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ACPD 8, 9761-9782, 2008 **Particle formation** from OH+SO₂ T. Berndt et al. **Title Page** Introduction Abstract Conclusions References Figures Back Full Screen / Esc





SO₂ oxidation products other than H₂**SO**₄ as a trigger of new particle formation – Part 1: Laboratory investigations

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Abstract

Mechanistic investigations of atmospheric H₂SO₄ particle formation have been performed in a laboratory study taking either H₂SO₄ from a liquid reservoir or using the gas-phase reaction of OH radicals with SO₂. Applying both approaches for H₂SO₄
 ⁵ generation simultaneously we found that H₂SO₄ evaporated from the liquid reservoir acts considerably less effective for the process of particle formation and growth than the products originating from the reaction of OH radicals with SO₂. Furthermore, for NO_x concentrations >5×10¹¹ molecule cm⁻³ the formation of new particles from the reaction of OH radicals with SO₂ is inhibited. This suggests that substances other than H₂SO₄ (likely products from sulphur-containing peroxy radicals) trigger lower tropospheric new particle formation and growth. The currently accepted mechanism for SO₂ gas-phase oxidation does not consider the formation of such substances making a revision necessary.

1 Introduction

¹⁵ For more than a decade the formation of new aerosol particles in the atmosphere has been the subject of intense studies in the field and in the laboratory. The formation mechanisms and the participating substances, however, have not been resolved yet (Kulmala, 2003). Large discrepancies between model-predicted nucleation rates for the favoured binary system H₂SO₄/H₂O and much higher atmospheric nucleation data
 were explained by various supportive additional participants such as ammonia (Kulmala et al., 2000), organic molecules (Yu et al., 1999; Hoffmann et al., 1998) or ions (Lee et al., 2003; Lovejoy et al., 2004).

In a previous investigation of our laboratory under near-atmospheric conditions, experimental evidence for the formation of new particles in the system H_2SO_4/H_2O was

²⁵ found for H₂SO₄ concentrations of ~10⁷ molecule cm⁻³ if "H₂SO₄" was produced insitu via the reaction of OH radicals with SO₂ ("H₂SO₄" here stands for all products of



converted SO₂). As the OH radical source served UV-photolysis of O_3 or ozonolysis of alkenes in a dark reaction. The experiments have been performed in the presence and absence of organics. No clear difference concerning new particle formation was observed using the different experimental conditions (presence or absence of UV)

- light, presence and absence of organics) pointing clearly at the SO₂ oxidation products ("H₂SO₄") being the exclusive precursors for the newly formed particles observed (Berndt et al., 2004, 2005, 2006). A significant increase of the nucleation rate for increasing r.h. was measurable (Berndt et al., 2005, 2006). Similar observations are reported from another flow-tube experiment (Friend et al., 1980) as well as from in vestigations in a chamber (Burkholder et al., 2007). In contrast, taking H₂SO₄ from a
- liquid reservoir concentrations of $10^9 10^{10}$ molecule cm⁻³ are needed for new particle formation (Ball et al., 1999, Zhang et al., 2004). This observation is roughly in line with the prediction from the binary nucleation theory H₂SO₄/H₂O (Kulmala et al., 1998).

In this study mechanistic investigations on H_2SO_4/H_2O particle formation are reported explaining the different threshold H_2SO_4 concentrations needed for nucleation, i.e., ~10⁷ molecule cm⁻³ ("H₂SO₄" produced via gas-phase reaction of OH radicals with SO₂) vs. 10⁹-10¹⁰ molecule cm⁻³ (H₂SO₄ via evaporation from a liquid reservoir).

2 Experimental

The experiments have been performed in the atmospheric pressure flow-tube *IfT*-LFT (i.d. 8 cm; length 505 cm) at 293±0.5 K. The first tube section (56 cm) includes an inlet system for gas entrance (humidified air premixed with SO₂, O₃, NO or hydrocarbons or H₂SO₄ vapour from a liquid reservoir premixed in dry air). The middle section (344 cm) is equipped with 8 UV lamps (Hg-lamps made of quartz-glass PN235 with a cut-off wavelength of 210 nm) for a homogeneous irradiation of the tube if needed. A nonirradiated end section (105 cm), which also holds the sampling outlet, is attached.

