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Nitrogen isotope fractionations among gaseous and aqueous NH⁺₄, NH₃, N₂, and metal-ammine complexes: Theoretical calculations and applications

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Abstract

Ammonium (NH_4^+), ammonia (NH_3) and N_2 are key nitrogen species in geological nitrogen recycling. NH_3 has also been proposed to play an important role in mobilizing base metals in the form of metal-ammine complexes in hydrothermal fluids. The nitrogen isotope fractionation factors among these nitrogen species in aqueous and gaseous phases are essential parameters to trace source signatures and geochemical properties in geological processes. However, the nitrogen isotope fractionation factors for metal-ammine complexes are largely absent, and the few existing nitrogen isotope fractionation factors for the aqueous NH_4^+ – aqueous NH_3 pair show large discrepancy between experimental results and theoretical calculations. In this study, we employed the density functional theory to systematically calculate the nitrogen isotope fractionation factors among the nitrogen species that may occur in a hydrothermal system, i.e., gaseous N₂, gaseous and aqueous NH₄⁺ and NH₃, and ammine complexes of Co, Zn, Cu, Cd, Ag, Au, and Pt. Based on these new results, the large nitrogen isotope fractionations for the aqueous NH_4^+ – aqueous NH_3 pair observed in previous experimental studies can be well explained by a combined effect of an equilibrium isotope fractionation between aqueous NH_4^+ and aqueous NH_3 and a kinetic isotope fractionation during NH₃ degassing from the solution. This suggests that the nitrogen isotopic behavior during NH₃ degassing in natural hydrothermal system can be more complicated than previous thought. A numeric model is thus established here to quantify the combined isotopic effect on partial NH₃ degassing. Using the new results of metal-ammine complexes, we also tested the hypothesis that nitrogen mobilization could be controlled by copper-ammine complex based on the copper concentration- δ^{15} N relationship previously observed in meta-gabbros.

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Keywords: Nitrogen isotope fractionation; Ammonium; Metal-ammine complex; Ammonia degassing; Hydrothermal fluid

1. INTRODUCTION

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https://doi.org/10.1016/j.gca.2020.12.010 0016-7037/© 2020 Elsevier Ltd. All rights reserved. Geological nitrogen cycle, which is commonly mediated by hydrothermal fluids (e.g., Busigny and Bebout, 2013; Li et al., 2007, 2014; Halama et al., 2010, 2017), involves transformation of various nitrogen species within or between Earth's reservoirs, e.g., the atmosphere, crust, and mantle

(e.g., Halama et al., 2014; Mikhail and Sverjensky, 2014; Bebout et al., 2016). In particular, the nitrogen transfer between the atmosphere, in which nitrogen occurs mainly as N₂, and the lithosphere, in which nitrogen mainly occurs as NH₄⁺ substituting K⁺ in mineral lattices (Honma and Itihara, 1981), may pass through an intermediate nitrogen species of NH₃ (e.g., Brandes et al., 1998; Li et al., 2007, 2009, 2014). NH₃ may also play an important role in alkaline fluids occurring in a variety of geological settings, such as the deep aquifer in ophiolites (e.g., the Oman ophiolite, the Coast Range ophiolite: see Holm et al., 2006 and reference therein), the deep subsurface fracture waters in Precambrian cratons (e.g., South Africa; Onstott et al., 2006), ridge flank hydrothermal systems (e.g., Lost City, Rainbow; see the discussion in Li et al., 2012 and reference therein), subduction zones (e.g., Mariana forearc; Wheat et al., 2008), hot spots (e.g., Yellowstone; Holloway et al., 2011), and alkaline lakes (e.g., Lake Bosumtwi, Ghana; Talbot and Johannessen, 1992). This is because NH_4^+ in alkaline fluids can be dissociated into NH₃, which can be further removed from the fluid by degassing. The relative proportions of NH_4^+ , NH_3 , and N_2 in hydrothermal fluid are strongly dependent on the redox and pH conditions (e.g., Duit et al., 1986; Li et al., 2012; Li and Keppler, 2014; Mikhail and Sverjensky, 2014).

Besides being a key species in geological nitrogen cycle, NH₃ may also play an important role in hydrothermal enrichment and mobilization of base metals because it is an effective ligand to form metal-ammine complexes with transition metals, such as Cu (Hathaway and Tomlinson, 1970; Han et al., 1974; Chu et al., 1978), Ni (Gupta and Sarpal, 1967), Co (Meek and Ibers, 1970), Zn (Eßmann, 1995), and Ag (Geddes and Bottger, 1969; Widmer-Cooper et al., 2001; Fox et al., 2002). This property of NH₃ has been applied in industry to recover transition metals from ore deposits (e.g., Meng and Han, 1996; Katsiapi et al., 2010). In natural hydrothermal system, a possible coupling between NH₃ and Cu has been proposed based on geochemical signatures of hydrothermally altered gabbros (Busigny et al., 2011). If NH₃ can promote the solubility and mobility of base metals in hydrothermal system, it may potentially act as an important agent for ore genesis (Irving and Williams, 1953; Martell and Hancock, 1996).

Nitrogen isotopes have been used as a robust tool to trace nitrogen remobilization (e.g., Bebout et al., 1999; Busigny et al., 2005; Li et al., 2007) and geological nitrogen recycling (e.g., Bebout and Fogel, 1992; Busigny et al., 2003; Svensen et al., 2008; Halama et al., 2010; Li et al., 2009, 2014). In order to apply nitrogen isotope system to constrain nitrogen sources and fluxes in geological nitrogen recycling pathways, the nitrogen isotope fractionation factors between involved nitrogen species are crucial prerequisite parameters. However, the nitrogen isotope fractionation factors between aqueous NH₃ and metalammine complexes have not been well constrained yet, despite some early efforts (e.g., Gupta and Sarpal, 1967; Ishimori, 1960a). Several previous studies (Urey, 1947; Scalan, 1958; Hanschmann, 1981; Petts et al., 2015) have

investigated the equilibrium nitrogen isotope fractionations among NH_4^+ , NH_3 and N_2 by theoretical calculations and given very different results. These calculations were based on vibrational frequencies of nitrogen species in gas phases. However, in natural systems, particularly in hydrothermal systems, NH₄⁺ and NH₃ mostly exist in aqueous phases (hereafter referred as $NH_4^+ \cdot nH_2O$ or $NH_{4aqueous}^+$), and NH_3^- ·nH₂O or NH_{3(aqueous)}, respectively). One previous laboratory experimental study (Li et al., 2012) showed that, under hydrothermal condition, partial dissociation of $NH_4^+ \cdot nH_2O$ coupled with complete degassing of the produced NH₃ induced large ¹⁵N enrichments in the remaining $NH_4^+ \cdot nH_2O$, which cannot be explained by the theoretically predicted equilibrium fractionation factors between the NH⁺_(gaseous) - NH_{3(gaseous)} pair (Urey, 1947; Scalan, 1958; Hanschmann, 1981). To solve the discrepancy between the experimental and theoretical results, Li et al. (2012) proposed that the $NH_4^+ \cdot nH_2O$ dissociation – NH_3 degassing process involved an intermediate step that NH₄⁺·nH₂O was first equilibrated with NH3·nH2O, from which NH3 was further exsolved and degassed (Li et al., 2012). Such a process can be described as Eq. (1):

$$NH_{4}^{+} \cdot nH_{2}O + OH^{-} \rightleftharpoons NH_{3} \cdot nH_{2}O + H_{2}O$$

$$\rightarrow NH_{3(gaseous)} + (n + 1)H_{2}O$$
(1)

Given that the produced NH₃·nH₂O was completely removed by NH₃ degassing in the experiments, and more importantly, the ¹⁵N enrichments in the remaining NH₄⁺-·nH₂O apparently fitted well to a batch model assuming equilibrium isotope fractionation between NH_4^+ nH_2O and NH₃·nH₂O, Li et al. (2012) interpreted the strong ¹⁵N enrichments observed in the remaining NH_4^+ nH_2O as a result of large equilibrium isotope fractionations (e.g., +45.4‰ at 23 °C and + 33.5‰ at 70 °C) between NH₄⁺-·nH₂O and NH₃·nH₂O without considering kinetic isotopic effect from NH₃ degassing. However, a recent laboratory experimental study (Deng et al., 2018) found significant kinetic nitrogen isotopic effect (-8.2% at 21 °C, and -5.2‰ at 70 °C) during degassing of NH_{3(gas)} from NH₃-·nH₂O. In addition, a recent theoretical calculation (Walters et al., 2019) using relatively simple HF/6-31G(d) and B3LYP/6-31G(d) levels of theory yielded significantly different nitrogen isotope fractionation factors between $NH_4^+ \cdot nH_2O$ and $NH_3 \cdot nH_2O$. Thus, it is necessary to reassess the isotopic behavior during NH₃ degassing process described by Eq. (1).

To fill these knowledge gaps, we employed theoretical calculations to determine the equilibrium isotope fractionations among gaseous N_2 and several other nitrogen species related to NH_3 in hydrothermal fluids, including NH_4^+ and NH_3 in both gaseous and aqueous phases and metalammine complexes of several important base metals, i.e., Co, Ni, Cu, Zn, Cd, Ag, Au, and Pt. Theoretical calculation is a robust and efficient way to estimate the equilibrium isotope fractionation factors among these species, given that they are difficult to be characterized by laboratory experiments.

