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A cloud chemistry module for the 3-D cloud-resolving mesoscale model Meso-NH with application to idealized cases

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Abstract

A complete chemical module has been developed for use in the Meso-NH three-dimensional cloud resolving mesoscale model. This module includes gaseous and aqueous phase chemical reactions that are analysed by a pre-processor generating the Fortran90 code automatically. The kinetic solver is based on a Rosenbrock algorithm, which is robust and accurate for integrating stiff systems and especially multiphase chemistry. The exchange of chemical species between the gas phase and cloud droplets and raindrops is computed kinetically by mass transfers considering non-equilibrium between the gas and the condensed phases. Microphysical transfers of chemical species are considered for the various cloud microphysics schemes available, which are based on one-moment or two-moment schemes. The pH of the droplets and of the raindrops is diagnosed separately as the root of a high order polynomial equation. The chemical concentrations in the ice phase are modelled in a single phase encompassing the two categories of precipitating ice particles (snow and graupel) of the microphysical scheme. The only process transferring chemical species in ice is retention during freezing or riming of liquid hydrometeors. Three idealized simulations are reported, which highlight the sensitivity of scavenging efficiency to the choice of the microphysical scheme and the retention coefficient in the ice phase. A two-dimensional warm, shallow convection case is used to compare the impact of the microphysical schemes on the temporal evolution and rates of acid precipitation. Acid wet deposition rates are shown to be overestimated when a one-moment microphysics scheme is used compared to a two-moment scheme. The difference is induced by a better prediction of raindrop radius and raindrop number concentration in the latter scheme. A two-dimensional mixed-phase squall line and a three-dimensional mixed-phase supercell were simulated to test the sensitivity of cloud vertical transport to the retention efficiency of gases in the ice phase. The 2-D and 3-D simulations illustrate that the retention in ice of a moderately soluble gas such as formaldehyde substantially

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decreases its concentration in the upper troposphere. In these simulations, retention of highly soluble species in the ice phase significantly increased the wet deposition rates.

1 Introduction

More than 50 % of the Earth's surface is under cloud and several studies have shown that clouds interact with chemical species in many ways, over a wide range of scales, from micrometres up to thousands of kilometres. On global and regional scales, clouds have a major impact on the composition of the troposphere through multiphase removal processes (Tost et al., 2007). The impact of clouds on the ozone budget and on simple soluble compounds, such as hydrogen peroxide, has been well assessed (Lelieveld and Crutzen, 1991; Monod and Carlier, 1999). However, there are still uncertainties concerning the impact of cloud vertical transport of chemical species by deep convection on the Upper Troposphere (UT) composition, and about the impact of the chemical reactivity of organic compounds in clouds on the formation of Secondary Organic Aerosol (SOA). For instance, the source of HO_x in the deep convective tropical cloud outflow needs more investigation since the production of ozone in the UT is almost proportional to the HO_x mixing ratio (Wennberg et al., 1998; Jaeglé et al., 2001). Moreover, the nucleation of new particles observed downwind of the anvil of deep convective tropical clouds is still uncertain. Discrepancies are observed between in-situ measurements and theoretical calculations pointing out the possible role of volatile organic compounds associated with sulphur dioxide as precursors of aerosol particles (Waddicor et al., 2012). Organic aerosols affect the earth's radiative budget by their role in both direct and indirect aerosol forcing (Kanakidou et al., 2005). The majority of the organic fraction of aerosols is suspected to be of secondary origin. However, the sources, chemical composition and formation mechanisms of SOA remain one of the least well understood processes relevant to the atmosphere (Hallquist et al., 2009). In particular, the potential contribution of the aqueous phase reactivity is highly uncertain,

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as is the chemical nature of the aqueous phase products that are precursors of SOA (Hallquist et al., 2009; Ervens et al., 2011).

The importance of the vertical transport of chemical species by convection has been underlined by many authors (Dickerson et al., 1987; Prather and Jacob, 1997; Lawrence and Crutzen, 1998; Mari et al., 2000 for instance). In particular, local convection is a major source of HO_x in the UT (Jaeglé et al., 1998). This production of HO_x in the UT perturbed by deep convection is mainly due to photochemical reactions of hydrogen peroxide, methyl hydroperoxide, and formaldehyde, which are transported from the boundary layer to the UT by convection or arise from secondary production in the UT (Jaeglé et al., 1997; Cohan et al., 1999). As these species are soluble, they are also impacted by cloud microphysical processes and aqueous phase chemistry (Barth et al., 2007b). Another important topic concerns new routes of SOA formation through the condensation precursors having low volatility that are formed in cloud droplets or raindrops and released in the clear atmosphere as the clouds or rain evaporate (Chen et al., 2007; Lim et al., 2010; Ervens et al., 2011). Assessing the budgets of HO_x or SOA thus requires a detailed understanding of the coupling between cloud venting, microphysics and aqueous chemistry.

However, the parameterization of convective transport and gas scavenging at the global scale still remains approximate (Tost et al., 2010) due to the huge number of non-linear processes and the high variability of the solubility and reactivity of the chemical compounds. Meanwhile, at convective-resolved scale, current computational power enables Cloud Resolving Models (CRM) to be run, where interactions between the cloud microphysics and the chemistry and the advective/turbulent transport of chemical species can be reasonably well detailed (Flossmann and Wobrock, 1996; Barth et al., 2001; Yin et al., 2002). As a result, there is a great need to study and develop an efficient gaseous and aqueous chemical scheme tightly coupled to the microphysics of mixed-phase clouds in order to evaluate the budget of chemical compounds after a perturbation caused by convective events.

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5 A CRM is a powerful tool for studying the complex interactions between transport, chemistry and cloud microphysics. Whether they are used for improving the representation of the vertical redistribution of gases and aerosol particles by convective clouds, or to contribute to the assessment of SOA formation by cloud processes, CRMs have to integrate a cloud chemistry module, as well as an aerosol module and detailed cloud microphysics.

10 In this study, a cloud chemistry module for the three-dimensional meteorological model Meso-NH (Lafore et al., 1998) is developed and tested. Among the three ingredients: transport, microphysics and chemistry, the coupling between microphysics and chemistry is the most original part of the package, in particular in cases of mixed-phase cloud. The new module takes advantage of the resolved and turbulent transport schemes and the mixed phase cloud microphysical scheme, which have been continuously improved, in the host model. Moreover, an aerosol module is available in Meso-NH (ORILAM, for Organic Inorganic Lognormal Aerosol Model, Tulet et al., 2005, 15 2006), although its coupling with the cloud chemistry module is not presented in the present work.

20 First, the chemical module is described, including the detailed treatment of the temporal integration of the chemical production and destruction terms and of the cloud microphysics transfer terms. The diagnostic computation of the pH is also detailed. Then, three applications of the model are presented. The first one is a warm, idealized, two-dimensional precipitating case to focus on the sensitivity of aqueous phase chemistry to the cloud microphysics scheme: one-moment versus two-moment. The second case corresponds to an idealized, two-dimensional squall line to underline the effect of the ice phase on cloud chemistry via the retention of chemical species when riming or freezing occurs. The last case is the simulation of a mixed-phase, three-dimensional supercell, which has been widely studied in the framework of a model intercomparison exercise (Barth et al., 2007a). Finally, some perspectives concerning the use of the full package for simulating complex three-dimensional cloud situations are discussed, 25 together with possible extensions of the module.

2 Description of the cloud chemistry module

The cloud chemistry module was implemented in the meteorological Meso-NH model. Meso-NH is a non-hydrostatic mesoscale atmospheric model, which was jointly developed by CNRM (Météo France) and Laboratoire d'Aérodynamique (CNRS) (Lafore et al., 1998). Meso-NH simulates small-scale (LES with horizontal resolution of a few metres) to synoptic-scale (km scale resolution up to several tens of kilometres) phenomena. Different sets of parameterizations have been introduced for convection (Bechtold et al., 2001; Pergaud et al., 2009), cloud microphysics (Pinty and Jabouille, 1998; Cohard and Pinty, 2000a; Khairouidinov and Kogan, 2000), turbulence (Cuxart et al., 2000), surface processes (Noilhan and Mahfouf, 1996; Masson, 2000), gaseous chemistry (Suhre et al., 2000; Tulet et al., 2003), aerosol chemistry (Tulet et al., 2005, 2006) and cloud electricity including lightning flash production (Barthe et al., 2012). The new cloud chemistry module represents chemistry processes in both warm clouds and mixed-phase clouds.

In nested mode, it is possible to activate the cloud chemistry module in only the inner domain to save computing time while the coupling models (“father” models) treat the gas phase chemistry only.

Continuity equations for a chemical species X (mol per volume of dry air) in the gas phase and in the aqueous phase are of the form:

$$\frac{\partial X_g}{\partial t} = \frac{\partial X_g}{\partial t} \Big|_{\text{dyn}} + \frac{\partial X_g}{\partial t} \Big|_{\text{chem}} + \frac{\partial X_g}{\partial t} \Big|_{\text{others}} \quad (1)$$

$$\frac{\partial X_w}{\partial t} = \frac{\partial X_w}{\partial t} \Big|_{\text{dyn}} + \frac{\partial X_w}{\partial t} \Big|_{\text{chem}} + \frac{\partial X_w}{\partial t} \Big|_{\text{others}} \quad (2)$$

In Eq. (2), the subscript “w” stands for either cloud droplets or raindrops. In both equations, the term dyn refers to dynamical tendencies by advection and turbulence, which applies to all prognostic scalars in the model. The term others describes emission, dry

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deposition and release from the aqueous phase when evaporation, freezing or riming occurs. This point is detailed in Sect. 2.5. For the aqueous phase species, the term others represents the cloud microphysical processes, which depend on the cloud microphysics scheme and are detailed in Sect. 2.2 for warm clouds and in Sect. 2.5 for mixed-phase clouds. Finally, the term chem includes gas-liquid transfer and chemical reactions and will be explained in the following subsection. To avoid numerical problems, Eq. (2) is solved only when the liquid water content of cloud water or rainwater goes beyond a threshold value fixed by the user. A typical value is $1 \times 10^{-8} \text{ vol vol}^{-1}$. If the liquid water content is below this value, the concentrations of the chemical species in water are set to zero after transfer to the gas phase. For ions, which are not linked to a gas phase species by dissociation equilibria, the concentrations are set to zero and the associated mass is lost. However, this concerns very few species with very low concentrations (intermediate sulphur species, see below and Table 3).

2.1 Chemical kinetic scheme

The evolution of the chemical concentrations in the gas and liquid phases, denoted by the term *chem* above, of a chemical species *X* is given by the generic set of differential equations:

$$\left. \frac{\partial X_g}{\partial t} \right|_{\text{chem}} = P_g - D_g X_g - k_{tw} \left(L_w X_g - \frac{X_w}{H_{\text{eff}} RT} \right) \quad (3)$$

$$\left. \frac{\partial X_w}{\partial t} \right|_{\text{chem}} = P_w - D_w X_w - k_{tw} \left(L_w X_g - \frac{X_w}{H_{\text{eff}} RT} \right) \quad (4)$$

X_g and X_w are the concentration of *X* in the gas phase and the liquid phases, respectively. P_g and P_w , and D_g and D_w are the gaseous and the aqueous production terms in mol per volume of dry air s^{-1} ; and the gaseous and aqueous destruction terms in s^{-1} , respectively. L_w is the liquid water volume ratio (volume of the drops per volume of air). H_{eff} is the effective Henry's law constant in mol atm^{-1} . T is the temperature in K and

$R = 0.08206 \text{ atm M}^{-1} \text{ K}^{-1}$ is the universal gas constant. The rate constant of transfer between the gas phase and the aqueous phase k_{tw} in s^{-1} is the inverse of the characteristic times for gaseous diffusion and for the interfacial mass transport according to Schwartz (1986):

$$k_{\text{tw}} = \left(\frac{a_w^2}{3D_{\text{gas}}} + \frac{4a_w}{3\bar{v}\alpha} \right)^{-1} \quad (5)$$

a_w is the cloud droplet or raindrop radius in m, D_{gas} is the gaseous diffusion coefficient taken equal to $10^{-5} \text{ m}^2 \text{ s}^{-1}$ for all the species, \bar{v} is the mean quadratic speed in m s^{-1} of the soluble species (see Leriche et al., 2000) and α is the accommodation coefficient of the soluble species. For cloud droplet and raindrop populations, a_w is taken as the mean radius of the respective size distributions. If a one-moment scheme is used, the cloud droplet radius is fixed ($a_w = 10 \mu\text{m}$) and the mean raindrop radius is diagnosed. Both a_w are computed if a two-moment scheme is chosen.

