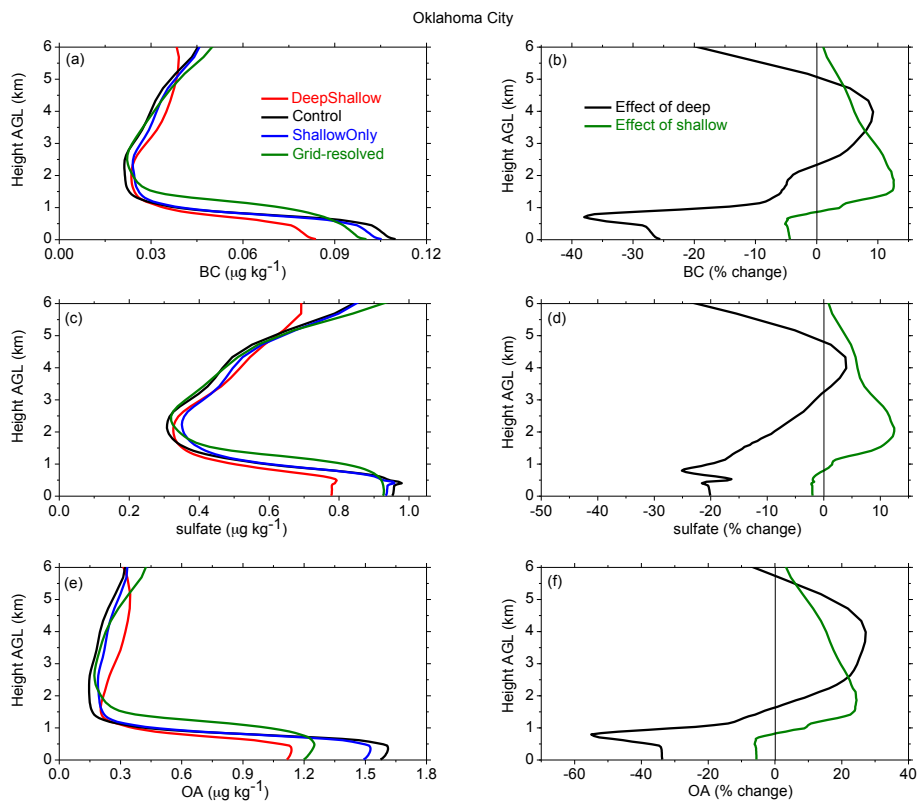


Fig. 8. As in Figs. 6 and 7 for the OKC box. The green line in (a, c, and e) represents results from the high-resolution simulations of Shrivastava et al. (2013).

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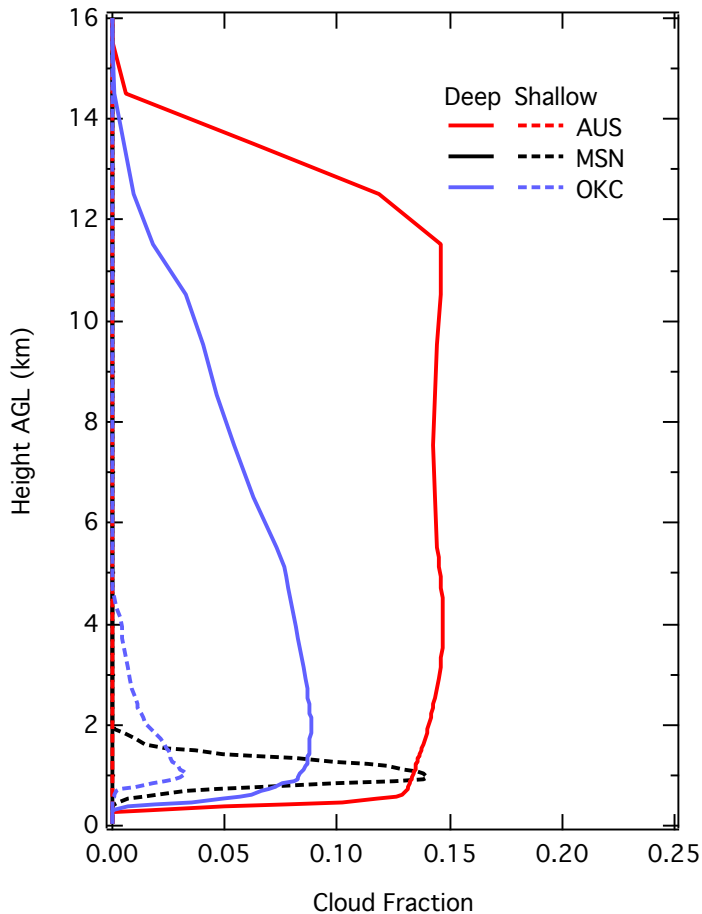


Fig. 9. Convective cloud fraction associated with deep (solid) and shallow (broken) convective clouds: Averages over the entire AUS (red), MSN (black), and OKC (blue) boxes at 20:00 UTC on 25 June 2007.

