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Surface adsorption in strontium chloride ammines

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An adsorbed state and its implications on the ab- and desorption kinetics of ammonia in strontium chloride ammine is identified using a combination of ammonia absorption measurements, thermogravimetric analysis, and density functional theory calculations. During thermogravimetric analysis, ammonia desorption originating from the adsorbed state is directly observed below the bulk desorption temperature, as confirmed by density functional theory calculations. The desorption enthalpy of the adsorbed state of strontium chloride octa-ammine is determined with both techniques to be around 37-39 kJ/mol. A simple kinetic model is proposed that accounts for the absorption of ammonia through the adsorbed state. © 2013 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4800754]

I. INTRODUCTION

Metal halides are useful materials for absorption of ammonia, forming ammine complexes in the solid state. Applications of the materials include thermochemical heat pumps,1 ammonia separation,2 and ammonia storage and delivery3, 4 for fuel cells5 or selective catalytic reduction of NOx gasses.6 Salts that form complexes with ammonia are very common and show a wide spectrum in binding energies and ammonia capacities.1 For reasons of ammonia storage capacity and availability, alkaline Earth halides are very attractive, and for practical reasons of operating temperature and pressure, CaCl2 and SrCl2 are often preferred. Since SrCl2 binds 7 out of 8 ammonia molecules with low binding energy (1-7: 41.4 kJ/mol, 8: 48.1 kJ/mol) compared to 6 out of 8 for CaCl2 (1-4: 41.0 kJ/mol, 5-6: 42.3 kJ/mol, 7: 63.2 kJ/mol, 8: 69.1 kJ/mol), SrCl2 is often the chosen material for practical applications.7 The stable ammines of SrCl2 are the monoammine Sr(NH3)Cl2, the di-ammine Sr(NH3)2Cl2, and the octa-ammine Sr(NH3)8Cl2. Depending on temperature and pressure, transitions can go between mono-, di-, and octa-ammine or directly between mono- and octa-ammine.8

For all the above mentioned applications, the absorption/desorption dynamics of ammonia is essential for the performance. In the literature, effective kinetic models of the form

\[ r_a = k_0 \exp(-E_a/RT) \left(1 - s\right)^M f(s, T, p) \]  

have previously been investigated.9-12 Here, \( E_a \) is the activation energy, \( R \) is the gas constant, \( k_0 \) and \( M \) are constants, \( T \) is the absolute temperature, and \( s \) is the ammonia saturation degree. Different forms of \( f \) have been proposed including a linear term \( f = p_{rel}(s, T) \), a power term \( f = p_{rel}(s, T)^N \), and a logarithmic term \( f = \log p/p_{eq} \), where \( p \) is the pressure and \( N \) is another constant. The relative pressure \( p_{rel} \) is given by

\[ p_{rel}(s, T) = \frac{p - p_{eq}(s, T)}{p_{eq}(s, T)} \]

and the equilibrium pressure \( p_{eq} \) is calculated from the van ’t Hoff relationship

\[ \log K = \log \frac{p_{eq}}{p_0} = -\frac{\Delta H}{RT} + \frac{\Delta S}{R} \]

In the present study, absorption rates are measured as a function of pressure at different temperatures, and a deviation from the effective kinetic model (1) is observed at pressures approaching the equilibrium pressure. It is proposed that this indicates the existence of an adsorbed state that is filled at pressures significantly higher than the equilibrium pressure, but less occupied closer to the equilibrium pressure.

Desorption from the adsorbed state is directly demonstrated using thermogravimetric analysis (TGA) and confirmed by density functional theory (DFT). Finally, a simple microkinetic model based on the adsorbed state is analyzed and shown to fit the absorption rate data well.

II. MEASUREMENT OF ABSORPTION RATE

The overall principle used to measure absorption rates is: (i) to prepare a reactor of known volume with an amount of strontium chloride, (ii) almost instantly fill the reactor to a high pressure with ammonia, and (iii) follow the pressure as ammonia is absorbed in the salt until the pressure stabilizes close to the equilibrium pressure. The rate, at which the pressure drops, is proportional to the absorption rate. This measurement principle is dynamic rather than a steady state. The main advantage is that the amount of ammonia in the gas phase is small compared to the amount of strontium chloride. This implies that the temperature can be considered constant.

\[ \frac{\Delta p}{\Delta t} = k_{abs} \]

where \( k_{abs} \) is the absorption rate constant.

