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# Unraveling the electronic structure of LuH, LuN, and LuNH: building blocks of new materials†

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Advances in superconductor technology have been pursued for decades, moving towards room temperature models, such as a postulated nitrogen-doped lutetium hydride network. While experimental observations have been contradictory, insight into the building blocks of potential new superconductor materials can be gained theoretically, unravelling the fascinating electronic structure of these compounds at a molecular level. Here, the fundamental building blocks of lutetium materials (LuH, LuN, and LuNH) have been examined. The structures, spectroscopic constants for the ground and excited states, and the potential energy curves have been obtained for these species using complete active self-consistent field (CASSCF) and multireference configuration interaction with Davidson's correction (MRCI+Q) methods. For LuNH, the energetic properties of its isomers are determined. The bond dissociation energies of the three building blocks are calculated with the state-of-the-art *f*-block *ab initio* correlation consistent composite approach (*f*-ccCA) and the high accuracy extrapolated *ab initio* thermochemistry (HEAT) scheme. As well, an analysis of different formation pathways of LuNH has been provided.

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## 1 Introduction

From the concept of superconductivity and its first observation in mercury at 4.2 K, scientists have strived to create the first room temperature superconductor.<sup>1</sup> The first ambient temperature superconductor would not only revolutionize electronics, but also impact industries, such as quantum computing, and transform the chemical industry. Ambient superconductors will allow for the development of energy efficient computer chips and fusion reactors, and aid in the development of magnets, which can be used in applications such as maglev trains. One very important application would be for particle accelerators, such as the Large Hadron Collider (LHC), which requires 96 tons of liquid helium to cool their current system.<sup>2</sup> If this extreme level of cooling was not necessary, the engineering would be simplified.<sup>2</sup> An added layer of difficulty in building an ambient superconductor are the required pressures. In the solid state these materials form a lattice that offers no resistance to conducting current. However, large pressures are required to maintain the stability and structure of the material lattices. Under lower pressure conditions (below 200–300 kPa), the lattice can break down, due to the coupling of the electrons and phonons.

Early in 2023, a nitrogen-doped lutetium hydride lattice was proposed as an ambient temperature superconductor (294 K and 1–2 GPa).<sup>3</sup> It was reported that the lattice structure is composed of: LuH<sub>2±x</sub>N<sub>y</sub>, depicting a dark-blue color, according to the authors consistent with the novel ambient temperature superconductivity. However, this finding did not achieve consensus among the chemistry community, and the publication on the proposed superconducting properties of this structure was retracted in November 2023. Ming *et al.*, synthesized the same compound (LuH<sub>2±x</sub>N<sub>y</sub>) utilizing high pressure and high temperature conditions, which was then confirmed by X-ray diffraction.<sup>4</sup> The authors found the same color change from dark blue to violet and then pink/red, however superconductivity was not observed above 2 K. Peng *et al.*, have also performed a quantitative temperature-dependent resistance comparison of Lu–N–H and pure lutetium before reaction and concluded that the change is resistance mentioned by Dasenbrock-Gammon *et al.*, is likely caused by a metal-to-poor conductor transition.<sup>5</sup> Xingzhou *et al.* also reported that they could not find evidence of a superconducting transition in all phases from temperatures ranging from 1.8 to 300 K.<sup>6</sup> The aforementioned and several other studies have also shown, that the superconductivity properties described by Dasenbrock-Gammon *et al.*<sup>3</sup> cannot be reproduced, hence the original manuscript was retracted.<sup>4,5,7–28</sup> However, the electronic properties of the building blocks of these materials have not been explained or understood. As Xingzhou *et al.* mentioned in their work: “...we identify a notable temperature-induced resistance

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anomaly of electronic origin in  $\text{LuH}_{2\pm x}\text{N}_y$ , which is most pronounced in the pink phase and may have been erroneously interpreted as a sign of superconducting transition".<sup>6</sup> In addition, the electronic structure of the lattice building blocks has not been detailed properly.

In terms of the building blocks of the nitrogen-doped lutetium hydride lattice, there is little known about it from an electronic structure standpoint. In 1974, Efantin *et al.* calculated the electronic spectra of LuH.<sup>29</sup> The ground state was established as  $1^1\Sigma^+$ , the bond length,  $R_e = 1.9111 \pm 0.0001 \text{ \AA}$ , and the  $\omega_e$  of the ground state is  $1520 \pm 20 \text{ cm}^{-1}$ . More recently, gas phase spectroscopic constants have been calculated by Wang *et al.* by laser ablation with the  $\omega_e$  of the ground state calculated to be  $1476 \pm 20 \text{ cm}^{-1}$ .<sup>30</sup>

Theoretically, LuH has been studied (mainly for ground state properties), while LuN and LuNH have received less attention. Pantazis *et al.* calculated the  $R_e$  of the LuH ground state ( $1^1\Sigma^+$ ) using density functional theory (DFT), specifically PBE0 with segmented all-electron relativistically contracted (SARC) basis sets, paired with a Douglas-Kroll-Hess second order Hamiltonian (DKH2) with counterpoise corrections, and rendered the ground state bond length  $1.904 \text{ \AA}$ .<sup>31</sup> In addition, the authors estimated the dissociation energy ( $D_e$ ) to be  $3.12 \text{ eV}$ . Later, Jalbout *et al.* utilized a panoply of DFT functionals, which were then fitted, and predicted the ground state bond length to range between  $1.911$  to  $1.940 \text{ \AA}$ .<sup>32</sup> In this investigation, B3LYP obtained the closest value to experiment, which led to the following spectroscopic constants:  $1.911 \text{ \AA}$  ( $R_e$ ),  $1517 \text{ cm}^{-1}$  ( $\omega_e$ ), and  $20.48 \text{ cm}^{-1}$  ( $\omega_e \chi_e$ ). In terms of wavefunction methods, Møller-Plesset second order (MP2) perturbation theory with an all-electron basis set was utilized to investigate ground state properties of LuH resulting in  $1.883 \text{ \AA}$  and  $1540 \text{ cm}^{-1}$  for  $R_e$  and  $\omega_e$ , respectively.<sup>33</sup> Dolg and Stoll also calculated spectroscopic constants for the  $1^1\Sigma^+$  ground state,<sup>34</sup> obtaining  $1.95 \text{ \AA}$ ,  $1445 \text{ cm}^{-1}$ , with CI(SD + Q) and the Wood-Boring pseudopotential for Lu.<sup>34</sup> In addition, Cao and Dolg utilized SCF, CCSD, and CCSD(T) paired with natural orbital valence basis sets (small-core pseudopotentials for the lanthanide) to calculate the  $R_e$ ,  $D_e$  and  $\omega_e$  for the  $1^1\Sigma^+$  ground state of LuH.<sup>35</sup> At the CCSD(T) level, the values were estimated from  $1.882$ – $1.914 \text{ \AA}$ , and the  $\omega_e$  from  $1507$  and  $1577 \text{ cm}^{-1}$  (depending upon the level of core-correlation considered). For  $D_e$ , the values ranged from  $3.35$  to  $3.64 \text{ eV}$ .<sup>35</sup> Assaf *et al.* calculated the ground and first excited states of LuH and provided spectroscopic constants for this molecule.<sup>36</sup> The authors show the importance of accurately detailing the spin-orbit properties of this system to understand the energetics of the complex.