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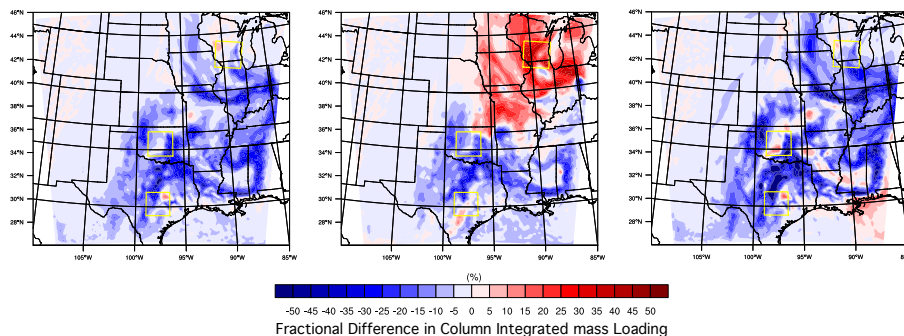


Fig. 10. Fractional differences in column integrated aerosol mass loading between DeepShallow and control simulations for size bins 1 through 4, including both interstitial and activated aerosol in the cloudy grid cells, for BC (left), sulfate (center) and OA (right), valid at 20:00 UTC on 25 June 2007. Yellow boxes indicate boxes used in the analysis.

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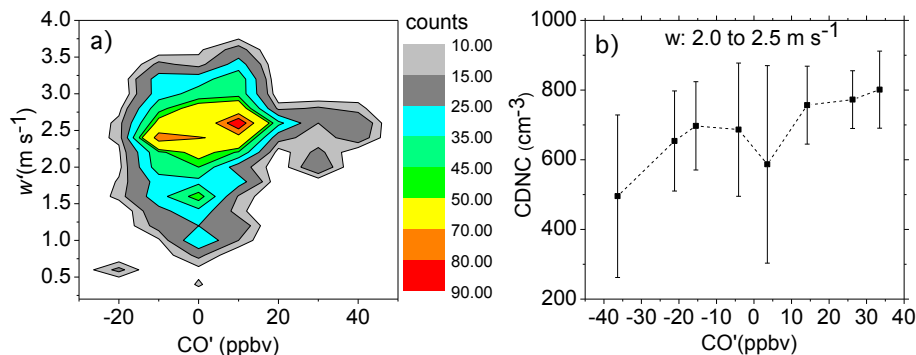


Fig. 11. PDF of simulated cloud updraft speed and CO loading in cloudy updrafts **(a)**, change in CDNC with perturbation values of CO (CO') for perturbation values of w (w') between 2.0 and 2.5 m s^{-1} **(b)**. Error bars in **(b)** indicate the standard deviation.

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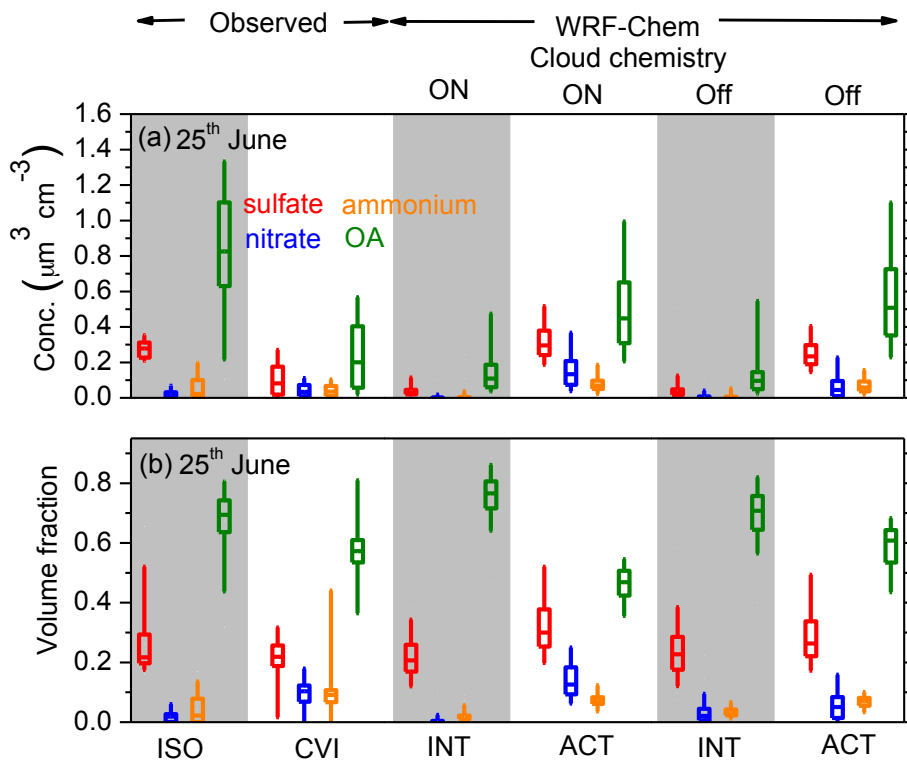


Fig. 12. Aerosol mass concentration (top) and volume fraction (bottom) for observed interstitial (sampled via an isokinetic inlet (ISO; grey areas)) and activated (sampled via a counter-flow virtual impactor inlet (CVI; white areas)) aerosol; and simulated interstitial (INT; grey areas) and activated (ACT; white areas) aerosol at 20:00 UTC on 25 July 2007. Colors indicate sulfate (red), ammonium (orange), nitrate (blue), and organic aerosol (green) in size bins 1 through 4. Box-and-whisker plots indicate 90th, 75th, 50th, 25th, and 10th percentiles.

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