\[ k_{abs} = \frac{\Delta V}{\Delta t} \]

where \( \Delta V \) is the volume change due to absorption and \( \Delta t \) is the time.

\[ \Delta V = V_0 - V_f \]

where \( V_0 \) is the initial volume and \( V_f \) is the final volume. The absorption rate is calculated from the slope of the pressure versus time graph.

\[ \frac{\Delta p}{\Delta t} = \frac{\Delta V}{\Delta t} \]

where \( \frac{\Delta V}{\Delta t} \) is the slope of the pressure versus time graph.
during absorption, because the heat evolved during absorption of the limited amount of ammonia gas is small. Also, the saturation degree can be considered constant. In this way, the measurement principle has the unique feature that absorption rates as function of ammonia pressure can be measured at different and independently chosen saturation degrees and temperatures.

The experimental setup is shown in Figure 1 (for a detailed description, see Ref. 8). The experiment consists of a sample chamber containing SrCl₂ (China Newcent, purity 99.7%, anhydrous, further dried at 200 °C for 24 h to less than 0.05% water content) and a measurement chamber that can be evacuated or filled with ammonia to a certain pressure through the valve V2. The measurement volume is kept at a fixed temperature in a climate chamber. For each cycle of the experiment, the two volumes are first separated by closing the valve V1 between them. This is followed by evacuating the measurement volume and pressurizing it with ammonia to a specified initial pressure. Hereafter, the valve V1 to the sample volume is opened. The pressure equilibrates between the sample chamber and the measurement chamber within milliseconds. From then on, the sample and measurement chambers behave as a batch reactor, and the ammonia pressure, \( P(t) \), measured with the pressure sensor P, decreases as ammonia is absorbed in the strontium chloride. After a while, the pressure becomes almost constant in time, which indicates an approach to equilibrium. By repeating the procedure, the salt can be prepared at any saturation degree. During the experiment, the absorption rate is determined from \( \frac{dp}{dt} \) and the equation of state

\[ r_a = \frac{dn}{dt} = \frac{V}{RT} \frac{dp}{dt}. \] (4)

From the initial pressure and final pressure in each cycle, the increase in absorbed ammonia can be calculated. In the measured range, the saturation degree changes less than a few percent during each cycle. Also, the sample is highly dispersed between a high number of steel balls8 both to maximize the heat capacity, thus reducing the temperature changes during absorption to less than a few Kelvin, and to minimize any mass transport limitations. If the steel balls were not introduced, the temperature of the SrCl₂ would approach the equilibrium temperature instead of being an independently controllable parameter. The saturation degree is verified by measuring the mass of the sample chamber.

A new sample of SrCl₂ will break up and form a porous structure upon ammonia absorption and desorption.13 Before the experiment is started, the sample is saturated with ammonia and degassed several times to prepare a stable sample with a non-changing structure. The experiment contains 5.1 g SrCl₂ and the gas volume in the two volumes V1 and V2 together is 90 ml.

All together, absorption rates as function of ammonia pressure can be obtained for specific saturation degrees and temperatures.

In Figure 2, measured absorption rates \( r_a \) are plotted as function of the relative pressure \( (2) \) for four temperatures.

The best fit of the proposed model (1) to the data in Figure 2 is obtained with the linear expression for

\[ r_a = k_0 \exp\left(-\frac{E_a}{RT}\right)(1-s)p_{rel}(s, T), \]

with \( k_0 = 9.5 \times 10^3 \) mol/s, \( E_a = 32.9 \) kJ/mol, and \( s = 0.3 \). The relative pressure, \( p_{rel} \), is calculated from (2) using literature values of \( \Delta H = 41.4 \) kJ/mol and \( \Delta S = 228.8 \) J/(mol K).1

The model fits well at high relative pressures, but deviates at low relative pressures.

It is proposed, that this indicates the existence of an adsorbed state in the absorption dynamics: The linear regime at high pressures, where the absorption rate is high, is interpreted to belong to a high ammonia surface coverage. At lower pressures, the surface coverage will shift to lower values and the rate will drop.

III. THERMOGRAVIMETRIC ANALYSIS OF THE ADSORBED STATE

Thermogravimetric analysis is a technique where a sample can be heated while monitoring the mass change with a balance. In a standard configuration, the sample is heated in a (typically) non-corrosive gas stream at ambient pressure. For measurements with corrosive gasses at varying pressures, the sample and gas stream need to be isolated from the balance and electronics. The present investigations are performed on a high pressure TGA (HP-TGA, Isosorp Gas, LP-flow) from Rubotherm, where the sample and sample holder are magnetically coupled to the balance and electronics. The HP-TGA
works in an ammonia pressure of 0-5 bars and a temperature range of 0 °C-400 °C with a mass resolution of 10 μg.