As a part of the source code solving the chemistry depends on the chemical mechanism, Meso-NH includes a kinetic pre-processor similar to KPP (Damian et al., 2002), which can be applied to any chemical reaction mechanism to automatically generate the Fortran90 code to compute the reaction rates and the Jacobian for the kinetic solver. In the present study, the chemical mechanism for the gas phase chemistry was an updated version of the ReLACS scheme (Regional Lumped Atmospheric Chemical Scheme, originally with 37 prognostic gaseous species and 128 reactions, see Crassier et al., 2000). Compared to the original ReLACS scheme, four prognostic gaseous species were added for the purposes of the aqueous chemistry application (ammonia, sulphuric acid, hydroxyl radical and formic acid), leading to 41 prognostic species in the gas phase. The chemical mechanism developed for the aqueous phase considers 20 soluble compounds, which are exchanged between the gas phase and liquid phases (cloud water and rainwater), and five intermediate ions in the liquid phases describing the sulphur chemistry. The resulting chemical scheme includes

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gaseous chemical reactions, reactions of mass transfer for the exchange of soluble species between the gas phase and liquid phases, and aqueous chemical reactions. This new scheme is called ReLACS-AQ and includes 91 prognostic species. In case of mixed phase clouds, 19 additional prognostic species were added to describe the concentration of soluble species in ice-precipitating hydrometeors (see Sect. 2.5).

Table 1 lists the chemical species exchanged between the gas and liquid phases together with the values of their Henry's law constants and accommodation coefficients. Table 2 gives the aqueous phase equilibria, which are used to compute the effective Henry's law constant and the total species mixing ratio in the aqueous phase. The chemical species involved in equilibrium are treated as total species to prevent losing mass within an aqueous phase (see Leriche et al., 2000). This approach allows us to treat the pH as a diagnosed variable (see Sect. 2.4). Finally, Table 3 presents the aqueous phase chemical mechanism of ReLACS-AQ. This mechanism was developed consistently with the gas phase mechanism ReLACS and based upon two existing reduced aqueous phase mechanisms (Tost et al., 2007; CAPRAM2.4 (Chemical Aqueous Phase RADical Mechanism) from Ervens et al., 2003). The resulting reduced ReLACS-AQ mechanism was tested by comparison with an explicit mechanism and with CAPRAM2.4 in the box model M2C2 (Model of Multiphase Cloud Chemistry; Leriche et al., 2003) using the three scenarios from Ervens et al. (2003): urban, rural and marine. Results of the time evolution of chemical species concentrations showed no significant differences between the full mechanism, the CAPRAM2.4 mechanism and the ReLACS-AQ one (the largest difference was less than 5%). As the only source of acetic acid in the aqueous phase in ReLACS-AQ is its mass transfer from the gas phase (no source of its precursors in the aqueous phase), which is of minor importance, its aqueous phase reactivity was neglected.

2.2 Cloud microphysics transfer terms

Once dissolved or produced in the drops, the aqueous chemical species are redistributed between the cloud droplets and the raindrops by the cloud microphysical

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processes that govern the formation, duration and dissipation of clouds. These processes, grouped in the term *others* of Eq. (2), are autoconversion (AUTO), accretion (ACCR) and sedimentation (SEDI). The transfers of chemical species between liquid drops and gas phase due to condensation/evaporation processes are considered by the transfer terms in Eqs. (3) and (4). Autoconversion and accretion are processes leading to raindrop formation and growth, respectively. Autoconversion and accretion processes transfer chemical materials from the cloud droplets to the raindrops. The sedimentation process describes the vertical fluxes of raindrops due to their significant terminal velocity, which ultimately lead to wet deposition of the soluble species by rain. The mass transfer of chemical species due to microphysical processes is computed using the classical assumption that it is proportional to that of the microphysical reservoirs. These microphysical transfer terms are computed as follows:

$$\left. \frac{\partial X_c}{\partial t} \right|_{\text{AUTO}} = - \left. \frac{\partial X_r}{\partial t} \right|_{\text{AUTO}} = -X_c \frac{1}{r_c} \left. \frac{\partial r_c}{\partial t} \right|_{\text{AUTO}} \quad (6)$$

$$\left. \frac{\partial X_c}{\partial t} \right|_{\text{ACCR}} = - \left. \frac{\partial X_r}{\partial t} \right|_{\text{ACCR}} = -X_c \frac{1}{r_c} \left. \frac{\partial r_c}{\partial t} \right|_{\text{ACCR}} \quad (7)$$

$$\left. \frac{\partial X_r}{\partial t} \right|_{\text{SEDI}} = -X_r \frac{1}{r_r} \left. \frac{\partial r_r}{\partial t} \right|_{\text{SEDI}} = -X_r \frac{1}{r_r} \frac{\partial}{\partial z} F_{\text{SEDI}}(r_r) \quad (8)$$

r_c and r_r are the mass mixing ratios of the cloud droplets and of the raindrops in kg of water per kg of dry air, respectively. F_{SEDI} is the sedimentation flux of the rain mixing ratio.

The sedimentation flux and the autoconversion and accretion rates depend on the cloud microphysical scheme. In Meso-NH, there are four different ways to calculate these terms. The simplest one computes the cloud microphysics with a one-moment scheme following Kessler (1969) and assuming a Marshall-Palmer distribution for the raindrops. The second one uses the one-moment scheme for mixed phase cloud, ICE3 (Pinty and Jabouille, 1998), which is equivalent, for the warm part of the scheme, to a Kessler (1969) parameterization assuming a generalized gamma distribution for

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all hydrometeors including rain. The third one, the C2R2 scheme (Cohard and Pinty, 2000a), is a two-moment scheme for warm cloud, in which the parameterization of the CCN activation follows the diagnostic and integral approach of Twomey (1959) as improved by Cohard et al. (1998). The last one is a two-moment scheme for LES or stratocumulus applications (Khairouidinov and Kogan, 2000). The detailed expressions of each term for the four cloud microphysics schemes available in Meso-NH can be found in the scientific documentation of Meso-NH (<http://mesonh.aero.obs-mip.fr/mesonh/>).

Equations (6)–(8) are integrated in the chemical tendencies (Eqs. 1 and 2) before the kinetic solver is called to resolve the chemical ODE system at each grid point of the computational domain.

2.3 Kinetic solver

The set of non-linear differential equations describing the evolution of chemical species forms a stiff ODE system (Eqs. 3 and 4). Moreover, including an aqueous phase in the chemical scheme increases the stiffness of the numerical ODE (Audiffren et al., 1998) so we substituted the kinetic solvers available in Meso-NH by the Rosenbrock family of solvers described in Sandu et al. (1997).

Rosenbrock solvers are based on multistage implicit methods with an adaptive sub-timestep to achieve high order accuracy. Each stage of the method needs to solve a system of linear equations by inverting a matrix. This is done with an **LU**-decomposition method (where **L** is a lower triangular matrix and **U** is an upper triangular matrix) and efficient index coding rules that exploit the sparsity of the Jacobian matrix of the chemical system (only the non-zero coefficients are stored).

Finally, the Rosenbrock solver suite including multiple order accuracy of the publicly available code of Sandu and Sander (2006) was coupled to the chemistry monitor of Meso-NH. For the sake of efficiency, the solver was adapted in a “vectorized” form such that several independent chemical systems (one per grid point) were solved simultaneously. Mixing the treatment of cloud-free and aqueous-phase chemistry was made possible by first ordering the gas-phase species.

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2.4 pH solver

As the solubility of some important gaseous pollutants depends on the pH (by definition $\text{pH} = -\log_{10}[\text{H}^+]$) of the drops as well as the aqueous phase reactivity. E.g. in sulphate formation, it is crucial to predict the evolution of the pH of the cloud droplets and raindrops. There are two ways to solve the pH in a cloud chemistry module.

The first method explicitly considers all the ionic species by including dissociation equilibria in the aqueous phase chemical mechanism (see Table 2) as backward and forward reactions (e.g. Ervens et al., 2003). Here, the concentration of H^+ is clearly a prognostic variable. The drawback of this method is the possibility of numerical instabilities. Because backward and forward reactions of dissociation equilibria are very fast, a very small displacement from equilibrium leads to the sporadic occurrence of abnormally high or low concentrations of species, which increases the stiffness of the system.

The second method treats the chemical species involved in equilibria as total species. For instance, formic acid in aqueous phase is the sum of the concentration of the dissolved formic acid plus the concentration of formate ions. Using this formulation, it is possible to set up an electroneutrality equation of the system and then to simplify it by keeping only the main acids and bases. The complex electroneutrality equation was further developed into a high-order polynomial equation for which the concentration of H^+ was the physical root to be selected. This operation was carefully conducted using a formal calculus software to avoid errors of manipulation. The simplified form of the electroneutrality equation deduced from the ReLACS-AQ mechanism is:

$$[\text{H}^+] + [\text{NH}_4^+] = [\text{OH}^-] + [\text{HCO}_3^-] + 2[\text{CO}_3^{2-}] + [\text{HSO}_3^-] + 2[\text{SO}_3^{2-}] + [\text{NO}_3^-] + 2[\text{SO}_4^{2-}] + [\text{HCOO}^-] + \sum [\text{ions}] \quad (9)$$

The ions in the last term are the intermediate ions of sulphur chemistry (see Table 3), which exist only in aqueous phases and are explicitly represented in the mechanism.

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Using the dissociation constants (Table 2) to replace ionic concentrations by concentrations of total species and assuming that strong acids (nitric and sulphuric) are completely dissociated, Eq. (9) leads to a polynomial equation in H^+ concentration of degree 8. A Laguerre method (Press et al., 2007) is used to extract the physical root of Eq. (9). A flag allows the pH to be computed or a constant value to be preset for the pH.

2.5 Extension to the ice phase

In mixed phase clouds, additional processes need to be considered for the soluble chemical species. These include direct gas uptake by ice crystals, partitioning during the freezing or riming of liquid hydrometeors, and the surface and bulk reactions in/on ice hydrometeors. Gas uptake by ice crystals is a complex process because the surface of ice crystals grows continuously by vapour deposition or evaporates (Pruppacher and Klett, 1997). A simple way of parameterizing this process is to introduce a single parameter, the burial coefficient, to describe different efficiencies of gas trapping in growing ice hydrometeors (Yin et al., 2002). The partitioning of soluble gases during the freezing or riming of liquid hydrometeors is classically described by a retention coefficient that partitions the fraction of a dissolved trace gas that is retained in hydrometeors during freezing/riming. Finally, information is almost non-existent for the surface and bulk reactivity of chemical species in the ice crystals and concerns mainly stratospheric conditions (Sander et al., 2006); ice reactivity is classically not considered in mixed-phase cloud modelling. However, this reactivity in ice has to be considered in long-term cirrus cloud chemistry modelling (Pruppacher and Klett, 1997).

A very recent modelling study of interactions between chemistry and mixed-phase cloud microphysics (Long et al., 2010) confirms the results obtained by Yin et al. (2002): the main process to be considered in the evolution of chemical species concentrations in mixed-phase clouds is the retention of soluble gases when liquid hydrometeors freeze/rime. This is why the process of gas trapping in growing ice hydrometeors is not considered in the present version. The mixed-phase cloud microphysics scheme

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available in Meso-NH, ICE3, considers three ice hydrometeor categories: pristine ice, newly formed crystals, snow and graupel, defined by an increasing degree of riming. The pristine ice category is not concerned by the riming processes. This means that the amount of liquid water transferred in this category by microphysical processes is very small, so we consider no source of chemical species in the cloud chemistry module for this category of ice.

In order to limit the number of prognostic variables, the concentrations of chemical species in snow and graupel are not differentiated but treated as a single set of species X_{ice} :

$$X_{ice} = X_{snow} + X_{grau} \quad (10)$$

X_{ice} , X_{snow} and X_{grau} are the global concentration of species X in total ice, the concentration of species X in snow and the concentration of species X in graupel, respectively. X_{ice} is the prognostic variable with a continuity equation analogous to Eqs. (1) and (2).