For LuN and LuNH, computational studies have been done, through mainly to characterize their behavior within a lattice, utilizing DFT, and considering potential solid-state applications. Oualdine *et al.* investigated the structural, elastic, and electronic properties of LuN using *ab initio* (solid state) techniques. The authors found that LuN has a semiconductor nature and calculated its band structure and gap.<sup>37</sup> Yurdaşan *et al.* also found LuN to be a promising material for optoelectronic applications.<sup>38</sup> More recently, in 2022, Devese *et al.* probed the

defects of  $\text{LuN}_{1-\delta}$  experimentally and computationally.<sup>39</sup> The authors calculated the optical bandgap of this material to be  $1.7 \text{ eV}$ . In addition, that its conductivity can be controlled by nitrogen vacancy doping.

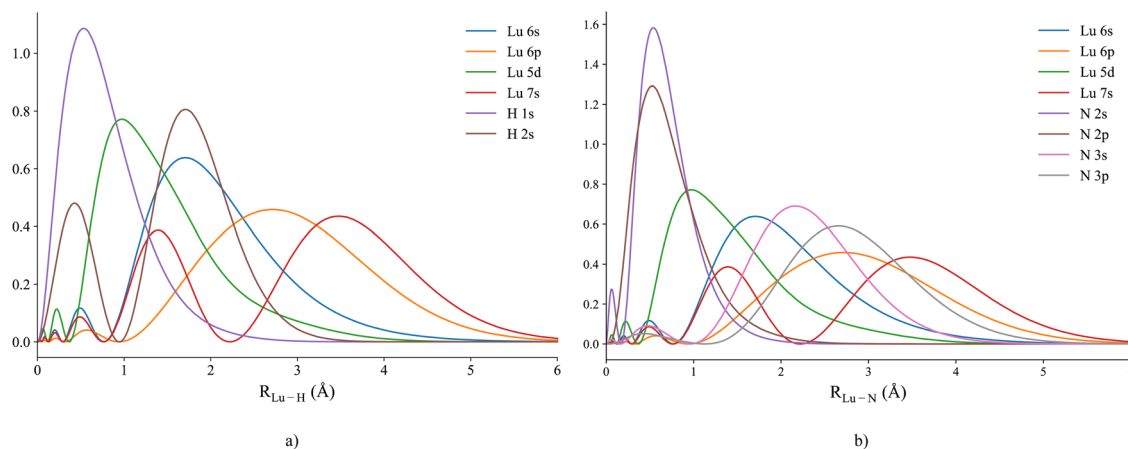
Herein, the first detailed electronic structure calculations for LuH, LuN, and LuNH (and its isomers) are reported, detailing the ground and excited states of the molecules, along with the different routes of formation of the molecules. Bond dissociation values are also provided with state-of-the-art methodologies: *f*-block *ab initio* correlation consistent composite approach (*f*-ccCA)<sup>40</sup> and a type of high accuracy extrapolated *ab initio* thermochemistry (HEAT).<sup>41–44</sup>

## 2 Computational details

Due to the complexity of the electronic structure of lanthanides, multireference wavefunction approaches are typically necessary to accurately describe the fundamental properties of LuH, LuN and LuNH, and also to construct potential energy curves (PECs). MRCI+Q was utilized to describe the ground and several excited states of these molecules. For LuH, the active space considered was: four electrons in ten orbitals. The orbitals included at the atomic level are: five of  $a_1$  symmetry ( $1s$  (H);  $6s$ ,  $5d_{z^2}$ ,  $5d_{x^2-y^2}$ ,  $6p_z$  (Lu)), two of  $b_1$  symmetry ( $5d_{xz}$  and  $6p_x$  (Lu)), two of  $b_2$  symmetry ( $5d_{yz}$  and  $6p_x$  (Lu)) and one of  $a_2$  symmetry ( $5d_{xy}$  (Lu)). Due to the smaller active space utilized for this molecule a DKH3 Hamiltonian was employed in MOLPRO 2020.<sup>45</sup> Paired with DKH3, a cc-pVQZ-DK<sup>46</sup> basis set was used for lutetium, along with aug-cc-pVQZ-DK<sup>47</sup> for hydrogen. For LuN, an expansion of the active space to six electrons in fifteen orbitals was necessary: six of  $a_1$  symmetry ( $2p_z$ ,  $3p_z$  (N);  $6s$ ,  $5d_{z^2}$ ,  $5d_{x^2-y^2}$ ,  $6p_z$  (Lu)), four of  $b_1$  symmetry ( $2p_x$ ,  $3p_x$  (N);  $5d_{xz}$  and  $6p_x$  (Lu)), four of  $b_2$  symmetry ( $2p_y$ ,  $3p_y$  (N);  $5d_{yz}$  and  $6p_x$  (Lu)) and one of  $a_2$  symmetry ( $5d_{xy}$  (Lu)). Since the active space was expanded to fifteen orbitals, it was not possible to employ a fully relativistic Hamiltonian (DKH3), instead, a non-relativistic Hamiltonian was utilized. For Lu, the ECP28MWB segmented pseudopotential was employed at a triple- $\zeta$  level.<sup>35,48</sup> For nitrogen, aug-cc-pVTZ was utilized.<sup>49</sup> Radial distribution functions (RDFs) were calculated for LuH and LuN, showing the importance of the inclusion of  $3p_{x,y,z}$  orbitals for the LuN PECs (see Fig. 1(a) and (b)). The  $3p_{x,y,z}$  orbitals overlap significantly at the equilibrium region of LuN (and probably LuNH). For both molecules, core correlation was also employed, by including the  $4f$  orbitals (seven extra orbitals). By including these orbitals at the “core” level in the MRCI expansion, single and double excitations are allowed to the active space. Moreover, in previous studies, it was shown that these core orbitals are necessary to reproduce experimental quantities accurately.<sup>40,50</sup>

For LuNH PECs, the same orbitals employed for LuN were considered at the CASSCF level, with one more electron originating from the  $1s$  of hydrogen. In addition, the  $2p_z$  of nitrogen hybridizes with the  $1s$  of hydrogen, so no additional orbitals were necessary in the active space. At the MRCI level, the three





**Fig. 1** (a) Radial distribution functions for LuH, with the bond distance of the molecule plotted along the z-axis, obtained using CCSD(T), the cc-pVTZ-DK3 basis set for Lu, and the cc-pVTZ-DK basis set for H (and using a DKH3 Hamiltonian). (b) Radial distribution functions for LuN, with the bond distance of the molecule plotted along the z-axis, obtained using CCSD(T), the cc-pVTZ-DK3 basis set for Lu, and the aug-cc-pVTZ-DK basis set for N (DKH3 Hamiltonian).

