Atmos. Chem. Phys. Discuss., 5, 10369–10408, 2005 www.atmos-chem-phys.org/acpd/5/10369/ SRef-ID: 1680-7375/acpd/2005-5-10369 European Geosciences Union



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The heterogeneous chemical kinetics of N₂O₅ on CaCO₃ and other atmospheric mineral dust surrogates

F. Karagulian, C. Santschi, and M. J. Rossi

Ecole Polytechnique Fédérale de Lausanne, Laboratoire de Pollution Atmosphérique et Sol (EPFL, LPAS), Bâtiment CH H5, Station 6, CH-1015 Lausanne, Switzerland

Received: 7 September 2005 - Accepted: 12 September 2005 - Published: 24 October 2005

Correspondence to: M. J. Rossi (michel.rossi@epfl.ch)

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Abstract

Uptake experiments of N_2O_5 on several mineral dust powder samples were carried out under continuous molecular flow conditions at 298±2K. At $[N_2O_5]_0 = (4.0 \pm 1.0) \times 10^{11} \text{ cm}^{-3}$ we have found γ_{ss} values ranging from $(3.5 \pm 1.1) \times 10^{-2}$ for CaCO₃ to (0.20±0.05) for Saharan Dust with γ_{ss} decreasing as [N₂O₅]₀ increased. We have observed delayed production of HNO₃ upon uptake of N₂O₅ for every investigated sample owing to hydrolysis of N₂O₅ with surface-adsorbed H₂O. At high and low $[N_2O_5]$ Arizona Test Dust and Kaolinite turned out to be the samples to produce the largest amount of gas phase HNO₃ with respect to N_2O_5 taken up. In contrast, the yield of HNO₃ for Saharan Dust and CaCO₃ are lower. On CaCO₃ the disappearance 10 of N_2O_5 was also accompanied by the formation of CO_2 . For CaCO₃ sample masses ranging from 0.33 to 2.0 g, the yield of CO₂ was approximately 42-50% with respect to the total number of N_2O_5 molecules taken up. The reaction of N_2O_5 with mineral dust and the subsequent production of gas phase HNO₃ leads to a decrease in $[NO_{*}]$ which may have a significant effect on global ozone. 15

1. Introduction

Dinitrogen pentoxide, N_2O_5 , is part of the family of active nitrogen, NO_y , owing to its equilibrium with NO_2 and NO_3 according to Reaction (1), which releases photochemically active NO_3 and NO_2 free radicals.

²⁰
$$N_2O_5 \leftrightarrow NO_2 + NO_3$$

Under atmospheric pressure and 298 K the thermal lifetime of N_2O_5 is approximately 20 s and is a strong function of temperature (Atkinson et al., 1997; Wängberg et al., 1997). The observation of the buildup of the concentration of nitric acid, HNO₃, in the polluted urban PBL during the night could only be explained by the heterogeneous

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conversion of N_2O_5 to HNO_3 according to Reaction (2) (Hjorth et al., 1987; Tuazon et al., 1983)

 $N_2O_5 + H_2O_{(ads)} \rightarrow 2HNO_3$

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The substrate for the heterogeneous Reaction (2) has not been unambiguously identified but is thought to be urban H₂O-containing aerosols or humid surfaces on the ground whose purpose is to make available adsorbed H₂O in order to enable Reaction (2).

Together with HNO₃ and NO₃, N₂O₅ contributes to the formation of particulate nitrate on the dust particles by surface processes in the troposphere (Zhang et al., 1994). These processes represent an important sink for nitrogen oxide species, with decreases of daytime NO_y levels reaching up to 60% in the presence of dust at a loading of about 1.8–11.5 µg m⁻³ corresponding to a particle surface area of (0.11–0.7)×10⁻⁶ cm² cm⁻³. During the nighttime, the conversion of NO₃ and N₂O₅ to particles dominates the overall nitrate formation accounting for 80% of total particulate nitrate formation while the heterogeneous hydrolysis pathway leading to HNO₃ accounts for only about 20% (Zhang et al., 1994).

In addition, the present study shows that gas phase HNO_3 is a potential reaction product formed by heterogeneous hydrolysis of N_2O_5 with adsorbed water present on mineral dust. Previous laboratory studies performed using a Knudsen cell flow reactor have extensively supported the reactivity of HNO_3 on $CaCO_3$, marble, mineral dust surrogates and metal oxides (Fenter et al., 1995; Frinak et al., 2004; Hanisch and Crowley, 2001a; Underwood et al., 2001).

Mineral aerosols provide reactive surfaces in the atmosphere where heterogeneous chemical reactions may take place. Dentener showed how the interaction of N_2O_5 ,

O₃ and HO₂ radicals with dust will affect the photochemical oxidant cycle, with ozone concentrations decreasing by up to 10% in and nearby the dust source areas (Dentener et al., 1996).

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Field observations, laboratory (Hanisch and Crowley, 2001a, b, c, d; Usher et al., 2003) and modelling studies have established the interaction of trace gases with mineral dust aerosol as well as the quantitative impact of the latter on the composition of the atmosphere. Box, regional and global scale models have shown the importance of dust on both the photochemical rates of oxidant formation as well as the loss of trace gases regarding atmospheric composition (Bauer et al., 2004; Bian and Zender, 2003; Dentener et al., 1996; Tabazadeh et al., 1998; Zhang et al., 1994).

The comparison of the global scale models of Bian and Zender with the one presented by Bauer et al. reveals significant quantitative differences of the effect of mineral dust on O_3 and several important trace gases such as HNO_3 , N_2O_5 and H_2O_2 . However, both studies agree that the direct interaction of O_3 with mineral dust is of

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- However, both studies agree that the direct interaction of O_3 with mineral dust is of minor importance, whereas the uptake of its precursor HNO₃ is responsible for most of the ozone decrease in the areas affected by dust. In the modelling studies of both Bian and Zender (2003) and Bauer et al. (2004) an uptake coefficient $\gamma_{N_2O_5}=10^{-3}$ at a relatively humidity (*RH*) of 30% has been used. In the modelling simulation performed
- by Bauer et al. the uptake of N_2O_5 resulted in a reduction of 11% of its mass in the gas phase whereas the simulations of Bian and Zender indicate a global reduction of 2%.

Despite the published results of field observations and modeling studies (Bauer et al., 2004; Bian and Zender, 2003; Bonasoni et al., 2004; de Reus et al., 2000), there are no reports on laboratory experiments that deal with the heterogeneous reactivity of N₂O₅ on mineral dust aerosol surrogates. Only a very recent laboratory study has been performed with N₂O₅ on Saharan Dust using a combination of Knudsen and Drifts cell (Seisel et al., 2005). We have therefore embarked on a laboratory program to measure some of the heterogeneous reactions involving N₂O₅ and NO₃ (Karagulian

²⁵ and Rossi, 2005) as relevant trace atmospheric gases. In the present work, we have used a Knudsen flow reactor in order to investigate the uptake and reaction of N_2O_5 on selected authentic mineral dust samples such as Kaolinite, Saharan Dust from Cape Verde Islands, Arizona test dust and natural limestone as well as on samples of pure CaCO₃. 5, 10369–10408, 2005

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2. Experimental set up and detection

All experiments were performed in a TEFLON^(R) coated Knudsen flow reactor operating in the molecular flow regime. This technique has been described in detail in the literature (Caloz et al., 1997). Briefly, N₂O₅ was introduced into the Knudsen flow reactor ⁵ from the gas handling system using a fine needle valve as a flow control device. An isolation plunger allows the separation of the reactive surface of interest from the reactor volume. The gases leave the Knudsen reactor through an escape orifice whose diameters (1, 4, 8, 14 mm) determine the residence time ($\tau_g = 1/k_{esc}$) and molecular concentration at a given flow rate $F_0^M = I_M \cdot C_{(M)}$, where I_M is the mass spectrometric ¹⁰ signal amplitude (*MS*) and $C_{(M)}$ is a calibration factor for the species *M* of interest

that depends on instrumental parameters. The concentration $[M]_{MS} = N/V_{cell} = \frac{F_0^M}{k_{esc} \cdot V_{cell}}$ is related to the flow of molecules leaving the reactor F_0^M . Mass spectrometry (*MS*) provides a direct measure of the flow rate F_0^M which is proportional to the concentration of a species escaping the Knudsen flow reactor. The rate constant for the effusive loss k_{esc} of N₂O₅ is given by the kinetic theory of gases and was measured for every used orifice. The characteristic parameters and relevant kinetic expressions used in this work are reported in Table 1.