A SrCl₂ sample of 226 mg is first saturated and degassed several times in the HP-TGA to create a stable structure of the sample with high porosity and surface area. The measurement cycle hereafter is (i) to fully saturate the sample for 120-180 min at 0 °C and an ammonia pressure of 3 bars. (ii) Stabilize the start temperature, \( T_s \), and pressure, \( p \), for the HP-TGA measurement. (iii) Ramp the temperature at 1 K/min to the desired final temperature \( T_f \).

A representative desorption measurement in 2 bars of ammonia is shown in Figure 3. The temperature is ramped from \( T_s = 273 \) K to \( T_f = 323 \) K at 1 K/min. The plotted data are \( \frac{dm}{dt} \), the mass change per time unit. The fast desorption range of 0-5 bars and a temperature from the surface begins, are listed. The asterisk denotes the dataset plotted in Figure 3. The pressures and desorption temperatures \( T_{\text{decomposition}} \) of Table I are plotted in Figure 4 and the fitted enthalpy and entropy changes are

\[
\Delta H_{\text{surf}} = 39.2 \text{ kJ/mol and } \Delta S_{\text{surf}} = 228 \text{ J/(mol K)}.
\]

Desorption from bulk is well understood and expected, when the temperature exceeds the bulk equilibrium temperature at a given pressure. The general relation between pressure and temperature at equilibrium is given by (3). For desorption from fully saturated Sr(NH₃)₆Cl₂, the enthalpy and entropy changes are \( \Delta H_{6:1} = 41.4 \text{ kJ/mol and } \Delta S_{6:1} = 228.8 \text{ J/(mol K)} \), where the subscript denotes going from the octa-ammine phase to the mono-ammine phase. From this, the equilibrium temperature related to desorption from bulk can be calculated as plotted in Figure 4.

It is clearly seen from Figure 4 that desorption starts at a significantly lower temperature than expected from the equilibrium temperature of ammonia absorbed in bulk Sr(NH₃)₆Cl₂. This is consistent with desorption from an adsorbed state with lower binding energy than the bulk states.

To estimate the possible amount of ammonia on the surface, the surface area of saturated SrCl₂ is assumed to be of the order of 50 m²/g, which has been reported for similar metal ammine chlorides. Setting the area of a surface bound ammonia molecule to 2.5 × 10⁻¹⁰ m² (a reasonable assumption as shown by DFT calculations in Figure 5), it is possible to adsorb 2 mg of ammonia on the surface, which is roughly the mass change observed from the onset of desorption at 309 K until bulk desorption kicks in at around 320 K as seen in Figure 3. (0.226 g of strontium chloride can absorb a total of 0.19 g of ammonia meaning that the ratio between surface sites and bulk sites is about 1:100 in the experiment.)

### IV. DFT CALCULATIONS

Density functional theory calculations were performed using the ASE framework with the GPAW version 0.7.6383 DFT code, GPAW is a grid-based real space implementation of the Projector Augmented Wave (PAW) method. A van der Waals corrected exchange correlation functional (vdW-DF) that includes long range correlation effects, important for systems, where hydrogen bonds are present, is used. The Brillouin zone was k-point sampled using a Monkhorst-Pack scheme. For each structure, the k-point sampling, grid spacing, and added vacuum were tested for convergence of the energy. A grid spacing of 0.18 Å, a k-point sampling of (4,2,1) (1 in the direction normal to the

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**TABLE I.** List of TGA experiments: For each experiment the pressure, the temperature interval, and the temperature, \( T_{\text{decomposition}} \), at which desorption from the surface begins, are listed. The asterisk denotes the dataset plotted in Figure 3.