2. METHOD

2.1. Equilibrium isotope fractionation theory

The equilibrium isotope fractionation factor between a species and its atomic form can be described by the β factor (Urey, 1947; Bigeleisen and Mayer, 1947). The details of the Urey-Bigeleisen-Mayer model for theoretical calculation of equilibrium isotope fractionation factor have been intensively reviewed in the literature (e.g., Richet et al., 1977; Schauble et al., 2004; Liu et al., 2010; Young et al., 2015; Dauphas and Schauble, 2016; Blanchard et al., 2017). In brief, for an isotope exchange reaction between species A and B:

$$\mathbf{A} + \mathbf{B}^* \rightleftharpoons \mathbf{A}^* + \mathbf{B} \tag{2}$$

where the species with * contain the rare isotope (i.e., ^{15}N in this case) and the ones without * contain the most abundant isotope (i.e., ^{14}N in this case). The nitrogen equilibrium isotope fractionation factor between A and B (i.e., α_{A-B}) can be expressed as the ratio of the $^{15}\beta$ factors of A and B:

$$\alpha_{A-B} = \frac{\left[\frac{15}{N}/\frac{14}{N}\right]_{A}}{\left[\frac{15}{N}/\frac{14}{N}\right]_{B}} = \frac{\frac{15}{\beta_{A}}}{\frac{15}{\beta_{B}}}$$
(3)

in which the ${}^{15}\beta$ factors can be estimated in harmonic approximation (Richet et al., 1977; Cao and Liu, 2012) using the Urey-Bigeleisen-Mayer model (Urey, 1947; Bigeleisen and Mayer, 1947):

$${}^{15}\beta = \prod_{i}^{N} \left(\frac{u_{i}^{*}}{u_{i}}\right) \left(\frac{e^{-u_{i}^{*}/2}}{e^{-u_{i}/2}}\right) \left(\frac{1-e^{-u_{i}}}{1-e^{-u_{i}^{*}}}\right)$$
(4)

$$u_i = \frac{hv_i}{k_B T} \tag{5}$$

in which v_i denotes the *i*th harmonic vibration frequency; *h* denotes the Planck constant; k_B denotes the Boltzmann constant; *T* denotes the temperature in Kelvin; and *N* denotes the harmonic vibrational modes (for non-linear molecules, N = 3n - 6; n is the total number of atoms).

2.2. Calculation methods

Density functional theory (DFT) calculation with the Urey-Bigeleisen-Mayer equation offers a reliable approach to theoretically estimate a β factor (Liu and Tossell, 2005). It is based on quantum chemical theories to predict the optimized molecular structure of a given system and calculate its vibrational frequencies. All optimization and harmonic vibrational frequency calculations for the ground states were performed by the software Gaussian 16 (Frisch et al., 2016). All calculations have been carried out with the B3LYP exchange-correlation functional (Lee et al., 1988; Becke, 1993), which has been widely employed in vibrational frequency-related calculations (e.g., Rustad et al., 2010; Li and Liu, 2011; Eldridge et al., 2016; Zhang and Liu, 2018a,b). The 6-311++G (d,p) basis set (Frisch et al., 1984) was used for H, O, and N atoms. LANL2TZ basis set (Hay and Wadt, 1985; Roy et al., 2008) was used for Zn, Co, Cu, Ag, Au, Cd, and Pt atoms to lessen

computation time yet with negligible loss of accuracy. No scaling factor was applied because the systematic errors only influence the absolute β values but are largely cancelled during the α calculations (Schauble et al., 2006; Méheut et al., 2007).

2.3. Anharmonic effect on nitrogen species

The anharmonic correction on the Urey-Bigeleisen-Mayer equation has been discussed in previous studies (Richet et al., 1977; Liu et al., 2010; Petts et al., 2015; Zhang and Liu, 2018a). One general consent is that anharmonic effect plays a greater role in vibrations involving light elements (e.g., H, B), although the magnitude of isotope fractionations caused by anharmonic effect is still under debate (Liu et al., 2010; Petts et al., 2015). To assess the magnitude of anharmonic effect on the nitrogen species in this study, we also calculated the nitrogen isotope fractionations with and without anharmonic correction for gaseous NH₄⁺ and gaseous NH₃.

2.4. Solvent effect on nitrogen species

When a substance is dissolved in a solution, its weak interaction with solvent molecules can cause large uncertainty of local structural configurations, and therefore influence the estimated ${}^{15}\beta$ values. To assess this solvent effect, we added water molecules surrounding a target species to simulate in a realistic way the environment of an aqueous solution (e.g., Rustad et al., 2008, 2010; Zhang and Liu, 2014; He and Liu, 2015; Gao et al., 2018; Zhang and Liu, 2018a,b), using the commonly used explicit solvation modes (i.e., water-droplet method; Liu and Tossell, 2005; Li and Liu, 2011; Gao et al., 2018).

For NH₃ and NH₄⁺, we built four starting cluster models with 6 water molecules as the first step. The structures were optimized to the local energy minimum with zero imaginary frequency. Subsequently, additional 6 water molecules were added to the second shell of the optimized structures and were optimized to the local energy minimum again. This process was repeated until the calculated ¹⁵ β value reached convergence. For the NH₃·nH₂O models, the convergence cluster has 30 water molecules. For the NH₄⁺·nH₂O models, the convergence cluster has 36 water molecules.

For dissolved metal-ammine complexes, the solvent effect was also assessed (using $Zn(NH_3)_6^{2+}$ and $Ni(NH_3)_6^{2+}$ as examples), following previous studies (e.g., Rudolph et al., 2000; Hill and Schauble, 2008) by adding a hydration sphere (12H₂O) on the studied molecular cluster.

3. RESULTS

3.1. Anharmonic effect on isotope fractionation between gaseous NH_4^+ and NH_3

The results of harmonic vibrational frequencies (w_i) and anharmonicity constants (x_{ij}) for NH₃ and NH⁺₄ are listed in Table 1. Comparison between the $\ln \alpha_{NH^+_4(gameous)} - NH_3(gameous)}$ results with and without anharmonic corrections indicates

Table 1 Harmonic vibrational frequencies (ω_i) and anharmonicity constants (x_{ij}) for NH₃ and NH₄⁺.

	$^{14}\mathrm{NH}_3$	¹⁵ NH ₃		$^{14}\mathrm{NH_4^+}$	$^{15}\mathrm{NH_{4}^{+}}$
w1	1005.6647	1000.2964	w ₁	1489.3992	1482.4191
W2	1668.9006	1665.5001	W2	1489.5274	1482.5471
W3	1668.9599	1665.5593	W3	1489.5677	1482.5874
w4	3480.3335	3478.3005	W_4	1727.4986	1727.4986
W5	3607.0377	3597.1959	W5	1727.5563	1727.5563
w ₆	3607.3700	3597.5267	w ₆	3371.2896	3371.2895
x ₁₁	-44.5655	-44.0991	W7	3474.2659	3464.5960
x ₁₂	-7.4656	-7.3437	W8	3474.6356	3464.9637
x ₁₃	-60.6278	-60.1753	W9	3474.9376	3465.2644
x ₁₄	-99.6980	-99.5133	x ₁₁	-35.9750	-35.6432
x ₁₅	-15.1020	-14.9203	x ₁₂	4.9813	4.8697
x ₁₆	25.8310	25.4004	x ₁₃	-63.5821	-63.4649
x ₂₂	-6.8370	-12.9044	x ₁₄	-16.5959	-16.4789
x ₂₃	-14.9178	-14.7277	x ₁₅	-11.3300	-11.2686
x ₂₄	-22.0048	2.4055	X ₁₆	-11.1957	-11.0121
x ₂₅	-12.3932	-12.4064	x ₁₇	-68.8225	-68.4742
x ₂₆	-14.9338	-15.0119	X ₁₈	-13.5188	-13.3919
x ₃₃	-44.2150	-43.7427	x ₁₉	-11.9319	-11.9304
x ₃₄	-99.6208	-99.4716	x ₂₂	-35.9787	-35.6468
x ₃₅	-7.3950	-7.2582	x ₂₃	-64.3850	-64.3460
x ₃₆	26.1471	25.7100	x ₂₄	-16.6125	-16.4950
X44	-26.5479	-26.5878	X ₂₅	-11.1808	-10.9972
x ₄₅	-21.9967	2.2832	x ₂₆	-11.3447	-11.2835
X46	18.0323	18.1441	X ₂₇	-68.0429	-67.6153
X ₅₅	-6.8509	-12.8954	x ₂₈	-13.5173	-13.3903
X56	-14.7680	-14.8366	X29	-11.9177	-11.9169
x ₆₆	-61.1998	-60.5714	X ₂₉ X ₃₃	-15.9055	-15.9056
	0111770	0010711	x ₃₄	-23.4745	-23.4746
			X34 X35	-4.8519	-5.3443
			x ₃₅ x ₃₆	-4.9011	-5.3913
			X ₃₇	-63.9414	-63.8669
			X ₃₈	-23.4652	-23.4654
			X39	-4.8868	-5.3762
				-1.6234	-1.6235
			X ₄₄	-3.5137	-3.5376
			x ₄₅	-3.5439	-3.5675
			x ₄₆	-12.0062	-11.8748
			X ₄₇	-3.4073	-3.4073
			X48	-18.4950	-18.4922
			X49		
			X55	-6.7531 4.4641	-6.5817
			x ₅₆		4.5306
			x ₅₇	-11.9000	-11.9027
			X58	-13.4279	-13.4342
			X59	-6.1422	-5.8856
			X66	-6.7632	-6.5921
			x ₆₇	-11.8785	-11.8801
			X68	-13.4427	-13.4488
			x ₆₉	-6.1608	-5.9043
			X ₇₇	-17.5888	-17.3853
			X78	-18.0933	-17.9816
			X79	-10.6254	-10.3797
			X88	-1.6219	-1.6220
			X89	1.3786	1.3461
			X99	-4.1523	-4.0280
ZPE _{harm}	7519.1332	7502.1895	ZPE _{harm}	10859.3390	10834.3611
ZPE _{anh}	7394.8057	7378.5208	ZPE _{anh}	10683.4961	10659.4508

that anharmonic effect on $\ln \alpha_{NH_4^+(gaseous)}^{-NH_3(gaseous)}$ is small, i.e., -1.4% at 0 °C, -0.6% at 400 °C, and -0.3% at 1000 °C (Table 2). Therefore, we did not apply the anharmonic

corrections for the other nitrogen species in this study. For consistency, we used all the harmonic results to calculate the $1000 \ln \alpha$ values.