The additional microphysical transfer terms due to the retention of soluble gases during freezing/riming are computed following:

$$\left. \frac{\partial X_g}{\partial t} \right|_{FREEZ} = (1 - RET) X_w \frac{1}{r_w} \left. \frac{\partial r_w}{\partial t} \right|_{FREEZ} \quad (11)$$

$$\left. \frac{\partial X_w}{\partial t} \right|_{FREEZ} = -X_w \frac{1}{r_w} \left. \frac{\partial r_w}{\partial t} \right|_{FREEZ} \quad (12)$$

$$\begin{aligned} \left. \frac{\partial X_{ice}}{\partial t} \right|_{FREEZ} &= (RET) X_w \frac{1}{r_w} \left. \frac{\partial r_{ice}}{\partial t} \right|_{FREEZ} \\ &= (RET) X_w \frac{1}{r_w} \left(\left. \frac{\partial r_{snow}}{\partial t} \right|_{FREEZ} + \left. \frac{\partial r_{grau}}{\partial t} \right|_{FREEZ} \right) \end{aligned} \quad (13)$$

FREEZ refers to freezing and riming processes. RET is the retention coefficient ($0 \leq RET \leq 1$). $RET = 0$ means that the soluble gas is completely released to the gas phase

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and is not retained in the ice phase at all. $RET = 1$ means that the soluble gas is fully incorporated into the ice phase. For the chemical species present in the aqueous phase only as ionic species (intermediate ions involved in the oxidation scheme of sulphur dioxide), Eqs. (12) and (13) are applied with $RET = 1$.

5 Table 4 gives the values of retention coefficients for the soluble gases considered in the ReLACS-AQ scheme. Very few experimental studies exist on the phase partitioning during liquid-to-solid freezing and riming (Iribarne and Pyshnov, 1990a,b; Lamb and Blumenstein, 1987; Snider et al., 1992, 1998; Voisin et al., 2000; von Blohn et al., 2011). A recent theoretical study on chemical retention during dry growth riming (Stuart and Jacobson, 2004) shows that the effective Henry's law constant is a particularly important forcing factor. According to this work, chemical species with very high effective Henry's law constants (e.g. strong acids) are likely to be fully retained in the ice hydrometeor under all conditions. Highly soluble gases such as strong acids are almost completely dissociated in water so ions are hardly able to leave the liquid phase 15 (von Blohn et al., 2011). This is consistent with the available experimental data on nitric acid (Iribarne and Pyshnov, 1990a; von Blohn et al., 2011), which show a retention coefficient of 1. For chemical species with lower effective Henry's law constants (e.g. SO_2 and H_2O_2), the pH, temperature, drop size, and air speed around the hydrometeor become important factors in the retention fraction (Stuart and Jacobson, 2004). 20 Following the conclusion of this theoretical approach and using the experimental data available, we chose to estimate the retention coefficient of chemical species using data for SO_2 and H_2O_2 and according to the value of the effective Henry's law constant. For the strong acids, a value of 1 was taken as recommended. Additionally, some data for SO_2 and NH_3 , deduced from in-situ measurements, were available (Voisin et al., 2000). 25 Finally, it was assumed that the retention coefficient of a slightly soluble gas was zero. Thus these species were not present in the ice phase.

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An important microphysical transfer term of ice phase chemical species was added for the sedimentation of graupel and snow, which contributes to wet deposition:

$$\begin{aligned} \left. \frac{\partial X_{\text{ice}}}{\partial t} \right|_{\text{SEDI}} &= -X_{\text{ice}} \left(\left. \frac{1}{r_{\text{snow}}} \frac{\partial r_{\text{snow}}}{\partial t} \right|_{\text{SEDI}} + \left. \frac{1}{r_{\text{grau}}} \frac{\partial r_{\text{grau}}}{\partial t} \right|_{\text{SEDI}} \right) \\ &= X_{\text{ice}} \left(\frac{1}{r_{\text{snow}}} \frac{\partial}{\partial z} F_{\text{SEDI}}(r_{\text{snow}}) + \frac{1}{r_{\text{grau}}} \frac{\partial}{\partial z} F_{\text{SEDI}}(r_{\text{grau}}) \right) \end{aligned} \quad (14)$$

As shown in Table 4, 14 + 5 prognostic variables were added to the cloud chemistry module for soluble chemical species and for intermediate ions in ice-precipitating hydrometeors, respectively.

3 Application to idealized test cases

Three cases were run to assess the new module in Meso-NH. For all cases, the highly accurate PPM (Piecewise Parabolic Method) scheme was used for the transport of the meteorological and scalar fields. The first case, an idealized, warm, two-dimensional precipitating case, was used to study the sensitivity of the cloud chemistry to available warm cloud microphysical schemes. The second case, an idealized, mixed-phase, two-dimensional squall line, allowed us to assess the sensitivity of the cloud chemistry to the ice phase via the retention of chemical species when riming or freezing occurred. The last one, concerning mixed-phase, three-dimensional supercells initialized with warm bubbles, was used to compare results with other CRMs, in particular the WRF-chem model (Barth et al., 2007b).

3.1 The HaRP case: a 2-D warm shallow convection case

The “HaRP” test case was an idealized 2-D kinematic simulation from “The Hawaiian Rainband Project” that took place in 1990 (Szumowski et al., 1998). It aimed at simulating a highly precipitating cell forced by an idealized, time-varying, non-divergent

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circulation. The duration of the simulation was 3000 s corresponding to the mean life cycle of the HaRP precipitating cell, which was produced by a narrow (1 km) wave with a slightly tilting updraught peaking at 8 m s^{-1} after 1500 s. Only advection, cloud microphysics and cloud chemistry were considered in the following simulations. The computational domain extended over 180×60 grid points with a spacing of 50 m in the x and z directions (3000 m in z and 9000 m in x). The concentrations of the chemical species were continuously replenished through the open lateral boundaries in the lowest 250 m, where inflow conditions occurred. Two simulations were performed to test the sensitivity of the cloud chemistry module to the cloud microphysics scheme. These simulations have already been performed and described in Cohard and Pinty (2000), who studied the implementation of the C2R2 cloud microphysics scheme in Meso-NH; no chemistry was run in their study. The first case used the classical Kessler scheme, which is a one-moment scheme. The second one used the C2R2 scheme, which includes a realistic parameterization of droplet nucleation described in Cohard et al. (1998). This parameterization is based on a four-parameter CCN activation spectrum taking account of the physicochemical properties of the accumulation mode in a natural aerosol population.

Numerous differences were observed between the Kessler and C2R2 simulations for the HaRP precipitating cell. As mentioned in Cohard and Pinty (2000), after 1200 s, the rain began to reach the ground for the C2R2 case, unlike the Kessler case. At 1500 s, heavy precipitation at sea level was observed for the C2R2 case while the Kessler case produced only a very sharp precipitating band. After 1800 s, the cell began to decay in the C2R2 case while it achieved a maximum development in the Kessler case. Very high radar reflectivity was observed for both cases at the end of the simulation. During the course of the simulation, the rain water content was very high for the Kessler run with values up to $9 \times 10^{-6} \text{ vol vol}^{-1}$ whereas the maximum value was $2.5 \times 10^{-6} \text{ vol vol}^{-1}$ in C2R2 run. This difference was explained by the presence of an unrealistically small raindrop radius with the Kessler scheme (Fig. 1).

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The initial conditions for chemistry are shown in Table 5. These values correspond to a tropical marine atmosphere (Seinfeld and Pandis, 1998; Finlayson-Pitts and Pitts, 2000). Three different types of vertical profiles were allowed. For the “stratospheric vertical profile”, the initial mixing ratio was multiplied by 1 from 0 to 2 km and by 0.5 at 3 km. For the “boundary layer vertical profile”, the initial mixing ratio was multiplied by 1 from 0 to 1 km and by 0.1 from 2 to 3 km. Linear interpolation was used between values given at two levels. Simple DMS chemistry in gas phase following Mari et al. (1999) was added to the ReLACS chemical scheme.

Results are shown as the time evolution of vertical profiles in the centre of the cloud cell, i.e. the fields have been x-averaged over 200 metres in the middle of the domain. For hydrogen peroxide, which is a soluble gas, it is easy to see the effect of wet deposition in the Kessler case (Fig. 2). At the beginning of the simulation, hydrogen peroxide is scavenged by cloud water and then transferred to rain by microphysical conversion. The sedimentation of the raindrops and the simultaneous scavenging by raindrops lead to its efficient wet deposition. Moreover, during the course of the run, hydrogen peroxide was involved in the oxidation of sulphur dioxide leading to sulphuric acid. This was the main chemical sink of sulphur dioxide occurring in the liquid phase. Contrasting behaviour can be observed for the C2R2 case (Fig. 2): the microphysical transfer from cloud water to rainwater is limited due to the two-moment approach in C2R2. Such a scheme is able to produce large raindrops in a reasonable concentration, thus with a moderate rainwater content (Fig. 1). Due to the large size of raindrops, the equilibrium time of the mass transfer between the gas phase and the aqueous phase is very long, leading to inefficient scavenging of hydrogen peroxide by raindrops (cf. Eq. 5). This is why the mixing ratio of hydrogen peroxide in rainwater is limited inside the cloud, where collision/coalescence processes transfer chemical species from cloud water to rainwater. These processes are the main source of hydrogen peroxide inside raindrops for the C2R2 case. For sulphur dioxide, the same disparity can be observed between the Kessler and the C2R2 simulation. Additionally, the form of the initial vertical profile (boundary layer type) is visible (Fig. 3). Also, the mixing ratio of sulphur dioxide

exceeds its initial value (50 pptv, see Table 5) in the low levels due to a production by DMS oxidation in the gas phase. Finally, its scavenging by cloud water is delayed compared to hydrogen peroxide because of its lower solubility. Because the initial mixing ratio of hydrogen peroxide is larger than that of sulphur dioxide (1 ppbv versus 50 pptv, see Table 5), the oxidation of S(IV) into S(VI) is not limited. No sulphuric acid is present initially. Results for the Kessler case show a production of sulphuric acid up to 38 pptv in rainwater (Fig. 4). However, due to limited transfer of chemical species in rainwater for the C2R2 case, the production of sulphuric acid in rainwater is only up to 16 pptv and is limited inside the cloud (Fig. 4). The resulting pH simulated for rainwater (Fig. 5) is near the atmospheric equilibrium value considering only the scavenging of carbon dioxide in water (i.e. 5.6). This value is consistent with a remote marine atmosphere (Seinfeld and Pandis, 1998) and is explained by a low production of sulphuric acid, which is counterbalanced by the scavenging of ammonia (initial concentration 50 pptv, see Table 5). Figure 6 summarizes these results: below the cloud, for the Kessler case, rainwater is able to efficiently scavenge hydrogen peroxide and sulphur dioxide, which is converted into sulphuric acid in rain. For the C2R2 case, the scavenging of gases below the cloud is inefficient and the production of sulphuric acid in rainwater is negligible.

This simple idealized simulation highlights the role of the cloud microphysics scheme used in cloud chemistry. In particular, using a two-moment scheme for cloud microphysics allows a better prediction of raindrop number concentration (Fig. 1), leading to a realistic mean raindrop diameter, which greatly changes the assessment of strong acid wet deposition, as the temporal evolution of acid wet deposition rates is very sensitive to the values of raindrop radius. Another important point is the below-cloud aerosol particle scavenging by rain, which is another significant source of strong acid wet deposition. This source is not considered here and should contribute mainly to the acidity of precipitation in a marine atmosphere, where aerosol particles are in the coarse mode, which is the mode more efficiently scavenged by falling raindrops (Berthet et al., 2010.)

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3.2 The COPT case: a 2-D tropical squall line

A tropical squall line is composed of two circulation features: a convective region and a stratiform region. The convective region is characterized by the mesoscale boundary layer convergence feeding deep convective updrafts and mid- to upper-level divergence associated with mass flux outflow from individual cells. The stratiform region is characterized by mid-level convergence feeding both a mesoscale downdraft below the anvil and a mesoscale updraft within the stratiform part. The “COPT” test case (Caniaux et al., 1994) is typically a 12-h simulation of a tropical squall line with kilometre-scale-resolved internal circulations, a 3-D turbulence scheme and mixed-phase microphysics (ICE3 scheme). This case was built up from a tropical squall line observed in West Africa during the Convection Profonde Tropicale (COPT81) campaign on 23 June 1981. The domain contains 320×44 grid points unevenly spaced in the vertical ($z = 70$ m at ground level and $z = 700$ m above 12 km). The horizontal resolution is 1.25 km. The model is integrated with a time step of 10 s. A gravity wave damping layer is inserted between 17 km and the model top at 22.5 km. A constant speed translation is used to compensate for the motion of the squall line. No fluxes are considered in the surface layer. Convection is initiated by forming a -0.01 K.s^{-1} artificial cold pool in the low levels of a small domain for 10 min. Measurements available from the AMMA campaign (Stone et al., 2010) were used to build the initial profiles for chemical species. For unavailable species, the initial profiles were estimated either from simulation by the MOCAGE CTM model (Bousserez et al., 2007; Teysédre et al., 2007) at the location corresponding to the COPT campaign and for June; or from measurements averaged over June and July for the years 1998 to 2007 and taken from the IDAF database (Adon et al., 2010, <http://idaf.sedoo.fr>) at Lamto, Ivory Coast. In order to allow short-lived species with unknown concentrations to form and adjust their mixing ratios to an equilibrium state, a simulation with a box model corresponding to 24 h of spin-up was performed using the set of data composed by AMMA and IDAF measurements as initial conditions, as well as MOCAGE simulation results. The resulting initial mixing ratios in

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the boundary layer for the COPT case are indicated in Table 6. For the “stratospheric vertical profile”, the initial mixing ratio was multiplied by 1 from 0 to 2 km, by 0.5 from 3 to 13 km, by 0.75 at 14 km and by 1 at 15 km. For the “boundary layer vertical profile”, the initial mixing ratio was multiplied by 1 from 0 to 1 km, by 0.1 from 2 to 13 km and by 0.05 from 14 km to 15 km. Linear interpolation was used between values given at two levels.

