Uptake of N_2O_5 by aqueous aerosol unveiled using chemically accurate many-body 1 potentials 2

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The reactive uptake of N_2O_5 to aqueous aerosol is a major loss channel for nitrogen oxides in the troposphere. Despite its importance, a quantitative picture of the uptake mechanism is missing. Here we use molecular dynamics simulations with a data-driven many-body model of coupled-cluster accuracy to quantify thermodynamics and kinetics of solvation and adsorption of N_2O_5 in water. The free energy profile highlights that N_2O_5 is selectively adsorbed to the liquid-vapor interface and weakly solvated. Accommodation into bulk water occurs slowly, competing with evaporation upon adsorption from gas phase. Leveraging the quantitative accuracy of the model, we parameterize and solve a reaction-diffusion equation to determine hydrolysis rates consistent with experimental observations. We find a short reaction-diffusion length, indicating that the uptake is dominated by interfacial features. The parameters deduced here, including solubility, accommodation coefficient, and hydrolysis rate, afford a foundation for which to consider the reactive loss of N_2O_5 in more complex solutions.

11 12 13 14 15 16 17 18 19 20 21 tive on gaseous uptake, free of underlying rate limita- ⁵⁴ merically, the simplified models can be refined. 22 tion assumptions.[14] However, studying such processes 23 theoretically imposes challenges, since uptake coefficients 55 24 26 28 29 30 31 32 34 35 36 take.

37 38 39 40 the appropriate reaction diffusion equations is not typi- 72 of the model employed. As direct experimental confir-41 cally tenable, approximate models are commonly postu-73 mation of the importance of the interface is difficult, an

The uptake of trace gases from the air into aerosol par- a_3 kinetic properties.[17] For example, the uptake of N₂O₅ ticles impacts a wide range of environmental systems. [1, 44 in aqueous aerosol has been assumed to follow such a 2] Among other things, such multiphase processes help 45 model, determined by bulk accommodation followed by to determine the oxidative power of the atmosphere by 46 bulk phase hydrolysis and parameterized by its bulk solacting as sinks for nitrogen oxides. [3, 4] Of particular 47 ubility and hydrolysis rate. [18] Such kinetic models typilong-standing interest is the reactive uptake of N_2O_5 in 46 cally lack molecular details, neglecting the finite width of aqueous aerosol, which is estimated to account for 15- 40 the liquid-vapor interface and its potential unique prop-50% of the loss of NO_x in the troposphere. [5, 6] Despite 50 erties. With molecular dynamics simulations these assignificant study, basic questions remain concerning the 51 sumptions can be relaxed, and the relevant parameters mechanism of N_2O_5 uptake. [7–13] Molecular dynamics s_2 extracted to inform an atomistic kinetic model. [19–21] simulations can be used to obtain a molecular perspec- 53 Further, by solving the reaction diffusion equations nu-

The validity of the traditional resistor model for the are exponentially sensitive to free energy differences and 56 reactive uptake of N₂O₅ has been recently called into the simulations involve large systems and long times to 57 question due to the difficulty of reconciling the kinetmodel the complex dynamics. While qualitative predic- 55 ics with field measurements, combined with theoretical tions of mechanisms can be typically studied with con- 59 work providing indications of interfacial stability and ventional empirical force fields or density functional the- or reactivity. [15, 22] The mechanism of uptake has been reory based models, [13, 15] quantitative predictions require 61 cently explored directly using a neural network based rehigher levels of accuracy. To address this challenge, a 62 active model, and it was found that interfacial rather many-body potential, MB-nrg, [16] has recently been pa- 63 than bulk phase processes dictate the observed uptake rameterized from coupled-cluster calculations, providing 64 coefficient.[13] Using training data obtained from denthe capability of making quantitative predictions of the 65 sity functional theory, this study found that the hydrolthermodynamics and kinetics leading to the N_2O_5 up- $_{66}$ ysis rate was sufficiently fast at the interface that bulk 67 phase partitioning cannot kinetically compete, and the ⁶⁸ uptake was determined by a competition between interfa-Gaseous uptake into fluid particles couples thermody- 60 cial hydrolysis and evaporation. These calculations found namic constraints of solubility with kinetic details of re- 70 modest agreement with experimental uptake coefficient action and diffusion. As a complete analytical analysis of 71 values, consistent with the expected qualitative accuracy 42 lated employing a small number of thermodynamic and 74 alternative means of validating it is to employ models 75 with higher chemical accuracy. This is the aim of the 128 current work, to apply a quantitatively accurate poten-76 tial to extract the thermodynamic and kinetic properties 77 underpinning the uptake of N_2O_5 into water. 78