Gas-phase analysis was carried out using a GC-FID (HP 5890) with a cryoenrichment unit for hydrocarbons, a humidity sensor (Vaisala), analyzers for O_{3} , SO_{2} ,



and NO_x (Thermo Environmental Instruments: 49C, 43C, and 42S), a butanol-based UCPC (TSI 3025) as well as a H₂O-based UCPC (TSI 3786) for integral particle measurements, and a differential mobility particle sizer (Vienna-type DMA with UCPC, TSI 3025) for monitoring of size distributions.

- ⁵ The carrier gas consisted of synthetic air (99.9999999%, Linde and further purification with Gate Keeper, AERONEX). A small part of the total gas flow was flushed through an ozone generator (UVP OG-2) for O₃ generation outside the flow tube. SO₂ was taken from a 1ppmv or 10ppmv calibration mixture in N₂ (Messer) and NO from 0.5 vol% mixture in N₂ (Messer). In the case of experiments with H₂SO₄ vapour from a
- ¹⁰ liquid reservoir a part of the total gas stream of dry air was flushed through a saturator (maintained at 288 K) filled with concentrated sulphuric acid (95–98%, Aldrich). The outlet tube of the saturator served directly as the inlet for the H_2SO_4 vapour containing gas stream. In the first tube section this H_2SO_4 gas stream was brought together with humidified air.
- ¹⁵ The total gas flow inside the *IfT*-LFT was set at 3.33, 10, 20, or 30 standard litre min⁻¹ resulting in a bulk residence time (middle and end section) of 378, 126, 63, or 42 sec., respectively. It is to be noted that the given bulk residence time stands for the upper limit of the nucleation time in the flow tube. All gas flows were set by means of calibrated gas flow controllers (MKS 1259/1179) and the pressure in the tube was measured using a capacitive manometer (Baratron).
 - 2.1 Determination of H₂SO₄ taken from a liquid reservoir

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For experiments using H_2SO_4 vapour from a liquid reservoir, H_2SO_4 concentrations were measured at the outlet of the saturator by means of a denuder system (Amanda, ECN). The working fluid was a 10^{-4} M NaOH solution of de-ionised water and sampling times were in the range of 2–10 h. The subsequent analysis of SO_4^{2-} ions has been done by means of ion chromatography (761 compact IC, Metrohm).

ACPD 8, 9761-9782, 2008 **Particle formation** from OH+SO₂ T. Berndt et al. **Title Page** Abstract Introduction Conclusions References **Figures** Þ١ Back Full Screen / Esc **Printer-friendly Version** Interactive Discussion



2.2 Determination of "H₂SO₄" (products of converted SO₂) and OH concentrations

Starting from the reaction of OH radicals with SO_2 "H₂SO₄" concentrations were calculated using a model according to the following reaction scheme used in the case of photolysis experiments (Berndt et al., 2005, 2006):

$$_{5} O_{3} \rightarrow 2 OH$$
 (1)

$$OH + furan/CO \rightarrow products$$
 (2)

$$OH + SO_2 \rightarrow \ldots \rightarrow "H_2SO_4"$$
(3)