Three simulations were performed. The first one (GAS) considered only gas phase chemistry; the cloud chemistry module was not activated. The second one (AQ-NOICE) neglected the retention of soluble chemical species in the ice phase, assuming that mixing ratios of soluble species in the liquid phase were fully transferred to the gas phase when freezing or riming occurred, which corresponds to $RET = 0$ for all species. The third one (AQ-ICE) considered the retention of soluble chemical species in the precipitating ice hydrometeors, i.e. prognostic scalar variables were considered for the mixing ratio of soluble species in ice. Results are shown as averages obtained between 7 and 8 h of simulation and corresponding to the mature stage of the squall line (Carniaux et al., 1994), which was characterized by a convective zone 40 km wide giving large precipitation, a well developed stratiform zone stretching over 150 km and giving moderate precipitation over an area 80 km wide, and a forward anvil at the tropical easterly jet level near 12 km.

In this section, we focus on three chemical species with increasing solubility and retention coefficient in the ice phase: formaldehyde HCHO, formic acid HCOOH and sulphuric acid H₂SO₄.

The budget of formaldehyde in the upper troposphere (UT) is still uncertain although HCHO is a potentially a significant source of HO_x via its photolysis in the UT (Cohan et al., 1999). HCHO observed in the UT is due to direct transfer of boundary layer HCHO and chemical secondary production by transported VOCs. Fried et al. (2008) and Stickler et al. (2006) showed an HCHO enhancement in the convective outflow, for summertime deep convection over North America and the North Atlantic, and in the middle and upper troposphere over Europe, respectively. Borbon et al. (2012) observed

a moderate enhancement of HCHO in the convective outflow for a mesoscale convective system over West Africa. Fried et al. (2008) and Borbon et al. (2012) estimated that 70 % and 60 %, respectively, of the HCHO observations in the UT after convection were related to an enhanced production from precursors rather than an upward transport from the boundary layer.

The initial mixing ratio of HCHO in the boundary layer was 500 pptv (Table 6) for the mature stage of the squall line. This mixing ratio was enhanced due to secondary production of HCHO by the oxidation of VOC (Volatile Organic Compounds) in the front of the squall line whereas it was reduced behind the passage of the convective front (Fig. 7a–c). This reduction was also observed for CO (not shown), which is a good tracer for convective transport, and was explained by the rear-to-front flow in the low levels of the stratiform area of the squall line. The production of secondary HCHO in the front of the squall line reached about 100 pptv and came mainly from the photo-oxidation of alkanes (ETH and ALK ReLACS species) and of methyl hydroperoxide (OP1 ReLACS species) in the gas phase. Indeed, the maximum of the total mixing ratio of HCHO for AQ-NOICE and AQ-ICE simulations (Fig. 7b, c) and of the gas phase mixing ratio of HCHO for the GAS simulation (Fig. 7a) was the same, about 600 pptv. A comparison between Fig. 7a and b shows that the liquid cloud chemistry reduced the vertical transport of HCHO by the convective updraft in the UT, due to the scavenging of HCHO by the cloud droplets and the raindrops. In the GAS simulation, a mixing ratio of HCHO up to 350 pptv was observed in the forward anvil and in the stratiform part of the cloud while it fell to less than 150 pptv for the AQ-NOICE case. The comparison between the two simulations including cloud chemistry (Fig. 7b, c) is interesting: for AQ-ICE case (Table 4, RET = 0.64 for HCHO), values of HCHO in the anvil and in the stratiform part of the cloud were very low, less than 100 pptv with some areas having values less than 50 pptv. The mixing ratio of HCHO in the ice phase during the mature stage of the cloud was less than 30 pptv (Fig. 7d) so, alone, it cannot explain the difference between Fig. 7b and c. Along the squall line development, the HCHO in the ice phase was partly eliminated by rain due to the melting of snow and graupel during

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their sedimentation. This was the main reason for the depletion of HCHO in the gas phase in the UT observed in the AQ-ICE simulation. Non-soluble precursors of HCHO, such as ETH and ALK (ReLACS species), were transported in a similar way to CO in the forward anvil, the stratiform part of the cloud and the convective outflow. However, their contribution to the HCHO mixing ratio in the UT was very weak, as shown by the comparison between Fig. 7c and a. No signature of a secondary production of HCHO in the UT was observed for the AQ-ICE case. In conclusion, the effect of the retention of HCHO in ice is to enhance its scavenging by the tropical squall line but it leads to a negligible perturbation of HCHO mixing ratio in the convective outflow. The COPT81 squall line was a very intense convective system with heavy precipitation. The peak of the simulated precipitation rate located below the convective part of the system was about 90 mm h^{-1} . Similarly to previous model studies on the transport of soluble species by deep convection using CRM models (Barth et al., 2007a, b; Marécal et al., 2006), results obtained for the AQ-ICE case show a depletion of HCHO in the convective outflow in comparison with unperturbed UT. This result is consistent with the hypothesis made by previous works based on aircraft observations (Stickler et al., 2006; Fried et al., 2008; Borbon et al., 2012) that direct transport of formaldehyde from the boundary layer by convective updrafts is unlikely. These authors explain the observed HCHO increase in the UT by secondary production from vented precursors. In the COPT81 simulation however, only 50 pptv of HCHO were produced by transported VOC in the convective outflow, which is not sufficient to compensate for the depletion due to the scavenging by mixed phase processes (Fig. 7c).

Formaldehyde is the major precursor of formic acid in the aqueous phase, which is a good tracer of cloud-processed air since its sources in the gas phase are weak (Barth et al., 2007b). Formic acid is a secondary species, both in gas and aqueous phases. In the gas phase, formic acid is mainly produced in the ReLACS mechanism by the ozonolysis of alkenes (ALKE species in ReLACS) and of biogenic VOC (BIO species in ReLACS). In the aqueous phase, it is produced by the oxidation of hydrated formaldehyde by OH radical (Table 3). Its main destruction pathway is oxidation by OH in either

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the gas or aqueous phase. No formic acid is present initially. As for formaldehyde, a gas phase production is observed in the front of the squall line (Fig. 8a–c). However, in contrast to HCHO production, the maximum of HCOOH mixing ratio for the GAS simulation is about 10.5 pptv whereas it is about 11.6 pptv for the AQ-NOICE and AQ-ICE cases. This difference is due to a weak contribution of aqueous phase chemistry to the gas phase mixing ratio in the front of the squall line. Although a greater production of formic acid via aqueous phase chemistry was expected (the initial mixing ratio of its main precursor in aqueous phase is 500 pptv in the boundary layer), results actually show a very weak contribution of cloud reactivity in the production of formic acid. This is partly explained by the destruction of formate ion by OH in the aqueous phase, which is more efficient than the oxidation of formic acid by OH (Table 3). This pathway dominates due to high pH values in cloud water and rainwater ($\text{pK}_a(\text{formic acid}) = 3.75$ at 298 K; Table 2). Values of pH were between 6 and 6.5 (not shown) due to a very high mixing ratio of ammonia (almost 3 ppbv in the boundary layer, see Table 6) and low mixing ratio of sulphur dioxide. The effect of the retention of HCOOH in ice is the same as simulated for HCHO with a depletion of formic acid in the UT (Fig. 8c), which cannot be directly explained by its amount in ice precipitating hydrometeors (Fig. 8d).

Sulphuric acid is one of the precursors for the nucleation of new aerosol particles in the UT (e.g. Waddicor et al., 2012). Sulphuric acid is a strong acid with a very high Henry law constant (Table 1). It is completely retained in the ice phase when freezing or riming occurs, with a retention coefficient equal to 1 (Table 4). No sulphuric acid is present initially. Figure 9a shows a very weak production of sulphuric acid of less than 1.5 pptv in the gas phase by oxidation of sulphur dioxide by OH radical. When liquid phase chemistry is activated (Fig. 9b AQ-NOICE case), the total mixing ratio of sulphuric acid shows two maxima: one located at the base of the squall line, with a maximum value of about 280 pptv, and another located in the forward anvil of the squall line with a maximum value of about 8 pptv. The first maximum is due to the high mixing ratio of sulphuric acid inside cloud water and rainwater, which is produced by the oxidation of sulphur dioxide by ozone and hydrogen peroxide in the aqueous phase.

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A similar maximum is observed for the AQ-ICE simulation (Fig. 9c). The upper level maximum is in the gas phase and can only be explained by the degassing of super-cooled droplets when riming or freezing occurs throughout the AQ-NOICE simulation. For the AQ-ICE case, the mixing ratio in the ice phase at the mature stage of the cloud is about 2 pptv (Fig. 9d) and cannot explain the high maximum located in the forward anvil simulated for the AQ-NOICE case. Only complex interactions between sulphuric acid degassing from supercooled droplets and its successive transport throughout the simulation can explain this result: the released gaseous sulphuric acid is entrained by the convective updraft inside the forward anvil while the convective cell evolves in a squall line. As already pointed out for formic acid, the AQ-ICE simulation shows a very weak mixing ratio of soluble species (here sulphuric acid) inside the forward anvil and the stratiform part of the squall line (Fig. 9d) due to efficient trapping of soluble species inside the ice-precipitating hydrometeors, then contributing to wet deposition.

For this particular simulation, the results show that the effect of the retention of soluble species in the ice phase is to enhance the scavenging efficiency of soluble species and thus to increase the wet deposition of the soluble species. This is consistent with the very high accumulated precipitation due to melting of snow and graupel in the convective part of the cloud.

3.3 The STERAO case: a 3-D continental storm

The STERAO (Stratospheric-Tropospheric Experiment: Radiation, Aerosol, and Ozone) convective system observed on 10 July 1996 took place near the southern border of Wyoming and Nebraska (see Dye et al., 2000 for an overview of the experiment). The storm developed during the late afternoon, the main cells propagated south-south-eastward into north-eastern Colorado before dissipating in the evening. Radar reflectivity observations showed an evolution of the storm from a multicell stage followed by a quasi supercellular one (Dye et al., 2000). The simulation of this storm performed with the Meso-NH model was a 3-D idealized run based upon Skamarock et al. (2000) with some modifications. The convection was initiated with three warm

not considered. For the interaction with the ice phase of the cloud, two simulations were performed in Barth et al. (2007b), either dissolved species were completely retained in the ice phase when cloud droplets or raindrops froze ($RET = 1$) or all dissolved species were degassed during freezing ($RET = 0$).

As shown in Barth et al. (2007b), CO is transported from the boundary layer to the anvil of the cloud where mixing ratios are > 110 ppbv, greater than the unperturbed UT at the same level (Fig. 10a, b). As in Barth et al. (2007b) the comparison of CO and ozone with measurements across the anvil were in good agreement (not shown). Results are now discussed for HCHO and H_2O_2 , two soluble and reactive gases. For HCHO, the results are shown at 1 h, corresponding to the multicell stage of the cloud.