MB-nrg potentials can serve to make quantitative 79 prediction of gaseous uptake as they can be accurate 80 yet computationally amenable to the large system sizes 81 and long timescales required to simulate interfacial pro-82 cesses just like the MB-pol water model. [23, 24] Con-83 trary to common neural network models, these many-84 body potentials have an explicit representation for long-85 ⁸⁶ range interactions, which can be important at extended interfaces. [25] For example, it has been shown that MB-87 pol[26, 27] yields quantitative accuracy for a variety of 88 molecular properties across water's phase diagram[26-89 43] including at the water-vapor interface [44]. Exten-90 91 sions of this modeling framework to describe mono-92 atomic ions and small molecules in aqueous solutions 93 as well as generic mixtures of molecules have been re-94 cently realized. [45–49] These MB-nrg models include a ⁹⁵ model of N₂O₅ that has been developed using analogous approaches.[16] This MB-nrg model was demonstrated to vield comparable accuracy with respect to the coupled 97 cluster reference data it was parameterized on, enabling highly accurate simulations of N₂O₅ in aqueous environ-99 ments. While not able to describe reactions with water, 100 the model nevertheless is capable of quantifying the pro-101 cesses that establish the physical uptake of N_2O_5 . 102

Here we employ this MB-nrg model to study the phys-103 ical uptake of N₂O₅ into water using molecular dynamics 104 simulations and enhanced sampling techniques, making 105 106 quantitative predictions of the thermodynamics and kinetics of N_2O_5 uptake. We subsequently leverage the 107 quantitative accuracy of the model to parameterize and 108 solve a reaction-diffusion equation and infer hydroly-109 sis rates consistent with experiment, providing a com-110 plete quantitative picture of the reactive uptake of N_2O_5 157 111 $_{112}$ by aqueous aerosol. We find a short reaction-diffusion $_{158}$ ubility of N_2O_5 in liquid water. In dilute solution at ¹¹³ length, indicating that the uptake is dominated by inter- c_l in contact with a solute with partial

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RESULTS

In order to extract the thermodynamic and kinetic 116 properties that determine the uptake of N_2O_5 , we have 117 simulated a system containing a slab of liquid water in 162 and computable from the solvation free energy defined 118 119 121 122 123 124 consistent with emergent capillary waves. [50] The bulk 168 of N_2O_5 . [52, 53] This value is higher than recent esti-125 density of the MB-pol model $\rho_{\rm B}$ is 1.007 g/cm³.[29] We 169 mates employing fixed charge force fields and neural net- $_{126}$ take the origin of z to be coincident with the Gibbs di- $_{170}$ work potentials, each of which found a value closer to 0.5 127 viding surface of the interface.

Thermodynamics of adsorption and solvation

Using this simulation setup, we first considered the $_{130}$ thermodynamics of N_2O_5 solvation in liquid water. Fig-¹³¹ ure 1c shows the free energy profile for moving a gaseous ¹³² N₂O₅ into liquid water. Supplementary Figure 1 depicts 133 the free energy profile in more detail and the Supple-¹³⁴ mentary Notes and Supplementary Table 1 contain an 135 overview of all thermodynamic and kinetic parameters 136 discussed in this text. For both $z \gg 0$ and $z \ll 0$, the free ¹³⁷ energy profile is flat, reflecting the translationally invari-138 ant bulk liquid and vapor on either side of the interface. 139 We define the offset between these asymptotic values as 140 $\beta \Delta F_{\rm s} = -4.3 \pm 0.1$, the solvation free energy for the gas 141 phase N₂O₅, where $\beta = 1/k_{\rm B}T$ and $k_{\rm B}$ is Boltzmann's 142 constant. In between these two extremes, the free energy 143 is non-monotonic and exhibits a global minimum approx-144 imately centered at the Gibb's dividing surface and a $_{145}$ barrier to move the N_2O_5 molecule from this interfacial position into the bulk liquid. Relative to the gas phase, 147 the global minimum corresponds to a favorable interfaial adsorption free energy of $\beta \Delta F_{\rm a} = -6.2 \pm 0.1$. The 149 globally favorable interfacial adsorption indicates N_2O_5 150 is relatively hydrophobic, consistent with previous obser-¹⁵¹ vations of its relatively weak solvation.[13, 15, 51] This ¹⁵² free energy profile dictates that the equilibrium density $_{153}$ profile of N_2O_5 would be inhomogeneous in the vicinity of 154 the liquid-vapor interface, a feature neglected in typical 155 kinetic models.