 $\text{``H}_2\text{SO}_4\text{''} \rightarrow \text{wall}$

The effective photolysis rate coefficient k_1 was determined in each experiment measuring the O₃ decay. Rate coefficients $k_{2,\text{furan}} = 4.0 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, $k_{2 \text{ CO}} = 2.4 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, and $k_3 = 1.2 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, were taken from literature (Finlayson-Pitts and Pitts, 2000, and references therein). For description of the assumed, diffusion controlled wall loss of "H₂SO₄", k_4 =0.017 s⁻¹, the same data as for H₂SO₄ were taken (Hanson and Eisele, 2000). The concentration of furan (or CO) was chosen such that $k_{2 \text{ furan}} \times [\text{furan}] > 20 \text{ s}^{-1}$ ($k_{2 \text{ furan}} \times [\text{furan}]$ 15 $\sim k_{2 CO} \times [CO]$) was fulfilled. Modelling studies showed that under these conditions the reactions of OH radicals with $\mathrm{O}_3,\,\mathrm{HO}_2$ or any impurities (impurity concentration $\leq 10^{10}$ molecule cm⁻³, $k_{OH+impurity} = 10^{-10}$ cm³ molecule⁻¹ s⁻¹) as well as the wall loss of OH radicals were negligible allowing also the determination of OH radicals from the measured amount of reacted furan. The ratio "reacted O₃"/"reacted furan" was exper-20 imentally found to be 0.5±0.2 as expected from the simple reaction scheme given above. This finding confirms that OH radical concentration, and consequently the "H₂SO₄" concentrations, are well described by the model applied. For normally used initial reaction conditions (O_3 : ~1.5×10¹¹ with a conversion <25%, furan: 6.2×10¹¹

(4)

with a conversion $\leq 6\%$ or CO: 1.0×10^{14} with a conversion <1% and SO₂: (6.7–350)×10¹⁰ with a conversion <2.5%; concentration unit: molecule cm⁻³) and measured effective photolysis rate coefficient k_1 in the order of 10^{-3} s^{-1} , modelling of the reaction system yielded OH radical concentrations of about 10^7 molecule cm⁻³ being close to atmospheric levels. Given "H₂SO₄" concentrations in figures and explanations represent averaged values for the irradiated middle section.

In the case of added NO_x in the carrier gas, NO and NO₂ can undergo reactions with O₃ and OH radicals. An additional pathway for OH radical generation represents the reaction of NO with HO₂ forming OH and NO₂. Therefore, OH radical concentrations were determined experimentally using the titration procedure with furan. The OH profile in the tube was calculated from the measured furan decay assuming an exponential function for the overall decay of O₃ and for the decay of furan. With the knowledge of the axial OH radical profile "H₂SO₄" concentrations represent also averaged values for the irradiated middle section.

3 Results and Discussion

3.1 Measurement of consumed SO₂

It was attempted to measure the amount of consumed SO₂ to evaluate the reliability of calculated "H₂SO₄" concentrations via pathways (1)–(4). In Fig. 1 the temporal behaviour of SO₂ concentration and particle number for an experiment with CO for OH radical titration is given. SO₂ measurement using Thermo Environmental Instruments 43C in the absence of OH radicals (UV off) yielded: $(6.78\pm0.24)\times10^{10}$ molecule cm⁻³ (full line) and in the presence of OH radicals (UV on): $(6.63\pm0.24)\times10^{10}$ molecule cm⁻³ (dashed line) resulting in a SO₂ consumption of 1.5×10^9 molecule cm⁻³ with an uncertainty by a factor of ~2.3. Modelling of this example (including wall loss for "H₂SO₄")



shows an average " H_2SO_4 " concentration in the tube of 2.46×10^8 molecule cm⁻³ and a SO₂ consumption of 1.46×10^9 molecule cm⁻³. Simulated and measured SO₂ consumption are in good agreement. The particle number exceeded the counting range of the TSI 3025 (particle number >10⁵ cm⁻³). Integration of simultaneously performed size distribution measurement yielded a particle number of 3.6×10^5 cm⁻³.

The determination of reacted SO₂ stands for a complementary measure of "H₂SO₄" and the good agreement between measured and calculated SO₂ conversion shows explicitly that the "true" concentration of "H₂SO₄" in the experiment cannot exceed the modelling data considerably.