Despite its solubility and its reactivity in the aqueous phase, HCHO is efficiently transported from the boundary layer to the anvil (Fig. 10c, d) during the multicell stage of the cloud. However, the total mixing ratio of HCHO in the anvil is greater for the simulation considering retention in ice (Fig. 10c) than in the one with no retention (Fig. 10d). HCHO and CO have the same initial vertical profile but with different magnitudes. It is thus interesting to compare the redistribution by the cloud of HCHO, a soluble and reactive species, and that of CO, an insoluble passive species. Consistently with results for the total mixing ratio of HCHO, the ratio of total HCHO to CO shows a transport of HCHO from the boundary layer to the anvil which is efficient in the simulation with retention in ice. The ratio of total HCHO to CO also reveals that vertical transport inside the cloud is less efficient for HCHO than for CO, indicating that a fraction of HCHO has reacted or precipitated to the ground. This effect described in Barth et al. (2007b) is similar. Looking at the gas phase mixing ratio of HCHO (Fig. 11a, b), the simulated gas phase mixing ratio of HCHO in the anvil is low in the simulation with retention in ice whereas an efficient transport of HCHO in the anvil is observed for the simulation without retention. As for the COPT81 case, the result from the simulation with retention in ice shows a depletion of HCHO in the convective outflow in comparison with unperturbed UT, in agreement with Barth et al. (2007b). The mixing ratios of HCHO in cloud water and in rainwater (Fig. 11c–f) for both simulations are very close, with a difference

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at the top of the cloud water zone due to less HCHO being available in the gas phase at this altitude in the simulation with retention. In comparison to Barth et al. (2007b), the same order of magnitude is retrieved for the HCHO mixing ratio in cloud water but lower values are observed in rainwater. In Barth et al. (2007b), the mixing ratios of soluble species are explicitly represented in each of the ice phase hydrometeor categories. However, the comparison of the sum of HCHO mixing ratios in snow and in hail categories from Barth et al. (2007b) with the HCHO mixing ratio in precipitating ice from our simulation (Fig. 11g), shows the same order of magnitude. Thus, the global approach for the ice phase species mixing ratio representation used in Meso-NH seems to be comparable to a more explicit approach. The main general remarks concern results of HCHO mixing ratio during the supercell stage of the cloud (after 2.5 h of simulation). In particular, the depletion of gas phase mixing ratio of HCHO in the convective outflow is again observed for the simulation with retention in ice, whereas the simulation without retention shows enrichment in the anvil (not shown). This difference is due to the capture of HCHO in the ice phase. The greater value of total HCHO mixing ratio observed for the simulation with retention in ice whatever the stage of the storm in comparison to simulation without retention indicates the formation of a reservoir of HCHO in the ice phase in the simulation with retention in ice, leading to less HCHO remaining available to react in gas and aqueous phases.

The same effect is observed for hydrogen peroxide (Figs. 12a, b and 13e). Moreover, as hydrogen peroxide is more soluble than HCHO in liquid water and is very reactive in aqueous phase with sulphur dioxide, its vertical transport from the boundary layer to the anvil is less efficient than for HCHO. A significant depletion is observed in the convective cores for hydrogen peroxide in gas phase for the simulation with retention in ice (Fig. 12c) underlying its capture in the ice phase (Fig. 13e). This depletion is also observed for the simulation without retention in ice (Fig. 12d), whereas this effect is not observed for HCHO (Fig. 11b). This difference stems from the very efficient transfer of hydrogen peroxide from the gas phase to the liquid phase due to its high reactivity and its high solubility in the aqueous phase.

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As in Barth et al. (2007b), integrated results are shown as percentages of HCHO and H_2O_2 in each water chemical reservoir relative to the total amount of the species in the condensed phase (Fig. 14). The total amount of each of the hydrometeor reservoirs relative to the total amount of condensed water is also indicated (Fig. 14a). The transition between the multicell and the supercell stages of the storm is marked by the minimum of the percentage of cloud water between 90 and 120 min. The percentage of the pristine ice reservoir is of the same order of magnitude as the graupel reservoir, whereas the proportion of ice is very small in Barth et al. (2007b), less than 5%. Moreover, the amount of cloud water is greater than that of rainwater, in contrast to Barth et al.'s (2007b) findings. These differences come from the microphysical schemes used in the two models with different parameterizations of microphysical processes and different ice phase categories. For instance, Barth et al. (2007b) used pristine ice, snow and hail as ice phase categories whereas, in Meso-NH, the ICE3 scheme uses pristine ice, snow and graupel. Results for the vertical cross section across the anvil of ice particle concentration shown in Barth et al. (2007a) reflect these differences in the cloud microphysics schemes of the models participating in the intercomparison exercise, including WRF-Chem and Meso-NH. Looking at the simulation without retention in ice (Fig. 14d, e), the distributions of HCHO and H_2O_2 between cloud water and rainwater are seen to be very similar, in agreement with Barth et al. (2007b). For the simulation with retention in ice, the percentages of HCHO and H_2O_2 in rainwater are close. The percentage of HCHO in ice-precipitating hydrometeors is of the same order of magnitude as in cloud water (Fig. 14b), whereas the percentage of H_2O_2 in ice-precipitating hydrometeors is less than half those in cloud water (Fig. 14c). This difference comes from the reactivity of the hydrogen peroxide in aqueous phase leading to its destruction in cloud water and, thus, limiting its transfer to the ice phase in comparison to HCHO.

This 3-D idealized simulation already run by several models shows consistency of the results obtained by Meso-NH with similar features observed by Barth et al. (2007b) for instance. Moreover, this particular simulation underlines the potential role of ice

phase as a reservoir of soluble species that are unavailable to react in either gas or aqueous phases.

4 Conclusions

A cloud chemistry module has been developed in the mesoscale atmospheric model Meso-NH. This module is a complete suite including aqueous phase reactivity, exchange of chemical species between the gas phase and the condensed phase (liquid and ice), a diagnostic estimation of pH in cloud water and rainwater, and the redistribution of chemical species by mixed phase cloud microphysical processes. The aqueous phase reactivity was developed for the gas phase using the chemical mechanism ReLACS, which is a reduced form of the RACM mechanism. The exchange of chemical species between gas phase and aqueous phase has been considered using the theory of mass transfer kinetics including the limitation of the solubility inside droplets by gas phase diffusion and accommodation at the air/water interface. The redistribution of chemical species by cloud microphysical processes has been developed for both one-moment and two-moment cloud microphysical schemes. Cloud microphysical processes include the autoconversion, the accretion and the sedimentation of all precipitating hydrometeors (rain, snow, graupel), and the freezing and the riming of supercooled drops. For these last processes, retention coefficients have been estimated to compute the amounts of soluble species retained in the ice and released in the gas phase. If the effect of an ice phase on chemical budgets is taken into account, prognostic scalar variables are added for the global mixing ratio of soluble species in ice-precipitating hydrometeors to save computing time. It is possible to run a simulation with mixed phase microphysics and without these additional prognostic variables, in which soluble chemical species are completely released to the gas phase when freezing or riming occurs.

The new cloud chemistry module has been tested on three cases. The first one was a 2-D simulation of a warm cumulus cloud formed over Hawaii Island. This case was

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used in particular to focus on the impact of a 1-moment cloud microphysics scheme vs. a 2-moment cloud microphysics scheme. Results show significant differences between the two microphysical approaches in terms of mixing ratios of chemical species. In particular, the 2-moment approach led to smaller raindrops, thus limiting sulphuric acid formation inside the rain and leading to less acidic raindrops than for the 1-moment scheme.

The second case was a 2-D simulation of a tropical squall line observed in West Africa. When no retention in ice was considered, this case underlined the complex interactions between the degassing of soluble species from supercooled droplets and the transport of chemical species throughout the simulation. This led to a release of soluble species, which were entrained by the convective updraft inside the anvil while the convective cell evolved in a squall line. When the retention in ice was taken into account, very low mixing ratios of soluble species inside the forward anvil and the stratiform part of the squall line were observed due to efficient trapping of soluble species inside the ice-precipitating hydrometeors, which then contributed to wet deposition.

The last test case was a 3-D idealized simulation of a storm observed in Colorado. First, the comparison of our results with those of Barth et al. (2007b), who used an explicit representation of the mixing ratio of soluble species in each ice phase hydrometeor category, shows that the global approach for the ice phase species mixing ratio representation used in Meso-NH is comparable to a more explicit approach. The comparison of simulations considering or neglecting the retention of soluble species in ice shows a depletion of the gas phase mixing ratio of soluble species in the convective outflow for the simulation with retention in ice, whereas the simulation without retention shows enrichment in the anvil. This difference is due to the capture of soluble species in the ice phase. Also, a greater value of the total mixing ratio of the soluble species observed in the simulation with retention in ice, in comparison to simulation without retention, indicates a reservoir of the soluble species in the ice phase leading to lower concentration of soluble species available to react in gas and aqueous phases. This effect is the inverse of the one observed for the COPT test case, showing inefficient

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wet deposition of soluble species due to the transfer of soluble species in rain from the ice-precipitating hydrometeors.

Results for COPT and STERAO cases on the amount of formaldehyde in the convective outflow show a depletion in comparison to the unperturbed UT. This result is consistent with previous modelling studies (Barth et al., 2007b, Marécal et al., 2006) and agrees with the hypothesis of an inefficient direct convective transport of HCHO to the upper troposphere (Stickler et al., 2006; Fried et al., 2008; Borbon et al., 2012). Unlike the observations reported in these works, however, the model does not show an increase of HCHO in the upper troposphere due to secondary production by vented VOC precursors. Moreover, the aqueous phase chemistry mechanism used in this study includes simple organic chemistry, and thus overestimates the concentration of the OH radical in the aqueous phase (Herrmann et al., 2005), leading to an overestimation of the oxidation of HCHO in cloud droplets and raindrops. In order to try to understand the discrepancy between models and measurements for the amount of HCHO in the convective outflow, additional studies are needed based on well documented case studies (e.g. the AMMA experiment described in Borbon et al., 2012).

An interesting extension of the cloud chemistry module of Meso-NH would consider coupling the present scheme with the explicit electrical scheme of Meso-NH (Barthe et al., 2012) in order to include the sporadic source of NO produced by the simulation of lightning flashes in deep convective storms.

The next step for the cloud chemistry module will be to include more complex organic chemistry in the aqueous phase coupled to the ReLACS2 gas phase mechanism (Tulet et al., 2006) available in Meso-NH. The ReLACS2 mechanism describes the gas phase chemistry of the ozone-NO_x-VOC system including the formation of the gas phase semi-volatile precursors of SOA. The objective is to simulate the formation of the aqueous phase precursors of SOA in droplets, and to study the effect of aqueous phase reactivity on SOA formation in the troposphere.

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Table 1. Values of Henry's law constants at 298 K (H_{298}) and associated temperature dependencies ($\Delta H/R$) and values of mass accommodation coefficients (α).

Chemical species	H_{298} (Matm ⁻¹)	$\Delta H/R$ (K)	Ref.	α	Ref.
O ₃	1.0×10^{-2}	-2830	Sander et al. (2006)	0.05	Sander et al. (2006)
OH	3.9×10^1		Sander et al. (2006)	0.05	Estimated
HO ₂	6.9×10^2		Sander et al. (2006)	0.2	Sander et al. (2006)
H ₂ O ₂	7.73×10^4	-7310	Sander et al. (2006)	0.11	Davidovits et al. (1995)
NO	1.92×10^{-3}	-1790	Sander et al. (2006)	0.0001	Sander et al. (2006)
NO ₂	1.4×10^{-2}		Sander et al. (2006)	0.0015	Sander et al. (2006)
NO ₃	3.8×10^{-2}		Sander et al. (2006)	0.05	Estimated
N ₂ O ₅	2.1×10^0	-3400	Fried et al. (1994)	0.0037	George et al. (1994)
HNO ₃	2.1×10^5	-8700	Schwartz and White (1981)	0.054	Davidovits et al. (1995)
HNO ₂	5.0×10^1	-4900	Becker et al. (1996)	0.05	Bongartz et al. (1994)
HNO ₄	1.2×10^4	-6900	Régimbal and Mozurkewich (1997)	0.05	Estimated
NH ₃	6.02×10^1	-4160	Sander et al. (2006)	0.04	Sander et al. (2006)
SO ₂	1.36×10^0	-2930	Sander et al. (2006)	0.11	Sander et al. (2006)
H ₂ SO ₄	2.1×10^5	-8700	Estimated as HNO ₃	0.07	Davidovits et al. (1995)
CO ₂	3.4×10^{-2}	-2710	Sander et al. (2006)	0.0002	Sander et al. (2006)
CH ₃ O ₂	2.7×10^0	-2030	Estimated ^a	0.05	Estimated
CH ₃ OOH	3.0×10^2	-5280	Sander et al. (2006)	0.007	Sander et al. (2006)
HCHO	3.23×10^3	-7100	Sander et al. (2006) ^b	0.04	Sander et al. (2006)
HCOOH	8.9×10^3	-6100	Sander et al. (2006)	0.012	Davidovits et al. (1995)
CH ₃ COOH	4.1×10^3	-6300	Sander et al. (2006)	0.03	Sander et al. (2006)

^a Estimated from the empirical relation: $H_{\text{ROO}_2} = (H_{\text{ROOH}}H_{\text{HO}_2}/H_{\text{H}_2\text{O}_2})$;

^b effective value.

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Table 2. Aqueous phase equilibria at 298 K (K_{298}) and temperature dependencies associated ($\Delta H/R$).