Solubility

From the free energy profile we can calculate the solfacial features in the vicinity of the liquid/vapor interface. p_{160} pressure p_{160} , this solubility is traditionally reported as a 161 Henry's law constant defined as [52]

$$H = \frac{c_l}{p} = \beta e^{-\beta \Delta F_{\rm s}} \tag{1}$$

contact with its vapor and a single N_2O_5 molecule as de- 163 operationally from our free energy profile. This estimate scribed in the Methods section and illustrated in Figure 164 gives a Henry's law constant $H = (3.0 \pm 0.4)$ M/atm. 1a. The corresponding density profile of water along the 165 This value is in line with typical inferences from experdirection perpendicular to the interface, $\rho(z)$, is shown 166 iment, which range between 1 to 10 M/atm, though its in Figure 1b, exhibiting the expected sinusoidal profile 167 direct measurement is hindered by the facile hydrolysis 171 M/atm. [13, 15]

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173 174 ics, we have also characterized the dynamical processes 227 to equilibration at the interface followed by subsequent 175 176 178 180 ¹⁸¹ N₂O₅ by computing mean-squared displacements. The ²³⁴ kinetic theory.[17] Specifically, the rates are given by value obtained for the self-diffusion constant of N₂O₅ de-182 rived from the average mean-squared displacement was 183 $(1.53 \pm 0.06) \times 10^{-5} \text{ cm}^2/\text{s}$. Hydrodynamic effects are 184 known to suppress the diffusion constant for finite systems employing periodic boundary conditions.[54] Using 186 the known experimental viscosity of liquid water at am-187 bient conditions, [29] we can correct for these finite size 188 effects resulting in a diffusion constant in the thermody-189 namic limit of $(1.89 \pm 0.06) \times 10^{-5}$ cm²/s. We have also 237 190 estimated the change in the diffusion constant at the liq-191 uid vapor interface, [55] and find an increase from its bulk 192 value to $(5.3 \pm 0.1) \times 10^{-5} \text{ cm}^2/\text{s}.$ 193

Adsorption and evaporation rates 194

Evaporation from the liquid-vapor interface and sol-195 vation into the bulk are both activated processes with 196 barriers estimated from the free energy in Figure 1 to 197 be larger than typical thermal values. As such, they are 198 rare events and difficult to sample with straightforward 199 simulations. However, as we consider a system whose 200 dynamics satisfies detailed balance, we can alternatively 201 study comparatively typical events like desolvation and 202 adsorption, and infer their reverse using the previously 203 evaluated free energy profile. [56] Definitions for these dif-204 ferent dynamical processes are well described in Ref. 17. 205 To compute the adsorption and therefore evapora-206 tion rates, we have sampled 250 scattering trajectories 207 whereby an initially gas phase N_2O_5 placed at z = 1.2208 nm, shown in the right panel of Figure 2a, is evolved to-209 ward the liquid slab. To do this we take 10 distinct equi-210 $_{211}$ librium configurations of N_2O_5 generated by constrain- $_{212}$ ing its center of mass to z = 1.2 nm, and draw 25 re-²¹³ alizations of a Maxwell-Boltzmann distributed velocity 214 at 300 K for each. Figure 2b reports the trajectories of ²¹⁵ the center of mass of the N₂O₅ as it impinges on the liq- $_{216}$ uid slab. Overwhelmingly, the incipient gas phase N_2O_5 ²¹⁷ molecule meets the interfaces and sticks, with only 11 218 out of the 250 scattering trajectories exhibiting a back 259 where $\beta \Delta F_{\rm b} = 0.8$ is the barrier to move from the bulk 219 scattering event, with N₂O₅ bouncing off of the inter- 260 liquid to the interface and $\ell = 0.6$ nm is the width of the 220 face and going back into the gas phase within the 100 ps 261 interface. The width of the interface is determined by fit-²²¹ observation time employed. The scattering rate is quan-²⁶² ting the free energy minima to a parabola and integrating 222 tified with the so-called thermal accommodation coeffi- 263 the resultant Gaussian distribution from $1 \text{ nm} > z > z^{\dagger}$. 223 cient, $S = 0.96 \pm 0.06$, relating the probability of being 264 We find $k_d = 340/\text{ns}$ and $k_s = 51/\text{ns}$.