Table 2 $\ln \alpha_{\rm NH4+(gaseous) - NH3(gaseous)}$ for harmonic and anharmonic calculations at 0–1000 °C.

T (°C)	Harmonic	Anharmonic
0	35.2	33.8
20	32.4	31.0
40	29.8	28.5
60	27.5	26.4
80	25.5	24.4
100	23.8	22.7
200	17.3	16.5
400	10.2	9.6
600	6.7	6.2
800	4.7	4.4
1000	3.6	3.3

3.2. Isotope fractionations between NH_4^+ and NH_3 in gaseous and aqueous phases

Fig. 1 illustrates the optimized geometries for gaseous NH_3 and NH_4^+ , and their aqueous phases represented by $NH_3 \cdot 30H_2O$ and $NH_4^+ \cdot 36H_2O$, respectively. The coordinates of these optimized geometries are listed in the Supplementary Data. Our calculations yielded the N-H bond length and H-N-H bond angle as 1.01(4) Å and 107.9° , respectively, for gaseous NH_3 , and 1.02(6) Å and 109.5° , respectively, for gaseous NH_4^+ . These values are consistent with previous theoretical calculation and experimental results, e.g., 1.01(2) Å for the N-H bond length and

106.67° for the H-N-H bond angle in gaseous NH₃ (Haynes, 2014), and 1.02(7) Å for the N-H bond length and 109.5° for the H-N-H bond angle in gaseous NH₄⁺ (Chen and Davidson 2001). The NH₃ molecule is hydrated in all explicit solvent models, in which N forms a 1.65–1.69 Å hydrogen bond with H of a water molecule.

Table 3 lists the calculated ${}^{15}\beta$ results of NH₃ and NH₄⁺ in gaseous phases as well as aqueous phases hydrated by 6-30 H₂O molecules in 4 different configurations at 25 °C. The results show that the ${}^{15}\beta$ values increase significantly from gaseous NH₃ ($^{15}\beta = 1.0687$) to hydrated NH₃, whose $^{15}\beta$ values vary strongly with an increase in the surrounding H₂O molecule numbers until converge at 1.0776 when the H₂O molecule number reaches 30. This solvent effect results in large equilibrium isotope fractionations between $NH_{3(aqueous)}$ and $NH_{3(gaseous)}$ (e.g., $ln\alpha_{NH_{3(aqeous)}-NH_{3(gaseous)}} =$ +8.3‰ at 25 °C). In contrast, the ¹⁵ β value of NH₄⁺ is not significantly shifted after hydration (Table 3), resulting in negligible isotope fractionation between gaseous and aqueous NH₄⁺, e.g., the $ln\alpha_{NH_{4(ageous)}^{+}-NH_{4(gaeous)}^{+}}$ value is +0.3‰ at 25 °C, and mostly within the calculation uncertainty of $\pm 0.5\%$ at other temperatures (Tables 4–5). Therefore, we will not specifically distinguish between aqueous and gaseous NH_4^+ in the discussions below.

The 1000ln β values at selected temperatures are given in Table 4, and the general equations describing the temperature-dependent ¹⁵ β values for individual species are given in Table 5 and plotted in Fig. 3. The temperature-dependent equilibrium isotope fractionations

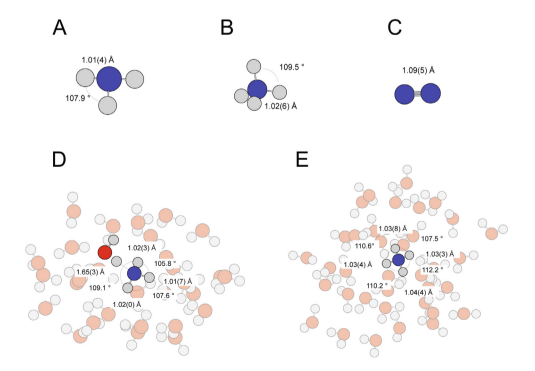


Fig. 1. Optimized geometries for gaseous NH_3 (A), NH_4^+ (B), N_2 (C), and representative local configuration of NH_3 ·30H₂O (D) and NH_4^+ ·36H₂O (E).

Table 3 Calculated ¹⁵ β factors (¹⁵N/¹⁴N) of ammonia and ammonium in gaseous and aqueous phases at B3LYP/6-311G++(d,p) level at 25 °C.

Species	¹⁵ β	$1000 \ln^{15} \beta$		$^{15}\beta$	$1000 \ln^{15} \beta$
Gaseous phase					
NH ₃	1.0687	66.4	NH_4^+	1.1031	98.1
Aqueous phase					
$NH_3(H_2O)_6A$	1.0766	73.8	$NH_4(H_2O)_6^+A$	1.1032	98.2
$NH_3(H_2O)_6_B$	1.0767	73.9	$NH_4(H_2O)_6^+B$	1.1038	98.8
$NH_3(H_2O)_6_C$	1.0757	73.0	$NH_4(H_2O)_6^+C$	1.1038	98.8
$NH_3(H_2O)_6_D$	1.0786	75.7	$NH_4(H_2O)_6^+D$	1.1039	98.8
Average*	1.0769 ± 0.0011	74.1 ± 1.0	Average	1.1037 ± 0.0003	98.6 ± 0.3
NH ₃ (H ₂ O) ₁₂ A	1.0799	76.9	$NH_4(H_2O)_{12}^+A$	1.1032	98.2
$NH_3(H_2O)_{12}B$	1.0766	73.8	$NH_4(H_2O)_{12}^+B$	1.1034	98.4
$NH_{3}(H_{2}O)_{12}C$	1.0781	75.2	$NH_4(H_2O)^+_{12}C$	1.1032	98.2
$NH_3(H_2O)_{12}$ D	1.0781	75.2	$NH_4(H_2O)_{12}^+$ D	1.1035	98.5
Average	1.0782 ± 0.0012	75.3 ± 1.1	Average	1.1033 ± 0.0001	98.3 ± 0.1
NH ₃ (H ₂ O) ₁₈ A	1.0787	75.8	$NH_4(H_2O)_{18}^+A$	1.1039	98.8
$NH_3(H_2O)_{18}B$	1.0781	75.2	$NH_4(H_2O)_{18}^+B$	1.1031	98.1
$NH_{3}(H_{2}O)_{18}C$	1.0778	74.9	$NH_4(H_2O)_{18}^+C$	1.1039	98.8
$NH_{3}(H_{2}O)_{18}D$	1.0762	73.4	$NH_4(H_2O)_{18}^+$ D	1.1026	97.7
Average	1.0777 ± 0.0009	74.8 ± 0.9	Average	1.1034 ± 0.0006	98.4 ± 0.5
NH ₃ (H ₂ O) ₂₄ A	1.0776	74.7	$NH_4(H_2O)^+_{24}A$	1.1039	98.8
$NH_3(H_2O)_{24}B$	1.0778	74.9	$NH_4(H_2O)^+_{24}B$	1.1041	99.0
$NH_3(H_2O)_{24}$ C	1.0776	74.7	$NH_4(H_2O)_{24}^+C$	1.1042	99.1
$NH_{3}(H_{2}O)_{24}D$	1.0773	74.5	$NH_4(H_2O)_{24}^+D$	1.1035	98.5
Average	1.0776 ± 0.0002	74.7 ± 0.2	Average	1.1039 ± 0.0003	98.9 ± 0.2
$NH_3(H_2O)_{30}$ A	1.0771	74.3	$NH_4(H_2O)_{30}^+A$	1.1032	98.2
$NH_3(H_2O)_{30}B$	1.0777	74.8	$NH_4(H_2O)_{30}^+B$	1.1044	99.3
NH ₃ (H ₂ O) ₃₀ C	1.0780	75.1	$NH_4(H_2O)_{30}^+C$	1.1037	98.7
$NH_{3}(H_{2}O)_{30}D$	1.0776	74.7	$NH_4(H_2O)_{30}^+D$	1.1028	97.9
Average	1.0776 ± 0.0003	74.7 ± 0.3	Average	1.1035 ± 0.0006	98.5 ± 0.5
Ū.			$NH_4(H_2O)^+_{36}A$	1.1028	97.9
			$NH_4(H_2O)_{36}^+B$	1.1030	98.0
			$NH_4(H_2O)_{36}^+C$	1.1034	98.4
			$NH_4(H_2O)^+_{36}D$	1.1031	98.1
			Average	1.1031 ± 0.0002	98.1 ± 0.2
Preferred value**	1.0776	74.7	Preferred value**	1.1035	98.5

* The "Average" data in bold are the mean values of 4 configurations (i.e., configurations A,B, C, and D).

^{**} The preferred ¹⁵ β value of aqueous NH₃(H₂O)_n are the average results of 8 configurations (4 × NH₃(H₂O)₂₄ and 4 × NH₃(H₂O)₃₀), and the preferred ¹⁵ β value of NH₄(H₂O)_n⁺ are the average results of 12 configurations (4 × NH₄(H₂O)₂₄, 4 × NH₄(H₂O)₃₀), and 4 × NH₄(H₂O)₃₆).

between species are given in Table 6 and plotted in Fig. 4. It is noted that the different solvent effects on NH_4^+ and NH_3 result in much larger isotope fractionation in the $NH_{4(aqueous)}^+ - NH_{3(gaseous)}$ pair (e.g., +32.1‰ at 25 °C) than the $NH_{4(aqueous)}^+ - NH_{3(aqueous)}$ pair (e.g., +23.8‰ at 25 °C).

