Modelling below-cloud scavenging of size resolved particles in GEM-MACHv3.1

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Abstract. Below-cloud scavenging (BCS), the process of aerosol removal from the atmosphere between cloud-base and the ground by precipitation, affects aerosol number/mass concentrations, size distribution, and lifetime. The impact of different representations of BCS was examined through implementing a new aerosol BCS scheme in the

- 10 Environment and Climate Change Canada (ECCC) air quality prediction model GEM-MACH and comparing it with the existing scavenging scheme in the model. Further, the current GEM-MACH employs a single-phase precipitation for BCS: total precipitation is treated as either liquid or solid depending on a fixed environment temperature threshold. Here, we consider co-existing liquid and solid precipitation phases as they are predicted by the GEM microphysics. GEM-MACH simulations, in a local-area domain over the Athabasca oil sands areas,
- 15 Canada, are compared with observed precipitation samples.

Evaluation against observed precipitation samples reveals enhanced GEM-MACH performance with the new multiphase approach and scavenging scheme. Notably, sulfate wet deposition showed improvement with reduced biases. In contrast, nitrate and ammonium improvements were less significant, showing episodic changes tied to specific locations and cloud events. Discrepancies in wet deposition patterns are traced to the complexities of below-cloud scavenging parameterizations. The GEM-MACH current BCS parameterization lacks representation of crucial physical processes, while the updated semi-empirical scheme displays distinct scavenging coefficients, notably important for aerosol sizes ranging from 0.1 to 1 µm and varying precipitation intensities. The changes in wet scavenging resulted in a higher formation rate and larger concentrations of atmospheric particle sulphate.

Model evaluation against precipitation chemistry and ambient air concentrations indicates the effectiveness of the new parameterization with multi-phase partitioning, improving overall model-observation agreement. Despite slight biases in wet-deposited nitrogen, the inclusion of partitioning and new scavenging enhances model performance.

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Introduction

Atmospheric aerosols impact air quality, and influence climate directly by absorbing and scattering sunlight, and indirectly by modifying cloud properties (e.g. Haywood and Boucher, 2000). Aerosol number/mass concentration, lifetime and distribution are regulated by the removal processes such as wet scavenging (Pruppacher and Klett, 1997).

- 35 Wet scavenging, including in-cloud and BCS, is defined as the removal of both aerosol particles and gases from the atmosphere by different hydrometeors such as rain or snow. Scavenging of pollutants by cloud droplets and by removal due to precipitation formation (also known as "rainout") occurs as a result of the solubility of gases and via aerosol activation. BCS is also known as the "washout" process, in which atmospheric material is exposed to falling hydrometeors and hence may be subject to collection and removal (e.g., Rogers and Yau, 1989). Ryu and Min
- 40 (2022) study reevaluated aerosol deposition in WRF-Chem using observation results. Below-cloud scavenging was found to play a more significant role (50%–60%) in wet deposition than traditionally understood, while in-cloud scavenging was revised up to 34%–37%. Their updated model improved wet deposition accuracy for various aerosols, particularly SO₄ and NH₄, enhancing surface PM2.5 and PM10 concentration predictions. Wet scavenging is the dominant removal pathway of aerosol particles at the global scale (Stier et al., 2005), however, it is worth
- 45 noting that recent observational studies, such as Emerson et al., 2020, highlighted the significance of aerosol dry deposition. In general, the study of the wet deposition process requires an understanding of cloud processes, including the chemical reactions occurring within hydrometeor. Details of these processes underlie some of the major sources of uncertainty in wet deposition modeling (Tost et al., 2007; Kajino and Aikawa, 2015). To study the BCS, it is necessary to consider some of the physical processes, such as condensation, Brownian motion,
- 50 thermophoresis, diffusiophoresis, turbulent inertial interception, rear-capture and gravitational/electric forces (Lemaitre, et al., 2017; Pruppacher and Klett 1978; Gong et al., 2003 and 2011). Brownian motion is more important for scavenging particles smaller than 0.1 μ m, while the gravitational forces are more efficient to remove particles larger than 1 μ m. Collision efficiency of hydrometeors with particles has a minimum for particles in the size-range of 0.1–2 μ m. This region of low collection efficiency due to Brownian motion and gravitational settling, with its
- 55 minima in this size range, is referred to as the Greenfield gap (Greenfield, 1957; Slinn and Hales, 1971; Andronache et al. 2006; Ladino, et al. 2011). The size distributions of aerosol particles and the diffusivity/solubility of gases also need to be specified in order to accurately represent the collection kernel for hydrometeor-aerosol collision efficiency (c.f. Cherrier et al, 2017). In this study, we focus on BCS of aerosols.

The BCS of particles is more efficient in the cold months due to snow scavenging, while gas scavenging is more
 efficient during the warm months (Cheng et al., 2017). Snow scavenging is an important removal mechanism in
 mid-latitude, polar regions and mountainous areas. Snow and ice crystals are more efficient scavengers than water
 drops, due to their larger surface-to-volume ratio (Wang et al., 2014). The study of snow scavenging is more
 complicated than rain scavenging, due to the wide variety of snow-particle shapes, sizes, and densities, which results
 in different fall speeds, cross-sectional areas, and flow patterns around snow particles (Pruppacher and Klett, 1997;

55 Jylhä, 1999, Wang et al., 2014), while it has been a common practice to assume spherical shapes for rain droplets. However, it is important to note that this assumption can introduce inaccuracies. This is particularly evident for

larger raindrops, which often deviate from perfect spherical shapes and exhibit more oblate forms. The modeling study of Croft et al., (2009) estimated around 30% of BCS of sulphate particles is due to snow scavenging, globally.

Some studies indicate a tendency of the precipitation phase to be moving towards the liquid (rain) due to the global warming (Trenberth, 2011). It is important to differentiate precipitation phases in model simulations, as scavenging efficiency differs between rain and snow (examples of aerosol scavenging coefficients for rain and snow are shown in Figure 3). The temperature cross-over characterizing the transition point between equal contributions of rain and snow towards particle collection also varies significantly, between -0.4 and 2.4 °C (a 29-year Northern Hemisphere observational dataset demonstrating this range may be found in Jennings et al., (2018)). Despite this temperature variation, and the importance of precipitation phase on wet scavenging and wet fluxes, some scavenging models use

75 variation, and the importance of precipitation phase on wet scavenging and wet fluxes, some scavenging models use partition precipitation phase based on a simple, uniform air temperature threshold (Harpold, et al., 2017; Feiccabrino et al., 2015; Jennings, et al., 2018).