The samples we have used in this study are the following: Kaolinite, poorly ordered (KGa-2, Warren County, Georgia, USA), CaCO₃ (Fluka), Arizona Medium Test ²⁰ Dust (Powder Technology Incorporated, Burnsville MN, USA), natural limestone (Transmat, La Sarraz, Switzerland) and Saharan Dust from deposits on Cape Verde Islands (SDCV).

Two kinds of sample holders were used: one consisted of a TEFLON[®] coated Pyrex holder of available sample surface of 19.6 cm², the other consisted of an internal reduction piece made out of DELRIN[®] leading to a sample surface of 4.9 cm². Both TEFLON[®] and DELRIN[®] did not show any reactivity towards N₂O₅ under the present

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experimental conditions.

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N₂O₅ was synthesized by the oxidation of NO₂ with excess ozone. The O₃/O₂ mixture at the outlet of the ozonator (Fisher 502) is passed through a P₂O₅ trap in order to eliminate residual moisture before being mixed with equally dried NO₂. The N₂O₅ is collected in a methanol/dry-ice bath at 195 K and is subsequently analyzed for purity by *MS*. Hydrolysis of N₂O₅ may occur on surfaces of the traps and of the line before admission into the Knudsen flow reactor generating HNO₃ as an impurity ranging from 10 to 15% as studied by *MS* at m/e 63, its molecular ion peak.

 N_2O_5 does not have a measurable parent and fragment peak at m/e 108 and 62, respectively, under the present experimental conditions; the most intense peaks are its fragment NO_2^+ at m/e 46 followed by the less intense fragment NO^+ at m/e 30. However, the HNO₃ impurity also contributes to the *MS* signal at m/e 46 and 30. Under the experimental conditions used HNO₃ has a detectable parent peak at m/e 63. Through a calibrated mass spectrum of pure HNO₃ we have accurately determined the effective contribution of HNO₃ at m/e 46 and 30 by using the fragmentation pat-

tern expressed as the ratios $f_{46} = \frac{I_0^{\frac{76(HNO_3)}{0}}}{I_0^{\frac{53(HNO_3)}{0}}} = 52 \pm 8 \text{ and } f_{30} = \frac{I_0^{\frac{30(HNO_3)}{0}}}{I_0^{\frac{63(HNO_3)}{0}}} = 33 \pm 4$. In the absence

of a substrate, f_{46} · $I_0^{63(\text{HNO}_3)}$ and f_{30} · $I_0^{63(\text{HNO}_3)}$ have been subtracted from the total *MS* signals I_0^{46} and I_0^{30} at m/e 46 and 30, respectively, in order to assign the remaining *MS* amplitude to the NO₂⁺ and NO⁺ fragments to N₂O₅: $I_0^{46(N_2O_5)} = I_0^{46} - f_{46} \cdot I_0^{63(\text{HNO}_3)}$ and $I_0^{30(N_2O_5)} = I_0^{30} - f_{30} \cdot I_0^{63(\text{HNO}_3)}$.

The ratio between the two most intense peaks of N₂O₅ therefore was $r = \frac{I_0^{46(N_2O_5)}}{I_0^{30(N_2O_5)}} = 1.4 \pm 0.2$. Mixtures of N₂O₅ and HNO₃ may thus be monitored using *MS* signal intensities at m/e 46, 30 and 63. However, in order to quantify N₂O₅ we chose its most intense peak corresponding to its fragment NO₂⁺ at m/e 46. The subscript 0

²⁵ and *r* refer to continuous gas uptake experiments in the absence and presence, respectively, of the solid sample.

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3. Uptake coefficient of N₂O₅ and identity of reaction products

When N₂O₅ is exposed to the sample, it is taken up and undergoes a heterogeneous reaction on the mineral dust surface which results in a decrease of the N₂O₅ concentration that is monitored using the *MS* signal $l_r^{46(N_2O_5)}$. As shown in previous studies on mineral dust, HNO₃ that is always present as an impurity, is taken up on the mineral dust surface without releasing any product that may contribute to the total *MS* signal l_r^{46} at m/e 46 (Hanisch and Crowley, 2001b). We therefore determined the rate constant k_{obs} for the disappearance of N₂O₅ following Eq. (E.1), assuming that the rate law is first order in N₂O₅:

$$k_{\rm obs} = (\frac{I_0^{46(N_2O_5)}}{I_r^{46(N_2O_5)}} - 1) \cdot k_{\rm esc}$$

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 $I_0^{46(N_2O_5)}$ and $I_r^{46(N_2O_5)}$ are the intensities of the NO₂⁺ fragment of N₂O₅ before and during heterogeneous reaction, respectively, and k_{esc} is the measured rate constant of effusion for N₂O₅ out of the flow reactor (see Table 1). In order to determine $I_r^{46(N_2O_5)}$, the raw *MS* signal at m/e 46 was corrected for HNO₃ generated through hydrolysis of N₂O₅ with water adsorbed on the substrate surface. Production of HNO₃ has in fact been observed at m/e 63 in every uptake experiment of N₂O₅. Figure 1 shows the raw *MS* signal at m/e 63 for a typical uptake experiment of N₂O₅ on CaCO₃. Therefore, we have corrected the *MS* signal of N₂O₅ at m/e 46 for the presence of HNO₃ according to Eq. (E.2) when the sample is exposed to N₂O₅:

$${}_{20} I_r^{46(N_2O_5)} = I_r^{46} - f \cdot I_r^{63(HNO_3)}$$
(E.2)

The ratio *r*, reported above, turned out to be the same before and during the exposure of the mineral dust sample to N₂O₅. Therefore, $l_r^{46(N_2O_5)}$ exclusively corresponds to N₂O₅ for this experiment after correction of the *MS* signal at m/e 46 for the contribution of HNO₃ desorbing from the mineral dust substrate.

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In the Knudsen flow reactor, reactant gas molecules either exit through an aperture into the mass spectrometer or are lost on the reactive surface. The net observed uptake coefficient for N₂O₅, γ_{obs} , is given by the following expression:

$$\gamma_{\rm obs} = \frac{k_{\rm obs}}{\omega}$$

⁵ Equation (E.3) is only valid if the rate law for uptake is first order in N₂O₅, where k_{obs} is the pseudo-first order rate constant at steady state conditions given in Eq. (E.1), ω is the collision frequency of the average molecule with the reactive surface area of the sample defined as ω = ^{c̄}/_{4V_{cell}} A_s, c̄ is the mean velocity of the gas-phase molecule, V_{cell} is the volume of the reactor and A_s is the surface area of the sample holder as will be justified below. In the following, we evaluate γ_{obs} at the initial and steady state values of the uptake rate leading to γ₀ and γ_{ss}, respectively.