3.3. Isotope fractionations between NH_4^+ and N_2

The optimized geometry of gaseous N_2 is also shown in Fig. 1. Our calculations yielded the N-N bond length as 1.09(5) Å, which is consistent with previous published data (e.g., 1.0975 Å; Sutton and Bowen, 1958).

The calculated 1000ln β results at selected temperatures are listed in Table 4 with a general description equation given in Table 5. The results show that N₂ is more enriched in ¹⁵N than both gaseous and aqueous NH₃, but more depleted in ¹⁵N than gaseous and aqueous NH₄⁺ (Table 4). Given that experimental studies have demonstrated that the isotope fractionation between gaseous N₂ and dissolved N₂ is very small (<1‰ at 0 °C; Klots and Benson, 1963), our results of gaseous N_2 can also be approximately used for aqueous N_2 . Accordingly, our calculation results self-consistently put aqueous N_2 in the right ¹⁵N enrichment order between aqueous NH₃ and aqueous NH₄⁺.

3.4. Metal – ammine complexes

The geometries of metal-ammine complexes are still not well constrained. In this study, we select the complex structure of the minimum-energy level from previous studies as the dominant metal-ammine complex species (see below). The optimized geometries are shown in Fig. 2, and their coordinates are listed in the Supplementary Data. The calculated ¹⁵ β values at selected temperatures are listed in Table 4 with general description equations being listed in Table 5. A brief summary for each metal is given below.

Zinc-ammine complexes $Zn(NH_3)_n^{2+}$ (n = coordination number) with n values varying from 4 to 6 have been reported (Kim et al., 1993; Fatmi et al., 2006; Fatmi et al., 2010). However, our calculations only obtained

() ()	\mathbf{Z}_{2}^{2}	$\rm NH_3$	$\rm NH_3$	$\rm NH_4^+$	$\rm NH_4^+$	Metal –	ammine complexes*	omplexes*									
	(gas)	(gas)	(aq)	(gas)	(aq)	Zn ²⁺ (6)	Ni ²⁺ (6)	Co ²⁺ (6)	Co ³⁺ (6) ^H	Co ³⁺ (6) ^L	Cd ²⁺ (6)	Cd ²⁺ (4)	Cu ²⁺ (4)	Cu ⁺	A_{g^+}	Au^+ (2)	Pt ⁺
	90.9	73.9	83.5	109.1	110.1	84.3	86.9	85.5	87.6	97.2	82.8	86.4	6.06	94.0	89.6	98.2	98.7
_	83.6	67.8	76.4	100.2	100.7	77.1	79.3	78.1	80.0	88.5	75.7	78.8	82.9	85.7	81.8	89.4	89.7
_	77.1	62.6	70.2	92.4	92.5	70.7	72.8	71.7	73.4	81.0	69.6	72.4	76.0	78.5	75.0	81.9	82.1
_	71.5	58.0	64.7	85.5	85.4	65.3	67.1	66.1	67.6	74.5	64.2	66.7	70.0	72.2	69.1	75.3	75.6
_	66.5	53.9	60.0	79.5	79.2	60.5	62.1	61.3	62.6	68.8	59.5	61.8	64.8	66.7	63.9	69.5	69.7
100	62.0	50.3	55.8	74.1	73.6	56.2	57.7	56.9	58.1	63.7	55.3	57.4	60.1	61.9	59.4	64.4	64.7
00	45.5	36.9	40.4	54.2	53.3	40.8	41.7	41.2	41.9	45.6	40.2	41.5	43.3	44.4	42.9	46.1	46.2
0	27.3	22.6	24.4	32.9	32	24.5	25.0	24.8	25.1	27.0	24.3	24.9	25.9	26.4	25.7	27.3	27.4
0	18.0	15.4	16.4	22.1	21.3	16.5	16.8	16.6	16.7	17.9	16.3	16.7	17.3	17.6	17.2	18.1	18.1
800	12.7	11.0	11.7	15.8	15.1	11.7	11.9	11.8	11.9	12.7	11.6	11.9	12.2	12.5	12.2	12.9	12.9
000	9.4	8.3	8.7	11.8	11.2	8.8	9.0	8.9	8.9	9.5	8.7	8.9	9.2	9.4	9.2	9.7	9.7

meaningful results for the 6-coordinated complex Zn $(NH_3)_6^{2+}$ with a multiplicity of 1. The Zn-N bond length in the yielded geometry is 2.31 Å, which is close to the previously suggested value of 2.291 Å (Kim et al., 1993). The yielded ¹⁵ β is 1.0783 for Zn(NH₃)²⁺ at 25 °C. The ¹⁵ β value shows an insignificant increase with hydration, to 1.0788 for $Zn(NH_3)_6^{2+}.6H_2O$, and 1.0792 for $Zn(NH_3)_6^{2+}.12H_2O$ at 25 °C.

Nickel-ammine complex can display various geometries with coordination numbers varying from 4 to 6. In this study, we focus on the 6-coordinated complex Ni(NH₃) $_{6}^{2+}$ with a multiplicity of 3, which is considered to be more stable (e.g., Paul et al., 2004; Varadwaj et al., 2008; Casanova and Head-Gordon, 2009; Varadwaj et al., 2009). The yielded Ni-N bond length of Ni(NH₃) $_{6}^{2+}$ is 2.20 Å, which is consistent with the previously suggested value of 2.205 Å and close to the crystallographically observed mean value of 2.135 Å (Varadwaj et al., 2008). The calculated $^{15}\beta$ value is 1.0807 for $Ni(NH_3)_6^{2+}$ at 25 °C, which slightly increases to 1.0817 when an additional hydration sphere of 12 H₂O is added (i.e., Ni $(NH_3)_6^{2+} \cdot 12H_2O).$

Cobalt can complex with NH₃ at valence states of II and III. The optimized structures of the complexes are 6-coordinated $Co(NH_3)_6^{2+}$ and $Co(NH_3)_6^{3+}$ (Barnet et al., 1966; Meek and Ibers, 1970; Müller and Kraus, 2015). Co $(NH_3)_6^{2+}$ is believed to be more stable at high-spin state (Schmiedekamp et al., 2002) and thus has a multiplicity of 4. While $Co(NH_3)_6^{3+}$ has been considered as a low-spin complex (e.g., Williams, 1979) and thus has a multiplicity of 1, our single-atom test yielded the lowest energy of Co³⁺ at high-spin state with a multiplicity of 5. Therefore, we reported the results of both low- and high-spin states for Co³⁺ here. Our calculation gave an average Co-N bond length of 2.26 Å for $Co(NH_3)_6^{2+}$, which is close to previously published data of 2.257-2.264 Å (Schmiedekamp et al., 2002; Varadwaj and Marques, 2010). The yielded $^{15}\beta$ value of Co(NH₃) $_6^{2+}$ is 1.0794 at 25 °C. The Co-N bond length for $Co(NH_3)_6^{3+}$ is 2.03 Å at low-spin state, which is consistent with previously calculation results (2.033 Å; Rotzinger, 2009), but increases to 2.20 Å at high-spin state. This results in a large difference in the ${}^{15}\beta$ values, e.g., 1.0814 at low-spin state and 1.0904 at high-spin state at 25 °C.

Cadmium-ammine complex is considered to be the stable in both 4-coordinated (i.e., $Cd(NH_3)_{4}^{2+}$; multiplicity = 1) and 6-coordinated (i.e., $Cd(NH_3)_6^{2+}$; multiplicity = 2) forms (Nilsson et al., 2007; Zeng et al., 2015). The yielded Cd-N bond lengths are 2.35 Å for Cd $(NH_3)_4^{2+}$, which is higher than the value of 2.02 Å reported by Zeng et al. (2015), and 2.48 Å for $Cd(NH_3)_6^{2+}$, which is higher than the value of 2.35 Å reported by Nilsson et al. (2007). The calculated ${}^{15}\beta$ values at 25 °C are 1.0802 for $Cd(NH_3)_4^{2+}$ and 1.0769 for $Cd(NH_3)_6^{2+}$.

Copper can complex with NH₃ at valence states of I and II. The 4-coordinated Cu(II)-ammine complex is suggested to be more stable than the 5- and 6-coordinated complexes (Pavelka and Burda, 2005). For the 4-coordinated Cu $(NH_3)_4^{2+}$ (multiplicity = 2), the calculated structure has a Cu-N bond length of 2.07 Å, which is consistent with exper-

Table 4

Table 5

Polynomial fit parameters of calculated ¹⁵ β factors in the form of 1000·ln(¹⁵ β) = C₀ + C₁*10⁻¹x - C₂*10⁻²x² + C₃10⁻³x³ - C₄*10⁻⁴x⁴ + C₅*10⁻⁵x⁵ - C₆10⁻⁶x⁶ in which x = 10⁶/T² and T is temperature in Kelvin (valid from 273 to 1273 K)

Compound	C_0	C_1	C_2	C_3	C_4	C ₅	C_6
N _{2(gaseous)}	0.27	160.78	230.62	319.21	285.31	140.65	28.766
NH _{3(gaseous)}	0.65	137.43	238.30	383.41	373.46	191.98	39.882
NH _{3(aqueous)}	0.75	142.11	214.55	315.96	286.52	139.19	27.668
NH ⁺ _{4(gaseous)}	1.13	189.79	280.17	400.11	360.31	176.23	35.521
NH ⁺ _{4(aqueous)}	0.51	190.47	299.05	465.60	448.62	230.51	48.098
$Zn(NH_3)_6^{2+2}$	0.74	142.77	216.00	323.78	300.98	150.18	30.635
$Ni(NH_3)_6^{2+}$	0.84	143.69	208.87	305.13	277.40	135.51	27.075
$Co(NH_3)_6^{2+}$	0.77	143.52	215.04	321.63	298.11	148.06	30.022
$Co(NH_3)_6^{3+}$ (H)*	0.79	144.13	209.51	308.78	282.83	138.96	27.903
$Co(NH_3)_6^{3+}(L)^*$	0.70	154.72	216.85	318.04	291.16	143.41	28.932
$Cd(NH_3)_6^{2+}$	0.66	142.64	219.61	326.76	298.19	145.28	28.875
$Cd(NH_3)_4^{2+}$	0.83	142.65	203.83	289.48	254.75	120.34	23.266
$Cu[(NH_3)_4]^{2+}$	0.73	148.83	212.48	301.05	260.08	119.14	22.149
$Cu[(NH_3)_2]^+$	0.70	153.55	229.98	355.89	339.85	172.99	35.831
$Ag(NH_3)^{2+}_2$	0.85	147.16	212.91	313.78	290.42	145.54	29.982
$Au(NH_3)^{2+}_2$	0.97	152.43	197.47	263.92	217.27	95.08	16.882
$Pt(NH_3)_2^{2+}$	0.85	154.95	211.42	302.56	268.89	127.71	24.704

* H and L in $Co(NH_3)_6^{3+}$ denote high spin state (multiplicity = 5) and low spin states (multiplicity = 1), respectively.