 $_{10}$ 3.1.1 New particle formation in the presence of in-situ produced "H $_2SO_4$ " and H $_2SO_4$ from a liquid reservoir

In a first set of experiments in-situ formation of "H₂SO₄" via OH+SO₂ was coupled with evaporation of H₂SO₄ from the liquid reservoir as background in order to investigate the role of H₂SO₄ for the new particle formation and growth process. Initial H₂SO₄ concen-¹⁵ tration from the liquid reservoir was (1–2)×10⁹ molecule cm⁻³, i.e., close to the threshold concentration needed for binary H₂SO₄/H₂O nucleation in our experiment. The bulk residence time in the reaction zone (irradiated section) was 97 s and the end-H₂SO₄ concentration (from liquid reservoir) was estimated to be (2–4)×10⁸ molecule cm⁻³ considering diffusion controlled wall loss only (Hanson and Eisele, 2000). In Fig. 2 results from a typical measurement are depicted. After turning on the flow through the H₂SO₄-saturator, a small increase of particle number above the noise level was visible with a subsequent slight rise up to 2–3 particles cm⁻³ after more than 5 h on stream. These particles are due to binary nucleation of H₂SO₄/H₂O (Kulmala et al., 1998). The particle number was not affected by the presence of O₃ and the photolysis products

²⁵ (OH radicals, HO₂, and H₂O₂) after switching on UV irradiation. With addition of SO₂ (time=235 min), in-situ particle precursor formation in the gas phase started and a constant particle number of 5×10^4 cm⁻³ was observed for in-situ "H₂SO₄" concentration



of 2.3×10^8 molecule cm⁻³. Immediately after switching off UV (time=330 min), particle number went down to the level being attributed to binary H₂SO₄/H₂O nucleation. At 360 min the supply of H₂SO₄ from the liquid reservoir was stopped resulting in a decrease of particle number down to background noise level. Switching on UV, i.e., restarting the in-situ "H₂SO₄" production without background H₂SO₄, led to a particle number of 5×10^4 cm⁻³ again. This behaviour is a strong indication that H₂SO₄ from the liquid reservoir (now absent) did not participate in the nucleation process. To illus-

- trate the influence of background H_2SO_4 on particle growth, Fig. 3 shows measured size distributions in the diameter range d=2-4 nm recorded in the time period 275 to 325 min (presence of background H_2SO_4) and between 410 and 460 min (absence of
- background H_2SO_4). To make sure that the measurements were not affected by memory effects, after a 3 day flushing of the flow tube with clean air, the experiment without background H_2SO_4 from the liquid reservoir was repeated. Figure 3 shows that the three measured size distributions are nearly identical. This indicates that H_2SO_4 from
- the liquid reservoir, its concentration being at least in the same order or one order of magnitude higher compared to that of in-situ produced "H₂SO₄" (cf. Fig. 4), does not significantly contribute to particle growth. Obviously, condensational growth by H₂SO₄ evaporated from the liquid reservoir is of less importance under the chosen conditions and cannot be resolved. A kinetically limited collision model assuming H₂SO₄-like
 substances as the particle precursor (probably products of HOSO₂O₂, see later) can roughly describe particle growth into the size range measured.

It is to be noted that integration of the size distributions ($d \ge 2$ nm) given in Fig. 3 yielded a particle number of $\sim 5 \times 10^3$ cm⁻³, i.e., 1/10 of the total number found in the integral measurement using the UCPC. Therefore, the majority of produced particles in this example were smaller than 2 nm in diameter.

25

To get an rough estimate concerning the H_2SO_4 /" H_2SO_4 " mass balance it is assumed i) that H_2SO_4 is the exclusive product from the reaction of OH radicals with SO_2 and ii) that the particles consist of H_2SO_4 (density of 1.85 g cm⁻³) only, i.e., contributions of water to particle mass are neglected. For the example described above



(assuming 5×10^3 particle with d=2 nm and 4.5×10^4 particle cm⁻³ with d=1.5 nm) it follows that about 0.5% of gas phase H₂SO₄/"H₂SO₄" can attributed to the newly formed particles.