4. Uptake of N₂O₅ on CaCO₃: results and discussion

Typical raw data from an uptake experiment of N_2O_5 on 510 mg of CaCO₃ are shown ¹⁵ in Fig. 1 using the 14 mm diameter-orifice. After a steady state flow of N_2O_5 has been established, the isolation plunger is lifted at t=92 s and the substrate is thus exposed to the N_2O_5 flow. *MS* signals of 18 (H_2O^+), 30 (NO^+), 44 (CO_2^+), 46 (NO_2^+), and 63 (HNO_3^+) were simultaneously monitored during the uptake. Because of the uptake of N_2O_5 on CaCO₃, the number of molecules exiting through the escape orifice into the

²⁰ *MS* immediately decreases which leads to a decrease of the *MS* signal $I_r^{46(N_2O_5)}$ at m/e 46. During the exposure of the sample to N₂O₅ both HNO₃ and CO₂ monitored at m/e 63 and 44, respectively, have been observed in the gas phase. As the exposure time increases, the *MS* signal at m/e 46 partially recovers, indicating a decrease in the rate of uptake that ultimately leads to steady state towards the end of the displayed uptake experiment. At *t*=1150 s the sample compartment is sealed by lowering the plunger 5, 10369–10408, 2005

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and the *MS* signal at m/e 46 approximately returns to its initial steady-state value. The slight decrease of $I_0^{46(N_2O_5)}$ over extended periods of time such as displayed if Fig. 1 can be explained by a slight decrease of the corresponding flow rate into the reactor.

- An ancillary experiment was performed in order to estimate the amount of water adsorbed on the CaCO₃ substrate surface after a given pumping time. In order to limit H₂O desorbing from the Pyrex sample holder it was replaced by a gold-coated all-metal sample holder. In this way most desorbing H₂O may be attributed to the mineral dust sample. Typically, a fresh sample of 1 g of CaCO₃ was pumped for 30 min at $T=298\pm 2$ K in the 14 mm-orifice reactor until the *MS* signal of H₂O at m/e 18 dropped
- ¹⁰ to the background level. Subsequently, the CaCO₃ sample was heated up to 470 K and the *MS* signal at m/e 18 recorded until it reached background as well indicating that no additional water desorbed at that temperature. The *MS* signal at m/e 18 was integrated in order to calculate the number of adsorbed water molecules H₂O_(ads) using the measured BET surface area of $3.7 \text{ m}^2 \text{ g}^{-1}$ for precipitated CaCO₃. A value of ap-¹⁵ proximately 3.0×10^{13} molecule cm⁻² has been found for the surface density of strongly
 - adsorbed H₂O_(ads) (Santschi, 2003).

When the CaCO₃ substrate is exposed to N_2O_5 at 92 s, we observed rapid formation of $CO_{2(g)}$ and $H_2O_{(g)}$ as displayed in Fig. 1 which were either generated in a chemical reaction or desorbing from a precursor state. In order to better understand the up-

take of N₂O₅ on solid CaCO₃ powder, we will briefly digress to the description of the chemical nature of a carbonate surface. From experimental and theoretical surface science studies, (De Leeuw and Parker, 1998; Kuriyavar et al., 2000; Stipp et al., 1994) there is clear evidence that under ambient conditions of pressure, temperature, and relative humidity, the surface of CaCO₃ is terminated by OH groups that persist even under ultrahigh vacuum conditions. The OH-terminated surface may be a result of the dissociative adsorption of water according to Reaction (3):

 $CaCO_3 + H_2O \rightarrow Ca(OH)(HCO_3)$

In a recent study (Al-Hosney and Grassian, 2004), the surface chemistry of CaCO₃ with

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trace atmospheric gases such as HNO_3 , SO_2 , HCOOH, and CH_3COOH was investigated using FTIR absorption spectrometry. This study has pinpointed adsorbed carbonic acid H_2CO_3 to be involved in the surface chemistry of $CaCO_3$ and was identified as a stable intermediate species on the $CaCO_3$ surface in the presence of H_2O vapor.

⁵ The vibrational spectrum of carbonic acid is well characterized by its C= \overline{O} stretching frequency at 1685 and 1705 cm⁻¹ corresponding to the adsorbed and condensed phase, respectively. In the following, adsorbed H₂CO₃ will occur as an intermediate in several instances.

Dissolution of CaCO₃ in the system H₂O-CO₂-CaCO₃ is controlled by three ratedetermining processes: the kinetics of dissolution at the mineral surface, mass transport by diffusion, and the slow kinetics of Reaction (4):

$$CO_{2(g)} + H_2O_{(ads)} \leftrightarrow H_2CO_{3(ads)} \leftrightarrow H_{(aq)}^+ + HCO_{3(aq)}^-$$
(4)

A theoretical model by Buhmann and Dreybrodt (1985) taking these processes into account predicts that, due to the slow kinetics of Reaction (4), precipitation rates on

¹⁵ the surface of CaCO₃ minerals critically depend on the ratio V/A of the volume V of the solution to the surface area A of the mineral in contact with it. They concluded that Reaction (4) is rate limiting (Dreybrodt et al., 1997).

Bicarbonate ion, HCO_3^- , may react with $CaCO_3$ to yield a surface intermediate that is proposed to be the active surface reactant for the heterogeneous reactions discussed below according to the well-known "Karst dissolution" mechanism (Dreybrodt et al., 1996):

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$$CaCO_{3(s)} + H^{+}_{(ads)} + HCO^{-}_{3(ads)} \rightarrow Ca(OH)(HCO_{3}) + CO_{2(g)}$$
(5)

When the $CaCO_3$ sample is exposed to N_2O_5 the surface intermediate may react as follows:

²⁵
$$N_2O_{5(g)} + Ca(OH)(HCO_3) \rightarrow Ca(NO_3)_2 + H_2O_{(s)} + CO_{2(g)}$$
 (6)

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and the net reaction resulting from Reactions (4), (5) and (6) will be:

 $N_2O_5 + CaCO_3 \rightarrow Ca(NO_3)_2 + CO_{2(g)}$

It is evident that H₂O_(g) and CO₂ resulting from Reaction (6) could be used again to generate additional Ca(OH)(HCO₃) according to Reactions (4) and (5) provided Re-⁵ action (5) is fast enough under flow reaction conditions. In this way H₂O is neither consumed nor generated and may therefore be viewed as a catalytic species.

On the time scale of the uptake experiments reported in Fig. 1 the calculated ratio CO_2/N_2O_5 of the product yields was 0.42 which is significantly smaller than 1.0 that is stoichiometrically expected according to Reaction (7) (see Table 2). The mass balance

¹⁰ between the adsorbed N_2O_5 and the reaction product CO_2 is therefore not satisfied. If Reactions (4) and (5) were fast and not rate limiting, the CO_2 yield could be 100%. We therefore have to consider another possible pathway for Reaction (6) that does not result in release of CO_2 :

 $N_2O_5 + Ca(OH)(HCO_3) \rightarrow Ca(NO_3)(HCO_3) + HNO_{3(a)}$

- In this case N₂O₅ may directly be converted into gas phase HNO₃ which is also observed at longer exposure times as displayed in Fig. 1. Reaction (8) may also help explain why the ratio CO₂/N₂O₅ differs from 1.0 as no CO₂ is released from Reaction (8). In addition, HNO₃ may also react with CaCO₃ as already observed by Hanisch and Crowley (2001). We routinely measure a yield of 51% of gas phase H₂O with respect to N₂O₅ consumed which is comparable to the CO₂ yield according to Eq. (6) and re-
- sults from the uptake of N_2O_5 on samples of different mass of powdered CaCO₃. In the wake of the observation of H₂O, CO₂ and HNO₃ we propose that Reactions (7) and (8) compete with each other. H₂O is formed in Reaction (6) on CaCO₃ that is in part preexisting as a surface intermediate Ca(OH)(HCO₃) from prior exposure to atmospheric CO₂ and H₂O following Reactions (4) and (5).