<table>
<thead>
<tr>
<th>Pressure (bar)</th>
<th>( T_s ) (K)</th>
<th>( T_f ) (K)</th>
<th>( T_{\text{decomposition}} ) (K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>273</td>
<td>303</td>
<td>285</td>
</tr>
<tr>
<td>1</td>
<td>293</td>
<td>373</td>
<td>296</td>
</tr>
<tr>
<td>2</td>
<td>273</td>
<td>308</td>
<td>296</td>
</tr>
<tr>
<td>2*</td>
<td>273</td>
<td>323</td>
<td>309</td>
</tr>
<tr>
<td>2</td>
<td>273</td>
<td>323</td>
<td>311</td>
</tr>
<tr>
<td>2.5</td>
<td>273</td>
<td>323</td>
<td>315</td>
</tr>
</tbody>
</table>
A. Modeling the surfaces

Surfaces were made from bulk crystals of the different amines (see Ref. 8 for more information about the bulk structures). Cuts were made along different low Miller index planes to simulate possible surface terminations and the energy cost of exposing a surface from the bulk was found. The relaxation usually led to a small contraction in the direction perpendicular to the surface. The facets with the lowest surface energy were used for studying adsorption energies of NH$_3$, as these surfaces are expected to dominate the real surface of the material. The surface indexes are indicated in Table II. For all surfaces, the general rule of thumb was that the surface termination should break the least bonds, thereby letting the surface atoms have the highest possible coordination number. Adsorbate NH$_3$ molecules were investigated in different positions that maximized the number of hydrogen bonds formed between the adsorbates and the surface.

B. DFT results

For Sr(NH$_3$)$_2$Cl$_2$, only one surface site is stable in the unit cell, the energy corresponding to half coverage ($\theta = 0.5$) is therefore calculated by doubling one side of the unit cell in the surface and filling only one of the equivalent sites. For the other phases, two stable surface sites exist. The energy for full coverage ($\theta = 1.0$) is defined as: $E_{\theta=1} = (E_{\theta=0} + 2E_{NH_3})/2$, i.e., the average of the adsorbed ammonia molecules.

The adsorbed NH$_3$ molecules form long hydrogen bonds to the Cl atoms in the surface. For the mono-ammine, the surface site is asymmetric and has one short H-bond on 2.72 Å at 161°, and one longer on 2.87 Å at 142°. For the di-ammine, the hydrogen bonds to Cl atoms are of equal length and angle, 2.85 Å and 142°, respectively. Furthermore, a hydrogen bond to the N on the adsorbed ammonia of only 2.32 Å and 156° is established; this is what stabilizes the di-ammine surface site more than the mono-ammine (see Figure 5(b)).

The second adsorption energy, $\theta = 1$, for the mono-ammine is slightly higher than the first, probably owing to interaction with the first adsorbed NH$_3$. For the di-ammine, the second adsorbs in almost equivalent environments as the first just rotated 180°, leading to almost equal adsorption energies. The surface sites on the octa-ammine are different in hydrogen bond length and angles giving energies with large differences. The strongly adsorbed NH$_3$ (indicated with dark grey hydrogen in Figures 5(c) and 5(d)) molecule forms two hydrogen bonds of 3.07 Å and 3.16 Å at 160° and 156°, respectively. The weakly adsorbed NH$_3$ (marked with light grey hydrogen in Figures 5(c) and 5(d)) has distances that can hardly be called hydrogen bonds of 3.67 Å at 161° and 3.54 Å at 163°. Since the hydrogen bonds are more than 3 Å, it is more reasonable to ascribe the stability of the adsorbed state to van der Waals interactions. It should be noted that the octa-ammine when forming a surface exposes one NH$_3$ molecule (marked with black hydrogen in Figures 5(c) and 5(d)) bound by only 37.8 kJ/mol almost the same as the first adsorbed NH$_3$. The other phases do not expose weakly bound NH$_3$ when the surface is formed.

V. MICROKINETIC MODEL

With the knowledge of the adsorbed state it is possible to develop a simple microkinetic model of absorption. The

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**TABLE II. Energies of the adsorbed NH$_3$ molecules in half of the stable sites ($\theta = 0.5$) and all of the stable sites ($\theta = 1.0$).**

<table>
<thead>
<tr>
<th>Material</th>
<th>Adsorbed state energy (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$\theta = 0.5$</td>
</tr>
<tr>
<td>Sr(NH$_3$)$_2$Cl$_2$ (010)</td>
<td>27.5</td>
</tr>
<tr>
<td>Sr(NH$_3$)$_2$Cl$_2$ (001)</td>
<td>35.9</td>
</tr>
<tr>
<td>Sr(NH$_3$)$_8$Cl$_2$ (100)</td>
<td>37.0</td>
</tr>
</tbody>
</table>