Experiments, during which OH radicals were produced via ozonolysis of *trans*-⁵ butene (dark reaction) yielded similar results.

These findings pose the question, why H_2SO_4 from the liquid reservoir exhibits a totally different behaviour (no clear contribution to particle formation and growth) compared to in-situ produced " H_2SO_4 "? A possible reason could be that H_2SO_4 is not the particle precursor produced from the reaction of OH radicals with SO_2 .

The currently accepted mechanism of atmospheric SO_2 gas-phase oxidation initiated by OH radicals is as follows (Finlayson-Pitts and Pitts, 2000, and references therein):

 $OH + SO_2 \rightarrow HOSO_2$

 $HOSO_2 + O_2 \rightarrow SO_3 + HO_2$

10

 $SO_3 + 2 H_2O \rightarrow H_2SO_4 + H_2O$

- ¹⁵ Here, HOSO₂ radicals produced from the primary reaction of OH radicals with SO₂ via pathway (5) react in the very fast consecutive step (6) with O₂ forming SO₃. SO₃ reacts with two water molecules or a water dimer producing H₂SO₄. According to this scheme, each attacked SO₂ molecule from pathway (5) is transformed to a H₂SO₄ molecule.
- As a result of their pioneering work in the eighties, Stockwell and Calvert (1983) found that pathway (6) has to be important for the atmospheric fate of $HOSO_2$ as via this reaction more than 80% of $HOSO_2$ regenerates HO_2 . However, an earlier way of thinking was based on a different pathway (6a) which considered the addition of O_2 to $HOSO_2$:

²⁵ HOSO₂ + O₂ + M
$$\rightarrow$$
 HOSO₂O₂ + M

(5)

(6)

(7)

(6a)



Interactive Discussion

Pathway (6) represents an endothermic reaction whereas pathway (6a) is exothermic (Benson, 1978, Majumdar et al., 2000), i.e., pathway (6a) is the thermodynamically favoured one. From the pressure-dependence of the kinetics of the $HOSO_2+O_2$ reaction, it was concluded that pathway (6a) accounts for <10% of the $HOSO_2$ removal

Gleason et al., 1987). Simultaneous measurements of HO₂ and HOSO₂ concentrations yielded a lower limit of 70% for pathway (6) (Howard, 1985). Taking all experimental findings into consideration, up to 30% of HOSO₂ can be converted to HOSO₂O₂.

It should be noted that a reduction of H_2SO_4 formation from SO_2 oxidation due to pathway (6a) is within the measurement uncertainty of the CIMS instruments (chemical ionization mass spectrometry) applied in laboratory and field measurements (Lovejoy et al., 1996).

 $HOSO_2O_2$ is a peroxy-type radical and, analogous to organic peroxy radicals, it can react with HO_2 , other peroxy radicals or with NO and NO_2 (Finlayson-Pitts and Pitts, 2000, and references therein). Also the reaction of $HOSO_2O_2$ or its hydrated species $HOSO_2O_2(H_2O)_n$ with SO_2 is imaginable (Wayne, 1991). If the reaction of $HOSO_2O_2$ with HO_2

 $HOSO_2O_2 + HO_2 \rightarrow HOSO_2O_2H + O_2$ (8)

and/or with other peroxy radicals (RO₂)

 $HOSO_2O_2 + RO_2 \rightarrow products$

²⁰ and/or with SO₂

10

15

 $HOSO_2O_2 + SO_2 \rightarrow HOSO_2OSO_3$

is responsible for the formation of particle precursors, addition of NO and/or NO₂ should reduce or inhibit new particle formation as a part of the available $HOSO_2O_2$ will be consumed by NO and/or NO₂. (For simplification pathway (10) is written without hydration.)