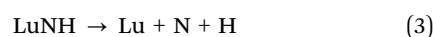
higher orbitals from  $a_1$ ,  $b_1$  and  $b_2$  symmetries were removed. Having fifteen orbitals and seven electrons in the active space, with additional core orbitals was not feasible at the MRCI level. In addition, at 7.0 Å, (the initial distance the potential energy curves were computed), the three aforementioned orbitals are a hybrid of the  $3p_{x,y,z}$  and the  $3d_{z^2}$ ,  $3d_{xz}$  and  $3d_{yz}$  of nitrogen. However, at equilibrium bond length, the  $b_1$  and  $b_2$  orbitals lose their  $3p_{x,y}$  character and become pure  $3d_{xz}$  and  $3d_{yz}$  respectively. As per LuH and LuN, the 4f orbitals were also included in the core. For the potential energy curves (PECs), the following basis sets were used: ECP28MWB segmented (Lu)<sup>35,48</sup> and aug-cc-pVTZ (N,H).<sup>49</sup> To optimize, the ground state of LuNH and respective isomers, CCSD(T) with a DKH3 Hamiltonian was utilized, paired with a triple- $\zeta$  level basis sets (cc-pVTZ for H and aug-cc-pVTZ for N). Analytical frequencies were calculated, ensuring a minimum in the potential energy surface.<sup>49</sup>

The spin-orbit corrections for LuH, LuN, and LuNH were calculated with the Breit–Pauli Hamiltonian within the MRCI method. The active space used was the same as previously mentioned for the non-spin orbit calculations, *i.e.*, four electrons in ten orbitals for LuH, six electrons in fifteen orbitals for LuN, and seven electrons in fifteen orbitals for LuNH. Core-correlation was utilized for LuH and LuN, the f-orbitals were included in the “core card”. However, for LuNH the 4f orbitals were not included, due to the added computational cost. For LuH due to the added computational cost, a triple- $\zeta$  basis set was utilized for spin-orbit calculations (cc-pVTZ-DK3<sup>46</sup> (Lu) and aug-cc-pVTZ-DK<sup>47</sup> (H), paired with a DKH3 Hamiltonian). For the other species (LuN and LuNH), the same basis set as for the non-spin-orbit calculations was utilized.

For CASSCF and MRCI+Q, equilibrium bond lengths, harmonic, anharmonic and  $\Delta G_{1/2}$  were calculated from the PECs of LuH, LuN, and LuNH, and the Numerov/Cooley approach was utilized.<sup>51,52</sup> For spin-orbit curves and with fewer sampling energetic points, the Dunham approach was utilized.<sup>53</sup>

$f$ -ccCA and HEAT were used to obtain thermodynamic quantities for LuH, LuN, and LuNH. For LuN, even though the Hartree–Fock orbitals are not ideal (see Results and Discussion), the higher correlation contributions beyond the CCSD(T) terms for HEAT are able to provide much of the electronic correlation necessary to describe the ground state of this molecule.

HEAT is utilized rather than our own super-ccCA (s-ccCA), as s-ccCA includes a pentuples contribution, which though successful for transition metal species,<sup>54</sup> is too expensive computationally for the heavy elements, even at a double- $\zeta$  level. HEAT and s-ccCA are similar, though with some differences such as the pentuples contribution (s-ccCA), and the full complete basis set (CBS) extrapolation for the full coupled cluster triples contribution (HEAT).<sup>54</sup> Composite schemes such as  $f$ -ccCA, s-ccCA, and HEAT mimic the accuracy of computationally more expensive methods, utilizing a series of lower level *ab initio* calculations, allowing a lower computational cost.<sup>40–44,54–58</sup> (An explanation of the composite schemes utilized herein is provided in the ESI†). Due to the large number of valence electrons for these systems for the HEAT protocol, the higher order correlation only consists of CCSDT and  $\Lambda$ -CCSDT(Q). For LuH and LuN, bond dissociation energies and enthalpies of formations were obtained using atomization energies, and standard thermochemical quantities for the atoms. For LuNH, the bond dissociation energies were obtained for three different routes:



The enthalpy of formation was obtained with the atomization scheme (route (3)), as well as from routes (1) and (2) using



experimental quantities for Lu, and NH; the computed dissociation energy of LuN was used for route (2).

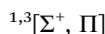
## 3 Results and discussion

### 3.1 Lutetium hydride (LuH)

The lutetium hydride curves are depicted in Fig. 2. The first two dissociation channels are considered for this molecule: Lu ( $^2D$ ) + H ( $^2S$ ), which render the following states:



and Lu ( $^2P^0$ ) + H ( $^2S$ ):



Attractive states of singlet and triplet multiplicity are depicted. The bond length ( $R_e$ ), harmonic ( $\omega_e$ ), anharmonic ( $\omega_e x_e$ ),  $\Delta G_{1/2}$ , and excitation energies ( $T_e$ ) are shown in Table 1. The CI vectors coinciding with the investigated states are displayed in Table S1 (ESI $^\dagger$ ). The ground state of LuH is a  $^1\Sigma^+$ . At equilibrium bond length, the valence 5d<sup>1</sup> electron from lutetium migrates to the 1s of hydrogen, filling the shell (1s<sup>2</sup>). MRCI + Q and CCSD(T) are both in agreement with predictions of the bond length of the complex to be 1.920 Å. In addition, the calculated  $\omega_e$ , from the PECs is 1503.0 cm<sup>-1</sup>, comparing favorably to our coupled-cluster predictions, 1487.1 cm<sup>-1</sup> (obtained with analytical gradients). In addition, both our equilibrium bond length and harmonic frequencies compare well to experimental values of Efanttin *et al.* and Wang *et al.*<sup>29,30</sup> Our  $\omega_e$  obtained with CCSD(T), with a DKH3 Hamiltonian and triple- $\zeta$  basis set is the closest value to experiment (1487.1 vs. 1476  $\pm$  20 cm<sup>-1</sup>). However, the DFT bond lengths shown in Table 1 (calculated by Pantazis *et al.* and Jalbout *et al.* with a panoply of functionals), range from 1.904 to 1.940 Å, demonstrating the sensitivity of the bond length predictions with respect to functional choice, which can dramatically impact

the bond length of lanthanide hydrides.<sup>31,32</sup> The MP2 bond length is also quite far from experiment (1.883 vs. 1.9111 Å).<sup>33</sup> Coupled-cluster predictions show sensitivity to core correlation, when considering different active and inactive orbitals within the core (below valence) set of orbitals. In addition, counterpoise corrections were also considered for these calculations.<sup>35</sup> One of the main conclusions of Cao *et al.*, based on the range of coupled-cluster predictions focused on the fact that the 4f orbitals are crucial to obtain correct results. The bond lengths obtained from Cao and Dolg fall between 1.882 to 1.914 Å.<sup>34</sup> Finally, the ic-MRCI prediction from Assaf *et al.* slightly underestimate the bond length of the ground state, though is on par with experiment.<sup>36</sup>