Table 6

Parameters in the general equations for temperature-dependent equilibrium nitrogen isotope fractionations (1000 $\ln \alpha = A \times 10^6/T^2 + B \times 10^3/T + C$; T is in Kelvin, valid in the range of 273–1273 K) of various NH₃-related species relative to NH_{3(gaseous)}, NH_{3(aqueous)}, and NH⁺_{tequeous)}, respectively.

	- NH3(gase	ous)		- NH _{3(aque}	eous)		$- \mathrm{NH}^+_{4(\mathrm{aquo})}$	eous)	
	А	В	С	A	В	С	A	В	С
N _{2(gaseous)}	-0.054	5.906	-3.78	-0.628	5.229	-3.30	-0.941	-1.850	0.16
NH _{3(gaseous)}	0	0	0	-0.574	-0.678	0.48	-0.887	-7.756	3.94
NH _{3(aqueous)}	0.574	0.678	-0.48	0	0	0	-0.314	-7.078	3.46
NH ⁺ (gaseous)	0.409	9.361	-4.34	-0.165	8.684	-3.87	-0.478	1.605	-0.40
NH ⁺ _(aqueous)	0.887	7.756	-3.94	0.314	7.078	-3.46	0	0	0
$Zn(NH_{3})_{6}^{2+}$	0.611	0.771	-0.56	0.038	0.094	-0.08	-0.276	-6.985	3.38
$Ni(NH_3)_6^{2+}$	0.748	0.997	-0.67	0.174	0.320	-0.19	-0.140	-6.759	3.27
$Co(NH_3)_6^{2+}$	0.683	0.858	-0.60	0.109	0.180	-0.12	-0.205	-6.898	3.34
$Co(NH_3)_6^{3+}$ (H)*	0.824	0.956	-0.71	0.250	0.279	-0.24	-0.064	-6.800	3.22
$Co(NH_3)_6^{3+}(L)^*$	1.293	2.044	-1.36	0.719	1.367	-0.89	0.406	-5.711	2.57
$Cd(NH_3)_6^{2+}$	0.510	0.707	-0.52	-0.064	0.030	-0.04	-0.378	-7.049	3.42
$Cd(NH_3)_4^{2+}$	0.720	0.968	-0.68	0.146	0.291	-0.21	-0.168	-6.788	3.26
$Cu(NH_3)_4^{2+}$	0.954	1.437	-0.95	0.381	0.760	-0.47	0.067	-6.319	2.99
$Cu(NH_3)_2^+$	1.138	1.660	-1.05	0.564	0.982	-0.57	0.250	-6.096	2.89
$Ag(NH_3)_2^+$	0.856	1.406	-0.86	0.282	0.729	-0.39	-0.032	-6.350	3.08
$Au(NH_3)_2^+$	1.321	2.212	-1.35	0.747	1.534	-0.88	0.434	-5.544	2.59
$Pt(NH_3)_2^+$	1.359	2.192	-1.36	0.785	1.515	-0.88	0.471	-5.563	2.58

* H and L in $Co(NH_3)_6^{3+}$ denote high spin state (multiplicity = 5) and low spin states (multiplicity = 1), respectively.

imentally determined value of 2.00 Å (Valli et al., 1996). The calculated $^{15}\beta$ value for Cu(NH₃)₄²⁺ is 1.0845 under 25 °C. For Cu(I)-ammine complex, the setup structure was adopted from Pavelka and Burda (2005) as a 2-coordinated complex Cu(NH₃)₂⁺ (multiplicity = 1). The yielded Cu-N bond length is 1.94 Å, which is consistent with the result (1.91 Å) of Pavelka and Burda (2005). The yielded ¹⁵β value for Cu(NH₃)₂⁺ is 1.0874 at 25 °C.

For the silver-ammine complex, previous studies have suggested a linearly 2-coordinated complex $Ag(NH_3)_2^+$ (multiplicity = 1) to be the most stable form (Geddes and Bottger, 1969; Shoeib et al., 2000; Widmer-Cooper et al., 2001; Fox et al., 2002). Our calculation gave the Ag-N bond length of $Ag(NH_3)_2^+$ as 2.17 Å, which is close to previously reported data (2.18 Å; Shoeib et al., 2001). The calculated ¹⁵ β value for $Ag(NH_3)_2^+$ is 1.0833 at 25 °C.

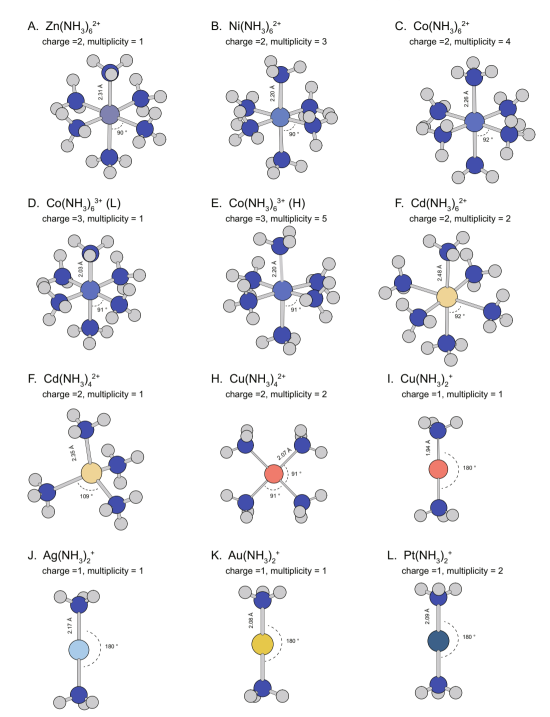


Fig. 2. Optimized geometries of $Zn(NH_{3})_{6}^{2+}(A)$, $Ni(NH_{3})_{6}^{2+}(B)$, $Co(NH_{3})_{6}^{2+}(C)$, $Co(NH_{3})_{6}^{3+}$ at low-spin state (D) or high-spin state (E), Cd $(NH_{3})_{6}^{2+}(F)$, $Cd(NH_{3})_{4}^{2+}(H)$, $Cu(NH_{3})_{2}^{+}(I)$, $Ag(NH_{3})_{2}^{+}(J)$, $Au(NH_{3})_{2}^{+}(K)$, and $Pt(NH_{3})_{2}^{+}(L)$.

For the gold-ammine complex, previous studies have also suggested the linearly 2-coordinated complex Au $(NH_3)_2^+$ (multiplicity = 1) to be most stable (Kryachko and Remacle, 2007). Our calculation gave the Au-N bond length as 2.08 Å, which is the same with the data reported by Kryachko and Remacle (2007). The calculated ¹⁵β value for Au(NH₃)₂⁺ is 1.0914 at 25 °C.

For the platinum-ammine complex, we also calculated the linearly 2-coordinated complex $Pt(NH_3)_2^+$ (multiplic-

ity = 2), which is considered to be the most stable form (Juhász et al., 2012). The results gave the Pt-N bond length as 2.09 Å, which is close to the value of 2.07 Å reported by Juhász et al. (2012). The calculated ¹⁵ β value for Pt(NH₃)⁺₂ is 1.0917 at 25 °C.

The equilibrium nitrogen isotope fractionations of these metal-ammine complexes relative to $NH_{3(gaseous)}$, $NH_{3(aqueous)}$, $NH_{4(aqueous)}^+$ are given in Table 6 and plotted in Fig. 3.

4. DISCUSSION

4.1. Factors controlling the N isotope fractionations in the $NH_4^+ - NH_3$ – metal-ammine complex system

Fig. 3 shows that ¹⁵N is most enriched in NH₄⁺ and most depleted in NH_{3(gaseous)}. This is consistent with a more stable tetrahedral structure of NH₄⁺ relative to the pyramidal structure of NH₃. The hydration of NH₃ induces significant ¹⁵N enrichment in NH_{3(aqueous)} relative to NH_{3(gaseous)}, because of the additional N-H bond formed in NH_{3(aqueous)} (Fig. 1). However, the hydration of NH₄⁺(aqueous) does not cause much more ¹⁵N enrichment in NH_{4(aqueous)} (+0.4%; Table 3), because the bonding environment does not change significantly for the N in NH₄⁺ (Fig. 1). Similarly, hydration of ammine complexes of Zn²⁺ (from Zn(NH₃)₆²⁺ to Zn(NH₃)₆²⁺ to 21(NH₃)₆²⁺ to 21(NH₃)₆²⁺ to 21(NH₃)₆²⁺ to 25 °C, suggesting that the solvent effect is insignificant on metal-ammine complexes either.