The rate of change (loss) of aerosol mass concentrations due to BCS by precipitation (rain and snow) is referred as the scavenging coefficient Λ (s⁻¹) (Seinfeld and Pandis, 2006):

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$$\frac{\partial C(t)}{\partial t} = -\Lambda \cdot C(t),$$

(1)

$$\Lambda(d_{p}) = \int_{0}^{\infty} \frac{\pi}{4} D_{p}^{2} U_{t}(D_{p}) E(D_{p}, d_{p}) N(D_{p}) dD_{p}$$
(2)

In Eq. 1, C(t) is the mass concentration of aerosol at time t, and the scavenging coefficient Λ is the loss frequency (s⁻¹) of mass concentration per unit time; their product in (1) is the rate of mass concentration loss per unit time associated with scavenging. In Eq. 2, d_p , D_p and $N(D_p)$ refer to aerosol diameter, droplet diameter and droplet number density at given droplet size, respectively. In this equation, $\Lambda(d_p)$ is expressed as a function of the particle collection efficiency, $E(d_p, D_d)$ and the hydrometeor's fall speed $U_t(D_d)$.

Eq. (1) is also dependent on the size of the aerosol (via d_p), hence requiring the aerosol size distribution to be incorporated into the equations.

- 90 Here, a semi-empirical, size-resolved Λ parameterization from Wang et al., (2014) was implemented in the Environment and Climate Change Canada (ECCC) air quality prediction model GEM-MACH v3.1 (Global Environmental Multi-scale model–Modeling Air quality and Chemistry). The Λ parameterization in Wang et al., (2014) study was developed based on the uncertainty analyses and follows power law relationships with precipitation intensity R (Wang et al., 2014). Furthermore, it is applicable to BCS of the rain and snow and over a wide range of
- 95 particle sizes and precipitation intensities. The default GEM-MACH scavenging scheme is based on Slinn 1984, and its collection efficiency is formulated as a linear combination of the collection efficiencies due to Brownian motion, interception, and impaction. Therefore, the Slinn parameterization misses important processes in the Greenfield gap, such as thermophoresis and electrostatic forces, which are included implicitly in the Wang et al. (2014) model. The semi-empirical Wang 2014 scheme was developed to provide an optimization of all available theoretical formulations
- 100 of scavenging coefficients in comparison with available observations at the time.

In the present study, we determine the impact of precipitation phase partitioning on aerosol BCS, and compare the Wang et al., (2014) scavenging scheme with the previous GEM-MACH scavenging scheme (based on Slinn 1984). The implementation details of both schemes are described below.

In what follows, BCS in the GEM-MACH model and the simulation set-up are described (Sections 2.1 - 2.3), followed by a brief description of the measurement data used for model evaluation (Section 2.4). Section 3 presents the results including (1) GEM-MACH BCS tests to examine the impact of precipitation partitioning and of the Wang et al., (2014) scheme, and (2) comparison of GEM-MACH simulations with observations. The latter work has a focus on the particulate base cation NH_4^+ , acidic anions NO_3^- , $SO_4^=$ and HSO_3^- , and particulate sulphate, ammonium and nitrate. The summary and conclusions of our work are reported in Section 4.

110 1 Methodology

2.1 GEM-MACH Model

The GEM-MACH base model consists of an online tropospheric chemistry module embedded within ECCC's GEM numerical weather forecast model (Côté *et al.* 1998a, b; Charron *et al.*, 2012, Moran et al., 2013). A fully-coupled version of GEM-MACH is used for this study; "fully-coupled" here refers to the aerosol chemistry and meteorology coupling, i.e., this version allows for the model to include the influence of on-line aerosols on modelled meteorology through radiation and cloud microphysics (Makar *et al.*, 2015a, b; Gong *et al.*, 2015; Makar *et al.*, 2021). The aerosols predicted by the chemistry module are used to calculate aerosol extinction in the GEM radiation code and droplet number concentration (CDNC) in turn affecting aqueous phase chemical processes within the model's chemistry modules.

- 120 The GEM-MACH chemistry module includes a comprehensive representation of air quality processes, such as gasphase, aqueous-phase, and heterogeneous chemistry and aerosol processes (e.g. Moran *et al.*, 2013; Makar *et al.*, 2015a, b; Gong *et al.*, 2015). The default gas-phase chemistry of GEM-MACH was based on the ADOM-II mechanism with 47 species and 114 reactions (Lurmann *et al.*, 1986; Stockwell and Lurmann, 1989). In the current study, we use gas-phase chemistry parameterized by the SAPRC 11 mechanism with 175 species and 837 reactions (Carter and Heo,
- 125 2013); inorganic heterogeneous chemistry represented by a modified version of ISORROPIA algorithm of Nenes *et al.* (1999) as described in Makar *et al.* (2003); secondary organic aerosol (SOA) formation parameterized using a two-product, overall or instantaneous aerosol yield formation (Odum *et al.*, 1996; Jiang 2003; Stroud *et al.*, 2018); aerosol microphysical processes, including nucleation and condensation (sulphate and SOA), hygroscopic growth, coagulation, and dry deposition/sedimentation parameterized based on Gong *et al.* (2003); and the representation of
- 130 cloud processing of gases and aerosols includes uptake and activation, aqueous-phase chemistry, and wet removal (Gong *et al.*, 2006, 2015). The default GEM-MACH model includes eight internally mixed aerosol components: sulphate, nitrate, ammonium, primary organic aerosol, secondary organic aerosol, elemental carbon, crustal material, and sea salt. For this study, crustal

material is further speciated into 6 elements: calcium, magnesium, sodium, potassium, iron, and manganese. A sectional approach is used for representing aerosol size distribution with a 12-bin (between 0.01 and 40.96 µm, logarithmically spaced) configuration.

2.2 Below-cloud scavenging

In GEM-MACH model, the default wet BCS schemes of aerosol by rain and snow are based on Slinn (1984). Wet scavenging is a removal process for the atmospheric tracers, and describes the uptake of gases and aerosols into cloud hydrometeors. Partial or complete hydrometeor evaporation subsequent to that uptake may result in the release of captured particles back to the atmosphere. In such cases, rather than a net removal (wherein the captured material reaches the ground within precipitation), the captured material may instead be transported to a different vertical level in the atmosphere from the point where the uptake occurred. The wet deposition flux of tracer i at a given level in a vertical column is calculated by Eq. (3) in GEM-MACH (Gong et al., 2006):

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$$F_{i}(z) = (F_{i}(z+1) + \Delta F_{i}(z))(1.0 - f_{evp}(z))$$
(3)

where f_{evp} is the fraction (0 to 1) of precipitation loss by evaporation. $\Delta F_i(z)$ is the input flux of aerosols to the hydrometeors at level z due to either precipitation production (e.g., cloud-to-rain) or precipitation scavenging, and $F_i(z + 1)$ is the flux arriving at that level from above.