The first excited state of the complex is a  $^3\Delta$ , followed closely by a  $^3\Pi$ , and these states lie 12738.3 and 12963.1 cm<sup>-1</sup>, respectively, above the ground state according to MRCI+Q. The  $^3\Delta$  has two electrons coupled in the 1s of hydrogen and two unpaired electrons on the 6s and 5d of lutetium. However, the  $^3\Pi$  state has one electron on the 6p<sub>x</sub> or 6p<sub>y</sub> of lutetium, while two electrons are also coupled in the hydrogen's 1s orbital. The first  $^3\Sigma^+$  has one electron on the 6s and one on 6p<sub>z</sub> of lutetium. The first singlet excited state only appears at 16528.1 cm<sup>-1</sup> according to MRCI+Q, and it is a  $^1\Delta$ . The  $^1\Delta$ , is an open shell singlet, *i.e.*, it has one electron in the 6s of lutetium of alpha spin and a beta spin electron on the 5d orbital. The next state,  $^2^3\Pi$  is  $\sim$ 2100 cm<sup>-1</sup> above  $^1\Delta$  and has its bond length shortened by  $\sim$ 0.1 Å (1.889 Å according to MRCI+Q). This is the first triplet state that places one electron in the 5d<sub>xz</sub> or 5d<sub>yx</sub> of lutetium. The last two bonding states are the  $^2^1\Sigma^+$  and  $^2^1\Pi$ . The  $^2^1\Sigma^+$  is at 19309.1 cm<sup>-1</sup> and is a multireference state, *i.e.*, placing one alpha electron in the 6p<sub>z</sub>, a beta electron on the 6s and *vice-versa*. The last plotted state ( $^2^1\Pi$ ) is also quite multireference, alternating between alpha and beta electrons on the 6p<sub>x</sub>, 6p<sub>y</sub>, and 6s orbitals of lutetium.

The ground and the first excited states were considered for spin-orbit contributions for LuH and are displayed in Fig. 3. The following  $^{2S+1}\Lambda$  states were selected:  $X^1\Sigma^+ \rightarrow X^1\Sigma_0^+$ ,  $^1^3\Pi \rightarrow ^3\Pi_{0+}, ^3\Pi_{0-}, ^3\Pi_1, ^3\Pi_2$ ,  $^1^3\Delta \rightarrow ^3\Delta_1, ^3\Delta_2, ^3\Delta_3$ ,  $^1^3\Sigma^+ \rightarrow ^3\Sigma_0^+, ^3\Sigma_1^+$ , and  $^1^1\Delta \rightarrow ^1\Delta_2$ . The spin orbit constants and composition of states are shown in Table S2 (ESI $^\dagger$ ).

The ground state is well separated from the first excited state,  $^1^3\Pi_{0-}$ . When corrected at a spin-orbit level, the  $^1^3\Delta$  is not the first excited state anymore. In fact, the  $^1^3\Pi_{0-}$  and  $^1^3\Delta$  are separated by 700 cm<sup>-1</sup>. Furthermore, the three components of the  $^1^3\Delta$  state are separated by 2138.8 cm<sup>-1</sup>, which shows the impact of spin-orbit for these species. Also, by comparing the composition of the  $^1^3\Delta_1$  and  $^1^3\Pi_1$ , it is noted that these states are severely mixed. The  $\Omega = 2$  states are also mixed between  $^1^3\Delta_2$  and  $^1^3\Pi_2$  but not to the same extent as  $\Omega = 1$  states. The first component of the  $^1^3\Pi$  is at 12133.2 cm<sup>-1</sup> and the four components of this state are also separated by 2370.1 cm<sup>-1</sup>. The  $^1^3\Sigma_1^+$  and  $^1^3\Sigma_0^+$  are at 15248.8 and 15401.1 cm<sup>-1</sup> respectively. The last considered states in Fig. 3 is the  $^1\Delta_2$ , which lies at 16944.2 cm<sup>-1</sup>. Even though the  $^1\Delta_2$  is well separated from the rest it is still mixed with the  $^1^3\Delta$  and  $^1^3\Pi$ .

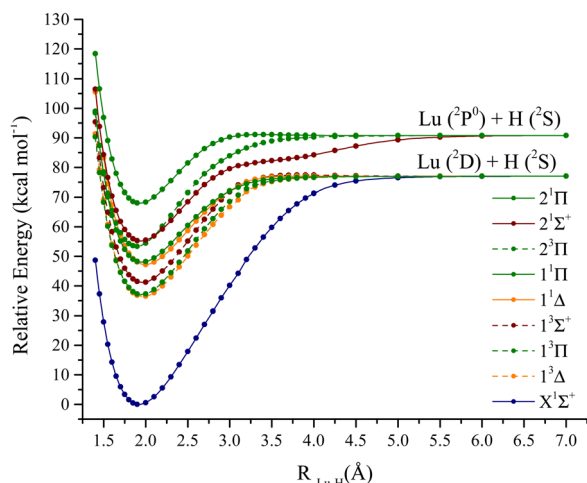


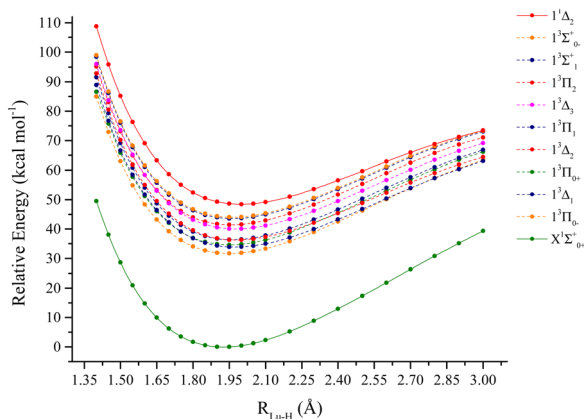
Fig. 2 LuH MRCI + Q potential energy curves (kcal mol<sup>-1</sup>) as a function of the Lu-H distance.