Relative to $NH_{3(aqueous)}$, most metal-ammine complexes are more enriched in ^{15}N (Fig. 3B), which is in general consistent with that the metal-N bond in the complexes is stronger than the N-H bond in NH3(aqueous). However, the varying magnitudes of ¹⁵N enrichment in the metalammine complexes relative to NH3(aqueous) suggest that more factors can affect the nitrogen isotope fractionations in the metal-ammine complexes. For example, compared with NH_{3(aqueous)}, Zn(NH₃)²⁺₆ has slightly higher ¹⁵ β values and Cd(NH₃)₆²⁺ has even slightly lower ¹⁵ β values (Table 5; Fig. 3B). Even taking into account a small increase from the solvent effect (<+1‰), nitrogen isotope fractionations between $Zn(NH_3)_6^{2+}$ or $Cd(NH_3)_6^{2+}$ and NH_{3(aqueous)} are very small (<+2% at 0 °C). In contrast, the ammine complexes of Cu⁺, Au⁺, and Pt⁺ have much higher ${}^{15}\beta$ values than NH_{3(aqueous)}. One speculation is that coordination of NH₃ in the metal-ammine complex may play an important role in determining the nitrogen isotope fractionations. As a fact, the ¹⁵ β values of metalammine complexes show an increasing trend following the decrease of coordination number from 6 (for Cd^{2+} , Zn^{2+} , Co^{2+} , Ni^{2+} , Co^{3+}), 4 (for Cd^{2+} and Cu^{2+}), to 2 (for Ag^+ , Cu^+ , Au^+ , Pt^+) (see Fig. 3). The effect of NH₃ coordination can even overrule the valence effect, which is indicated by the inversed order in isotopic enrichment between $Cu(NH_3)_2^+$ and $Cu(NH_3)_4^{2+}$.

4.2. Comparison with literature data

Our calculated ¹⁵ β values for NH_{3(gaseous)} are very close to those previous calculation results. For example, the calculations by Liu et al. (2010) using a similar method but a larger basis set gave a ¹⁵ $\beta_{NH3(gaseous)}$ value of 1.0685 (vs. 1.0687 in this study) at 25 °C. In a recent study, Walters et al. (2019) also calculated the ¹⁵ β values for both gaseous and aqueous NH₃ and NH₄⁺ in a temperature range of 250– 350 K by the HF/6-31G(d) and B3LYP/6-31G(d) levels of theory and recommended to use the HF/6-31G(d) results. Therefore, in the discussions below, all data of Walters et al. (2019) refer to the HF/6-31G(d) results. Our calculation results agree relatively well with their results for $NH_{3(gaseous)}$ (¹⁵ β value at 25 °C: 1.0687 in this study vs 1.0700 in Walters et al., 2019) and $NH_{4(gaseous)}^+$ (¹⁵ β value at 25 °C: 1.1031 in this study vs 1.1049 in Walters et al., 2019) (Fig. 4B). However, our calculation results show a larger discrepancy to their results for $NH_{3(aqueous)}$ (¹⁵ β value at 25 °C: 1.0776 in this study vs 1.0743 in Walters et al., 2019) and NH_{4 (aqueous)} ($^{15}\beta$ value at 25 °C: 1.1035 in this study vs 1.1073 in Walters et al., 2019). This discrepancy could be caused by two factors. First, it is noticed that, in the optimized geometry of Walters et al. (2019) for NH_{3(aqueous)}, the NH₃ molecule located at the edge of a water cluster, implying that this aqueous NH₃ model does not simulate a fully hydrated environment. Second, the calculations in Walters et al. (2019) only sampled one configuration for each explicit solvent model. However, previous studies have observed local configuration uncertainties caused by explicit solvent molecules and urged to sample multiple configurations (e.g., Table 3) to produce more accurate results (e.g., Zhang and Liu, 2014; He and Liu, 2015; Gao et al., 2018).

Nitrogen isotope fractionations between N_2 and NH_4^+ or NH_3 have been mostly investigated by theoretical calculations because it is difficult to reach isotope equilibrium between N_2 and NH_4^+ or NH_3 at experimental conditions (Li et al., 2009). Compared with previous theoretical calculations based on measured vibration frequencies (Urey, 1947; Scalan, 1958; Hanschmann, 1981; Petts et al., 2015), our results are closer to those of Scalan (1958) (see Fig. 4A).

Nitrogen isotope fractionations between NH_4^+ and NH3(gaseous) have been intensively studied by both theoretical calculations and experimental studies. The data are compiled in Fig. 4B. Our new results are broadly consistent with the results for a temperature range of 250–350 K by Walters et al. (2019). Compared with the diverse results in previous calculations based on measured vibration frequencies (Urey, 1947; Scalan, 1958; Hanschmann, 1981; Petts et al., 2015), our results fall between the results of Urey (1947) and Scalan (1958) at low temperature range (0-200 °C) but converge to those of Scalan (1958) and Hanschmann (1981) at temperatures higher than 200 °C (Fig. 4B). Among all these theoretical calculations, the results from Petts et al. (2015) are significantly lower than the others. This difference is mainly attributed to the large anharmonic effect from the calculations by Petts et al. (2015), which is however not observed in our calculations. Experimental determinations of nitrogen isotope fractionation between NH₄⁺ and NH_{3(gaseous)} were mostly carried out at low temperatures. The data of Thode et al. (1945), Kirshenbaum et al. (1947) and Heaton et al. (1997) are close to our calculation results (Fig. 4B), whereas the data of Urey and Aten (1936), Savard et al. (2017), Kawashima and Ono (2019) diverge from the theoretical predictions. This deviation may be attributed to two factors: (1) the equilibrium isotope fractionation could have not been achieved in those experiments; (2) some of the experiments were measured on solid ammonium salt, in which an additional isotope fractionation between solid NH_4^+ and gaseous NH_4^+ may exist (Ishimori, 1960b).

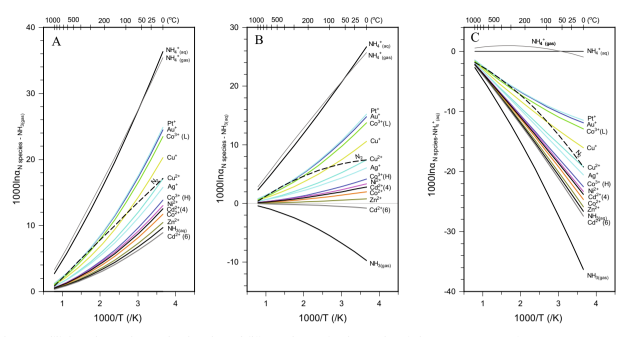


Fig. 3. Equilibrium nitrogen isotope fractionations of different nitrogen-bearing species relative to gaseous NH_3 (A), aqueous NH_3 (B), and aqueous ammonium (C). The letter "L" and "H" after Co^{3+} denote the low-spin state and the high-spin state, respectively. The number "4" and "6" after Cd^{2+} denote the coordination number of NH_3 .

Nitrogen isotope fractionations between NH₄⁺ and NH_{3(aqueous)} are relatively less studied. The solvent effect, which is strong for NH₃ hydration but negligible for NH_4^+ hydration, results in diminished isotope fractionation between the NH_4^+ – $NH_{3(aqueous)}$ pair relative to the NH_4^+ – $NH_{3(gaseous)}$ pair. This has been demonstrated by our calculation as well as those by Walters et al. (2019). However, our calculations yielded much smaller magnitudes of nitrogen isotope fractionations between NH_4^+ and $NH_{3(aqueous)}$ (Fig. 4C) than those of Walters et al. (2019), likely because the calculations in Walters et al. (2019) only incorporated a partial solvent effect, which can be inferred by the configuration of their hydrated NH₃. Our results fit well with the experimental results of Urey and Aten (1936), Thode et al. (1945), and Ishimori (1960a), but are slightly lower than the result of Kirshenbaum et al. (1947) and slightly higher than the result of Hermes et al. (1985). One striking phenomenon on Fig. 4C is that the experimental results by Li et al. (2012) are much higher than the theoretical predictions, which will be explored in detail in Section 5.1.

For the $NH_{3(aqueous)} - NH_{3(gaseous)}$ pair (Fig. 4D), our calculations considering a full hydration effect yielded larger nitrogen isotope fractionations than those of Walters et al. (2019). The relatively few experimental estimates (e.g., Wahl et al., 1935; Urey and Aten, 1936; Thode et al., 1945; Kirshenbaum et al., 1947; Deng et al., 2018) mostly fall between the line from this study and the one from Walters et al. (2019). The difference between our results and the experimental results may be attributed to the kinetic isotope effect associated with the movement of

 NH_3 gas (e.g., Deng et al., 2018), which could have interfered the quantification of the equilibrium isotope fractionations in laboratory experiments.