Diffusion in bulk and at the liquid/vapor interface 224 accommodated at the interface upon collision, consistent ²²⁵ with previous simulations. [15] This is likely a lower bound As gaseous uptake couples thermodynamics and kinet- 226 as some of the back scattering events can be attributable of N_2O_5 as it moves between phases across the liquid- 228 evaporation. The near unity value implies a lack of a barvapor interface. Before considering the rare events of 229 rier to adsorption, that subsequent evaporation is analoevaporation and solvation, we first discuss the diffusive 230 gously limited only by the free energy of adsorption, and properties of N_2O_5 in the bulk liquid. With a simula- ²³¹ that uptake is not significantly influenced by this initial tion of N_2O_5 immersed in a bulk liquid containing 272 ²³² thermal accommodation. The corresponding rates of adwater molecules, we have estimated the self-diffusivity of 233 sorption, $k_{\rm a}$, and evaporation, $k_{\rm e}$, can be computed from

$$k_{\rm a} = S \frac{v}{4} , \quad k_{\rm e} = k_{\rm a} e^{\beta \Delta F_{\rm a}}$$
 (2)

235 where $v = \sqrt{8/\beta\pi m}$ is the average molecular speed of 236 N₂O₅. These are $k_{\rm a} = 57 \text{ nm/ns}$ and $k_{\rm e} = 0.11 \text{ nm/ns}$.

Solvation and desolvation rates

738 We have also computed the rates of solvation and de-239 solvation following the Bennet-Chandler approach. [57] Identifying $z^{\dagger} = -0.42$ nm as the location of the pu-241 tative transition state for solvation into the bulk liquid ²⁴² from the interface (see Figure 1c), we can estimate the ²⁴³ rate of solvation and analogously desolvation by computing the transmission coefficient, κ , for committing to the ²⁴⁵ interface conditioned on starting at the transition state. 246 The transmission coefficient is defined as [57]

$$\kappa(t) = \frac{\langle v(0)\Theta(z(t) - z^{\dagger}) \rangle_{z^{\dagger}}}{\langle |v| \rangle/2} \tag{3}$$

²⁴⁷ where Θ is the step function and the brackets denote 248 an ensemble average where in the numerator it is condi-²⁴⁹ tioned on starting at the transition state. We have evalu-²⁵⁰ ated the transmission coefficient using 2000 trajectories. ²⁵¹ Like the scattering calculations, we have taken 80 differ-²⁵² ent equilibrium configurations of N₂O₅ at $z = z^{\dagger}$, and ²⁵³ compose 25 Maxwell-Boltzmann velocity distributions at ²⁵⁴ 300 K for each. Each trajectory was evolved for 10 ps. Figure 2c shows that κ decays to 0.08 over 1 ps, consis-²⁵⁶ tent with a diffusive barrier crossing. From the plateau 257 we can estimate the rates to solvate into the bulk, $k_{\rm s}$, **258** and desolvate into the interface, $k_{\rm d}$, as

$$k_{\rm d} = \kappa \frac{v}{2\ell} e^{\beta \Delta F_{\rm b}} , \quad k_{\rm s} = k_{\rm d} e^{\beta (\Delta F_{\rm a} - \Delta F_{\rm s})} \tag{4}$$