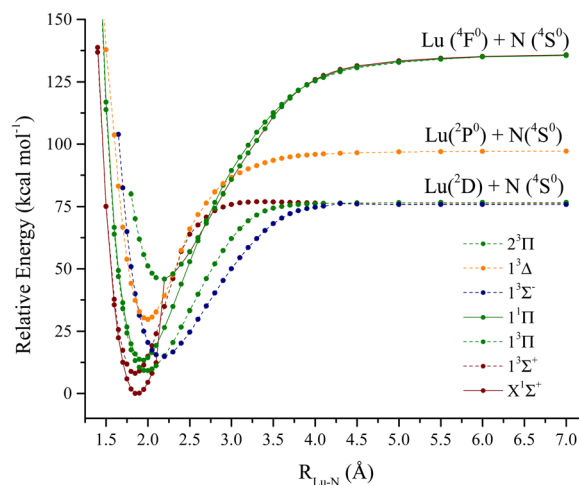
**Table 1** Bond lengths ( $R_e$ ), harmonic vibrational frequencies ( $\omega_e$ ), anharmonicity constants ( $\omega_e\chi_e$ ),  $\Delta G_{1/2}$ , and excitation energies ( $T_e$ ) for the lowest electronic excited states of LuH. CASSCF and MRCI + Q calculations were performed using cc-pVQZ-DK3 for Lu and aug-cc-pVQZ-DK with DKH3

States	Methods	$R_e$ (Å)	$\omega_e$ (cm $^{-1}$ )	$\omega_e\chi_e$ (cm $^{-1}$ )	$\Delta G_{1/2}$ (cm $^{-1}$ )	$T_e$ (cm $^{-1}$ )
$X^1\Sigma^+$	CASSCF	1.982	1404.8	17.6	1370.5	0
	MRCI+Q	1.920	1503.0	18.3	1467.9	0
	CCSD(T) <sup>a</sup>	1.920	1487.1	—	—	0
	PBE0 <sup>31,b</sup>	1.904	—	—	—	0
	DFT functionals <sup>32</sup>	1.911–1.940	1511.8–1557.6	14.0–20.48	—	0
	MP2 <sup>33</sup>	1.883	1540	—	—	0
	CCSD(T) <sup>35</sup>	1.882–1.914	1507–1577	—	—	0
	CI(SD+Q) <sup>34</sup>	1.95	1445	—	—	0
	ic-MRCI <sup>36</sup>	1.906	1539.24	18.69	—	0
	Exp. <sup>30</sup>	—	1476 ± 20	—	—	0
	Exp. <sup>29</sup>	1.9111	1520 ± 20	22 ± 1	—	0
$1^3\Delta$	CASSCF	2.035	1313.1	25.8	1256.4	14426.5
	MRCI+Q	1.966	1421.7	22.2	1375.0	12738.3
	ic-MRCI <sup>36</sup>	1.962	1436.42	21.71	—	12653
$1^3\Pi$	CASSCF	1.990	1386.5	23.7	1336.5	12763.0
	MRCI+Q	1.949	1448.4	22.8	1402.0	12963.1
	ic-MRCI <sup>36</sup>	1.938	1469.7	22.18	—	12828
$1^3\Sigma^+$	CASSCF	2.013	1336.5	24.6	1283.3	14435.0
	MRCI + Q	1.949	1426.6	22.5	1380.3	14418.3
	ic-MRCI <sup>36</sup>	1.945	1453.25	30.55	—	14116
$1^1\Delta$	CASSCF	2.056	1303.2	36.0	1220.7	17369.0
	MRCI + Q	1.994	1389.3	28.8	1327.6	16528.1
	ic-MRCI <sup>36</sup>	1.987	1401.07	27.60	—	16335
$1^1\Pi$	CASSCF	2.037	1271.9	57.5	1132.4	18954.1
	MRCI+Q	1.952	1396.0	32.9	1323.0	16808.8
	ic-MRCI <sup>36</sup>	1.942	1400.68	31.23	—	16843
$2^3\Pi$	CASSCF	1.988	1442.0	68.7	1262.8	19743.9
	MRCI+Q	1.889	1566.5	32.8	1492.7	18688.0
	ic-MRCI <sup>36</sup>	1.879	1575.38	25.42	—	18514
$2^1\Sigma^+$	CASSCF	2.056	1299.5	36.4	1218.5	17381.1
	MRCI+Q	1.941	1420.1	32.7	1346.1	19309.1
	ic-MRCI <sup>36</sup>	1.928	1427.95	30.17	—	19364
$2^1\Pi$	CASSCF	2.012	1479.5	—	—	23138.9
	MRCI+Q	1.941	1494.4	46.8	1384.6	23786.7
	ic-MRCI <sup>36</sup>	1.916	1488.29	35.34	—	23620

<sup>a</sup> Values calculated in this study. CCSD(T) with a DKH3 Hamiltonian. cc-pVTZ-DK3 for Lu and cc-pVTZ-DK for H. <sup>b</sup> The data from prior studies reports the exact number of digits as provided in the original sources.



**Fig. 3** LuH MRCI + Q spin-orbit potential energy curves (kcal mol $^{-1}$ ) as a function of the Lu–H distance.



**Fig. 4** LuN MRCI + Q potential energy curves (kcal mol $^{-1}$ ) as a function of the Lu–N distance.

### 3.2 Lutetium nitride (LuN)

The lutetium nitride molecule PECs are depicted in Fig. 4. From an electronic standpoint, this molecule is a lot more challenging

to analyze when compared to LuH. The lowest dissociation channels of LuN are:



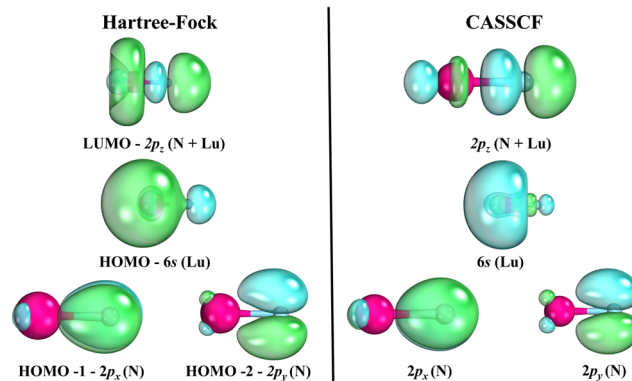
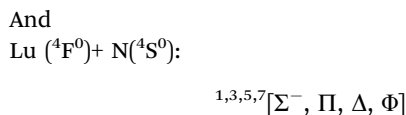
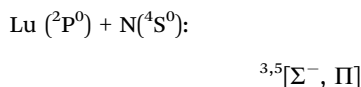
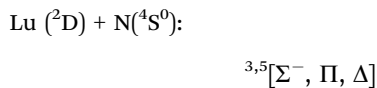


Fig. 5 Hartree–Fock orbitals versus selected CASSCF orbitals. For purposes of visualization, the Hartree–Fock orbitals were obtained with cc-pVDZ-DK3 (Lu) and aug-cc-pVDZ (N) at the equilibrium bond length for this method (2.010 Å). The CASSCF orbitals were collected from the PECs at 1.85 Å.