The study of nitrogen isotope fractionation involving metal-ammine complexes is very few in our knowledge. By ion-exchange experiments using cation exchange resins, Ishimori (1960a) investigated the equilibrium isotope fractionation factors during the single-stage separation of NH₃ from ammine complexes of Ni²⁺, Cu²⁺, Zn²⁺, Cd²⁺, and Ag²⁺ in 30 °C aqueous solutions, which yielded +7.6% between Ni(NH₃)_n²⁺ and NH₃, +11.6% between $Cu(NH_3)_4^{2+}$ and NH_3 , +11.5‰ between $Zn(NH_3)_4^{2+}$ and NH_{3} , +10.2‰ between Cd(NH_{3})²⁺ and NH_{3} , and +9.2‰ between $Ag(NH_3)_2^{2+}$ and NH_3 . Using similar experiments at 30 °C, Gupta and Sarpal (1967) obtained the equilibrium isotope fractionation factors of +6.2%, +7.9%, and +10.0% between Ni²⁺-ammine complexes and NH₃ in aqueous solutions during the single-stage separation of NH₃ from Ni(NH₃)²⁺₄, Ni(NH₃)²⁺₅, and Ni(NH₃)²⁺₆, respectively. These experimental and theoretical data are significantly higher than our calculated data between these metal-ammine complexes and NH3(aqueous) (e.g., +2.8% for $Ni(NH_3)_6^{2+}$, +2.3% for $Cd(NH_3)_4^{2+}$, +6.2% for Cu $(NH_3)_4^{2+}$, +0.6% for $Zn(NH_3)_6^{2+}$, +5.1% for $Ag(NH_3)_2^{+}$, at 30 °C), but mostly lower than those between metalammine complexes and NH3(gaseous) (e.g., +10.8% for Ni $(NH_3)_6^{2+}$, +10.3% for Cd(NH_3)_4^{2+} ,+14.2% for Cu $(NH_3)_4^{2+}$,+8.6% for Zn(NH_3)_6^{2+} ,+13.1% for Ag(NH_3)_2^{+}, at 30 °C). Because these experimental studies (Ishimori, 1960a; Gupta and Sarpal, 1967) did not describe the detailed controlling conditions in their experiments, in particular, whether other nitrogen species (e.g., NH_4^+)

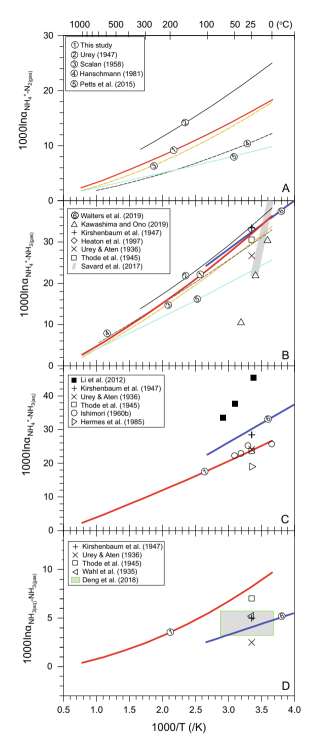


Fig. 4. Comparison of nitrogen isotope fractionation factors between our data and previous theoretical calculations and experimental results. The line numers in B, C and D are the same to those in A. See text for discussion.

coexisted in the solution or whether any NH_3 degassing was involved, it is difficult to assess the exact cause of the discrepancy between our calculation results and these experimental results. In a theoretical calculation based on vibrational frequency data, Jeevanandam and Gupta (1968) obtained the isotope fractionation factors (relative to NH₃) at 25 °C to be ~ +27‰ for Co(NH₃)₆³⁺, ~ +15‰ for Ni(NH₃)₆²⁺, and ~ +11‰ for Co(NH₃)₆²⁺, which are slightly higher than our calculated results between these complexes and gaseous NH₃, i.e., +11.8‰ and +20.0‰ for Co(NH₃)₆³⁺ at high-spin state and low-spin state, repsectively, +11.1‰ for Ni(NH₃)₆²⁺, and ~ +10.0‰ for Co (NH₃)₆²⁺.

5. APPLICATIONS

5.1. Complicated isotopic effect during NH3 degassing

NH₃ degassing may occur in alkaline fluids in a variety of geological settings (see discussion in Li et al., 2012; Deng et al., 2018). Li et al. (2012) carried out laboratory experiments to simulate NH₃ degassing in the field at a temperature range from 2 °C to 70 °C. The experiments were started by adding inadequate hydroxyl to partially dissociate NH₄⁺ in a solution to drive NH₃ degassing, i.e., the processes described by Eq. (1). The nitrogen isotopic compositions of the remaining NH⁺₄ after complete degassing of NH_3 produced by partial dissociation of NH_4^+ at various extents displayed a pattern close to batch equilibrium model rather than a Rayleigh model, which made the observed large isotopic effect (from +33.5% at 70 °C to +45.5% at 23 °C) be interpreted as a result mainly from equilibrium isotope fractionations between NH_4^+ and $NH_{3(-1)}^+$ aqueous), whereas the possible effect from NH₃ degassing was not fully assessed due to the lack of data (Li et al., 2012). Our calculations yielded significantly smaller nitrogen isotope fractionations between NH_4^+ and $NH_{3(aqueous)}$ (Fig. 4C) than the experimental results by Li et al. (2012). Recently, experimental investigations on the NH_{3(aqueous)} - NH₃ system by Deng et al. (2018) revealed that NH₃ degassing could have a significant kinetic nitrogen isotope effect. To assess the possible influence of this kinetic isotope effect on the experimental results in Li et al. (2012), we carried out numeric modeling of those isotopic results using the equilibrium isotope fractionation factors between NH_4^+ and $NH_{3(aqueous)}$ in this study and the kinetic isotope effect between NH_{3(aqueous)} and NH_{3(gaseous)} determined by Deng et al. (2018).

Following the methods from previous studies (e.g., Rees, 1973; Dauphas and Rouxel, 2006), a two-step reaction model was applied to characterize the nitrogen isotope fractionation involving an intermediate species. The nitrogen isotope transfer in the reaction network can be described by the equations below.

$${}^{14}NH_{4}^{+} + OH^{-} + (n-1)H_{2}O \stackrel{{}^{14}k_{1}}{\rightleftharpoons} {}^{14}NH_{3} \cdot nH_{2}O$$

$$\times \xrightarrow{{}^{14}k_{3}} {}^{14}NH_{3(gas)} + nH_{2}O$$

$${}^{15}NH_{4}^{+} + OH^{-} + (n-1)H_{2}O \stackrel{{}^{15}k_{1}}{\rightleftharpoons} {}^{15}NH_{3} \cdot nH_{2}O$$

$$\times \xrightarrow{{}^{15}k_{3}} {}^{15}NH_{3(gas)} + nH_{2}O \qquad (6)$$

in which ${}^{14}k_1$, ${}^{15}k_1$, ${}^{14}k_2$, ${}^{15}k_2$, ${}^{14}k_3$, ${}^{15}k_3$ represent reaction rates of ${}^{14}N$ and ${}^{15}N$ in the reversible reaction between NH_4^+ and $NH_{3(aqueous)}$ (denoted by subscripts 1 and 2)

1.

and the unidirectional reaction from $NH_{3(aqueous)}$ to $NH_{3(-gaseous)}$ (denoted by subscript 3), respectively. Accordingly, the concentration of reactants and products at time *t* can be obtained by:

$$\frac{dn_{[}^{14}NH_{4}^{+}]}{dt} = - {}^{14}k_{1} [{}^{14}NH_{4}^{+}] \cdot [HO^{-}] + {}^{14}k_{2} [{}^{14}NH_{3} \cdot nH_{2}O]$$
(7)

$$\frac{dn_{[1^{5}NH_{4}^{+}]}}{dt} = - {}^{15}k_{1} [{}^{15}NH_{4}^{+}] \cdot [HO^{-}] + {}^{15}k_{2} [{}^{15}NH_{3} \cdot nH_{2}O]$$
(8)

$$\frac{dn[14NH_3 \cdot nH_2O]}{dt} = {}^{14}k_1[{}^{14}NH_4^+] \cdot [HO^-] - {}^{14}k_2[{}^{14}NH_3 \cdot nH_2O] - {}^{14}k_3[{}^{14}NH_3 \cdot nH_2O]$$
(9)

$$\frac{dn_{[15NH_3\cdot nH_2O]}}{dt} = {}^{15}k_1 [{}^{15}NH_4^+] \cdot [HO^-] - {}^{15}k_2 [{}^{15}NH_3 \cdot nH_2O] - {}^{15}k_3 [{}^{15}NH_3 \cdot nH_2O]$$
(10)

$$\frac{dn_{[\ ^{14}NH_3 \cdot nH_2O]}}{dt} = \ ^{14}k_3 \left[\ ^{14}NH_3 \cdot nH_2O \right] \tag{11}$$

$$\frac{dn_{[1^{5}NH_{3}\cdot nH_{2}O]}}{dt} = {}^{15}k_{3} [{}^{15}NH_{3} \cdot nH_{2}O]$$
(12)

Finally, the ratio of the remaining NH_4^+ and $NH_{3(aqueous)}(R_{remaining})$ at time *t* can be derived from:

$$R_{remaining} = \frac{\begin{bmatrix} 1^5 NH_4^+ \end{bmatrix} + \begin{bmatrix} 1^5 NH_3 \cdot nH_2 O \end{bmatrix}}{\begin{bmatrix} 1^4 NH_4^+ \end{bmatrix} + \begin{bmatrix} 1^4 NH_3 \cdot nH_2 O \end{bmatrix}}$$
(13)

The nitrogen isotope fractionations in these reactions can be described as:

$$\alpha_1 = \frac{{}^{15}k_1 \cdot {}^{14}k_2}{{}^{15}k_2 \cdot {}^{14}k_1} \tag{14}$$

$$\alpha_2 = \frac{^{15}k_3}{^{14}k_3} \tag{15}$$

where α_1 is the equilibrium nitrogen isotope fractionation factor, which has been calculated above; α_2 is the kinetic nitrogen isotope fractionation factor, which has been determined by Deng et al. (2018). In our numeric modeling, values were arbitrarily assigned to the unknowns ${}^{14}K_1$, ${}^{15}K_1$, and ${}^{14}K_2$ to best fit the experimental data.