Owing to the fact that $CaCO_3$ has a specific surface area (BET) of 5.06 m²/g, the 510 mg sample from the experiment displayed in Fig. 1 has a total surface area of



(7)

(8)

 2.6×10^4 cm². N₂O₅ may be represented as a sphere with a projected surface area of 6.4×10^{-15} cm²/molecule or 1.56×10^{14} molecules cm⁻², assuming that N₂O₅ has a characteristic diameter of approximately 9 Å and a density of 2.93 g/cm³. After an exposure time of 1060 s 2.2×10^{18} molecules of N₂O₅ are taken up on 510 mg of CaCO₃ (Fig. 1) leading to a surface concentration of $\frac{2.2 \times 10^{18}}{2.6 \times 10^4} = 8.4 \times 10^{13}$ N₂O₅ molecule cm⁻² which corresponds to a coverage (θ) of approximately 54% based on the BET surface area after a reaction time of 1060 s. This assumes that at steady-state the total internal area of the powder sample is accessible to N_2O_5 in contrast to the initial uptake where only the surface layer is interacting with the gas. After N_2O_5 reacted on CaCO₃ powder for 1060s the substrate is apparently at steady state as displayed in Fig. 1. 10 This means that there is a sufficient number of intermediate species Ca(OH)(HCO₃) to allow Reactions (6) and (8) to occur. However, once reacted with N_2O_5 , the intermediate species cannot regenerate because Reactions (4) and (5) are too slow. Therefore, we observe an amount of $CO_{2(q)}$ smaller than the maximum yield of 100% at steady state conditions. On the other hand, N_2O_5 may react with adsorbed water $H_2O_{(ads)}$ that 15 still remains on the CaCO₃ substrate, thereby forming two molecules of nitric acid as

$$N_2O_{5(q)} + H_2O_{(ads)} \rightarrow 2HNO_{3(q)}$$

follows:

In this case solid CaCO₃ is just the support for the reactive adsorbed water $H_2O_{(ads)}$ and is not consumed in the chemical reaction. Initially, $HNO_{3(g)}$ is physically adsorbed on the surface to result in adsorbed HNO_3 which reacts with a surface OH-group and slowly forms surface nitrates and H_2O according to Reaction (10):

$$S-OH + HNO_{3(ads)} \rightarrow S-NO_3 + H_2O_{(ads,g)}$$
(10)

where S represents a surface site for physical adsorption. This mechanism has been
 proposed by Seisel et al. (2004) in a DRIFTS study of the heterogeneous reaction of HNO₃ on mineral dust where they observed the presence of free OH-groups located on the surface of mineral dust and the formation of surface nitrates.

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The fact that CO_2 is formed immediately after lifting the plunger (Fig. 1) whereas HNO₃ is formed later on after a certain delay clearly indicates that there are two competitive processes occurring during the reaction of N_2O_5 with $CaCO_3$. Immediately after the exposure of substrate, N_2O_5 reacts with the intermediate species $Ca(OH)(HCO_3)$ on the $CaCO_3$ sample (Reactions (6) and (8)). At the same time heterogeneous hydrolysis of N_2O_5 may occur due to the presence of $H_2O_{(ads)}$ on the substrate (Reaction (9)). However, the presence of a sufficient quantity of the intermediate formed in Reaction (5) may make Reactions (7) and (8) predominant at first with respect to Reaction (9). This explains the initial rapid formation of $CO_{2(g)}$ and the absence of any measurable trace of gas phase HNO₃ at t=92 s in Fig. 1. Once the surface intermediate starts to be consumed owing to the initial fast reaction with N_2O_5 the sample starts saturating and Reaction (9) becomes predominant compared to Reactions (7) and (8) which would explain the delayed formation of HNO₃ displayed in Fig. 1.

After having performed the uptake experiment shown in Fig. 1 on a substrate of CaCO₃, we sealed the sample and halted the inlet flow of N₂O₅ for approximately 10 min. Subsequently, we lifted the plunger again and observed small amounts of desorbing CO₂ (curve (a), Fig. 2) as well as a *MS* signal at m/e 30 without any measurable *MS* signal at m/e 46 (curve (b), Fig. 2). The total yield of desorbed CO₂ only represented 5% of N₂O₅ taken up on the substrate of CaCO₃ during the reaction time of 1058 s (Fig. 1). The small intensity of the *MS* signal at m/e 30 strongly suggests formation of NO as a decomposition product of N₂O₅.

Uptake experiments of N₂O₅ on 0.58 g of CaCO₃ powder were carried out at smaller orifice sizes, thus increased residence time τ_g at constant flow of N₂O₅ of 2.3×10¹⁵ molecule s⁻¹. The values of γ_0 and γ_{ss} decrease with increasing values of the residence time as displayed in Fig. 3 and Table 3. The strong dependence of γ_{ss} on τ_g suggests that the mechanism of N₂O₅ uptake is complex and does not correspond to a simple first order uptake reaction. These observations indicate that the reactivity of N₂O₅ on CaCO₃ decreases for long residence times as the heterogeneous reaction rate not only depends on the gas phase concentration but apparently also on interme-

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diates whose surface concentration depend on the extent of reaction that scales with τ_g akin to an effective second order reaction. Apparently, the active surface intermediate Ca(OH)(HCO₃) cannot regenerate sufficiently fast according to Reaction (5) so as to maintain the rate limiting Reaction (6) at higher partial pressure, that is at longer ⁵ residence time τ_a .

In order to unravel whether or not the effective available surface area for uptake is influenced by the internal surface area formed by interstitial voids between individual dust particles, the mass dependence of the N_2O_5 uptake on CaCO₃ was investigated in the Knudsen flow reactor at ambient temperature and at $[N_2O_5] = (4.0 \pm 1.0) \times 10^{11} \text{ cm}^{-3}$.

- ¹⁰ The mass of CaCO₃ which is a non-porous material ranged from 0.11 g to 1.8 g and the results are shown in Fig. 4. Table 4 reports values of γ_{ss} and γ_0 using the geometric surface area A_s=19.6 cm². The steady state and initial uptake coefficient γ_{ss} and γ_0 , respectively, of N₂O₅ were found to increase linearly at low masses of CaCO₃. Samples below 0.33 g were considered to be part of this linear mass-dependent regime.
- ¹⁵ Increasing the sample mass further had a negligible effect on the amount of adsorbed N_2O_5 because not the entire sample surface is apparently available for N_2O_5 adsorption. This maximum value is attributed to the inability of N_2O_5 to penetrate through all layers of the sample within the residence time of N_2O_5 in the gas phase, thus resulting in a constant number of molecules taken up despite the increasing sample mass. The limiting γ_{ss} value therefore represents the maximum amount of N_2O_5 able to interact
 - with $CaCO_3$ powder within the N₂O₅ residence time.

In order to better define the saturation behaviour of N_2O_5 on the substrate further uptake experiments were performed on CaCO₃ employing a pulsed valve to admit N_2O_5 into the reactor. The pulsed-valve experiments were carried out by using a solenoid valve, through which the gas was introduced in pulses with a duration of 5*MS* and at a dose of 2.5×10^{15} molecules per pulse (Caloz et al., 1997) corresponding to $[N_2O_5]_0=1.3 \times 10^{12}$ cm⁻³. We determined the total number of injected molecules per pulse and the value of k_{esc} in a reference experiment which was obtained by fitting the decaying *MS* signal to an exponential decay in the absence of reaction thereby

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obtaining k_{esc} . The "reactive pulse" was obtained by repeating the same pulsed experiment with the sample exposed to the gas. The total observed exponential decay in the presence of a reactive substrate was thus characterized by a new rate constant, k_{dec} , given by $k_{dec} = k_{obs} + k_{esc}$ that is used to determine the number of interest given by $\gamma_{obs} = k_{obs}/\omega$. The pulsed valve uptake experiments on CaCO₃ showed that there is a modest mass-dependence of the measured uptake γ_{obs} by approximately a factor of three to four. For a dose of about 2.5×10^{15} molecules per pulse we observed an unexpected increase of the uptake γ_{obs} with increasing mass of CaCO₃ as reported for steady state experiments in Table 4 displayed in Fig. 4. The increase of γ_{obs} saturates at large sample mass because the number of layers exceeds the depth of diffusion of gas into the internal voids. For all series of pulses the measurements are identical, within the uncertainty, to those for steady state experiments.