Equilibrium	K_{298} (M)	$\Delta H/R$ (K)	Ref.
$\text{HO}_2 \leftrightarrow \text{H}^+ + \text{O}_2^-$	1.6×10^{-5}		Bielski et al. (1985)
$\text{HNO}_2 \leftrightarrow \text{H}^+ + \text{NO}_2$	1.6×10^{-3}	1760	Park and Lee (1988)
$\text{HNO}_3 \leftrightarrow \text{H}^+ + \text{NO}_3$	2.2×10^1		Perrin (1982)
$\text{HNO}_4 \leftrightarrow \text{H}^+ + \text{NO}_4$	1.26×10^{-6}		Goldstein and Czapski (1997)
$\text{NH}_3 + \text{H}_2\text{O} \leftrightarrow \text{NH}_4^+ + \text{OH}$	1.7×10^{-5}	4350	Seinfeld and Pandis (1998)
$\text{SO}_2 + \text{H}_2\text{O} \leftrightarrow \text{H}^+ + \text{HSO}_3$	1.3×10^{-2}	-1965	Seinfeld and Pandis (1998)
$\text{HSO}_3 \leftrightarrow \text{H}^+ + \text{SO}_3^{2-}$	6.4×10^{-8}	-1430	Seinfeld and Pandis (1998)
$\text{H}_2\text{SO}_4 \leftrightarrow \text{H}^+ + \text{HSO}_4^-$	1.0×10^3		Seinfeld and Pandis (1998)
$\text{HSO}_4 \leftrightarrow \text{H}^+ + \text{SO}_4^{2-}$	1.0×10^{-2}		Seinfeld and Pandis (1998)
$\text{CO}_2 + \text{H}_2\text{O} \leftrightarrow \text{H}^+ + \text{HCO}_3$	4.3×10^{-7}	920	Seinfeld and Pandis (1998)
$\text{HCO}_3 \leftrightarrow \text{H}^+ + \text{CO}_3^{2-}$	4.7×10^{-11}	1780	Seinfeld and Pandis (1998)
$\text{HCHO} + \text{H}_2\text{O} \leftrightarrow \text{CH}_2(\text{OH})_2$	$2.5 \times 10^{3*}$	-4030	Bell (1966)
$\text{HCOOH} \leftrightarrow \text{H}^+ + \text{HCOO}$	1.8×10^{-4}	150	Serjeant and Dempsey (1979)
$\text{CH}_3\text{COOH} \leftrightarrow \text{H}^+ + \text{CH}_3\text{COO}$	1.74×10^{-5}		Serjeant and Dempsey (1979)

* K dimensionless

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Table 3. Aqueous phase chemical mechanism.

Reaction	k_{298} ($M^{-n+1} s^{-1}$)	E_a/R (K)	Ref.
$H_2O_2 + hv \rightarrow 2OH$	Calculated		Graedel and Weschler,1981; Zellner et al. (1990)
$OH + OH \rightarrow H_2O_2$	3.6×10^9	930	Elliot (1989)
$OH + HO_2 \rightarrow H_2O + O_2$	2.8×10^{10}	0	Elliot and Buxton (1992)
$OH + O_2^- \rightarrow HO^- + O_2$	3.5×10^{10}	720	Elliot and Buxton (1992)
$H_2O_2 + OH \rightarrow H_2O + HO_2$	3.2×10^7	1700	Yu and Barker (2003); Seinfeld and Pandis (1998)
$HO_2 + HO_2 \rightarrow H_2O_2 + O_2$	8.3×10^5	2700	Bielski et al. (1985)
$HO_2 + O_2 + H_2O \rightarrow H_2O_2 + O_2 + OH^-$	9.6×10^7	910	Christensen and Sehested (1988)
$O_3 + O_2^- + H_2O \rightarrow OH + 2O_2 + OH^-$	1.5×10^9	1500	Seinfeld and Pandis (1998)
$OH + HSO_3^- \rightarrow SO_3^- + H_2O$	2.7×10^9		Buxton et al. (1996)
$HNO_2 + OH \rightarrow NO_2 + H_2O$	1.0×10^{10}		Barker et al. (1970)
$NO_2 + HO_2 \rightarrow HNO_4$	1.8×10^9		Logager and Sehested (1993)
$NO_2 + O_2^- \rightarrow NO_4^-$	4.5×10^9		Logager and Sehested (1993)
$HNO_4 \rightarrow HO_2 + NO_2$	2.6×10^{-2}		Goldstein et al. (1998)
$NO_4^- \rightarrow NO_2^- + O_2$	1.1×10^0		Goldstein et al. (1998)
$HNO_4 + HSO_3^- \rightarrow SO_3^{2-} + NO_3^- + 2H^+$	3.3×10^5		Amels et al. (1996)
$NO_3^- + hv + H_2O \rightarrow NO_2 + OH + OH^-$	Calculated		Graedel and Weschler,1981; Zellner et al. (1990)
$N_2O_5 + H_2O \rightarrow 2HNO_3$	1.0×10^{15}		Estimated
$NO_3 + SO_4^{2-} \rightarrow NO_3^- + SO_4^-$	1.0×10^5		Logager et al. (1993)
$NO_3 + HSO_3^- \rightarrow SO_3^- + NO_3^- + H^+$	1.3×10^9	2200	Exner et al. (1992)
$CH_3O_2 + CH_3O_2 \rightarrow 2HCHO + 2HO_2$	1.7×10^9	2200	Herrmann et al. (1999)
$CH_3O_2 + HSO_3^- \rightarrow CH_3OOH + SO_3^-$	5.0×10^5		Herrmann et al. (1999)
$CH_2(OH)_2 + OH + O_2 \rightarrow HCOOH + HO_2 + H_2O$	7.8×10^8	1000	Chin and Wine (1994)
$HCOOH + OH + O_2 \rightarrow CO_2 + HO_2 + H_2O$	1.0×10^8	1000	Chin and Wine (1994)
$HCOO^- + OH + O_2 \rightarrow CO_2 + HO_2 + OH^-$	3.4×10^9	1200	Chin and Wine (1994)
$HSO_3^- + HCHO \rightarrow HOCH_2SO_3^-$	7.9×10^2	2900	Olson and Hoffmann (1989)
$SO_3^{2-} + HCHO + H_2O \rightarrow HOCH_2SO_3^- + OH^-$	2.5×10^7	2450	Olson and Hoffmann (1989)
$HOCH_2SO_3^- \rightarrow HSO_3^- + HCHO$	7.7×10^{-3}	9200	Seinfeld and Pandis (1998)
$HOCH_2SO_3^- + OH^- \rightarrow SO_3^{2-} + CH_2(OH)_2$	3.7×10^3		Seinfeld and Pandis (1998)
$HOCH_2SO_3^- + OH + O_2 \rightarrow HO_2 + HCOOH + HSO_3^-$	3.0×10^8		Herrmann (2003)
$SO_3^- + O_2 \rightarrow SO_5^-$	1.1×10^9		Das (2001)
$SO_5^- + HO_2 \rightarrow HSO_5^- + O_2$	1.7×10^9		Buxton et al. (1996)
$SO_5^- + SO_5^- \rightarrow 2SO_4^- + O_2$	2.2×10^8	2600	Buxton et al. (1996)
$HSO_5^- + HSO_3^- + H^+ \rightarrow 2SO_4^{2-} + 3H^+$	7.1×10^6		Seinfeld and Pandis (1998)
$SO_4^- + H_2O \rightarrow HSO_4^- + OH$	4.6×10^2	1100	Yu et al. (2004)
$HSO_3^- + O_3 \rightarrow HSO_4^- + O_2$	3.7×10^5	5500	Seinfeld and Pandis (1998)
$SO_3^{2-} + O_3 \rightarrow SO_4^{2-} + O_2$	1.5×10^9	5300	Seinfeld and Pandis (1998)
$HSO_3^- + H_2O_2 + H^+ \rightarrow SO_4^{2-} + 2H^+ + H_2O$	9.1×10^7	3600	Maaß et al. (1999)

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Table 4. Retention coefficients.

Chemical species	RET	Ref.
SO ₂	0.02	Voisin et al. (2000)
H ₂ O ₂	0.64	von Blohn et al. (2011)
NH ₃	1	Voisin et al. (2000)
HNO ₃	1	von Blohn et al. (2011)
H ₂ SO ₄	1	Stuart and Jacobson (2004)
O ₃ , NO, NO ₂ , NO ₃ , N ₂ O ₅ , CO ₂	0	Estimated
OH, CH ₃ O ₂ , CH ₃ OOH	0.02	Estimated, same as SO ₂
HO ₂ , HNO ₂ , HNO ₄ , HCHO, HCOOH, CH ₃ COOH	0.64	Estimated, same as H ₂ O ₂

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Table 5. Initial gas mixing ratios for the HaRP case. The form of the vertical profile is indicated in parentheses as: 1 for homogeneous, 2 for stratospheric and 3 for boundary layer profile.

Species	Mixing ratio	Species	Mixing ratio
O ₃	40 ppbv (2)	SO ₂	50 pptv (3)
H ₂ O ₂	1 ppbv (1)	CO	100 ppbv (3)
NO	100 pptv (2)	CH ₄	1.7 ppmv (1)
NO ₂	100 pptv (2)	ETH ^a	845 pptv (3)
NO ₃	50 pptv (1)	ALKA ^b	100 pptv (3)
N ₂ O ₅	100 pptv (1)	ALKE ^c	10 pptv (3)
HNO ₂	10 pptv (1)	BIO ^d	100 pptv (3)
HNO ₃	100 pptv (1)	HCHO	200 pptv (3)
HNO ₄	100 pptv (1)	KET ^e	100 pptv (3)
NH ₃	50 pptv (3)	PAN ^f	400 pptv (3)
DMS	100 pptv (3)	OP1 ^g	1 ppbv (3)

^a ETH = ethane;

^b ALKA = alkanes other than methane and ethane, together with alkynes, alcohols, esters and epoxides;

^c ALKE = anthropogenic alkenes;

^d BIO = biogenic species, i.e. isoprene, α -pinene and d-limonene;

^e KET = acetone and higher saturated ketones;

^f PAN = Peroxyacetal nitrate and higher saturated PANs;

^g OP1 = methyl hydrogen peroxide CH₃OOH.

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Table 6. Initial gas mixing ratios in the COPT case. The form of the vertical profile is indicated in parentheses as: 1 for homogeneous, 2 for stratospheric and 3 for boundary layer profile.

Species	Mixing ratio	Species	Mixing ratio
O ₃	42 ppbv (2)	CO	104 ppbv (3)
H ₂ O ₂	2 ppbv (1)	CH ₄	1.7 ppmv (1)
OH	0.5 pptv (1)	ETH	625 pptv (3)
HO ₂	30 pptv (1)	ALKA	767 pptv (3)
NO	120 pptv (2)	ALKE	122 pptv (3)
NO ₂	550 pptv (2)	BIO	186 pptv (3)
NO ₃	100 pptv (1)	HCHO	500 pptv (3)
N ₂ O ₅	35 pptv (1)	ALD	15 pptv (3)
HNO ₃	450 pptv (1)	KET	957 pptv (3)
NH ₃	2.95 ppbv (3)	PAN	50 pptv (3)
SO ₂	363 pptv (3)	OP1	750 pptv (1)

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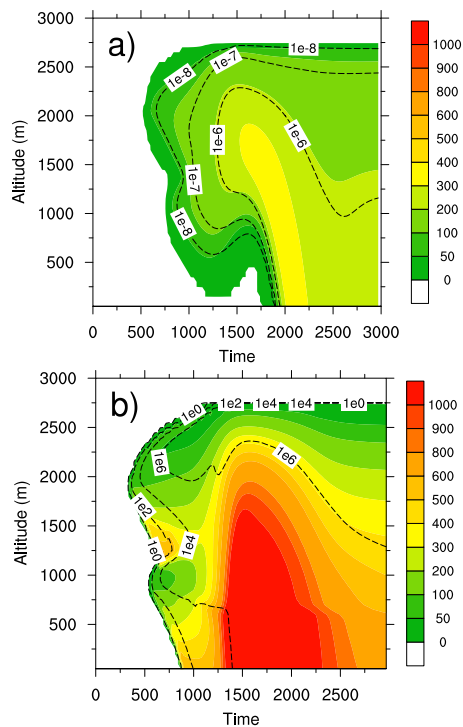


Fig. 1. HaRP case: time evolution of vertical profiles at the centre of the cloud cell for mean raindrop radius (μm): **(a)** using the Kessler scheme and **(b)** using the C2R2 scheme. For the Kessler scheme **(a)**, the rain liquid water content (vol vol^{-1}) is superimposed as isolines. For the C2R2 scheme **(b)**, the number concentration of raindrops (L^{-1}) is superimposed as isolines.