The spectroscopic constants calculated for LuN are shown in Table 2 and the calculated CI vectors are provided in Table S3 (ESI<sup>†</sup>). Even though, from the considered dissociation channels there are not any  $\Sigma^+$  states, when considering the lowest three dissociation channels of LuN (by Wigner–Witmer rules), the ground state is a  ${}^1\Sigma^+$ . There is crossing between 2.4–2.5 Å, from a higher dissociation channel (possibly the ionic channel), the  ${}^1\Delta$  state switches to the  ${}^1\Sigma^+$  ground state. This state is formed by nitrogen abstracting three electrons from lutetium, the ground state is  $\text{Lu}^{3+}\text{N}^{3-}$ , rather than  $\text{Lu}^{2+}\text{N}^{2-}$ . The latter is the electronic structure of the aforementioned  ${}^1\Delta$  state. As shown by the CI vectors in Table S3 (ESI<sup>†</sup>), the ground state is composed of three different components ( $\text{Lu}^{3+}\text{N}^{3-}$ ); Hartree–Fock can only describe one, which in this case is not one of the main CI vectors predicted by CAS/MRCI. In Fig. 5, the Hartree–Fock valence orbitals (except for the 2s of nitrogen) responsible for the bonding are shown. In addition, the LUMO orbital was also included for comparison. Furthermore, a comparison with the natural orbitals obtained with CASSCF is provided. At the Hartree–Fock level, the ground state determinant corresponds to filled  $2p_{x,y,z}$ , of nitrogen (HOMO–1 and HOMO–2), while the HOMO orbital is the distorted 6s of lutetium. This configuration has a very low probability (CI  $\sim$  0.11) when compared to the natural orbital occupations. Furthermore, an orbital rotation of the  $2p_z$  orbital with the 6s was performed; however, the energy of

the  $X^1\Sigma^+$  state became a lot higher. When CASSCF is utilized (and furthermore MRCI), there are three determinants responsible for the ground state. These consist of: six electrons in the  $2p_{x,y,z}$  orbitals from nitrogen, and two combinations of alpha and beta electrons on the  $2p_z$  of nitrogen and 6s of lutetium. The difference between CCSD(T) and CASSCF/MRCI+Q bond lengths arises from a multireference ground state ( ${}^1\Sigma^+$ ), along with incorrect orbitals, which cannot be described correctly by single reference methods. Despite the discrepant equilibrium bond lengths, the predicted frequencies by CASSCF, MRCI + Q and CCSD(T) are close in energy.

Regarding the excited states of LuN, the lowest six states were described with CASSCF/MRCI+Q and the  ${}^1\Sigma^+$  state was also described with CCSD(T). For LuN and in contrast to LuH, the excited states are not accessed from the internuclear region of the ground state, *i.e.* not the Franck–Condon region, which pertains to vertical transitions from the initial state (for example the ground state). In addition, there are also multiple crossings, which will be explained further. The first excited state is a  ${}^3\Sigma^+$ , also a product of a higher dissociation channel, which occurs between 2.5 Å and 2.4 Å. This state is constructed by placing an unpaired electron on the nitrogen  $2p_z$  orbital and lutetium 6s, which leads to a bond that is  $\text{Lu}^{2+}\text{N}^{2-}$  in nature. The bond length of the  ${}^1\Sigma^+$  is of the same magnitude of the  $X^1\Sigma^+$ , 1.844 Å according to MRCI+Q. For CCSD(T), the predicted bond length is 1.840 Å, which compares quite well to MRCI+Q. The main component of this state corresponds to a CI vector of 0.89, obtained from the CI vectors, making it largely single reference in nature as compared to the ground state, so it can also be described by single reference methods. The next excited state,  ${}^3\Pi$  is separated only by  $\sim 300\text{ cm}^{-1}$  from  ${}^1\Sigma^+$  according to MRCI + Q, however at the CASSCF level, the  ${}^3\Pi$  is the first excited state. Due to having unpaired electrons in the nitrogen’s  $2p_x$  or  $2p_y$  orbitals, the  ${}^3\Pi$  bond length is elongated to 1.978 Å. The next state in Table 2 is  ${}^1\Pi$ , which is an “open shell” singlet. The  ${}^1\Pi$  is the singlet version of the first  ${}^3\Pi$ , however, this state has alpha or beta

Table 2 Bond lengths ( $R_e$ ), harmonic vibrational frequencies ( $\omega_e$ ), anharmonicity constants ( $\omega_e\chi_e$ ),  $\Delta G_{1/2}$ , and excitation energies ( $T_e$ ) for the lowest electronic excited states of LuN. CASSCF and MRCI + Q calculations were performed using ECPMWB28–Seg for Lu and aug-cc-pVTZ for N

States	Methods	$R_e$ (Å)	$\omega_e$ ( $\text{cm}^{-1}$ )	$\omega_e\chi_e$ ( $\text{cm}^{-1}$ )	$\Delta G_{1/2}$ ( $\text{cm}^{-1}$ )	$T_e$ ( $\text{cm}^{-1}$ )
$X^1\Sigma^+$	CASSCF	1.908	773.5	6.6	765.6	0
	MRCI+Q	1.872	799.9	6.8	777.9	0
	CCSD(T)	2.010	757.2	—	—	0
${}^1\Sigma^+$	CASSCF	1.874	803.3	6.7	782.8	2898.4
	MRCI+Q	1.844	812.6	4.1	801.0	2865.2
	CCSD(T)	1.840	802.6	—	—	3452.5
${}^1\Sigma^-$	CASSCF	2.093	526.1	2.5	522.1	685.1
	MRCI+Q	1.978	562.7	1.5	556.9	3255.5
${}^1\Pi$	CASSCF	1.968	666.3	3.0	658.4	4476.2
	MRCI+Q	1.937	726.3	4.7	722.3	4716.4
${}^1\Sigma^-$	CASSCF	2.219	475.0	1.7	468.6	1111.5
	MRCI+Q	2.175	517.4	2.8	512.1	5199.7
${}^1\Delta$	CASSCF	2.134	339.1	6.2	415.1	17556.5
	MRCI+Q	1.993	743.6	4.6	733.2	10460.5
${}^2\Sigma^+$	CASSCF	1.977	737.0	3.5	724.1	8076.8
	MRCI+Q	2.167	533.0	2.6	528.2	16025.7



electrons in the 6s and 2p<sub>x</sub> or 2p<sub>y</sub> of lutetium and nitrogen, respectively. The bond length of the 1<sup>1</sup>Π is also elongated when compared to the ground state to 1.978 Å, at the MRCI + Q level. The first 3<sup>3</sup>Σ<sup>-</sup> state is only separated by 400 cm<sup>-1</sup> above the 1<sup>1</sup>Π. This state is the first one to have two unpaired electrons in nitrogen's p orbitals. This triplet is formed by placing one unpaired electron in each of the 2p<sub>x</sub> and 2p<sub>y</sub> orbitals. This state is quite single reference; its main CI component is 0.94. However, due to these two unpaired electrons in nitrogen orbitals, there is less orbital overlap, which elongates the Lu-N bond to 2.175 Å according to MRCI+Q. The last two states in Table 2 are triplets, the first of Δ symmetry and the 2<sup>3</sup>Π. CASSCF predicts the 2<sup>3</sup>Π to be lower in energy than the 1<sup>3</sup>Δ, however, MRCI + Q has the order reversed. The 2<sup>3</sup>Π has two main components in which two electrons are placed in the 2p<sub>z</sub> (nitrogen), two in the 6s of lutetium and one alpha and one beta electrons in the 2p<sub>x</sub> and 2p<sub>y</sub> orbitals of nitrogen. In addition, the 2<sup>3</sup>Π is also one of the longest states investigated for LuN, 2.167 Å at MRCI + Q level. In contrast, the 1<sup>3</sup>Δ six different CI contributions to the state, make this state extremely multi-reference in nature.