Similar behavior has been observed in a recent study carried out by Seisel et al. (2005). In that study uptake experiments of N₂O₅ on Saharan Dust obtained ¹⁵ from pulsed and steady state experiments were in good agreement, indicating that the steady state uptake coefficients are not influenced by saturation effects under their experimental conditions.

In order to determine the number of layers, the total volume of the powder was calculated from its true density (ρ_t =2.93 g/cm³) and the mass of the sample spread out across the geometric area of the sample holder. The number of formal layers calculated for an average sample grain diameter of 3.5 μ m from the average particle size and the height of the sample is reported in Table 4. The typical grain diameter of 3.5 μ m has been obtained from the manufacturer's specifications of the used CaCO₃ powder.

²⁵ Based on published microscopic images, it is reasonable to take into account a grain size diameter that is larger than $3.5 \,\mu$ m as suggested by electron microscopy (SEM) displayed in Fig. 5. Figure 4 shows that a mass of $0.33 \,\text{g}$ corresponds to one nominal layer of 57 μ m diameter CaCO₃ spread out over the geometric surface of the sample holder (19.6 cm²). Thus, one nominal layer of CaCO₃ will contain 330 mg of closely 5, 10369–10408, 2005

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packed spheres of "effective" grain diameter of 57 μ m knowing full well that the sample in reality is multidisperse and structurally heterogeneous. Therefore, the linear mass-dependent portion of γ_{ss} vs. mass for masses less that 330 mg corresponds to a sample holder that is partially covered with 57 μ m diameter CaCO₃ particles.

- ⁵ The use of the BET surface area and the application of the pore diffusion theory (Keyser et al., 1991) yields $\gamma_{pd,ss} = (7.4 \pm 1.7) \times 10^{-6}$ for steady state experiments using a grain diameter for CaCO₃ of 3.5 μ m. This value is lower by a factor of 10³ compared to $\gamma_{ss} = (2.8 \pm 0.5) \times 10^{-2}$ and $\gamma_0 = 0.16 \pm 0.02$ which were calculated on the basis of the geometrical surface area of the sample as displayed in Table 4. The use of the pore diffusion theory substantially underestimates the true uptake coefficient so that it may
- be interpreted as a lower limit for γ whereas γ_{ss} and γ_0 based on the geometrical surface area may be regarded as an upper limit, albeit close to the true value of γ . We take the fact that γ_0 and γ_{obs} obtained from pulsed valve experiments are identical within experimental uncertainty (Fig. 4) as evidence that N₂O₅ only interacts with the
- ¹⁵ surface layer of the solid sample during a typical pulse decay of 300 ms. We do not rule out that N₂O₅ will interact with deeper layers of the sample to some extent under steady state conditions. However, it is unlikely that the true value of γ will drop by more than three orders of magnitude during a steady state uptake experiment typically lasting tens of minutes. We will therefore not further discuss $\gamma_{pd,ss}$ also in view of the analysis of field data discussed below that are consistent with γ_{obs} and γ_0 values based
- on the geometric surface area.

5. Uptake of N₂O₅ on mineral dust substrates: results and discussion

Table 5 reports results on experiments performed on 1 g samples of surrogate mineral dust powder at a high initial concentration of $[N_2O_5]_0=(3.8\pm1.0)\times10^{12}$ cm⁻³. The steady-state uptake coefficients γ_{ss} of N_2O_5 range from $(2.2\pm0.6)\times10^{-3}$ for natural limestone to $(5.9\pm1.6)\times10^{-2}$ for Saharan Dust using the geometric surface area. At 5, 10369–10408, 2005

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a lower initial concentration of $[N_2O_5]_0 = (4.0 \pm 1.0) \times 10^{11} \text{ cm}^{-3}$ uptake experiments performed on 0.3 g have revealed larger values of γ_{ss} ranging from $(3.5\pm1.1)\times10^{-2}$ for CaCO₃ to 0.2±0.05 for Saharan Dust. We also report the observed initial uptake coefficients γ_0 for N₂O₅ on all the samples of mineral dust at low and high values of $[N_2O_5]$. For a concentration of $[N_2O_5]_0 = (3.8 \pm 0.5) \times 10^{12} \text{ cm}^{-3}$ the γ_0 values range from 5 $(6.4\pm1.9)\times10^{-3}$ for Arizona test dust to $(9.0\pm2.6)\times10^{-2}$ for Saharan Dust, whereas at $[N_2O_5]_0 = (4.0 \pm 1.0) \times 10^{11} \text{ cm}^{-3}$ the γ_0 values range from 0.12 ± 0.04 for CaCO₃ to 0.43±0.13 for natural limestone. For samples such as Saharan Dust, CaCO₃ and Arizona Test Dust values of γ_{ss} and γ_0 decrease between a factor of 3 and 7 from low to high $[N_2O_5]_0$. As indicated in Table 5, both the values γ_{ss} and γ_0 for CaCO₃ decrease 10 only by a factor 3.5 from low to high $[N_2O_5]$. On the other hand a particular case is represented by natural limestone which showed a decrease of γ_{ss} and γ_0 by a factor of 20 and 40, respectively, when increasing $[N_2O_5]_0$. CaCO₃ showed values of γ_{ss} and γ_0 higher by a factor of 3 with respect to natural limestone at high $[N_2O_5]_0$. This difference is reversed by the same amount for low $[N_2O_5]_0$ with γ of natural limestone 15 being highly sensitive to saturation by $[N_2O_5]$. Natural limestone is a sedimentary rock containing 97% CaCO₃ by weight and a small percentage of metal oxides (1.9% of SiO_2 , 0.5% of Al₂O₃, 0.3% of Fe₂O₃, and 0.3% of MgO) that may be responsible for the difference in the kinetic properties of CaCO₃ and natural limestone. In recent work Krueger et al. (2005) showed that dust containing calcium is very reactive with respect 20 to the uptake of nitric acid. However, because of differences in mineralogy of single

dust-particles, not all of the calcium-containing particles react similarly.

It is important to note that for Kaolinite γ_{ss} and γ_0 are independent of $[N_2O_5]_0$ over the investigated range. Typical raw data from an uptake experiment of N_2O_5 on 1 g of Kaolinite and Saharan Dust are shown in Figs. 6 and 7. In this series of experiments we did not succeed to saturate the samples during the present observation period. Uptake experiments of N_2O_5 on 0.2 g of Kaolinite powder were carried out by varying the initial flow of N_2O_5 into the reactor (Table 6). Figure 8 displays data for the 8 mm orifice corresponding to a residence time τ_g of 1.32 s for a variation of $[N_2O_5]_0$ by a factor

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9.5 that is between $(4.0\pm1.0)\times10^{11}$ cm⁻³ and $(3.8\pm0.5)\times10^{12}$ cm⁻³. Figure 8 shows that both γ_0 and γ_{ss} remain constant at 0.19 ± 0.05 and $(2.3\pm0.6)\times10^{-2}$, respectively, with increasing $[N_2O_5]$ and by changing the gas residence time τ_g . From this series of measurements it is evident that γ_{ss} follows a pseudo first order rate law in N_2O_5 in contrast to the other substrates which showed a decreasing trend from low to high $[N_2O_5]$ (see Fig. 3 and Table 5).