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Mass accomodation

266 $_{267}$ α , defined as the probability of a gas molecule striking $_{313}$ standard deviation of 0.05 nm was found to well approxi-268 269 $_{270}$ rameter determines the transfer rate of N₂O₅ across the $_{316}$ grid spacing of 0.02 nm and timestep 0.018 ps. 271 surface into the bulk liquid and can be computed from 317 $_{272}$ the sticking coefficient S and the free energy profile as

$$\alpha = \frac{S}{1 + e^{\beta(\Delta F_{\rm s} + \Delta F_{\rm b})}} \tag{5}$$

 $_{274}$ that infer a value larger than 0.4 for N₂O₅.[58]

Reactive uptake through interfacial and bulk 275 hydrolysis 276

277 278 drolysis with water and this reaction ultimately deter- 330 the overall reactive uptake, γ , can be computed by the mines the reactive uptake in aqueous aerosol. While we 331 portion of the loss through the reactive channel, 279 cannot simulate the reactive event with the MB-nrg po-280 tential employed directly, we can still make an inference 281 282 into the reactive uptake. Using the thermodynamic and 203 kinetic parameters evaluated from the molecular dynam- 332 We model the hydrolysis rate as having two characteristic 284 ics simulations, we can parameterize a molecularly de-333 values, one in the bulk for z < -0.5 nm, denoted $k_{\rm h}$, and 205 tailed reaction diffusion equation. Specifically, we con-334 one at the interface for -0.5 < z < 0.5 nm taken to 286 287 able to also undergo hydrolysis. 289

290 291 292 equilibrated at the interface. The subsequent evolution 341 Figure 3b, which rises from 0 to a plateau value at times 293 of its concentration profile, c(z,t), can be solved for using 342 much longer than the characteristic time associated with ²⁹⁴ a Smoluchowski equation[59] of the form,

$$\frac{\partial c(z,t)}{\partial t} = \frac{\partial}{\partial z} D(z) e^{-\beta \Delta F(z)} \frac{\partial}{\partial z} e^{\beta \Delta F(z)} c(z,t) -k_{\rm h}(z) c(z,t)$$
(6)

296 D(z) is the diffusion constant, and $k_h(z)$ is the unknown 349 believe these are the two likely extremes, as previous ex-297 hydrolysis rate, both of which in principle vary through 350 plicit calculations found that interfacial hydrolysis was 200 space. [60, 61] The first term is a drift diffusion encod- 351 suppressed relative to the bulk. [13] Experimentally, the 299 ing the stationary distribution implied by the free en- 352 range of reactive uptake coefficients on pure water has 300 ergy profile, while the second accounts for loss due to 353 been reported between 0.03 and 0.08, [7–9] which is con- $_{301}$ reaction. In practice we fit the free energy to an an- $_{354}$ sistent with a bulk hydrolysis rate between 0.01 ns⁻¹ 302 alytic function form, $\beta \Delta F(z) = a_1 \tanh[(z - a_2)/a_3] - 355$ and 0.07 ns⁻¹ in Figure 3c. These rates are slower than 303 $a_4 \exp[-(z-a_5)^2/a_6] + a_7 \exp[-(z-a_8)^2/a_9]$. Equa- 356 those computed directly from a previous neural network 304 tion 6 is valid only for the liquid and interface, not 357 model (0.2 ns⁻¹),[13] but faster than that typically in- $_{305}$ the vapor, since it is overdamped. [59] In order to model $_{358}$ ferred experimentally (0.002 ns⁻¹). [62] The disagreement 306 the vapor, we employ absorbing boundary conditions 359 with respect to the neural network model could likely be 307 c(z = 1 nm, t) = 0, and consider a domain that extends 300 a failure of the density functional used in the training 308 deeply enough into the liquid that the results are insensi- 361 data, by delocalizing the charge transfer accompanying 309 tive to the reflecting boundary condition employed there, 362 hydrolysis. [63–65] The disagreement with the rates in-

310 $\partial_z c(z = -30 \text{ nm}, t) = 0$. We solve Eq. 6 with a nor-³¹¹ malized Gaussian initial condition, localized in the free We can also evaluate the mass accomodation coefficient ³¹² energy minima near the Gibbs dividing surface where a the liquid surface to solvate into the bulk liquid phase in 314 mate the curvature of the interfacial minima. In practice, absence of surface reactions[17]. This fundamental pa- 315 we employ a simple finite difference scheme with constant