ACPD 8, 9761-9782, 2008 **Particle formation** from OH+SO₂ T. Berndt et al. **Title Page** Abstract Introduction Conclusions References **Tables Figures** Þ١ Back Full Screen / Esc **Printer-friendly Version** Interactive Discussion

(9)

(10)

3.1.2 New particle formation in the presence of NO/NO₂

New particle formation in the presence of NO/NO₂ using in-situ produced "H₂SO₄"

- To test the hypothesis that HOSO₂O₂ is a possible precursor for the nucleating species and that in the presence of NO and/or NO₂ new particle formation is inhibited, experiments were carried out, with NO_x (NO+NO₂) being present in the carrier gas. In this context it should be noted that generally NO and NO₂ can undergo reactions with O₃, OH radicals, HO₂ and other RO₂ radicals influencing the OH radical level significantly. Therefore, to determine the in-situ produced "H₂SO₄" concentration, OH radical titration with furan was applied during this set of experiments. Figure 5, as an example, shows the results of measurements performed for different initial NO concentrations. In the course of the overall reaction, 32% of NO reacted to NO₂. It
- is clearly seen that with increasing amounts of NO_x being present in the reaction gas, new particle formation is inhibited. Similar observations were made for relative humidities in the range 10–60 % or using NO₂ instead of NO. As a net inhibition of new particle formation is observed, the products from the NO_x reactions, if important at all, have to be less effective particle precursors than the products from the reaction with HO₂ and/or other peroxy radicals or with SO₂ via pathways (8), (9) or (10), respectively.
- NO_x cannot effect H_2SO_4 formation as described by pathways (5)–(7) because the $HOSO_2$ life-time regarding the O_2 reaction, pathway (6), as well as the SO_3 life-time regarding hydrolysis, pathway (7), are much shorter than those resulting from the corresponding NO_x reactions (A more detailed explanation is given in Appendix A).

25

New particle formation in the presence of NO/NO₂ using H_2SO_4 from the liquid reservoir



Furthermore, the influence of NO_x (NO+NO₂) for the process of new particle formation was studied using H₂SO₄ from the liquid reservoir. Figure 6 shows the temporal behaviour of O_3 concentration and the particle number for an experiment using an initial H_2SO_4 concentration (from the liquid reservoir) of ~5×10⁹ molecule cm⁻³. After turning on the flow through the H₂SO₄-saturator a slightly increasing particle number was observed for the whole time on stream. At 55 min NO with a concentration of 4.1×10^{12} molecule cm⁻³ was added and after turning on the external O₃ generation (O₃: 1.5×10^{11} molecule cm⁻³) NO was partly converted to NO₂ with an end concentration of ~1.1×10¹¹ molecule cm⁻³, O_3 +NO \rightarrow NO₂+O₂. Switching on UV lamps at 10 175 min OH radical formation started and beside OH radicals, HO_2 and H_2O_2 also HONO and HNO₃ were produced. After turning off the NO flow (time: 205 min) only the O_3 photolysis products (OH radicals, HO₂ and H₂O₂) remained. Under all conditions investigated, especially for NO_x addition, no impact of added trace gases on the number of newly formed particles was visible. The absence of a NO_x dependence on 15 the number of new particles is a clear disagreement regarding the findings using insitu produced "H₂SO₄". Consequently, these results represent a further indication that reaction products of SO₂ oxidation other than H_2SO_4 (most likely products of sulphurcontaining peroxy-type radicals like $HOSO_2O_2$) represent possible particle precursors.

20 4 Summary and conclusions

It was pointed out that, under lower tropospheric conditions, H₂SO₄ evaporated from a liquid reservoir did not significantly participate in the formation and growth of particles formed from the gas phase reaction of OH radicals with SO₂. Furthermore, it was shown that for NO_x concentrations >5×10¹¹ molecule cm⁻³ the formation of new particles from the reaction of OH radicals with SO₂ is inhibited. This behaviour is not in line with the currently accepted reaction mechanism describing an equal-molar formation of H₂SO₄ from SO₂. It is therefore suggested that in the course of SO₂ con-