Slinn (1984) formulations are parameterizations of the scavenging coefficient theoretically described in equation (2). Based on laboratory experiments and dimensional analyses, Slinn (1984) presented separate BCS schemes for rain and snow as functions of precipitation rate (P) and mean collection efficiency (E) (Gong *et al.*, 2003; 2011). For rain:

$$\Lambda(\mathbf{d}_{\mathrm{p}}) = \frac{c \,\mathrm{P}\,\mathrm{E}(\mathbf{d}_{\mathrm{p}},\mathbf{R}_{\mathrm{m}})}{\mathbf{R}_{\mathrm{m}}} \tag{4}$$

where the empirical constant *c* is 0.5, and R_m is the mean drop size. The precipitation rate *P* is also used to obtain the mean drop radius R_m . For snow:

$$\Lambda(\mathbf{d}_{\mathbf{p}}) = \frac{\gamma P \, \mathrm{E}(\mathbf{d}_{\mathbf{p}}, \lambda)}{\mathrm{D}_{\mathbf{m}}} \tag{5}$$

where λ is the characteristic capture length, D_m is the characteristic length, and γ, the empirically-derived collision efficiency, is 0.6. The values of λ and D_m vary depending on temperature and snow type. The temperature dependence
of Slinn's collection efficiency E for snow (solid precipitation) includes the consideration of hydrometeor forms (or shapes), which determine the characteristic capture length, settling velocity, etc. The different forms of precipitation are found in certain temperature ranges, e.g., needle snow (-8 °C < T < 0 °C), stellar snow (-25 °C < T < -8 °C), and graupel (T < -25 °C). The characteristic capture length, settling velocity, etc. are assigned with different values based on the types of solid hydrometeors (hence the temperature ranges). For both rain and snow, collection efficiency *E*, is a linear combination of three processes: Brownian diffusion, interception, and impaction (Slinn 1984).

In the pre-existing, base-case, GEM-MACH setup, the liquid and solid precipitation fluxes predicted from the GEM meteorological module are combined, and a single phase (*either* liquid or solid) is determined by a uniform environment temperature threshold, T = 0 °C (e.g. rain occurs for temperatures > 0 °C and snow for temperatures < 0 °C). In this study, a multi-phase partitioning is tested, by taking into account the liquid and solid hydrometeor phases

170 predicted by the microphysics in the meteorological module of GEM-MACH. Using this multi-phase partitioning approach, consistent with the GEM microphysics employed here, the *co-existence* of both liquid and solid phases can be treated (more information on this new scheme appears in section 2.3).

In addition to the above in-cloud scavenging parameterization comparisons, we compared the existing BCS methodology (Slinn, 1984) to a more recent semi-empirical methodology (Wang et al., 2014). In contrast to equations

(4,5) Wang *et al.* (2014) proposed separate formulae of Λ for rain and snow, as a function of aerosol diameter and rain rate:

$$\Lambda_{i} = A_{i} R^{B_{i}} \tag{6}$$

where A_i and B_i are empirical parameters and are polynomial functions of aerosol diameter, and the subscript *i* denotes a given particle size bin. *R* is the precipitation rate (Wang *et al.* (2014)).

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2.3 Model setup

This study focuses on the Oil Sands (OS) region, Alberta, Canada. Figure 1 shows the model domains and Canadian Air and Precipitation Monitoring Network (CAPMoN) and Alberta Precipitation Quality Monitoring Program (APQMP) observation stations used for model wet deposition evaluation. GEM-MACH simulations were carried out on a limited area model (LAM) domain with 2.5 km \times 2.5 km (red) resolution, nested from a 10 km \times 10 km (blue)

- horizontal resolution, for the months of April and July, 2018 (Fig. 1). The simulations incorporated a one-week "spinup" period prior to April and July 1st, 2018, to allow the model cloud fields to reach a steady-state prior to the monthlong simulations used for model comparison and evaluation. The model driving meteorology was updated every 24 hours, with each high resolution run duration of 30 hours incorporating a 6 hour spin-up, for the same reason. The
- 190 model set-up with regards to meteorological piloting, chemical lateral boundary condition, anthropogenic, biogenic, and wildfire emissions are similar to those presented in (Makar *et al.*, 2018, 2021), with the underlying version of GEM being updated to v5.0. The meteorology is piloted by the global GEM model and initialized daily (at 06 UTC) using the Canadian Meteorological Centre's regional objective analysis. Eighty vertical, unevenly spaced, hybrid coordinate levels were used to cover between the surface and 0.1 hPa, with the lowest terrain-following model layer
- 195 located about 20 m above the surface. The feedbacks between chemistry and meteorology were enabled so that the aerosols predicted by the chemistry module were allowed to influence the model radiative transfer (through aerosol extinction) and cloud microphysics (through droplet nucleation), as described above (further details can be found in Makar et al., 2015a,b, Gong *et al.*, 2015, and Makar *et al.*, 2021). For this study, three model runs were conducted:
- Base-case the existing GEM-MACH setup ("base-case"): The BCS of aerosol is parameterized based on Slinn (1984) ("Slinn1984") as described in 2.2; a single phase precipitation was employed at a given model

grid-cell: the liquid and solid precipitation fluxes at a given model grid-cell are combined to a total precipitation flux, and a uniform environment temperature threshold (T = 0 °C) is used to determine the precipitation phase, liquid or solid, for BCS (e.g. > 0 °C rain and < 0 °C snow).