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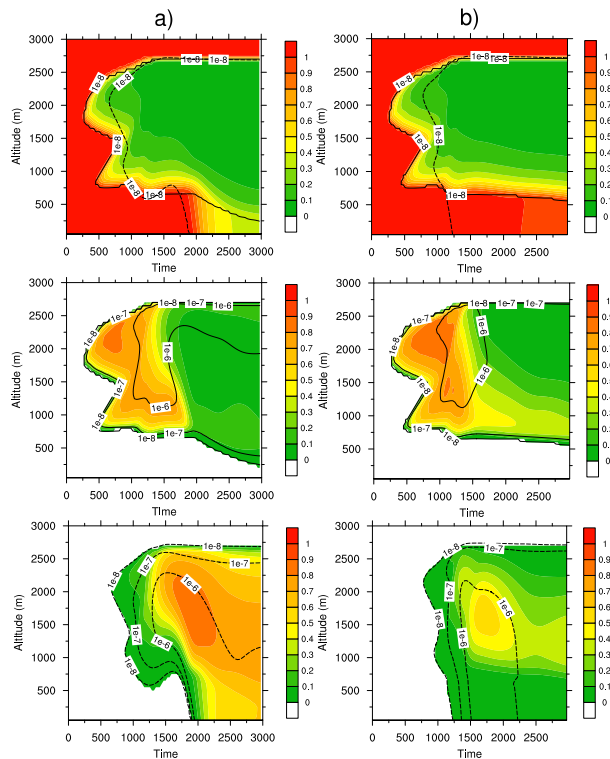


Fig. 2. HaRP case: time evolution of vertical profiles at the centre of the cloud cell for hydrogen peroxide mixing ratio (ppbv) in gas phase, in cloud water and in rainwater: **(a)** using the Kessler scheme and **(b)** using the C2R2 scheme. Isoline of cloud liquid water content (vol vol^{-1} , solid line) and of rain liquid water content (vol vol^{-1} , dashed line) are superimposed.

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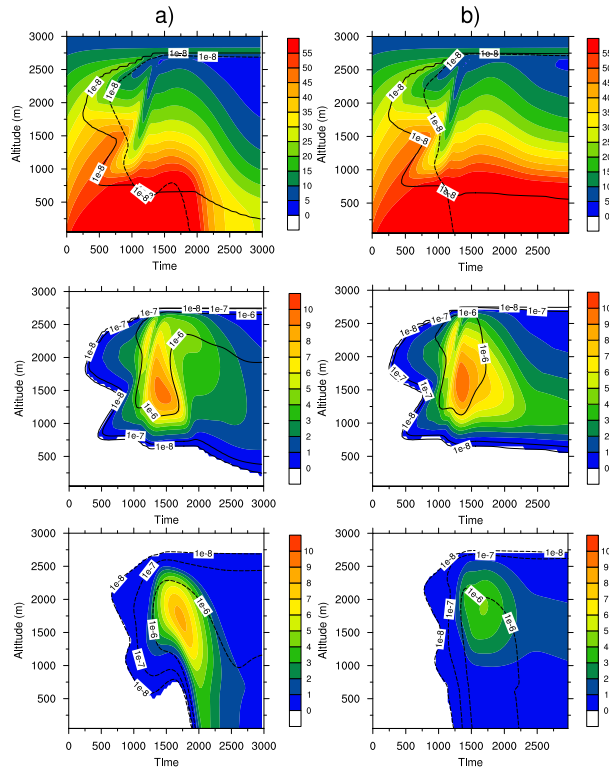


Fig. 3. HaRP case: same as Fig. 2 but for sulphur dioxide (pptv).

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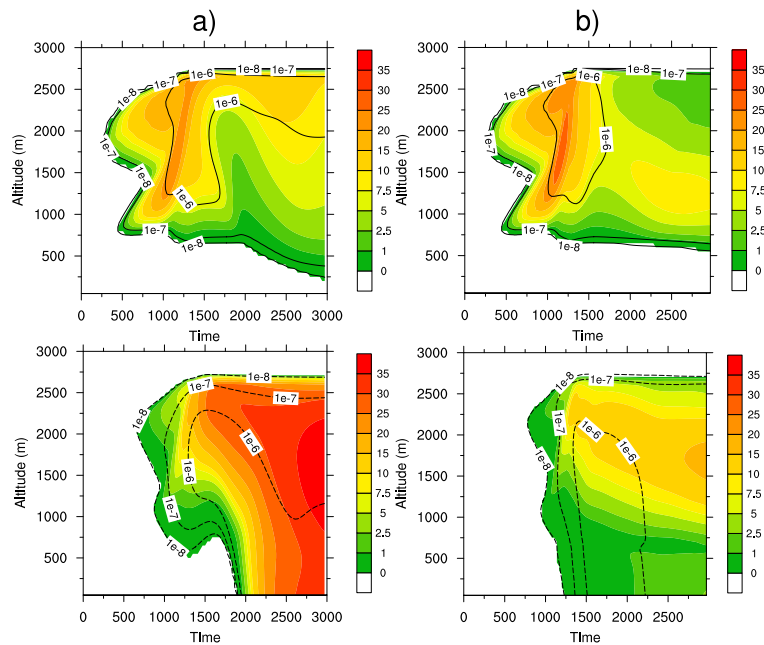


Fig. 4. HaRP case: same as Fig. 2 but for sulphuric acid in cloud water and in rainwater (pptv).

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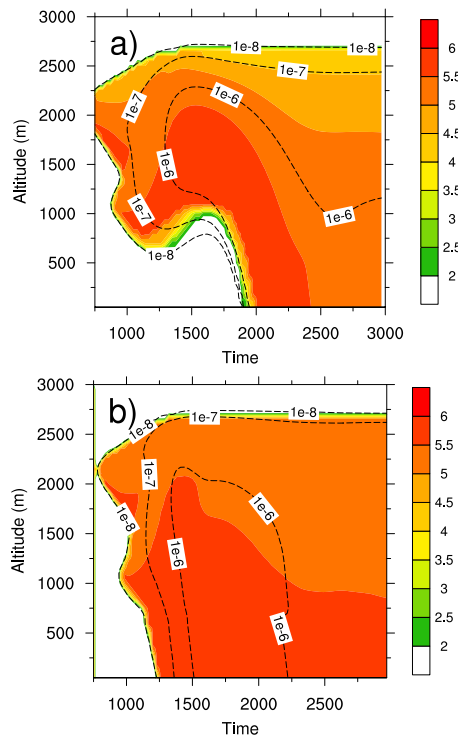


Fig. 5. HaRP case: time evolution of vertical profiles at the centre of the cloud cell for pH value in rainwater: **(a)** using the Kessler scheme, and **(b)** using the C2R2 scheme. The rain liquid water content (vol vol^{-1}) is superimposed as isolines.

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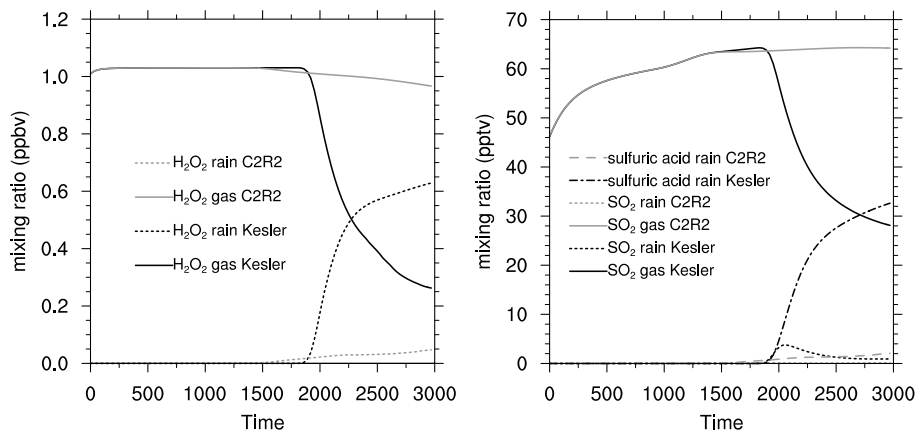


Fig. 6. HaRP case: time evolution of the mixing ratio below the cloud (250 m a.s.l.) for hydrogen peroxide in gas phase and in rainwater (left), for sulphur dioxide in gas phase and in rainwater, and for sulphuric acid in rainwater (right) for both cases: Kessler (black lines) and C2R2 (grey lines).

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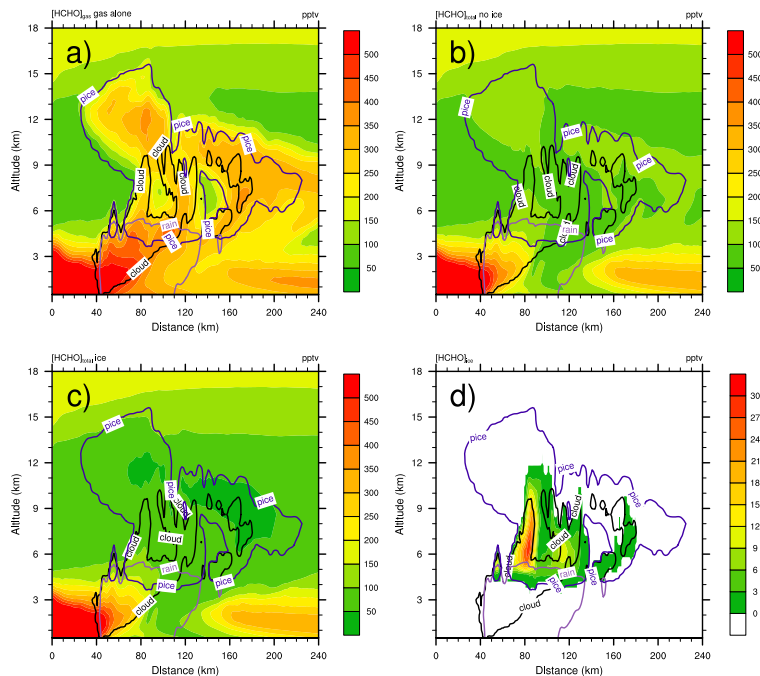


Fig. 7. COPT case: simulated mixing ratio of formaldehyde during the mature stage of the squall line: **(a)** simulated gas phase mixing ratio of HCHO for the GAS simulation, **(b)** simulated total mixing ratio of HCHO (gas + cloud water + rainwater) for the AQ-NOICE simulation, **(c)** simulated total mixing ratio of HCHO (gas + cloud water + rainwater + precipitating ice) for the AQ-ICE simulation, and **(d)** simulated mixing ratio of HCHO in precipitating ice for the AQ-ICE simulation. The black line is the 0.01 g kg^{-1} isoline cloud water mixing ratio, the mauve line is that for precipitating ice (snow + graupel) and the purple is that for rainwater.

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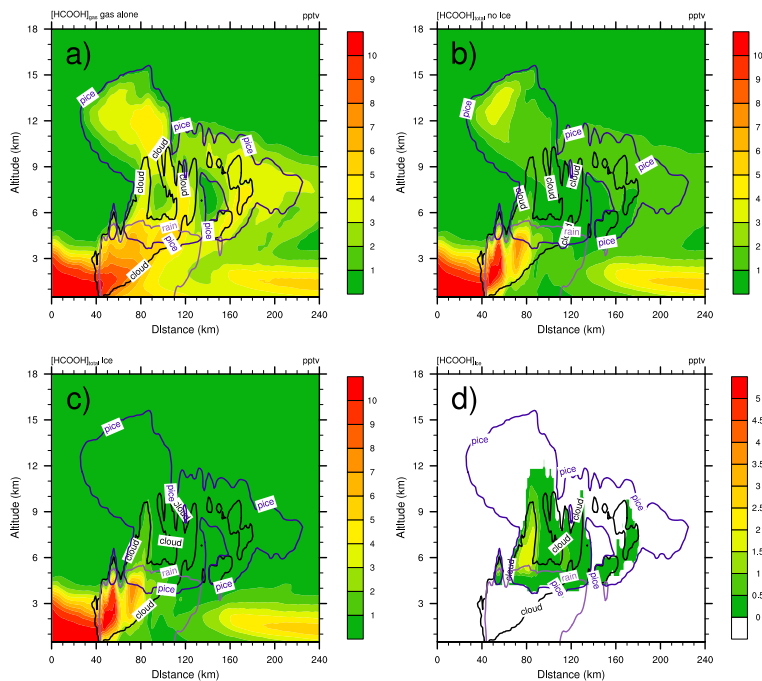
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**Fig. 8.** COPT case: same as Fig. 7 but for formic acid.