version, sulphur-containing peroxy radicals like HOSO₂O₂ can be formed in substantial fractions leading to products being different from H₂SO₄. It might be speculated, that under lower tropospheric conditions, HOSO₂O₂ could react with HO₂ and other peroxy radicals (including the self-reaction). These reactions, being of the RO₂+RO₂ type,
⁵ could result in low vapour pressure particle precursors. If, e.g., HOSO₂O₂ reacts with HO₂ (the most abundant RO₂ radical in atmosphere), peroxo sulphuric acid is formed. Peroxo sulphuric acid is a hydrophilic solid, which in the liquid phase (in water) is con-

verted to sulphuric acid and H₂O₂ (Kolditz, 1983). Another idea for particle formation starting from HOSO₂O₂ is given by Wayne (1991) describing the reaction of hydrated
 HOSO₂O₂ (HOSO₂O₂(H₂O)_n) with SO₂ producing HOSO₂O(H₂O)_nSO₃. For large "n" these hydrated radicals are regarded as aerosol particles (or at least as precursors). These scenarios, however, are highly speculative at the moment.

In conclusion, this study shows, that it is not necessarily sulphuric acid triggering atmospheric new particle formation and growth and that other compounds (like peroxo sulphuric acid or HOSO₂O(H₂O)_nSO₃ radicals) formed during the gas phase reaction

of OH radicals with SO₂ may play an important role.

In a second part a comparison of ambient and laboratory measurements are given and the atmospheric implications are discussed (Laaksonen et al., 2008).

Appendix A

20

Atmospheric life-times of HOSO₂ and SO₃

Assuming a concentration of 10^{12} molecule cm⁻³ for NO and NO₂ each, a HOSO₂ life-time regarding NO reaction of >2 s (Martin et al., 1986) and regarding NO₂ reaction of >10 milliseconds (assuming an upper limit for the rate coefficient of 10^{-10} cm³ molecule⁻¹ s⁻¹) follows. A comparison with the life-time regarding the overall O₂ reaction via pathway (6) of ~0.5 microseconds (Gleason et al., 1987) shows clearly that NO and NO₂ cannot influence the conversion of HOSO₂ via pathways (6) and (6a).

Assuming for the reaction of SO₃ with NO and NO₂ the rate coefficient measured for the NO₂ reaction (Penzhorn and Canosa, 1983), the SO₃ life-times regarding the NO and NO₂ reaction are in the order of 80 days. That means that also the hydrolysis of SO₃ via pathway (7) (with a life-time of ~20 microseconds assuming a H₂O con-⁵ centration of 3×10¹⁷ molecule cm⁻³; Jayne et al., 1997) cannot be affected by NO and NO₂. Consequently, NO and NO₂ for the concentrations considered here, cannot affect H₂SO₄ formation via pathways (5)–(7).

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ACPD		
8, 9761–9782, 2008		
Particle formation from OH+SO ₂		
T. Berndt et al.		
Title Page		
Abstract	Introduction	
Conclusions	References	
Tables	Figures	
I	۶I	
•	•	
Back	Close	
Full Screen / Esc		
Printer-friendly Version		
Interactive Discussion		

9775

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ACPD		
8, 9761–9782, 2008		
Particle formation from OH+SO ₂		
T. Berndt et al.		
Title Page		
Abstract	Introduction	
Conclusions	References	
Tables	Figures	
14	►I	
•	Þ	
Back	Close	
Full Screen / Esc		
Printer-friendly Version		
Interactive Discussion		



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ACPD		
8, 9761–9782, 2008		
Particle formation from OH+SO ₂		
Title Page		
Abstract	Introduction	
Conclusions	References	
Tables	Figures	
I	۶I	
•	•	
Back	Close	
Full Screen / Esc		
Printer-friendly Version		
Interactive Discussion		