- Multi-phase partitioning approach (or "multi-phase"): for this experiment, the BCS of aerosol is also based on Slinn (1984), however, coexisting multi-phase precipitation at a given model grid is considered. In contrast to the base-case, the liquid and solid precipitation fluxes are not combined prior to the BCS calculation. Rather, the predicted liquid and solid precipitation fluxes by the GEM microphysics are used in separate BCS calculations (according to the predicted phase), allowing for the co-existence of liquid and solid precipitation at a given model grid-cell for the treatment of BCS.
 - 3. Wang et al., (2014) scavenging scheme ("Wang2014"): this simulation also uses coexisting multi-phase precipitation similar to case 2 above, with the difference being the replacement of the Slinn (1984) formulation for the BCS coefficient for aerosols with the semi-empirical formulation from Wang et al., (2014).
- 215 In both the base-case and multi-phase approach, the formulae of Slinn (1984; Eqs. 4 and 5) are used to determine the scavenging coefficient (for liquid and solid precipitation, respectively). The main difference between the base-case and multi-phase approaches lies in whether a precipitation is treated as a single phase (base-case), or multiple phases of precipitation (multi-phase) are considered model grid cells containing clouds. The approaches consequently differ in how the precipitation phase is determined for BCS, i.e. through the use of a uniform temperature threshold which
- 220 determines whether the precipitation is in the form of rain or snow (base-case) or through the use of meteorological model (or parameterized) rain and snow (multi-phase). Note also that both methodologies make use of Eq. (3) to describe the flux between model layers. Figure 2 shows the average temperature and the rain and snow fluxes predicted by the GEM-MACH meteorological module for April 2018 at the model hybrid level of 0.98 (e.g. the level near the surface and at ~ 980 mb atmospheric pressure). The study area was relatively cold, and the average temperature is in the range (-15 to 15 °C). The model predicted both solid and liquid precipitation during the month of April, with solid precipitation dominating. There are areas (model grid-cells) where coexisting liquid and solid
- precipitation were predicted. With the base-case approach, all precipitation (liquid + solid) in the area where temperature was below 0° C was treated as solid phase for BCS. Similarly, all precipitation in the area where temperature was above 0° C was treated as liquid phase. This is illustrated in the lower panels of Figure 2 (Fig. 2d and
- 230 2e) by masking the areas with no liquid and solid precipitation. In contrast, the new multi-phase approach treats the liquid and solid precipitation as they are predicted in the GEM microphysics when applying the Slinn parameterization for liquid and solid precipitation scavenging coefficients. The comparison between the average temperature (Fig. 2a) and the solid and liquid fluxes (Fig. 2b and 2c) from these two approaches illustrates the difference between the assigned solid/liquid precipitation based on a uniform temperature (Fig. 2d and 2e liquid precipitation for T > 0 °C
- and solid for T < 0 °C) and the revised approach using the predicted solid and liquid phases from the GEM microphysics module (Fig. 2b and 2c). These results indicate the possible inconsistency between the model hydrometeor distributions generated with the use of a temperature threshold versus the "actual" precipitation phases,

which may further impact the aerosol scavenging/concentration and wet fluxes. Note that Figure S1, the same plot as Fig. 2, but for July 2018, shows the existence of solid phase precipitation at high altitudes.

- 240 Figure 3 compares the Slinn1984 and Wang2014 scavenging coefficients as a function of aerosol size. For liquid precipitation (Fig. 3a), the difference between the Slinn1984 and Wang2014 scavenging coefficients increases with increasing precipitation intensity. The two schemes differ the most for aerosol sizes between 0.1 to 1 μ m, particularly at high precipitation intensity (Fig. 3b). For the solid precipitation at lower intensity (R=0.01 mm/h), the two schemes diverge for aerosols smaller than 1 μ m, while at higher precipitation intensities (R=10 mm/h), the two schemes show
- 245 higher disparity for aerosols larger than 1 µm. Collection efficiency in the Slinn formula includes the effects of the processes of Brownian diffusion, interception, and impaction. However, Slinn's formulae does not include representation of some processes such as thermophoresis or diffusiophoresis, both of which may increase the collection efficiency for particles in the size range of 0.01-1 µm (e.g., Slinn and Hales, 1971; Wang et al., 1977; Mc-Gann and Jennings, 1991; Byrne and Jennings, 1993; Pranesha and Kamra, 1997; Tripathi and Harrison, 2001; Tinsley
- 250 et al., 2000; Andronache, 2004;2006; Wang et al., 2010). Wang2014 includes the consideration for these two additional processes. Jones et al. (2022) showed that the thermophoresis mostly enhances the collection of accumulation mode particles $(0.1 - 1 \mu m)$. This may explain the underestimation of scavenging coefficient by Slinn1984 in comparison to Wang2014 for particles below 1 µm diameter. Figure 4 shows snow scavenging coefficient magnitude versus aerosol size distribution with the intensity of 0.01 mm/h and different ambient atmospheric
- 255 temperatures. In the Slinn (1984) Eq. 5 (for snow or solid precipitation), collection efficiency E, characteristic capture length, λ , and characteristic length, D_m , vary with temperature. This temperature dependence is related to the assumption of the temperature regimes associated with types of hydrometeors and their shapes and sizes. However, the Wang2014 parameterization was developed assuming ambient temperatures of 15 °C for rain scavenging and -10 °C for snow scavenging, and an ambient pressure of 1013.5 hPa for both rain and snow scavenging. This introduces some uncertainties in the Wang et al (2014) approach, when the atmospheric conditions differ from these assumptions.
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2.4 Observation for model evaluation

The observation data used for model evaluation are from several surface monitoring networks, namely, precipitation chemistry data from the Canadian Air and Precipitation Monitoring Network (CAPMoN), Alberta Precipitation Quality Monitoring Program (APQMP) and air concentration of PM_{2.5} and speciated PM_{2.5} data from National Air Pollution Surveillance (NAPS).

2.4.1 **Precipitation samples**

Precipitation samples were collected at five sites by APQMP, and at three sites by CAPMoN (https://www.canada.ca/en/environment-climate-change/services/air-pollution/monitoring-networks-data/canadianair-precipitation.html). Measurement locations are shown in Figure 1. These samples are collected using wet-only

270 precipitation samplers. The samplers are designed to be operated only during precipitation; the sampling container lids open when the precipitation is detected by the heated precipitation sensors. The primary goal of the precipitation sample collection is analysis of the major ions. For the CAPMoN samples, the collector container was lined with a polyethylene bag which was removed, sealed, weighed, refrigerated, and shipped to the laboratory for major ion analysis. For the APQMP samples, the samples were transferred from the clean collection container to a smaller sample bottle, capped, refrigerated if stored on site, and shipped to the laboratory for analysis. The collection frequency varied between sites, with some sites collecting daily samples, others collecting weekly samples.

Using recommended methods and completeness criteria of WMO/GAW (2004, updated 2015), quality control procedures were performed by the collecting networks and precipitation-weighted mean concentrations of $SO_4^=$, NO_3^- and NH_4^+ were calculated from the samples.

280 Overall, the collecting method described above tends to underestimate the total precipitation amount, due to wind and evaporative loss, and delay in lid opening relative to the commencement of precipitation, so the flux of ions derived from the samples must be corrected using independent observations of total precipitation values. CAPMoN sites were equipped with separate standard rain and snow gauges for measuring the precipitation amounts in order to carry out this correction. These standard gauges were not available on APQMP sites, and hence deposition fluxes precipitation corrections were calculated using daily precipitation depth data from the nearest meteorological station (ECCC, 2022; AAF, 2022).