In recent work the uptake of N₂O₅ on Saharan Dust was found to be independent of $[N_2O_5]$ (Seisel et al., 2005). An initial uptake $\gamma_0 = (8.0 \pm 0.3) \times 10^{-2}$ was found, whereas the steady state value $\gamma_{ss} = (1.3 \pm 0.3) \times 10^{-2}$ was lower by a factor of five with respect to the present results at $[N_2O_5]_0 = (3.8 \pm 0.5) \times 10^{12} \text{ cm}^{-3}$.

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We have observed delayed production of HNO₃ upon uptake of N₂O₅ for every sample investigated. Gas phase HNO₃ formation may be due to the heterogeneous hydrolysis of N₂O₅ according to Reaction (9). In order to understand the gas phase production of HNO₃ we want to stress that all the investigated samples have a non negligible amount of adsorbed water available on the substrate surface. The quantities of H₂O_(ads) that still remain on the different mineral dust substrates at our experimental conditions as reported in Table 5 and were measured by gravimetric measurements. The hygroscopic properties of mineral aerosol samples have been examined in recent work (Gustafsson et al., 2005) which showed significant water adsorption on Arizona

- Test Dust compared to CaCO₃. In Table 5 we also report the percentage of gas phase HNO₃ produced with respect to N₂O₅ taken up during a reaction time of 200 s. At [N₂O₅]=(3.8±0.5)×10¹² cm⁻³, Arizona Test Dust and Kaolinite turned out to be the samples to produce the largest amount of gas phase HNO₃, that is 72% and 30%, respectively, with respect to N₂O₅ taken up. On the other hand, Saharan Dust and CaCO₃ have been the samples with a lower yield of absolute HNO₃ produced, namely 5% and 6%, respectively. At [N₂O₅]=(4.0±1.0)×10¹¹ cm⁻³ we obtained lower yields of
- 5% and 6%, respectively. At $[N_2O_5] = (4.0 \pm 1.0) \times 10^{-1}$ cm⁻² we obtained lower yields of gas phase HNO₃ compared to $[N_2O_5]$ ten times higher. In this case we may correlate the low yield of HNO₃ to the large rate of uptake of N_2O_5 and of HNO₃ on mineral dust.

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Adsorbed HNO₃ may then form salts such as $Ca(NO_3)_2$, $Fe(NO_3)_3$ and surface nitrates as shown in other experimental studies (Börensen et al., 2000; Goodman et al., 2001; Vogt and Finlayson-Pitts, 1994). A recent study on the reactivity of gaseous HNO₃ on atmospheric mineral dust samples reported values for the uptake of HNO₃ on CaCO₃,

⁵ Saharan Dust and Arizona Test Dust (Hanisch and Crowley, 2001a). In that work, a value of $\gamma_0=0.11$ was determined for Saharan Dust at $[HNO_3]=(5.6\pm0.4)\times10^{11}$ cm⁻³, 0.14 for CaCO₃ and 6.6×10^{-2} for Arizona Test Dust. These results are consistent with the trend of the present yields of HNO₃ measured for nominally the same mineral dust samples. Whenever the uptake coefficient of HNO₃ on the mineral substrates was low such as for Arizona Test Dust and Kaolinite, we find increased amounts of HNO₃ in the gas phase. Conversely, the reverse is true for samples that rapidly take up HNO₃ as for CaCO₃ and Saharan Dust (Börensen et al., 2000).

6. Conclusions and atmospheric implications

We have shown in this work that N_2O_5 undergoes a heterogeneous reaction with surrogate substrates of mineral dust aerosol at $T=298\pm 2$ K. The measured uptake coefficient showed different values for high and low N_2O_5 concentrations with the least differences for Saharan Dust. These γ values are generally larger than the ones used in a recent global modeling simulation of heterogeneous chemistry on mineral dust aerosol at dry conditions (Bauer et al., 2004) where $\gamma=3.0\times10^{-3}$ for N_2O_5 has been used. The γ values resulting from the present measurements are larger by at least a factor of 10. Therefore, the uptake of N_2O_5 on mineral dust aerosols may potentially have a greater influence on the reduction of the global ozone concentration compared to the estimated value of 0.7% (Bauer et al., 2004).

The measured uptake coefficient γ of N₂O₅ on sulfuric acid aerosols was reported to ²⁵ lie within the range 0.06–0.12 at a temperature between 230 and 300 K (Hanson and Lovejoy, 1994). Other measurements reported γ values of 0.05 on aqueous surfaces over a temperature range from 282 to 294 K (van Doren et al., 1990). To our knowledge

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no reaction probabilities of N_2O_5 on mineral aerosol have been determined to date. In a recent numerical modeling study the interaction of N_2O_5 (γ =0.1), O_3 and HO₂ radicals with dust resulted in a decrease of tropospheric ozone of up to 10% near the dust source areas (Dentener et al., 1996).

- ⁵ The photolytic rate of NO₃ (J(NO₃)=0.2 s⁻¹) is too fast to allow its recombination with NO₂ to N₂O₅ during daylight. Therefore, the heterogeneous chemistry of N₂O₅ is important only at night-time. The heterogeneous reaction of N₂O₅ is most effective during the night when *RH* is at a maximum in the boundary layer. Thus, under these conditions, dust particles are likely to contain significant quantities of adsorbed water and the assumed high values of γ seem justified (Dentener and Crutzen, 1993). In a recent global modeling study γ =0.02 (*RH*=70%) and γ =3.0×10⁻³ (*RH*=30%) for humid and dry conditions have been used as upper and lower limits, respectively (Bauer et al., 2004). The modeling results show that when applying the high value for the uptake coefficient 0.8% of the global ozone mass is removed by uptake of N₂O₅ on
- ¹⁵ aerosols. The observed reaction products of the heterogeneous reaction of N₂O₅ with mineral dust, mainly HNO₃, may also have an influence on the oxidizing potential of the atmosphere as well as on the atmospheric ozone balance. Previous laboratory work (Goodman et al., 2000; Hanisch and Crowley, 2001a; Seisel et al., 2004) has shown the importance of the reactivity of HNO₃ on mineral dust substrates. In addition, mod ²⁰ eling studies have quantitatively shown decreases in ozone concentration close to the
- area of HNO_3 destruction (Bauer et al., 2004).

The Saharan dust sample, from Cape Verde (SDCV), that we have used is mineralogically representative of atmospheric dust aerosol. Its composition has been described in the literature (Coudé-gaussen et al., 1994) and closely simulates atmo-

²⁵ spheric particles of crustal origin (Desboeufs et al., 1999). The clay fraction (<2 μ m) of dust from Cape Verde shows a kaolinite-illite-chlorite assemblage which is typical for central Saharan Dust. In the free troposphere mineral dust aerosol of a size less than 2 μ m have a settling velocity of approximately 50 cm h⁻¹ (Seinfeld and Pandis, 1998). Therefore it can remain in the atmosphere for several days, travel long distances and 5, 10369–10408, 2005

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undergo heterogeneous reactions with trace gases.