The spin-orbit curves of LuN are displayed in Fig. 6. Due to the complexity of these curves, and because the excitations are not vertical, only the ground and first three excited states are calculated. The spin-orbit spectroscopic constants and composition of the four considered states are shown in Table S4 (ESI<sup>†</sup>). The 2<sup>5</sup>+1Λ states split into the Ω-states as follows: X<sup>1</sup>Σ<sup>+</sup> → X<sup>1</sup>Σ<sup>+</sup><sub>0</sub>, 1<sup>3</sup>Σ<sup>+</sup> → 1<sup>3</sup>Σ<sup>+</sup><sub>0</sub>, 1<sup>3</sup>Σ<sup>+</sup><sub>1</sub>, 1<sup>3</sup>Π → 1<sup>3</sup>Π<sub>0</sub>, 1<sup>3</sup>Π<sub>1</sub>, 1<sup>3</sup>Π<sub>2</sub>, and 1<sup>1</sup>Π → 1<sup>1</sup>Π<sub>1</sub>. Even though the curves were not extended beyond 2.05 Å due to the multiple crossings, the four components of the 2<sup>3</sup>Π were also included in the calculation to have a more complete mixture of states and compositions. In addition, the 2<sup>3</sup>Π state was not tabulated, but its data was added to the state composition row in Table S4 (ESI<sup>†</sup>).

Contrary to LuH, the mixing of the Ω-states of LuN at the equilibrium bond length is much less extensive. However, there are multiple avoided crossings for this complex. The 1<sup>3</sup>Σ<sup>+</sup><sub>1</sub>/1<sup>3</sup>Π<sub>1</sub> and 1<sup>3</sup>Σ<sup>+</sup><sub>0</sub>/1<sup>3</sup>Π<sub>0</sub> avoid each other as shown in Fig. 6. Even though the

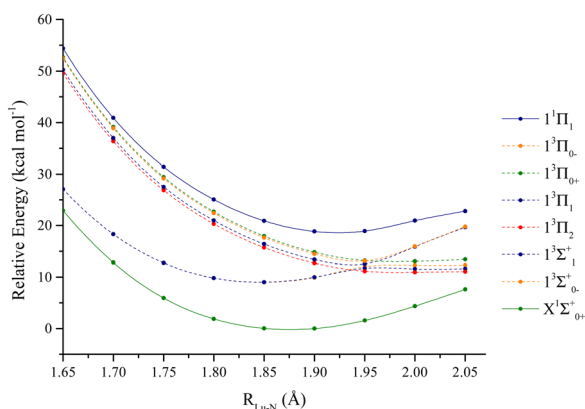


Fig. 6 LuN MRCI + Q spin-orbit potential energy curves (kcal mol<sup>-1</sup>) as a function of Lu-N distance.

ground state is separated by 3215.3 cm<sup>-1</sup> from the first excited state (1<sup>3</sup>Σ<sup>+</sup><sub>0</sub>), the rest of the considered states are quite close in energy. The 1<sup>3</sup>Π components are separated by 734 cm<sup>-1</sup>. It is interesting to note that the bond lengths of the different components also range from 2.002 Å (1<sup>3</sup>Π<sub>1</sub>) to 1.942 Å (1<sup>3</sup>Π<sub>0</sub>). Without spin-orbit estimations, it is not possible to accurately describe these states. Unfortunately, after 2.1/2.2 Å there are multiple crossings that do not allow for further extension of the PECs.

### 3.3 Lutetium imidogen (LuNH)

Combining the properties of LuH and LuN allows an understanding of the bonding properties of LuNH. To form this triatomic, all the possible isomers were considered. Herein, the following questions are raised: does lutetium bind to nitrogen and hydrogen, or just nitrogen? Which spin is the lowest, and how are the bonds of the triatomic formed? Is the lowest geometry linear or bent?

The electronic landscape of the bond formation of LuNH is shown in Fig. 7. The spectroscopic constants of the ground and first two excited states are shown in Table 3 and CI vectors in Table S5 (ESI<sup>†</sup>). The states considered for this complex at the CASSCF level comprise all of the excitations until the 6p<sub>x,y,z</sub> of lutetium of singlet and triplet spin is reached, which include 32 states. The lowest geometry consists of nitrogen as a central atom bonded to hydrogen and lutetium and it is perfectly linear (180° between Lu-N-H). The PECs were created with MRCI+Q for the ground and first excited state, and by fixing the N-H bond, utilizing the CCSD(T)/TZ with a DKH3 Hamiltonian bond length. The N-H bond length is 1.036 Å. The Lu-N distance was then stretched to seven angstroms, and the energy profile was analyzed. For this triatomic, as for LuN, there is a crossing from a higher ionic channel. However, for LuNH it occurs at ~3 Å, switching the 2Δ state to a 2Σ<sup>+</sup>. The ground state has an

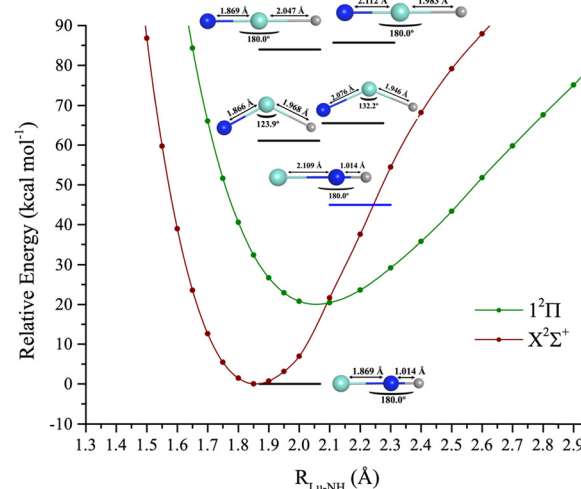


Fig. 7 Lu-NH MRCI + Q potential energy curves (kcal mol<sup>-1</sup>) as a function of the Lu-N distance. The blue horizontal bar corresponds to a CASSCF relative energy of the ground CASSCF state 2Σ<sup>+</sup>. The black horizontal bars correspond to CCSD(T)/DKH3 relative energies to the ground CCSD(T)/DKH3 state 2Σ<sup>+</sup>.