2.4.2 Speciated and total aerosol concentrations

PM_{2.5} speciation program of NAPS network includes the measurement of PM_{2.5} mass and speciated PM_{2.5} (e.g. sulphate, nitrate and ammonium) at the existing Canada-wide air quality monitoring sites (https://www.canada.ca/en/environment-climate-change/services/air-pollution/monitoring-networks-data/national-

<u>air-pollution-program.html</u>). Speciated PM_{2.5} samples were collected using a Partisol-Plus Model 2025-D sequential dichotomous particle samplers, by splitting the incoming PM₁₀ sample stream into fine and coarse fractions with a virtual impactor. The mass flow rates of the fine and coarse particle streams were maintained at 15 L min⁻¹ and 1.7 L min⁻¹, respectively. PM_{2.5} speciation samples were collected using a Partisol Model 2300 sequential speciation samplers, equipped with three Harvard designed, Thermo Scientific ChemComb[®] cartridges. The cartridges were designed to separate PM_{2.5} impactor inlets and maintain a constant flow rate of 10 L min⁻¹. More information regarding the data collection and analysis by NAPS is available from Dabek-Zlotorzynska *et al.* (2011).

3 Results and Discussion

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3.1 Modelled wet deposition

- 300 Makar *et al.*, (2018) evaluated the model total (wet and dry) deposition against observed precipitation data from the Athabasca oil sands region, and concluded that wet deposition dominates based on an earlier version of the GEM-MACH model. The contribution to April and July 2018 mean wet deposited flux for different components of sulfur and nitrogen for the base-case experiment, are shown in Figures 5 and 6, respectively (the domain-average values are shown in figures' captions). The wet deposition fluxes include both in-cloud and BCS. The wet scavenging of sulphate
- aerosol is lower during the cold season than in the warm season. HSO_3^- deposition mostly occurs close to the SO_2

emission sources as it is associated with wet scavenging of gas phase SO_2 , while the wet deposition of the oxidized form, SO_4^{-} , extends to a broader area downwind from the emission sources. Shown in Fig. S2 are maps of modelled average SO_2 concentration at the model hybrid level of 0.98 over the region for the periods of our simulations. The "hotspots" of SO_2 indicates the locations of major SO_2 emission sources in the oil sands area. The higher monthly total

- 310 precipitation in July (Fig. 7) affects the $SO_4^=$ deposition fluxes in July (Fig. 5). Furthermore, deposition of N species is higher in July than April due to higher total precipitation (Fig. 7), agricultural activities, and increased contributions of bi-directional fluxes when the ground is unfrozen. The comparison between wet deposition of N components (Fig. 6) shows higher contributions of NH_4^+ scavenging, especially in agriculture-rich areas during July, where reduced N is deposited closer to the emission source. The higher NO_3^- flux in winter (cold season) is likely driven by the uptake
- into cloud water of higher particulate nitrate concentrations. Both the temperature dependence of particle nitrate formation (which favours colder temperatures), and the shallower boundary layer in the winter, lead to higher particulate nitrate in wintertime, which may then be scavenged below clouds. This is further influenced by the elevated NOx emissions during the winter due to increased energy demand. Note that the NH₄⁺ wet flux includes contributions from scavenging of particulate NH₄⁺ and gaseous Ammonia (NH₃), and NO₃⁻ wet fluxes from both scavenging of particulate nitrate and gaseous Nitric acid (HNO₃).

Figure 8 shows the differences in wet deposition flux between the base and multi-phase experiments for SO_3^- , SO_4^- , NO_3^- and NH_4^+ , respectively. We computed the 90% confidence interval scores for each of the fields examined. The approach follows Makar et al. (2021) and Geer (2016), using a 90% confidence level in model predictions, with the statistical measures considered different at the 90% confidence level when the 90% confidence ranges do not

- 325 overlap. The 90% confidence interval scores for each of these difference fields are shown in Figure 9. Figure 8 shows a decrease of wet deposition of HSO₃⁻ when multi-phase partitioning is included. Although the mean wet deposition of SO₄⁼ is decreased for some areas, Figure 8 indicates overall enhancement of the scavenged sulphate particles mostly by precipitation partitioning (e.g. for multi-phase experiments). When multi-phase partitioning is included explicitly in the cloud processing parts of the model, a larger amount of solid phase precipitation occurs
 330 than would be the case using a temperature threshold. These changes in phase may result in the increase and
- than would be the case using a temperature threshold. These changes in phase may result in the increase and decrease of $SO_4^=$ and HSO_3^- , respectively. In the model, HSO_3^- is formed from SO_2 dissolved into liquid water (cloud droplets or rain droplet), and it is assumed that ice particles (cloud ice or snow) do not take up SO_2 . If solid precipitation dominates, there will be less uptake (or scavenging) of SO_2 into precipitation. This is in contrast to the case of particle $SO_4^=$ the process of particle scavenging (including particle sulphate) by precipitation is more
- efficient for solid precipitation (snow) than liquid precipitation (rain) as illustrated in Figure 2. NO_3^- and NH_4^+ do not experience uniform changes, and indicate both increases and decreases of wet scavenging, depending on location. The 90% confidence intervals show that the largest geographical area of the results between the two models differing at greater than 90% confidence is for HSO_3^- , with almost all of the difference field being significant above the 90% confidence level. Amongst SO_4^- , NO_3^- and NH_4^+ , the areas of confidence level above 90% for SO_4^-
- 340 are relatively more extensive and consistent with the areas of increased wet deposition in Fig. 8, while the areas of confidence level above 90% for the other two ions are small suggesting that these differences are occurring more sporadically, probably linked to differences in temperature and rainfall rates, as described above.

The comparison between the Multi-phase and Wang2014 runs for April 2018 shows the impact of using different BCS parameterizations, Slinn1984 vs. Wang2014 (Fig 10). The corresponding 90% confidence interval scores are shown

345 in Figure 11. Both formulations treat SO₂ scavenging in the same manner; differences are associated only with particle scavenging methodology. The differences in wet deposition flux of HSO₃⁻ between the two runs are due in part to the response of the meteorological system to changes in CDNC location and amount, and aerosol radiative properties (i.e. to the aerosols and meteorology feedbacks). Overall, the Wang2014 scheme has slightly lower HSO_3^- caused by the feedback in the model, and mixed changes of SO₄⁼, NO₃⁻ and NH₄⁺. These changes are not significant (refer to the mean domain values in the figures captions). July 2018 plots (Figs S3 and S4) show generally similar patterns to the

April plots (Figs 8 and 10) where the multi-phase and Wang2014 scheme are compared.