In the absence of any reactions, an initial interfacial $_{318}$ concentration of N_2O_5 will relax through a competition 319 between diffusion into the bulk liquid and evaporation 320 into the vapor towards the steady-state determined by ³²¹ the free energy profile. Figure 3a illustrates the relax-²⁷³ We find $\alpha = 0.93 \pm 0.06$, in agreement with experiments ³²² ation of this concentration profile. The initial Gaussian 323 distribution quickly looses amplitude and a diffusive front 324 propagates into the bulk liquid, while concentration is 325 irreversibly lost to the vapor. The initial rates to evapo-326 rate and solvate are consistent with our explicit molecular 327 simulation calculations. In the presence of hydrolysis, in 328 addition to loss from evaporation, there can be loss due Experimentally, N_2O_5 undergoes facile irreversible hy- 320 to reaction. Since the concentration is normalized to 1,

$$\gamma(t) = \int_0^t dt' \int dz \, k_{\rm h}(z) c(z, t') \tag{7}$$

sider the diffusive dynamics accompanying an initially 335 be a fraction of the bulk value, while it is set to zero in adsorbed N₂O₅ molecule as it enters the bulk liquid or $_{336}$ the vapor for z > 0.5 nm. The interfacial region was evaporates, and address what would happen if it were 337 determined from the inflection points in the free energy 338 curve and numerical tests have shown that the results are Consistent with the near unity thermal accommoda- 339 insensitive to the precise width of the interfacial region. tion S, we assume an initially adsorbed molecule locally $_{340}$ An example time series for the reactive uptake is shown in 343 the bulk hydrolysis rate. This asymptotic value is the 344 reactive uptake coefficient.

The reactive uptake as a function of the bulk hydrol-346 ysis rate is shown in Figure 3c. For each bulk hydroly-347 sis rate, we have computed γ setting the interfacial rate 295 where $\Delta F(z)$ is the free energy profile from Figure 1c, 348 equal to the bulk value, and also setting it to zero. We

³⁶⁴ tive uptake models that neglect interfacial reactivity and ⁴¹⁴ very weak aerosol particle size dependence. 365 stability. The confirmation of the importance of the in- 415 366 terface, reducing the diffusion into the bulk and account- 416 used as a new starting point for further modeling efforts $_{307}$ ing for a significant fraction of hydrolysis, agrees with the $_{417}$ to predict the reactive uptake of N₂O₅ in more complex 308 previous neural network model study, [13] and the need to 418 solutions. It is well known that the reactive uptake can 300 revise the standard resistor model. The hydrolysis rate 410 be modulated in the presence of inorganic salts, as in the 370 obtained with our modelling in addition to the other pa- 420 case of excess nitrate ions.[18] Further, the branching ra- $_{371}$ rameters relevant to reactive uptake are summarized in $_{421}$ tios to alternative less soluble products like to ClNO₂ in 372 Table 1.

Implications for the reactive uptake mechanism 373

374 375 376 active uptake. Specifically, the range of uptake values 430 with the full complexity of field measurements. 377 378 observed upon changing the interfacial hydrolysis rate 379 from 0 to $k_{\rm h}$ illustrates that, while interfacial reactivity 380 contributes to the reactive uptake coefficient, it accounts 431 for at most 20%. The interfacial contribution is lower 381 than recent estimates, [13] due to the increased solubility 432 382 predicted by the MB-nrg model and corresponding higher 383 accommodation coefficient relative to the previous neural 433 385 387 388 389 390 301 energy minima at the interface results in a propagation 440 corrections to thermodynamic properties are expected to $_{392}$ length of N₂O₅ into the bulk fluid of only around 2 nm $_{441}$ be minimal[51]. The system is embedded in a simula-303 (see Figure 3a). This implies that reactive uptake is af- 442 tion domain of the same cross-section and a length of 20 394 fected by interfacial characteristics, even though most 443 nm in order to accommodate periodic boundary condi-395 of the reaction is predicted to take place in the bulk. 444 tions. Simulations were executed with Amber 2020[67] 306 It also predicts a very weak aerosol particle size depen- 445 interfaced to the MBX[68] library. Ewald summation 307 dence to reactive uptake consistent with some experimen- 446 was employed to describe long-range electrostatics and ³⁹⁸ tal observations. [62]