---

**FIG. 5.** View of the surface atomic structures as calculated with DFT. Green: Cl, blue: N, white: H and Sr is in the center of the yellow coordination polyhedra. The adsorbed NH$_3$ are indicated by having grey hydrogen. The black line indicates the calculational unit cell. (a) The mono-ammine (010) surface with full coverage. The unit cell indicated is 4.69 × 7.72 Å$^2$. (b) The di-ammine (001) surface with half coverage. The unit cell is 6.23 × 8.45 Å$^2$. (c) Top view of the octa-ammine (100) surface with full coverage. Dark and light grey hydrogen indicates $\theta = 0.5$ and $\theta = 1.0$, respectively. The NH$_3$ molecule with black hydrogen is weakly bound as explained in the text. The unit cell indicated is 7.51 × 15.4 Å$^2$. (d) Side view of the octa-ammine surface structure. Half of the calculational vacuum between layers is indicated with the unit cell.
The simplest possible model is adsorption on the surface followed by crossing a barrier to the bulk as shown in Figure 6,

\[ \text{NH}_3 + * \leftrightarrow \text{NH}_3^* \quad \text{adsorption,} \quad (5a) \]

\[ \text{NH}_3^* + * \leftrightarrow \text{NH}_3^2 + * \quad \text{diffusion to bulk,} \quad (5b) \]

where "*" denotes a surface site and "€" a bulk site.

The simple model does not include any transport limitations by, e.g., bulk diffusion inside the crystals. From collision theory, it can easily be verified that for all realistic values, the time-scale to reach equilibrium between the gas phase and the surface is several orders of magnitude faster than all other time-scales in the system.\(^{23}\) As a consequence, the adsorption reaction (5a) can be considered in equilibrium, as in Langmuir adsorption, in which case the surface coverage can be written\(^{23}\)

\[ \theta = K_{\text{surf}} p_{\text{NH}_3} \frac{1}{1 + K_{\text{surf}} p_{\text{NH}_3}}, \quad (6) \]

where \(\theta\) is the surface coverage and the equilibrium constant \(K_{\text{surf}}\) is given from (3) with \(\Delta H_{\text{surf}} = 39.2 \text{ kJ/mol and } \Delta S_{\text{surf}} = 228 \text{ J/(mol K)}\) (Figure 4).

The rate equation for crossing the barrier between surface and bulk is

\[ \gamma \varphi'(t) = k^+ \theta(1 - \varphi) - k^- \varphi(1 - \theta), \quad (7) \]

\[ k^+ = v_{\text{in}} \exp\left\{ \frac{\Delta E_{\text{in}}}{k_b T} \right\}, \quad (8) \]

\[ k^- = v_{\text{out}} \exp\left\{ \frac{\Delta E_{\text{out}}}{k_b T} \right\}, \quad (9) \]

where \(\varphi\) is the occupancy of bulk sites and \(\gamma\) is the ratio between the number of bulk sites and surface sites.

To model the batch absorption experiments depicted in Figure 2, the mass balance between gas molecules and molecules on the surface and in the bulk is needed,

\[ p'(t) = -\frac{N_s k_b T}{V} (\gamma \varphi'(t) + \theta'(t)). \quad (10) \]

Here, \(N_s\) is the number of surface sites and \(V\) is the gas volume in the reactor.

The model can be integrated using the physical parameters of the reactor and the binding energy of the adsorbed state found from the TGA experiments. A complete list of values is found in Table III.

In Figure 7, the microkinetic model is plotted together with the experimental data of Figure 2. It is seen that absorption through an adsorbed state is consistent with the data. The only parameters for which there is no independent prediction are the effective attempt frequencies \(v_{\text{in}}, v_{\text{out}},\) and the barrier \(\Delta E_{\text{in}}\). Values for these parameters are chosen to make the best fit to data. A value of 30 Hz for the frequencies is too low to be a vibration frequency. Instead, it is understood as an effective value taking into account that the atomic configuration in the crystal change significantly when an ammonia molecule moves from one site to another. The barrier \(\Delta E_{\text{in}} = 10 \text{ kJ/mol}\) is the barrier an ammonia molecule has to surpass when going from the surface into the bulk.

It is a quite small barrier, but reflects the fact that the absolute absorption rates do not change much with temperature in the investigated temperature range as seen in Figure 7. The absorption rate, thus, mainly depends on the surface coverage. The model is not fitted to reach the exact equilibrium pressures. As a consequence, the relative pressure \(p_{\text{rel}}\) is not well defined, and the x-axis of Figure 7 linear in pressure in contrast to Figure 2.