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To illustrate the distinction between the Slinn1984 and Wang2014 schemes, and to explain the differences shown in Figure 10, the scavenging coefficients based on the domain averaged precipitation intensity from the two schemes are compared in Figure S5. In Figure S5a (rain scavenging), two different schemes (Slinn1984 and Wang2014) have 355 relatively similar scavenging coefficients during April, but the scavenging coefficients are more different during July, especially for aerosols at the size range of 0.1-1 μ m. The lower scavenging of the Slinn's scheme can be explained by its lack of processes such as thermophoresis in the Greenfield gap as discussed in section 2.3. For snow scavenging, Fig. S5b shows a significant difference between the two schemes for particles smaller than 1 µm, the Wang2014 scavenging coefficient being considerably higher than Slinn1984. Given the fact that the solid precipitation is dominating in the April precipitation, we expect to see higher particle wet deposition fluxes by using Wang2014 than

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by using Slinn1984 scheme (refer to Fig. 2 and Fig. S1).

3.2 Comparison with observations (precipitation chemistry)

Figure 12 shows the comparisons between measured and modelled precipitation amounts and wet deposition fluxes of SO₄⁼, NO₃⁻ and NH₄⁺, from the three different GEM-MACH experiments (e.g. Base-case, multi-phase and 365 Wang2014) for April. The statistical evaluations are also summarized in Fig. 12 for the model versus observation, and in Table 1 for the model versus APQMP and CAPMoN sites. Sample collection occurred daily at CAPMoN sites and weekly at the APQMP sites separately (given the difference in sample collection frequency and protocols). GEM-MACH captures the precipitation events/amounts well. The precipitation partitioning and the Wang2014 scheme do not directly affect model simulation of precipitation values. Some changes (especially local changes) are expected due 370 to the aerosol feedbacks included in the model; however the different GEM-MACH experiments show relatively similar precipitation amounts, overall (Fig. 12a-c and Table 1). The slight distinction between the precipitation amounts of three experiments is due to the feedbacks between modeled aerosols and cloud microphysics.

Comparison of the observed $SO_4^{=}$ data with the simulation results (Fig. 12d-f), suggests an overall better agreement with observations by including the multi-phase partitioning, and further improvement in agreement associated with

375 the use of the Wang et al. (2014) scavenging scheme. As shown in table 1, the normalized mean bias (NMB) values of SO_4^{-1} for the multi-phase and Wang2014 experiments are improved compared to the base-case (from 0.46 to -0.05) due to precipitation partitioning, and Wang2014 experiment has the best correlation (R = 0.86, compared to 0.83 for base run and 0.84 for multi-phase) and the best factor 2 score (0.64, compared to 0.57 for both base run and multiphase) at APQMP sites (Table 1). For the CAPMoN sites, the correlation values for $SO_4^=$ are slightly better for the

- 380 multi-phase and Wang2014 experiments (R = 0.92 and 0.93), however, the NMB value is smaller for the base experiment (NMB = 0.10, compared to 0.27 and 0.30 for the other two runs). All of these NMB values are improved compared to the earlier study of Makar *et al.* (2018) using a previous version of GEM-MACH, wherein wet sulphate precipitation fluxes were biased high (the slope of the linear fit of 2.2), although the low bias for the base experiment suggests this overall improvement may be due to other changes aside from wet scavenging. The latter reference
- 385 suggested the high bias may reflect an underestimation of the SO₂ dry deposition flux closer to the oil-sands sources, and a corresponding overestimation of in sulphate particles downwind. More recent work (Hayden et al, 2021) suggests that an underestimate in modelled SO₂ dry deposition fluxes relative to observations may be due to previously missing impacts of co-deposition of base cations on surface pH and hence SO₂ dry deposition velocity. Although CAPMoN and APQMP sites are located far from the oil-sands SO₂ emission sources; the plots seem to indicate a
- 390 positive bias only in modelled S wet flux from the base-case experiment at the APQMP sites, which has diminished in the other two experiments. Biases in S wet deposition were negative at the CAPMoN sites. We note that the current model emissions year (2018) has lower reported SO₂ emissions than the 2013 emissions year simulated in Makar et al (2018), though discrepancies between reported and satellite-derived SO₂ emissions have been noted (McLinden et al., 2020). The current model version also has higher particle dry deposition velocities than in Makar et al. (2018),
- following Emerson et al (2020), hence results in less particle sulphate being available for wet scavenging near the surface.

Overall, GEM-MACH estimates of wet deposited nitrogen are slightly biased low relative to observations. For both APQMP and CAPMoN, NO_3^- has better correlation values (R = 0.58 and 0.76, respectively) for the multi-phase and Wang2014 experiments relative to the base-case. GEM-MACH experiments have relatively similar statistical scores for the APQMP NH₄⁺ results, with the highest factor 2 value for the Wang2014 (Fac2 = 0.71). CAPMON NH₄⁺ shows

slightly a better correlation (R = 0.69) and a better NMB for the Wang2014 experiment.

3.3 Comparison with speciated PM data (NAPS)

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The impacts of partitioning and Wang2014 scavenging on modelled ambient concentration of speciated PM2.5 (sulphate (SU), nitrate (NI) and ammonium (AM)), near the surface, for April 2018 are shown in Figures 13, 14 and
15. The upper panels are April mean of SU2.5, NI2.5, and AM2.5 for the base-case, and the middle left panels show the net difference between multi-phase and base-case Slinn1984 experiments, while the middle right panels show the net difference between Wang2014 and multi-phase experiments. Corresponding 90% confidence interval scores for the difference plots are shown in the lower panels. Multi-phase partitioning leads to higher modelled concentration of particulate sulphate in the atmosphere, and the increases are statistically significant at the 90% confidence level. For particulate nitrate, partitioning leads to both increase in particulate nitrate over the OS source area and downwind is not significant at the 90% confidence level. For particulate ammonium, the increase of the concentration associated

with the partitioning approaches is the dominant change over the entire region, while a decrease occurs near the oilsands emissions sources.