The loss rate constant (k_{het}^{M}) due to heterogeneous uptake of a gas species M onto small aerosol particles is given by $k_{het}^{M} = \frac{\gamma A \bar{c}}{4}$ if the rate is not limited by diffusion, where γ is the uptake coefficient of M and is a function of the mineral dust aerosol composition, 5 A is the surface area density of the dust aerosol and \bar{c} is the mean molecular speed of M. Assuming a surface area density for Saharan Dust of about 1.5×10^{-6} cm² cm⁻³ (de Reus et al., 2000), we estimate $k_{het}^{N_2O_5} = 1.76 \times 10^{-3} \text{ s}^{-1}$ ($\tau_{het}^{N_2O_5} = 9.5 \text{ min}$) for N_2O_5 based on $\gamma = 0.2$ for Saharan dust at $[N_2O_5] < (4.0 \pm 1.0) \times 10^{11} \text{ cm}^{-3}$ (16 ppb). From the difference between the N_2O_5 formation and loss in the presence of the equilibrium with NO₃ we obtain the expression for the N_2O_5 steady state lifetime given in Eq. (11) (Brown et al., 2003):

$$\tau_{ss}^{N_2O_5} = \left(k_{het}^{N_2O_5} + \frac{k_{het}^{NO_3}k_1(T)}{k_{-1}[NO_2]} \right)^{-1}$$
(11)

 $k_1(T)$ and k_{-1} are the rate constant for thermal decomposition of N₂O₅ and its inverse at atmospheric pressure according to equilibrium (11), respectively. From recent work on the NO₃ heterogeneous reaction on mineral dust we have obtained γ =0.2 which leads to $k_{het}^{NO_3}$ =2.4×10⁻³ s⁻¹ ($\tau_{het}^{NO_3}$ =7 min) for NO₃ (Karagulian and Rossi, 2005). At *T*=273 K $k_1(273 \text{ K})$ =3.1×10⁻³ s⁻¹(τ =5 min) (Atkinson et al., 1997) and equilibrium (1) is shifted to the left. For [NO₂] a typical value is 10 ppb in polluted air so that the pseudo first-order recombination rate constant k_{-1} [NO₂] is 0.48 s⁻¹ and is thus much larger than the rate constant for heterogeneous loss $k_{het}^{NO_3}$ =2.4×10⁻³ s⁻¹. Therefore, the second term of expression (11) will be negligible and the steady state lifetime $\tau_{ss}^{N_2O_5}$ of N₂O₅ will be determined by its heterogeneous loss rate constant $k_{het}^{N_2O_5}$. At *T*=293 K equilibrium (1) is shifted to the right with $k_1(293 \text{ K})$ =4.6×10⁻² s⁻¹ (τ =20 s) which is an order of magnitude larger than at *T*=273 K. Therefore, both terms

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in Eq. (11) will be of comparable magnitude which will decrease the steady state life-time of N_2O_5 with respect to 273 K somewhat.

The calculated overall steady state lifetime for N₂O₅ at 293 K ($\tau_{ss}^{N_2O_5}$ =8.5 min) is significantly larger than the thermal dissociation lifetime (τ =20 s) of N₂O₅. At the same surface area concentration for dust aerosol of 1.5×10⁻⁶ cm² cm⁻³, the heterogeneous loss rate k_{het}^{hydr} due to hydrolysis (Reaction (2)) is 2.25×10⁻⁴ s⁻¹ corresponding to a lifetime of 74 min. This means that some N₂O₅ may be irreversibly converted to HNO₃ during the night by hydrolysis. During January 2004, Wood et al. (2005) performed in situ measurement of N₂O₅ in Contra Costa Country, California, and derived a steady state lifetime for N₂O₅ that ranged from 5 to 30 min at [N₂O₅]=200 ppt (5.0×10⁹ cm⁻³) in the temperature range 275–285 K which is comparable to $\tau_{ss}^{N_2O_5}$ calculated above. The measured lifetime for N₂O₅ measured in the present work.

Acknowledgements. We gratefully acknowledge OFES for funding this work in the context of the EU subproject MINATROC which is part of the EU Environment and Climate Program.

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 Table 1. Characteristic parameters and relevant kinetic expressions.

Definition	Value
Reactor volume, V	2000 cm ³
Reactor surface area, A_r	1500 cm ³
Sample surface area, A_s	19.5 cm ² (TEFLON holder), 4.5 cm ² (DELRIN holder)
Collision frequency ω for N ₂ O ₅ with A _s (M=108)	$\omega = 1.81(T/M)^{1/2}A_s$
Orifice diameters (nominal)	1, 4, 8 and 14 mm
Rate constant of effusion	$k_{\rm esc} ({\rm s}^{-1})$
$k_{\rm esc}$ (Experimentally determined values	$0.02(T/M)^{1/2} s^{-1}$ for 1 mm orifice
for orifice diameters of nominal diameter)	$0.25(T/M)^{1/2} s^{-1}$ for 4 mm orifice
	$0.8(T/M)^{1/2} \text{s}^{-1}$ for 8 mm orifice
	$1.9(T/M)^{1/2} \text{s}^{-1}$ for 14 mm orifice
Observed rate constant for heterogeneous reaction from steady state experiments, S_{I} = initial <i>MS</i> -signal; S_{f} = final <i>MS</i> -signal	$k_{\rm obs} = (S_i/S_f - 1)k_{\rm esc}$
Measured uptake coefficient	$\gamma_{\rm obs} = k_{\rm obs} / \omega$

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Table	2.	Uptake	experiments	with	N_2O_5	on	CaCO ₃	as	а	function	of	sample	mass	at
$[N_2O_5]_0$)=(4	.0±1.0)×	$(10^{11} \mathrm{cm}^{-3})$.											

CaCO ₃ Mass in g	γ ₀	γ_{ss}	Yield of HNO ₃ ^c	Yield of CO ₂
0.33 ^a	0.16±0.03	$(2.0\pm0.6)\times10^{-2}$	5.2%	47.8%
0.51 ^{a,b}	0.2 ± 0.05	(2.2±0.5)×10 ⁻²	5.4%	42.4%
0.58 ^a	0.18±0.05	(2.3±0.5)×10 ⁻²	-	-
0.73 ^a	0.22 ± 0.04	(2.4±0.4)×10 ⁻²	5.4%	50%
2 ^a	0.18±0.025	(1.6±1.6)×10 ⁻²	5.4%	50%

^a Uptake measurements performed with a surface sample area A_s =19.6 cm².

^b Uptake experiment displayed in Fig. 1.

 $^{\rm c}$ The yield is given as a percentage with respect to the total number of molecules of $\rm N_2O_5$ taken up during a reaction time of 550 s.

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Table 3. Relationship between γ_0 and γ_{ss} at different residence times τ_g (s⁻¹) of N₂O₅ interacting with 580 mg of CaCO₃ powder for data plotted in Fig. 2, surface sample area A_s =19.6 cm²; $F_0^{N_2O_5}$ =2.3×10¹⁵ molecule s⁻¹.

Orifice (mm)	γ ₀	γ_{ss}	$ au_g$ (s)
14	$(1.6\pm0.13)\times10^{-1}$	$(6.0\pm1.2)\times10^{-2}$	0.34
8	(8.5±0.8)×10 ⁻²	(3.0±0.7)×10 ⁻²	0.81
4	(3.4±1.0)×10 ⁻²	$(1.8 \pm 1.1) \times 10^{-2}$	2.72
1	(1.1±1.0)×10 ⁻²	$(1.0\pm1.0)\times10^{-2}$	30

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Mass (g)	γ_{ss}	γ_0	$\gamma_{0(\text{pulsed valve})}$	^a Number of	^b Number of	
			u /	formal layers	nominal layers	Title
0.11	$(8.0\pm5.0)\times10^{-3}$	$(5.5\pm2.0)\times10^{-2}$	$(4.7 \pm 1.5) \times 10^{-2}$	5	0.3	Abstract
0.15	(1.5±0.5)×10 ⁻²	(6.4±3.0)×10 ⁻²		7	0.5	
0.33	$(2.0\pm0.6)\times10^{-2}$	0.16±0.03	0.13±0.018	16	1	Conclusions
0.51	(2.2±0.5)×10 ⁻²	0.2±0.02		25	1.5	Tablas
0.58	$(2.3\pm0.5)\times10^{-2}$	0.18±0.02		30	1.8	Tables
0.73	$(2.4\pm0.5)\times10^{-2}$	0.22±0.02	0.21±0.016	36	2.2	• •
1.0	$(2.7\pm0.4)\times10^{-2}$	0.15±0.025		50	3	14
1.3	$(2.8\pm0.5)\times10^{-2}$	0.16±0.024		64	4	•
1.6	$(2.8\pm0.5)\times10^{-2}$	0.15±0.024	0.15±0.015	80	5	
1.8	$(2.8\pm0.5)\times10^{-2}$	0.17±0.03		90	5.5	Back

Effective diameter of ^a $3.5 \,\mu$ m and ^b $57 \,\mu$ m.