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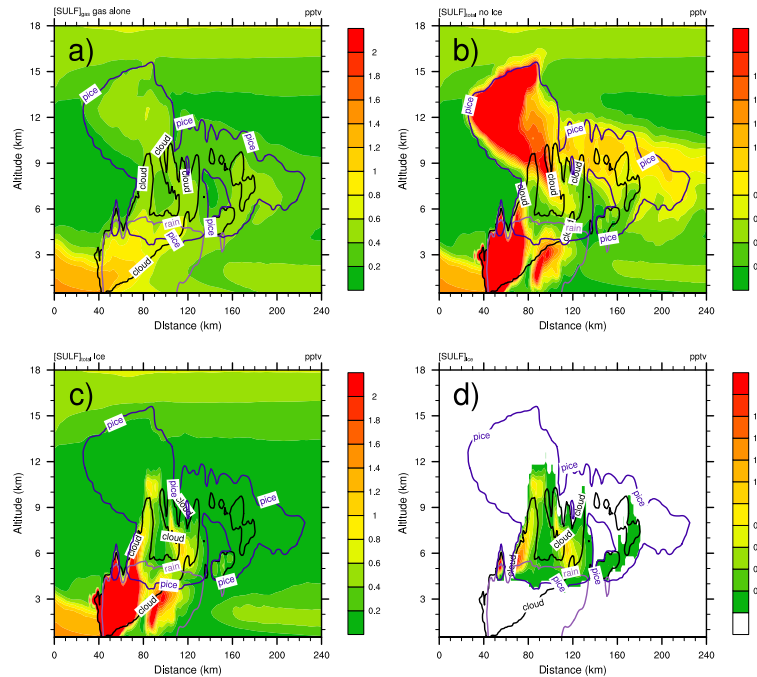


Fig. 9. COPT case: same as Fig. 7 but for sulphuric acid.

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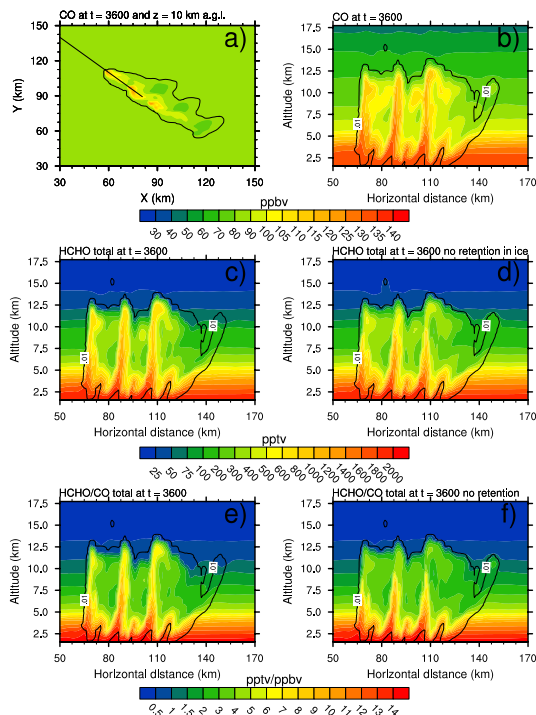


Fig. 10. STERAO case: results at $t = 3600$ s for (a) horizontal cross section of CO mixing ratio in ppbv at 10 km a.g.l., (b) vertical cross section along the segment indicated on (a) for CO mixing ratio in ppbv, (c) same vertical cross section for HCHO total mixing ratio (gas + cloud water + rainwater + precipitating ice) in pptv for simulation with retention in ice, (d) same as (c) for simulation without retention in ice, (e) same vertical cross section for the ratio of total HCHO to CO in pptv ppbv⁻¹ for simulation with retention in ice, (f) same as (e) for simulation without retention in ice.

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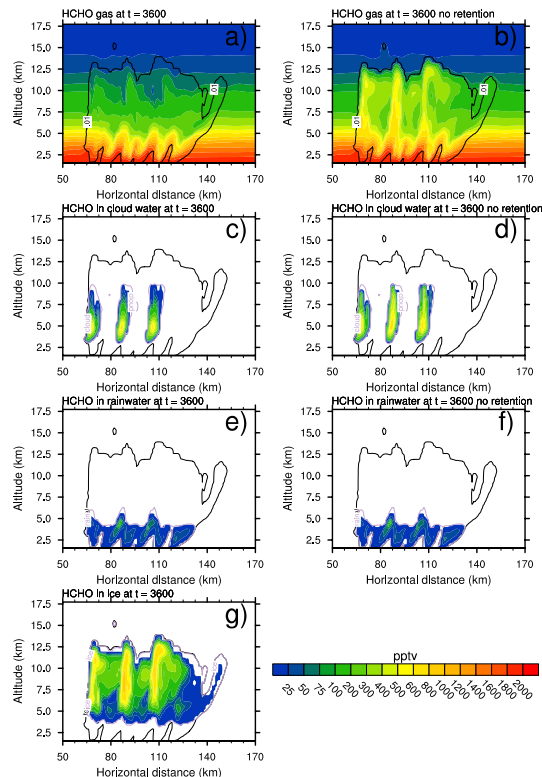


Fig. 11. STERAO case: results at $t = 3600$ s for vertical cross section of HCHO mixing ratio in pptrv along the segment indicated on Fig. 10a, at the left hand side for simulation with retention in ice and at the right hand side for simulation without retention: **(a)** and **(b)** in gas phase, **(c)** and **(d)** in cloud water, **(e)** and **(f)** in rainwater, **(g)** in precipitating ice. The black line represents the total hydrometeors mixing ratio equal to 0.01 kg kg^{-1} , the grey line on **(c)** and **(d)** is the same for cloud water, the grey line on **(e)** and **(f)** is the same for rainwater and the grey line on **(g)** is the same for precipitating ice.

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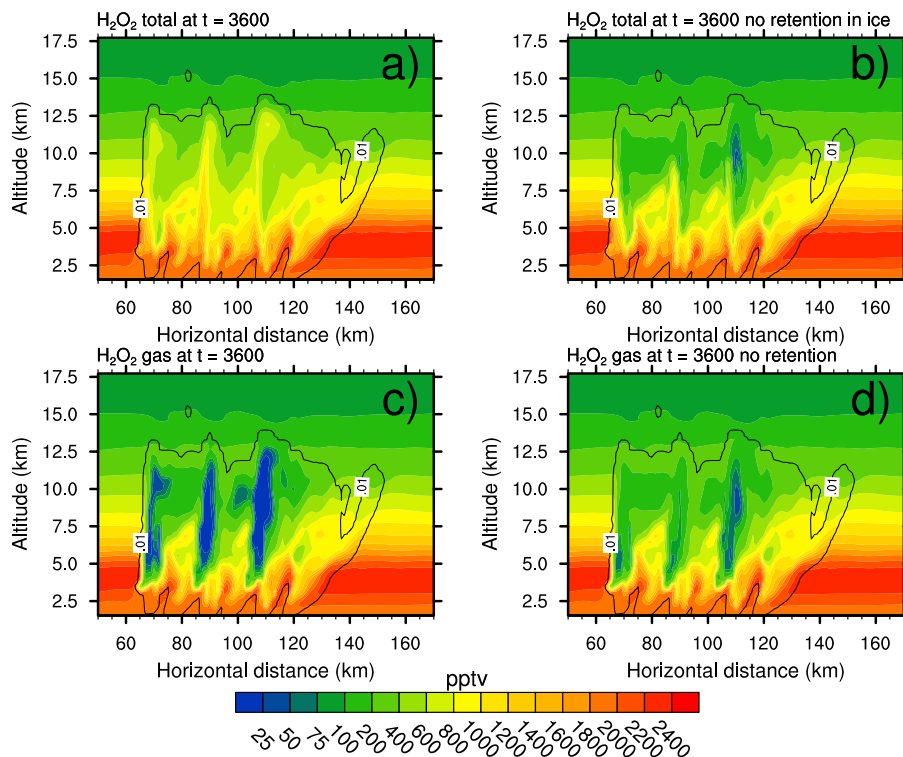


Fig. 12. STERAO case: results at $t = 3600$ s for vertical cross section of H_2O_2 mixing ratio in pptv along the segment indicated on Fig. 10a, on the left hand side for simulation with retention in ice and on the right hand side for the simulation without retention: **(a)** and **(b)** total mixing ratio, **(c)** and **(d)** in gas phase. The black line represents the total hydrometeors mixing ratio equal to 0.01 g kg^{-1} .

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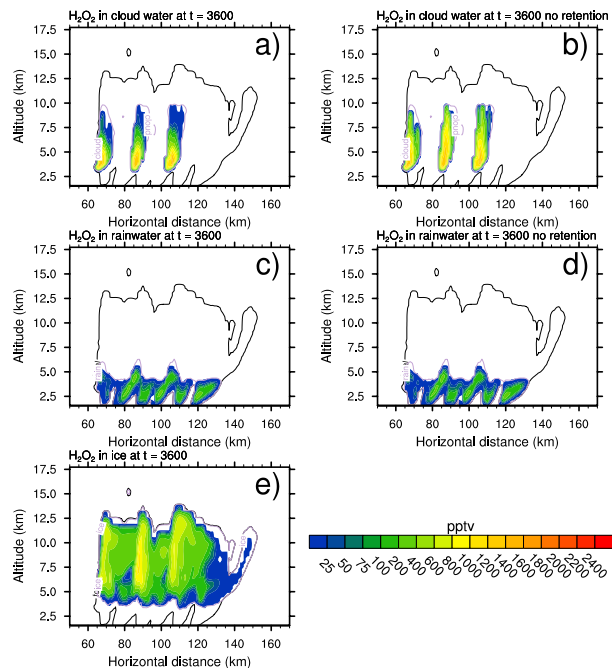


Fig. 13. STERAO case: results at $t = 3600$ s for vertical cross section of H_2O_2 mixing ratio in pptv along the segment indicated on Fig. 10a, on the left hand side for simulation with retention in ice and on the right hand side for the simulation without retention: **(a)** and **(b)** in cloud water, **(c)** and **(d)** in rainwater, **(e)** in precipitating ice. The black line represents the total hydrometeors mixing ratio equal to 0.01 g kg^{-1} , the grey line on **(a)** and **(b)** is the same for cloud water, the grey line on **(c)** and **(d)** is the same for rainwater and the grey line on **(e)** is the same for precipitating ice.

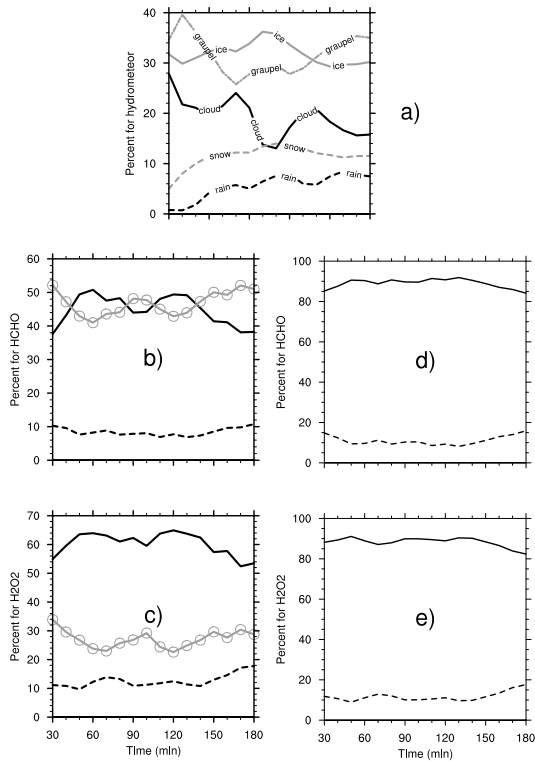


Fig. 14. STERAO case: results over the model domain as a function of time in the simulation sampled every 10 min: **(a)** percentage of each hydrometeor reservoir relative to the total amount of condensed water, **(b)** percentage of HCHO in each water category (cloud, rain, precipitating ice) to the total amount of HCHO in all the water reservoirs for simulation with retention in ice, **(c)** the same for H_2O_2 , **(d)** the same for HCHO but for simulation without retention in ice, **(e)** the same for H_2O_2 but for the simulation without retention in ice. Solid and dashed black lines **(b–e)** are for cloud water and rainwater reservoirs, respectively. Grey line with open circles **(b, c)** is for the ice-precipitating reservoir.