- 415 Figure 16 and Table 2 show GEM-MACH simulation results of speciated aerosols, in the 2.5-km domain, compared with the daily NAPS observation data for April 2018. For particulate sulphate, both multi-phase and Wang2014 experiments have lower NMB compared to the base-case, and the correlation value is improved for the Wang2014. Particulate nitrate model outputs show almost the same statistical evaluation results for all three experiments, and the correlation between the measured and model using Wang2014 scheme is the highest for the ammonium. Overall the
- 420 results show enhanced performance in the multi-phase experiment particularly over the source area and downwind. Model simulations making use of Wang2014 show mostly enhanced performance over OS facilities area and downwind, and a reduction over the area upwind of the OS, and these results are consistent with the changes in the wet scavenging.

4. Conclusions

- 425 To examine cloud processes and precipitation chemistry, we considered the co-existence of multi-phase precipitation, as predicted by the GEM microphysics, in GEM-MACH's BCS representation. Further, we implemented a new aerosol BCS scheme (Wang et al., 2014) and compared it with the GEM-MACH's existing scavenging scheme, based on Slinn (1984).
- An accurate representation of the precipitation phase is important in modelling the wet scavenging of atmospheric
 tracers particularly in cold environments. Considering the coexistence of multi-phase precipitation in BCS has a more consistent impact on the precipitation scavenging of SO₂. Here we have noted overall improvements in model performance based on the model-observation performance scores associated with multi-phase partitioning and in comparison to wet deposition evaluations carried out in the region in previous work (Makar et al., 2018). For example, the multi-phase approach resulted in the most significant improvement in modelled SO₄⁼ wet deposition flux over Alberta (at APQMP sites, reducing NMB from 0.46 to -0.05), as well as improvement in modelled ambient particulate
- sulfate concentration at NAPS sites.

As shown in this study and other existing studies, there is a considerable uncertainty in the various existing parameterizations for BCS of aerosol particles. Of the two schemes examined, the Slinn (1984) parameterization is theoretically based and lacks representation of several physical processes involved in the particle scavenging by

- falling hydrometeors (such as thermophoresis and diffusiophoresis). The Wang *et al.* (2014) scheme is based on a semi-empirical approach, and implicitly accounts for electrostatic forces, which are shown to be more important than diffusiophoresis in Jones *et al.* (2022). This scheme provided an overall best fit to an ensemble of existing models, although there is still a significant disparity between the scavenging coefficients based on Wang *et al.* (2014) and some of the observation-based scavenging coefficients (*e.g.*, Jones *et al.* 2022). The resulting scavenging
- 445 coefficients from the two schemes (Slinn1984 and Wang2014) show the greatest difference for aerosol sizes between 0.1 to 1 μm, particularly at high precipitation intensity. For the solid precipitation at lower intensity, the

two schemes diverge for aerosols smaller than 1 μ m, while at higher precipitation intensities, the two schemes show higher disparity for aerosols larger than 1 μ m. This resulted in the varied differences in modelled wet deposition (April – winter, July – summer), especially, higher Wang2014 scavenging coefficient for particles smaller than 1 μ m during the April, and higher particle wet deposition fluxes from using Wang2014 than using Slinn1984.

The model evaluation against observations (precipitation chemistry and ambient air concentration of speciated PM) seems to indicate that the use of Wang2014 parameterization along with the consideration for co-existence of multiphase precipitation results in the best scores overall, with the most significant improvement from the multi-phase partitioning in BCS. The multi-phase partitioning and Wang2014 scheme improve the comparison between observation and modeled results. Comparison of the SO₄⁼ data from the APQMP sites with the simulation results suggests better agreement by including the multi-phase partitioning and further improvement in agreement associated with the Wang et al. (2014) scavenging scheme. GEM-MACH estimates of wet deposited nitrogen are slightly biased low, and the simulated results improve by adding the partitioning and Wang2014 scheme (relative to the base-case). GEM-MACH experiments have relatively similar statistical scores for NH₄⁺ results. Including the partitioning and

- 460 Wang2014 scavenging impacts the modelled ambient concentration of speciated PM2.5 (sulphate, nitrate and ammonium). It leads to improvements in the model performance scores, with higher modelled concentration of particulate sulphate, and both increase and decrease of particulate nitrate concentration with an overall increase over, the OS source area and downwind. For particulate ammonium, the increase of the concentration is the dominant change over the entire region.
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Code Availability

The GEM-MACH code can be downloaded from this Zenodo site: <u>GEM-MACH | Zenodo</u>. GEM-MACH model output data are available by email request from Roya Ghahreman: <u>roya.ghahreman@ec.gc.ca</u>. The model output requires a large amount of storage space and is in a binary format specific to Environment and Climate Change Canada's modelling systems. Conversion to other formats may be possible on request by email.

Author contributions

WG and RG designed the study. RG implemented the new scavenging scheme in the model, ran the experiments, and analysed the output under the supervision of WG and PM and with assistance from AL, CL and AA. AC and KB provided the measurement data. RG prepared the paper with contributions from all co-authors.

Competing interests

The authors declare that they have no conflict of interest.

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Figure 1: Model domains - 2.5 km \times 2.5 km in red and 10 km \times 10 km in blue. CAPMoN and APQMP observation stations are shown in green and yellow dotes, respectively.







Figure. 2: Mean April: a. temperature, b. solid precipitation flux, c. liquid precipitation flux at the model hybrid level of 0.98 (e.g. at ~980 mb). d and e: as in a, b but showing approximate limits masking out the predominantly snow flux (d) and predominantly rain flux (e). The black masked regions are the portion of the solid precipitation that would be treated as liquid in the base-case due to the above-zero environment temperature (2c), and the portion of the liquid precipitation that would be treated as solid in the base-case due to sub-zero environment temperature (2e).



Figure. 3: Slinn (1984) and Wang2014 rain (left panel) and snow (right panel) scavenging coefficients (s⁻¹) versus the aerosol sizes and different precipitation intensities (R).



Figure. 4: Slinn (1984) and Wang *et al.* (2014a) snow scavenging coefficient versus aerosol size distribution with the intensity of 0.01 mm/h at three different ambient atmospheric temperatures (T = 240, 250 and 270 K). Note that the three curves associated with Wang *et al.* (2014a) collapse into a single curve due to their independence to temperature.



Figure 5: Hourly accumulated sulfur components deposition flux (μmol/m²) from the base-case experiment, averaged over April 2018 (upper panels) and July 2018 (lower panels) – SO₄⁼ (left panels), SO₃⁻ (right panels). Domain-mean deposition values are SO₃⁻ = 0.123 (9.84), SO₄⁼ = 0.083 (7.97) μmol/m² (μg/m²) for April, and SO₃⁻ = 0.027 (2.16), SO₄⁼ = 0.187 (17.95) μmol/m² (μg/m²) for July.