BY



Fig. 1. Temporal behaviour of SO₂ concentration and particle number for an experiment with "H₂SO₄" formation from the reaction of OH radicals with SO₂ using CO for OH radical titration, r.h.=22%, gas flow: 3.33 sl min⁻¹. Initial concentrations were (unit: molecule cm⁻³); O₃: 3.8×10^{11} ; CO: 5.3×10^{13} . SO₂ concentration in the absence of OH radicals (UV off): $(6.78 \pm 0.24) \times 10^{10}$ molecule cm⁻³ (full line) and in the presence of OH radicals (UV on): $(6.63 \pm 0.24) \times 10^{10}$ molecule cm⁻³ (dashed line) results in a SO₂ consumption of 1.5×10^9 molecule cm⁻³ (uncertainty by a factor of ~2.3).





ACPD 8,9761-9782,2008 **Particle formation** from OH+SO₂ T. Berndt et al. **Title Page** Introduction Abstract Conclusions References **Figures** Back Full Screen / Esc **Printer-friendly Version**

Fig. 2. Temporal behaviour of O_3 and SO_2 concentration and particle number for an experiment using H_2SO_4 from the liquid reservoir as well as in-situ formation of " H_2SO_4 " (SO₂ products) with CO for OH radical titration, r.h.=22%, gas flow: 10 sl min⁻¹, bulk residence time (middle and end section): 126 sec. Initial concentrations were (unit: molecule cm⁻³); H_2SO_4 (saturator): $(1-2)\times10^9$; O_3 : 1.6×10^{11} ; CO: 1.0×10^{14} ; SO_2 : 3.5×10^{11} .

Interactive Discussion





ACPD

8, 9761-9782, 2008

Particle formation from OH+SO₂

T. Berndt et al.



Fig. 3. Measured size distributions in the range d=2-4 nm from the experiment depicted in Fig. 2. The distribution with H₂SO₄ from the saturator was recorded between 275 and 325 min and that without H₂SO₄ from the saturator between 410 and 460 min, cf. Fig. 2. After a 3 day flushing of the flow tube with pure air the distribution with black symbols in absence of H₂SO₄ from the saturator was measured.



ACPD

8, 9761-9782, 2008

Particle formation from OH+SO₂

T. Berndt et al.



Fig. 4. CFD-Modelling: Calculated concentration profiles of H_2SO_4 from the liquid reservoir and from in- situ produced " H_2SO_4 " via the reaction of OH radicals with SO_2 , r.h.=22%, gas flow: 10 sl min⁻¹. Initial concentrations in the case of in situ " H_2SO_4 " formation were (unit: molecule cm⁻³); O_3 : 1.6×10^{11} ; CO: 1.0×10^{14} ; SO_2 : 3.5×10^{11} .



Fig. 5. Experimentally observed particle number as a function of in-situ produced " H_2SO_4 " (SO₂ products) concentration for different NO additions in the carrier gas using furan for OH radical titration, OH radical formation via O₃ photolysis, r.h.=22%, gas flow: 30 sl min⁻¹, bulk residence time (middle and end section): 42 s Initial concentrations were (unit: molecule cm⁻³); O₃: (1.5–1.6)×10¹¹; furan: 6.2×10¹¹; furan conversion: 2.1–5.3%; SO₂: (1.2–16)×10¹¹.

ACPD 8,9761-9782,2008 **Particle formation** from OH+SO₂ T. Berndt et al. **Title Page** Introduction Abstract Conclusions References **Figures** Back Full Screen / Esc **Printer-friendly Version** Interactive Discussion



ACPD

8, 9761–9782, 2008

Particle formation from OH+SO₂

T. Berndt et al.



Fig. 6. Temporal behaviour of O_3 concentration and the particle number for an experiment using H_2SO_4 from the liquid reservoir, r.h.=22%, gas flow: 30 sl min⁻¹, bulk residence time (middle and end section): 42 s Initial concentrations were (unit: molecule cm⁻³); H_2SO_4 (saturator): ~5×10⁹; O_3 : 1.5×10¹¹; CO: 1.0×10¹⁴; NO: 4.1×10¹².