When hydroxyl is added into an ammonium solution, the extent of the overall reaction is dependent on the initial $[OH^-]/[NH_4^+]$ ratio. As a combined effect of equilibrium isotope fractionation between the remaining NH_4^+ and $NH_{3(aqueous)}$ and kinetic isotope fractionation of NH_3 degassing from $NH_{3(aqueous)}$, the remaining NH_4^+ in the solution is progressively enriched in ¹⁵N with the proceeding of the reaction until the produced $NH_{3(aqueous)}$ is consumed by NH_3 degassing. The magnitude of ¹⁵N enrichment in the remaining NH_4^+ is a function of (1) temperature, which determines the magnitudes of the two involved isotope fractionations, and (2) the initial $[OH^-]/[NH_4^+]$ ratio, which determines the extents of the total reaction. The progressive ¹⁵N enrichment patterns for reactions

at room temperature, 50 °C and 70 °C, are illustrated in Fig. 5 for a variety of initial $[OH^{-}]/[NH_{4}^{+}]$ ratios. The results show that, when the initial $[OH^{-}]/[NH_{4}^{+}]$ ratio is large enough (e.g., \geq 2) to drive complete conversion of NH₄⁺ to NH_{3(aqueous)}, the ¹⁵N enrichment in the remaining NH_{3(aqueous)} along progressive NH₃ degassing is only controlled by the kinetic isotope fractionation from NH₃ degassing and follows the red curves in Fig. 5A-C. These scenarios resemble the laboratory experiments by Deng et al. (2018), and consistently, the data from Deng et al. (2018) fall closely to these curves. In contrast, when the initial $[OH^{-}]/[NH_{4}^{+}]$ ratio is small enough (e.g., < 1) to only induce partial conversion of NH_4^+ to $NH_{3(aqueous)}$, the isotope evolution pattern of the remaining nitrogen in the solution (a mixture of NH_4^+ and $NH_{3(aqueous)}$ in this case) along progressive NH₃ degassing is controlled by both equilibrium and kinetic isotopic fractionations as well as the initial $[OH^{-}]/[NH_{4}^{+}]$ ratio. The progressive ¹⁵N enrichments for a number of initial [OH⁻]/[NH⁺₄] ratios are illustrated by the grey curves in Fig. 5. Interestingly, after complete degassing of NH₃ from the solution, the $\delta^{15}N$ of the remaining NH₄⁺ from varying initial [OH⁻]/[NH₄⁺] ratios, i.e., the data points at the ends of the grey curves on Fig. 5, form the blue curves which show very small curvatures that mimic batch equilibration lines as initially thought in Li et al. (2012). When plotted on Fig. 5, the experimental data of Li et al. (2012) align closely along the blue lines. Therefore, the large isotope fractionations observed in the experiments by Li et al. (2012) more likely reflect the overall isotope effect combining the equilibrium isotope fractionation during the conversion of NH_4^+ to $NH_{3(aqueous)}$ (this study) and the kinetic isotope effect of NH₃ degassing (Deng et al., 2018). Nevertheless, the experiments in Li et al. (2012) represent an open-system scenario that is more likely to occur in the field, and thus can contribute to interpret the field data in a first order. More accurate data modeling and interpretation should follow the quantitative modeling described here (Eqs. (7)–(15); Fig. 5).

5.2. Metal-ammine remobilization in hydrothermal system?

Busigny et al. (2011) observed a linear relationship between Cu concentration and $\delta^{15}N$ values in metagabbros from the western Alps. To explain this correlation, the authors proposed that Cu in the protoliths of the metagabbros was hydrothermal leached and remobilized by fluid in which ammonia is complexed with Cu. However, because the nitrogen isotope fractionation factors between Cuammine complex (the species in the fluid) and ammonium (the species in the (meta-)mafic rocks; Busigny et al., 2005, 2011; Li et al., 2007, 2014) were not available at that time, Busigny et al. (2011) modeled their data using the nitrogen isotope fractionation factors between NH₄⁺ and NH₃ from Scalan (1958), which are now demonstrated to be very different from those between NH₄⁺ and copperammine complex (see Fig. 3). Our new data provide an unprecedently available opportunity to revisit this hypothesis.

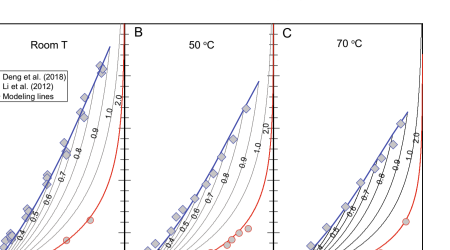


Fig. 5. Diagram showing the progressive ¹⁵N enrichment in the remaining nitrogen in a solution along progressive degassing of NH_3 as a result of partial dissociation of NH_4^+ . Number on individual line refers to the initial ratio of $[OH^-]/[NH_4^+]$ in the solution, which determines the fraction of NH_4^+ that can be dissociated and eventually degassed. The end point on each curve represents the isotopic composition of the final remaining NH_4^+ in the solution (if there is any) after complete NH_3 degassing. The blue lines linking these end points fit well with the experimental data of Li et al. (2012). The red lines represent the case of complete dissociation of NH_4^+ , which fit well with the experimental results of Deng et al. (2018). See text for discussion.

04

1-f

0.6

0.8

0

0.2

0.4

0.6

0.8

10

In our modeling (see Fig. 6), the valence of copper cation is considered to be either I or II. The coordination number for copper cation is strongly dependent on solution environments. Cu(II) can have coordination numbers of 4, 5 and 6, among which $Cu(NH_3)_4^{2+}$ is the most thermodynamically stable species; whereas Cu (I) can have coordination numbers of 2, 3 and 4, among which the most thermodynamically stable species has been suggested to be $Cu(NH_3)_2^+$ (Pavelka and Burda, 2005). Using the calculated nitrogen isotope fractionations of these two species relative to ammonium (Table 6; Fig. 3C), we modeled the meta-gabbro data from Busigny et al. (2011) by a batch model. Rayleigh distillation model is not employed here because it is apparently inconsistent with the observed linear relationship between Cu concentration and $\delta^{15}N$ of the low-strain meta-gabbros samples.

50

40

30

20

10

0 -

0.2

0.4

0.6

0.8

0

0.2

 $\delta^{15}N_{final}\delta^{15}N_{initial}$ (%)

A

Fig. 6 illustrates our modeling results. It shows that the relationship between nitrogen concentrations and δ^{15} N values can be easily explained by a leaching model (as well as a mixing model or a batch devolatilization model), but requires a large temperature range, e.g., 250–650 °C if in form of Cu(NH₃)₂⁺ or 300–700 °C if in form of Cu (NH₃)₄²⁺ (Fig. 6A). Applying the same temperature ranges, the Cu concentration and δ^{15} N data should fall in the triangular area labeled by Cu(NH₃)₂⁺ or Cu(NH₃)₄²⁺ in Fig. 6B, which however cannot explain the observed Cu data because of the low efficiency of NH₃ in mobilizing Cu in these two forms, i.e., 2:1 and 4:1, respectively (Fig. 6B). If

copper mobilization was indeed coupled with NH₃ in those samples, it had to be in a copper complex species containing only one NH₃ (Fig. 6B) in order to efficiently leach out > 90% of the Cu as observed in some samples (Fig. 6B). In this case, the Cu concentration – δ^{15} N relationship can be explained by a batch model (the triangular area labeled by $Cu(NH_3)^{+/2+}$ in Fig. 6B) with $\ln \alpha_{NH_4^+ - Cu(NH_3)^{+/2+}}$ values of +7.5% to + 10.1%. Even applying the isotope fractionations of $Cu(NH_3)_4^{2+}$ or $Cu(NH_3)_2^{+}$, which have larger coordination numbers than the expected species of $Cu(NH_3)^{2+}$ or $Cu(NH_3)^{+}$, respectively, these large isotope fractionations correspond to a small temperature range of 200–320 °C for Cu^{2+} or 150–270 °C for Cu^{+} , both are significantly lower than the large and high temperature ranges to explain the N concentration – δ^{15} N relationship (Fig. 6A). The real isotope fractionations between NH_4^+ and $Cu(NH_3)^{+/2+}$ would be smaller than these between NH_4^+ and $Cu(NH_3) \frac{2^+}{4}$ or between NH_4^+ and Cu $(NH_3)^{+}_{2}$ given that ¹⁵N is expected to be more enriched in species with smaller coordination numbers. Consequently, it requires even lower temperature range to explain the data. This self-inconsistency between the temperature ranges yielded from the N content- δ^{15} N relationship (Fig. 6A) and from the Cu content- $\delta^{15}N$ relationship (Fig. 6B) implies that the observed nitrogen signature of the meta-gabbros (Busigny et al., 2011) might not be controlled by leaching during seafloor hydrothermal alteration, or at least have been overprinted by other geochemical processes.

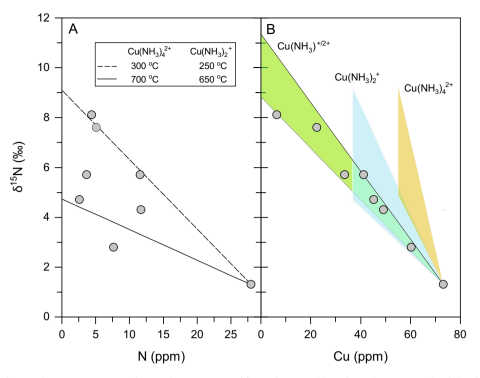


Fig. 6. Modeling of Cu and N contents versus nitrogen isotope compositions of meta-gabbros from the western Alps following the model of a coupled leaching of Cu and NH₃ by Busigny et al. (2011). See text for discussion.

6. CONCLUSION

Theoretical calculations of equilibrium nitrogen isotope fractionation factors between gaseous and aqueous ammonium, ammonia, N₂ and metal-ammine complexes indicate that ¹⁵N is enriched following the order of $NH_4^+ > N_2 - NH_{3(aqueous)} > NH_{3(gaseous)}$, with all but one metal-ammine complexes lying between NH_4^+ and $NH_{3(aqueous)}$. Our calculation suggests anharmonic effect is not significant on the isotope fractionation between NH_4^+ and NH_3 . In the metal-ammine complexes, coordination number may play an important role in controlling the isotope fractionations. Our new calculation results verify that nitrogen isotope behavior in a natural system involving multiple nitrogen species may be very complicated. The propagation of isotope fractionations along a reaction network may result in a much larger isotopic effect than expected.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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APPENDIX A. SUPPLEMENTARY MATERIAL

Supplementary data to this article can be found online at https://doi.org/10.1016/j.gca.2020.12.010.

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