Figure 6: Hourly accumulated nitrogen components deposition flux (μ mol/m²) from the base-case experiment, averaged over April 2018 (upper panels) and July 2018 (lower panels) – NO₃⁻ (left panels) and NH₄⁺ (right panels). Domain-mean deposition values are NO₃⁻ = 0.139 (7.65), NH₄⁺ = 0.377 (4.15) μ mol/m² (μ g/m²) for April, and NO₃⁻ = 0.169 (9.295), NH₄⁺ = 1.950 (21.45) μ mol/m² (μ g/m²) for July.





Figure 7: Daily accumulated precipitation (PR), averaged over April and July 2018. Domain-mean values of PR are 0.90 and 3.00 mm for April and July, respectively.



Figure 8: The net differences of hourly accumulated mean deposited flux of SO₃⁻, SO₄⁼, NH₄⁺ and NO₃⁻ for the multi-phase and base-case experiments (e.g. multi-phase – base-case) over April 2018. Domain-mean deposition values are SO₃⁻ = 0.123 (9.84), SO₄⁼ = 0.083 (7.970), NO₃⁻ = 0.139 (7.645), NH₄⁺ = 0.377 (4.147) μ mol/m² (μ g/m²) for base-case, and SO₃⁻ = 0.015 (1.20), SO₄⁼ = 0.089 (8.54), NO₃⁻ = 0.135 (7.42), NH₄⁺ = 0.371 (4.081) μ mol/m² (μ g/m²) for multi-phase experiment.



Figure 9: The corresponding 90% confidence interval scores for Fig. 8 (following Makar *et al.*, 2021, the differences are significant at or above 90% confidence level when the score \geq 1; red regions identify > 90% confidence differences).



Figure 10: The net differences of hourly accumulated mean deposited flux of SO_3^{1-} , $SO_4^=$, NH_4^+ and NO_3^- for the Wang2014 and multi-phase experiments (e.g. Wang2014 - multi-phase) for April 2018. Domain-mean deposition values are $SO_3^- = 0.014$ (1.120), $SO_4^= = 0.089$ (8.540), $NO_3^- = 0.135$ (7.425), $NH_4^+ = 0.371$ (4.081) μ mol/m² (μ g/m²) for Wang2014, and $SO_3^- = 0.015$ (1.20), $SO_4^= = 0.089$ (8.544), $NO_3^- = 0.135$ (7.425), $NH_4^+ = 0.371$ (4.081) μ mol/m² (μ g/m²) for multi-phase experiment.



Figure 11: The corresponding 90% confidence interval scores for Fig. 10 (following Makar *et al.*, 2021, the differences are significant at or above 90% confidence level when the score \geq 1).



Nitrate Wet Deposition (μ mol m⁻²) – APQMP + CAPMoN



Figure 12: Scatter plots for the precipitation amounts (a-c), SO₄⁼ (d-f), NO₃⁻ (g-i), and NH₄⁺ (j-l) wet deposition fluxes (μmol m⁻²) for April. CAPMoN and APQMP versus GEM-MACH simulations: base-case vs observation (left panels), multi-phase vs observation (middle panels) and Wang2014 vs observation (right panels).









Figure 13: April 2018 mean fine particulate sulphate (SU2.5) concentration near the surface (a). The net difference between multi-phase and base-case (b) and the net difference between Wang2014 and multi-phase (c). The corresponding 90% confidence interval scores are shown in the lower panels.

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950 Figure 14: April 2018 mean fine particulate nitrate (NI2.5) concentration near the surface (a). The net difference between multi-phase and base-case (b) and the net difference between Wang2014 and multi-phase (c). The corresponding 90% confidence interval scores are shown in the lower panels.

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Figure 15: April 2018 mean fine particulate ammonium (AM2.5) concentration near the surface (a). The net difference between multi-phase and base-case (b) and the net difference between Wang2014 and multi-phase (c). The corresponding 90% confidence interval scores are shown in the lower panels.



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Figure 16: GEM-MACH simulation results of speciated aerosols - Sulphate (upper panels, a-c), nitrate (middle panels, d-f) and Ammonium (lower panels, g-i) - compared with the daily NAPS observation data for April 2018. Base-case (or Slinn (1984) scheme) vs observation (left panels), multi-phase vs observation (middle panels) and Wang2014 vs observation (right panels).

Table 1: Performance scores for PR and deposited SO₄⁻, NO₃⁻ and NH₄⁺ with the different scavenging approaches the comparisons between the measured and modelled precipitation amounts and wet deposition fluxes of SO4⁼, NO3⁻ and NH4⁺. Highest scores are indicated in **bold face type**.

		APQMP		CAPMoN			
		R	NMB	FAC. 2	R	NMB	FAC. 2
PR							
	Base-case	0.73	0.39	0.29	0.68	0.20	0.34
	multi-phase	0.73	0.37	0.29	0.71	0.19	0.32
	Wang2014	0.73	0.37	0.29	0.69	0.18	0.32
SO4=							
	Base-case	0.83	0.46	0.57	0.90	-0.10	0.33
	multi-phase	0.84	-0.05	0.57	0.92	-0.27	0.47
	Wang2014	0.86	-0.05	0.64	0.93	-0.30	0.33
NO ₃ -							
	Base-case	0.51	-0.09	0.43	0.73	-0.68	0.27
	multi-phase	0.58	-0.13	0.43	0.76	-0.68	0.33
	Wang2014	0.58	-0.11	0.57	0.76	-0.68	0.27
$\mathrm{NH_{4}^{+}}$	5	0.02	0.1.4		0.60	0.47	0.40
	Base-case	0.93	-0.14	0.64	0.68	-0.47	0.40
	multi-phase	0.92	-0.14	0.64	0.68	-0.46	0.53
	Wang2014	0.93	-0.14	0.71	0.69	-0.44	0.47

Table 2: Performance scores for the near surface concentrations of speciated aerosols, sulphate, nitrate and ammonium, in the 2.5-km domain, compared with the daily NAPS observation data for April 2018.

		NAPS				
		R	NMB	FAC. 2		
Sulphate	Base-case	0.65	-0.41	0.54		
	multi-phase	0.69	-0.38	0.58		
	Wang2014	0.70	-0.38	0.58		
Nitrate	Base-case	0.70	7.18	0.04		
	multi-phase	0.70	7.32	0.04		
	Wang2014	0.70	7.07	0.04		
Ammonium	Base-case	0.48	2.27	0.46		
	multi-phase	0.45	2.35	0.46		
	Wang2014	0.50	2.26	0.46		