VI. DISCUSSION

In the TGA experiments, the sample is fully saturated and comparison to DFT calculations should be done for the

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**TABLE III. Parameters used for the microkinetic model.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(v_{\text{in}})</td>
<td>30 Hz</td>
</tr>
<tr>
<td>(v_{\text{out}})</td>
<td>30 Hz</td>
</tr>
<tr>
<td>(\Delta E_{\text{in}})</td>
<td>10 kJ/mol</td>
</tr>
<tr>
<td>(\Delta E_{\text{out}})</td>
<td>(\Delta E_{\text{in}} - \Delta E_{\text{surf}} + \Delta E_{\text{in}})</td>
</tr>
<tr>
<td>(\Delta E_{\text{surf}})</td>
<td>41.4 kJ/mol</td>
</tr>
<tr>
<td>(\Delta E_{\text{surf}})</td>
<td>39.2 kJ/mol</td>
</tr>
<tr>
<td>(V)</td>
<td>(9 \times 10^{-5} \text{ m}^3)</td>
</tr>
<tr>
<td>(\gamma)</td>
<td>100</td>
</tr>
</tbody>
</table>

---

**FIG. 7.** The desorption measurements at 235 K with blue circles, 277 K with green diamonds, 306 K with red squares and 328 K with cyan triangles. The full lines represent the simple microkinetic model at the corresponding temperature.
octa-ammine phase only. The measured energy of the adsorbed state of 39.2 kJ/mol is very close to the predicted 37.0 kJ/mol for octa-ammine (Table II) for the expected \( \theta = 0.5 \) coverage at the experimental conditions.

In the absorption rate experiments, the saturation degrees correspond to mostly mono-ammine or di-ammine and some octa-ammine. For the saturation degree of 30\%, the sample will be 20\% octa-ammine, if only a stable mono-ammine phase is assumed to exist, or 6\% octa-ammine assuming a stable di-ammine phase to exist.\(^3\) The experimental data should therefore be dominated by either the mono-ammine phase or the di-ammine phase, and the comparison between absorption rate experiments and DFT calculations should therefore be done using the DFT calculations for mono- and di-ammine.

However, fitting the model to the data in Figure 7 with the predicted energy of the adsorbed state of the mono- and di-ammine, give poor fits – especially for the lowest temperature. Only the high surface energy of the octa-ammine gives good fits for all temperatures. This can be explained by the dynamics of adsorption: The ammonia molecules adsorb on the surface, cross the barrier to the bulk sites closest to the surface, and then diffuse deeper into the material. This will induce a concentration gradient, where the bulk sites close to the surface can be dominated by octa-ammine.

Different mechanisms of ammonia diffusion in MgCl\(_2\) ammines have been investigated recently.\(^1\) The diffusion between layers in Ref. 24 can be considered equivalent to the diffusion of adsorbed NH\(_3\) molecules into the bulk of the mono- and di-ammine, as the layers are constructed almost identically in MgCl\(_2\) and SrCl\(_2\). The barriers for diffusion between layers in the mono- and di-ammines were found to be more than 1 eV (\(~96.5\) kJ/mol) that is a much larger value than observed here thereby ruling out diffusion as the mechanism giving rise to the different absorption rates at high and low pressures.

Barriers in the MgCl\(_2\) hexamine are approximately 0.5 eV (~48 kJ/mol), much higher than the estimate of \( \Delta E_{in} \) in our microkinetic model, however, the octa-ammine is a more open structure as evident in Figures 5(c) and 5(d), and more space leads to smaller barriers. If the weakly bound NH\(_3\) (marked with black in Figures 5(c) and 5(d)) is responsible for the adsorbed state, it is easy to imagine a very small barrier for diffusion from the adsorbed state into the bulk as this would just necessitate a move of the NH\(_3\) molecule to an adjacent position in the same Sr–NH\(_3\) complex, and no Sr–N bond would be broken and the barrier \( \Delta E_{in} \) would thus be very small.

Assuming that only a stable octa-ammine phase dominates in the experiments, the DFT surface site energy to compare with the experimental desorption enthalpy would be 37.0 kJ/mol (half coverage). That is in good agreement with the experimental finding of 39.2 kJ/mol, leading us to conclude that stable NH\(_3\) adsorbed states exist and play an important role in the ammonia absorption in metal halide ammines.

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