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DISCUSSION

400 with many-body potentials of coupled cluster accuracy 401 that quantify the thermodynamics and kinetics of ad-402 sorption and solvation of N_2O_5 by aqeuous aerosol. The ⁴⁵³ 403 404 hydrolysis rate of N_2O_5 is determined by numerically solving a molecularly detailed reaction-diffusion equation $_{454}$ We computed the free energy to move a gaseous N_2O_5 406 407 408 409 of the reactive uptake of N₂O₅ by aqueous aerosol. Al- 458 ployed harmonic biasing potentials of the form $\Delta U(z) =$ 410 though most of the hydrolysis is predicted to take place 459 $k/2(z-z^*)^2$ with a spring constant k of 2.5 kcal mol⁻¹ 411 in bulk water, our results highlight the importance of in- 460 A^{-2} and 52 independent windows with minima z^* spaced 412 terfacial features at the liquid/vapor interface leading to 461 evenly between -1.36 and 1.19 nm. Three separate sets of

303 ferred experimentally is because those are based on reac- 413 a relatively short reaction diffusion length and thus only

The framework and parameters determined here can be 422 the presence of NaCl have been well studied. [11, 66] By 423 quantifying the changes to both the thermodynamic and $_{424}$ kinetic properties of N_2O_5 in the presence of these al-425 ternative solutions, advanced molecular models such as 426 the ones used in this work in combination with similar Apart from quantifying a likely range of experimen- 427 analysis of generalized reaction-diffusion equations incortal hydrolysis rates, the analysis of the reaction diffusion 428 porating alternative loss mechanics, can be exploited to model provides insight into the likely mechanism of re- 420 provide a complete picture of reactive uptake of N₂O₅

METHODS

Molecular dynamics simulations

To simulate the uptake of N_2O_5 in water we employed network model study. Nevertheless, a significant adsorp- $_{434}$ the MB-nrg model of N_2O_5 in MB-pol water with extion free energy reduces diffusion into the bulk liquid, 435 plicit one-body, two-body and three-body short-range resulting in an effective renormalized reaction-diffusion 436 interactions. [16] The simulation system illustrated in Figlength. Absent barriers to solvation, the reaction dif- 437 ure 1a is made up of 533 water molecules forming a liquid fusion length, $\ell_r = \sqrt{D/k_h}$, would be around 15 nm. 438 slab measuring 2.416 nm x 2.416 nm in cross-sectional However, the barrier to solvation and corresponding free 439 area and 2.772 nm in length. For this slab size, finite size 447 dispersion interactions using a real-space cutoff of 1.2 448 nm. Thermodynamic averages were computed within an ensemble of fixed N particles, V volume, and T =450 300 K temperature, using a Langevin thermostat and 451 a timestep of 0.5 fs. Kinetic properties were evaluated We report extensive molecular dynamics simulations $_{452}$ within a constant energy ensemble with fixed N and V.

Umbrella sampling

that incorporates these parameters to yield results con- 455 into the liquid water slab using umbrella sampling apsistent with the experimentally observed reactive uptake $_{456}$ plied to the center of mass distance along the z dicoefficient. This provides a complete quantitative picture $_{457}$ rection between the water slab and N₂O₅.[69] We em462 calculations were run, each consisting of 1 ns equilibra- 503 tion time followed by 2.5 ns production time to compute 504 463 ⁴⁶⁴ partial histograms. The individual histograms from each 505 506 window were combined using umbrella integration.[70] 465 507 ⁴⁶⁶ Error bars of the free energy profile were computed from 508 467 the standard deviation for the three independent calcu-509 468 lations. 510

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Diffusion coefficients

Diffusion coefficients were computed from simulations 516
of N₂O₅ in bulk liquid water. Details of the calculation of 517
diffusion coefficients from mean-squared displacements in 518
the molecular dynamics simulations are discussed in the 519
Supplementary Methods. 520

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DATA AVAILABILITY

476Data supporting the findings of this study are in-526477cluded in the article and a Source Data file is pro-527478vided with this paper. In addition, the data related529479to this publication including simulation input files can530480be accessed from the NSF-CAICE Data Repository[71]531481(https://doi.org/10.6075/J0FF3SHB).532