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Table 4. Summary of uptake experiments with N₂O₅ on CaCO₃ as a function of sample mass

 $([N_2O_5]=(4.0\pm1.0)\times10^{11} \text{ cm}^{-3}, \text{ orifice diameter} = 14 \text{ mm}, A_s=19.6 \text{ cm}^2).$

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Table 5. Summary of uptake experiments of N₂O₅ on mineral dust samples: initial (γ_0) and steady state (γ_{ss}) uptake coefficients.

Mineral dust sample	^a _{Y0}	$^{a}\gamma_{ss}$	^a HNO ₃	$H_2O_{(ads)} [mg g^{-1}]$
Kaolinite	0.14±0.04	$(2.2\pm0.6)\times10^{-2}$	30%	23
Natural Limestone	(1.1±0.3)×10 ⁻²	(2.2±0.6)×10 ⁻³	18%	7
Arizona Test Dust	(6.4±1.9)×10 ⁻²	$(1.6\pm0.4)\times10^{-2}$	72%	22
CaCO ₃	(3.3±1.0)×10 ⁻²	(6.2±1.8)×10 ⁻³	5%	4
Saharan Dust	(9.0±2.6)×10 ⁻²	$(5.9 \pm 1.6) \times 10^{-2}$	6%	20
	^b Y ₀	${}^{b} \gamma_{ss}$	^b HNO ₃	
Kaolinite	(0.16±0.04)	$(2.1\pm0.6)\times10^{-2}$	17%	
Natural Limestone	(0.43±0.13)	(4.3±1.3)×10 ⁻²	12%	
Arizona Test Dust	(0.2±0.06)	(0.11±0.03)	20%	
CaCO ₃	(0.12±0.04)	(2.1±0.6)×10 ⁻²	5%	
Saharan Dust	(0.3±0.08)	0.2 ± 0.05	4%	
	Mineral dust sample Kaolinite Natural Limestone Arizona Test Dust CaCO ₃ Saharan Dust Kaolinite Natural Limestone Arizona Test Dust CaCO ₃ Saharan Dust	$\begin{array}{llllllllllllllllllllllllllllllllllll$	$\begin{array}{llllllllllllllllllllllllllllllllllll$	$\begin{array}{c cccc} \mbox{Mineral dust sample} & \begin{tabular}{ c c c c } & \begin{tabular}{ c c } & t$

Uptake experiments were performed at ^a $[N_2O_5]_0 = (3.8\pm0.5)\times10^{12} \text{ cm}^{-3}$ using 1 g of sample powder for $A_s = 19.6 \text{ cm}^2$; ^b $[N_2O_5]_0 = (4.0\pm1.0)\times10^{11} \text{ cm}^{-3}$ using 300 mg of sample powder for $A_s = 4.9 \text{ cm}^2$. The yield of HNO₃ is given as a percentage with respect to the total molecules of N_2O_5 taken up after a given reaction time of 200 s at orifice diameter of 8 mm.

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Table 6. Summary of uptake experiments of N₂O₅ on 200 mg Kaolinite for data plotted in Fig. 8: initial (γ_0) and steady state (γ_{ss}) uptake coefficients (A_s =4.9 cm²).

$[N_2O_5]$ molecules cm ⁻³	γ ₀	γ_{ss}
$a(4.0\pm1.0)\times10^{11}$	0.16±0.04	$(2.1\pm0.6)\times10^{-2}$
^b (6.2±1.5)×10 ¹¹	0.23 ± 0.06	$(2.6\pm0.5)\times10^{-2}$
$a(9.0\pm0.5)\times10^{11}$	0.22 ± 0.06	$(2.2\pm0.5)\times10^{-2}$
^b (2.1±0.5)×10 ¹²	0.18 ± 0.05	$(2.4\pm0.7)\times10^{-2}$
$a(3.8\pm0.5)\times10^{12}$	0.16 ± 0.04	$(2.2\pm0.6)\times10^{-2}$

^a Orifice diameter = 8 mm; ^b Orifice diameter = 4 mm.

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Fig. 1. Typical N₂O₅ uptake experiment on a sample of 510 mg of CaCO₃. Curves (a), (b), (c), (d) and (e) correspond to the raw *MS* signals monitored at m/e 46, m/e 30, m/e 44, m/e 63 and m/e 18, respectively, using an orifice diameter of 14 mm, A_s =19.6 cm² and [N₂O₅]₀=(4.0±1.0)×10¹¹ cm⁻³.



Fig. 2. Desorption of CO₂ and NO after N₂O₅ uptake on a sample of 510 mg CaCO₃ displayed in Fig. 1. Curves (a) and (b) correspond to the raw *MS* signals monitored at m/e 44 and m/e 30, respectively, at an orifice diameter of 14 mm and A_s =19.6 cm².

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Fig. 3. Dependence of γ_0 (full squares) and γ_{ss} (open circles) for the uptake of N₂O₅ on 580 mg CaCO₃ powder on orifice size: A_s is 4.9 cm², $F_0^{N_2O_5}$ =2.3×10¹⁵ molecule s⁻¹.

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Fig. 4. Dependence of the uptake coefficient γ_0 (open squares) and γ_{ss} (full circles) on sample mass at $[N_2O_5]_0 = (4.0 \pm 1.0) \times 10^{11} \text{ cm}^{-3}$ at an orifice diameter of 14 mm for the uptake of N_2O_5 on CaCO₃ powder ($A_s = 19.6 \text{ cm}^2$). Full triangles represent pulsed valve experiments carried out at the same experimental conditions.



Fig. 5. SEM-image of CaCO₃ powder sample used in this work.

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1.6 reaction on 1.4 իկոու MS Signal (Volt) 1.2 1.0 判例师 0.8 (a) 0.6 0.4 (c) x 10 0.2 • (d)/10 0.0 1600 1800 2000 2200 time (s)

Fig. 6. Typical N₂O₅ uptake experiment on a sample of 1 g of Kaolinite. Curves (a), (b), (c) and (d) correspond to the raw *MS* signals monitored at m/e 46, m/e 30, m/e 18 and m/e 63, respectively, using an orifice diameter of 8 mm and $[N_2O_5]_0 = (3.8 \pm 0.5) \times 10^{12} \text{ cm}^{-3}$.

1.5 н reaction on MS Signal (Volt) 0.1 :地相 新州市 (d)/10 (a) Amananan mananan anan ang kanan (c) x 10 0.0 1600 1400 1800 2000 time (s)

Fig. 7. Typical N₂O₅ uptake experiment on a sample of 1 g of Saharan Dust. Curves **(a)**, **(b)**, **(c)** and **(d)** correspond to the raw *MS* signals monitored at m/e 46, m/e 30, m/e 18 and m/e 63, respectively, using an orifice diameter of 8 mm and $[N_2O_5]_0 = (3.8 \pm 0.5) \times 10^{12} \text{ cm}^{-3}$.



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Fig. 8. N₂O₅ on 200 mg Kaolinite: uptake coefficient γ of NO₃ as a function of [N₂O₅]: initial (γ_0 , open circles) and steady state (γ_{ss} , full triangles) uptake coefficients for A_s =4.9 cm². Full and empty symbols are referred to uptake experiments carried out with an orifice diameter of 4 and 8 mm orifice diameter, respectively.

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