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CODE AVAILABILITY

This study utilized Amber 20 (https://ambermd.org) 538 483 484 interfaced toMBX (http:// paesani- 539 software ${}^{\mathbf{540}}$ group.ucsd.edu/software/mbx.html). The 485 541 for solving the reaction diffusion model is included 486 542 in the Supplementary Material and can be ac-543 cessed from the NSF-CAICE Data Repositorv[71] 488 544 (https://doi.org/10.6075/J0FF3SHB). 545

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AUTHOR CONTRIBUTIONS

D.T.L. and A.W.G. conceptualized and designed the Mardirossian, N. & Head-Gordon, M. Thirty years of 777 work. V.W.D.C. and A.W.G. prepared the simulation density functional theory in computational chemistry: an 778 models with many-body potentials of coupled cluster ac-779 curacy. V.W.D.C. performed the molecular dynamics 780 simulations (umbrella sampling, scattering at the wa- $_{781}$ ter/vapor interface, and N_2O_5 in bulk water). D.T.L. functionals and plane waves for chemical reactions. The 782 developed the reaction-diffusion model and executed the 783 corresponding simulations. V.W.D.C, M.G., D.T.L., and Roberts, J. M. et al. Laboratory studies of products of 784 A.W.G. performed data analyses. D.T.L. drafted the N₂O₅ uptake on Cl⁻ containing substrates. Geophysical 785 manuscript with contributions from all authors. D.T.L. 786 and A.W.G. acquired funding and supervised this re-787 search.

COMPETING INTERESTS

The authors declare no competing interests. 780

TABLES

TABLE 1.	Physical	and chemical	properties	of N ₂ O ₅
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Henry's Law Constant	H	(3.0 \pm 0.4) M/atm
Diffusivity	D	$\begin{array}{l}(1.89\ \pm\ 0.06)\ \times\\10^{-5}\ {\rm cm}^2/{\rm s}\end{array}$
Sticking Coefficient	S	0.96 ± 0.06
Mass Accommodation	α	0.93 ± 0.06
Hydrolysis Rate^{a}	$k_{ m h}$	$(4\pm3)\times10^{-2} \text{ ns}^{-1}$

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FIGURE LEGENDS/CAPTIONS

792 793 ⁷⁹⁴ interface. b) Water density profile where z = 0 demarks ⁸¹⁸ out (orange squares) and with (blue circles) interfacial 795 ros density. c) Free energy to move N_2O_5 in the z direction, s20 ficients observed experimentally on pure water, and grey

798 bars. $\Delta F_{\rm a}$ and $\Delta F_{\rm s}$ are the free energies of adsorption and solvation, respectively. 799

Fig. 2 Kinetics of N₂O₅ adsorption and solvation. a) Snapshots of N_2O_5 . Adsorption and evapora-801 tion takes N_2O_5 between the vapor (right) and interface 802 803 (center), while solvation and desolvation takes it between the interface and the bulk (left), with corresponding rate constants. **b**) Scattering trajectories following the z com-805 ponent of the center of mass of N_2O_5 . c) Transmission coefficient for transitions between the liquid and the in-807 terface. The shaded region are one standard deviation error bars. The solid red line is an exponential fit. 809

Fig. 3 Reactive uptake from the reaction-810 diffusion equation. a) Relaxation of the initial con-811 centration profile. Blue lines are c(z,t) separated by 0.25 812 ⁸¹³ ns and the dashed black line is the equilibrium profile s14 computed from $\exp[-\beta\Delta F(z)]$. b) An example time des15 pendent reactive uptake coefficient $\gamma(t)$. Both a) and b) Fig. 1 Thermodynamics of N₂O₅ solvation. a) are is computed with $k_{\rm h} = 0.02 \text{ ns}^{-1}$ and the interfacial rate Characteristic snapshot of N_2O_5 near the water-vapor a_{17} equal to $k_h/5$. c) Asymptotic uptake coefficients withthe Gibbs dividing interface and $\rho_{\rm B}$ is the bulk water *** reactivity. Blue regions denote the range of uptake coef-797 where the shaded region are one standard deviation error 821 the corresponding likely range of bulk hydrolysis rates.