**Table 3** Bond lengths ( $R_e$ ), harmonic vibrational frequencies ( $\omega_e$ ), anharmonicity constants ( $\omega_e x_e$ ),  $\Delta G_{1/2}$ , and excitation energies ( $T_e$ ) for the lowest electronic excited states of LuN–H. CASSCF and MRCI + Q calculations were performed using ECPMWB28-Seg for Lu and aug-cc-pVTZ for N and H

States	Methods	$R_e$ (Å)	$\omega_e$ (cm <sup>-1</sup> )	$\omega_e x_e$ (cm <sup>-1</sup> )	$\Delta G_{1/2}$ (cm <sup>-1</sup> )	$T_e$ (cm <sup>-1</sup> )
$X^2\Sigma^+$	CASSCF	1.882	775.3 <sup>a</sup>	-17.8	821.0	0
	MRCI+Q	1.858	809.8 <sup>a</sup>	-5.7	835.3	0
	CCSD(T)*	1.869	552.8 <sup>b</sup>	—	—	—
		560.1 <sup>b</sup>	—	—	—	—
		816.1 <sup>a</sup>	—	—	—	
		3569.4 <sup>c</sup>	—	—	—	—
$1^2\Pi$	CASSCF	2.108	651.0	9.6	650.2	1240.1
	MRCI+Q	2.056	603.1	2.1	598.3	7021.6
$1^4\Sigma^+$	CASSCF	2.109	631.6	9.3	634.3	15801.2

<sup>a</sup> Lu–NH symmetric stretch frequency. <sup>b</sup> Lu–N–H bending frequency. <sup>c</sup> N–H asymmetric stretch frequency.

unpaired electron on the 6s of lutetium, due to the movement of one of the lutetium valence electrons to the hydrogen, filling its 1s orbital. The bond distance of the first excited state is elongated to  $\sim 2.1$  Å and it is formed by moving one unpaired electron to the 2p<sub>x</sub> or 2p<sub>y</sub> of nitrogen rather than to the 6s of lutetium.

Both structures are quite single reference as shown by the CI vector in Table S5 (ESI†). The first quartet ( $1^4\Sigma^+$ ) was not possible to determine at the MRCI level due to the time required to solve the different MRCI states, however, it was still estimated at the CASSCF level, represented by a blue horizontal bar in Fig. 7. The  $1^4\Sigma^+$  lies at  $\sim 45$  kcal mol<sup>-1</sup> and is the second excited state, after the  $1^2\Pi$ . The  $1^4\Sigma^+$  has two main components, the first by having unpaired electrons on the 2p<sub>x</sub> (N), 6p<sub>x</sub> and 6s of (Lu), and second on the 2p<sub>y</sub> (N), 6p<sub>y</sub> and 6s of (Lu). The Lu–N bond length is elongated to 2.109 Å.

The isomers of the ground state structure were also investigated. For this step, all of the isomers were optimized at CCSD(T), utilizing a DKH3 Hamiltonian, with a triple- $\zeta$  basis set. These include: (a) the linear structure, with lutetium as the central atom,  $S = 1/2, 3/2$ , and, (b) the bent structure ( $C_s$ ) also  $S = 1/2, 3/2$  (see Table S6, ESI† for geometry details). There is an interesting trend among the isomers. For the doublets ( $S = 1/2$ ), the Lu–N is  $\sim 0.3$  Å shorter than for the quartet ( $S = 3/2$ ) counterpart. The state above  $1^4\Sigma^+$  is the first state with  $C_s$  symmetry ( $1^2A'$ ), at 61.1 kcal mol<sup>-1</sup> or 21373.5 cm<sup>-1</sup>. The  $C_s$  quartet is 1542.4 cm<sup>-1</sup> above the doublet and the Lu–N bond length is elongated by  $\sim 0.2$  Å and the Lu–H shortened by 0.22 Å. For the linear structures, here treated as a  $C_{2v}$  point group, the doublet and quartet structures are only separated by 600 cm<sup>-1</sup>. However, the Lu–N bond is still quite elongated by  $\sim 0.3$  Å. The N–H bond is again shortened by  $\sim 0.06$  Å. The  $2S+1A$  spin-orbit states of the  $X^2\Sigma^+ \rightarrow X^2\Sigma^+_{1/2}$  and  $1^2\Pi \rightarrow 1^2\Pi_{3/2}, 1^2\Pi_{1/2}$  were also considered. These two states are displayed in Fig. S1 (ESI†) and their spectroscopic constants, and composition of states are shown in Table S7 (ESI†). The ground state is  $X^2\Sigma^+_{1/2}$ , followed by excited states at  $1^2\Pi_{3/2}$  and  $1^2\Pi_{1/2}$ . The  $1^2\Pi_{3/2}$  and  $1^2\Pi_{1/2}$  are separated by 417.9 cm<sup>-1</sup>.

### 3.5 Bond dissociation energies (BDEs) of LuH, LuN and LuNH

Until now, the different states and bond patterns of LuH, LuN, and LuNH have been elucidated. However, it is also important to calculate bond dissociation energies. The dissociation energies of LuH and LuN, along with the different energetics routes for breaking the LuNH bonds with  $f$ -ccCA and HEAT are shown in Table 4. The breakdown of the energetics arising from each of the different terms of  $f$ -ccCA and HEAT are shown in Tables S8 and S9 (ESI†). For LuH,  $f$ -ccCA and HEAT energies are similar, with a difference of only 1.29 kcal mol<sup>-1</sup>. However, for LuN, the difference in dissociation energy is much larger when comparing HEAT and  $f$ -ccCA, at 5.45 kcal mol<sup>-1</sup>. From Tables S8 and S9 (ESI†), the two main differences from  $f$ -ccCA and HEAT arise from the core–valence term and the contribution from the quadruple excitations in HEAT, calculated at the  $\lambda$ -CCSD(T)/double- $\zeta$  level. The core–valence correlation for HEAT is extrapolated to the CBS utilizing energies determined at the triple- and -quadruple- $\zeta$  basis set level, while for  $f$ -ccCA, the core–valence correlation is solely obtained at a double- $\zeta$  level. Another contributing factor for the energetic difference is attributed to the full triples contribution at the CBS (from triple- and quadruple- $\zeta$  basis sets) limit, with  $-10.03$  kcal mol<sup>-1</sup> for HEAT, while  $f$ -ccCA does not include full triple corrections, only perturbative triples. Furthermore, the quadruple excitation contribution is quite large at 3.38 kcal mol<sup>-1</sup>. Even though the LuN ground state is a  $1^1\Sigma^+$ , it is by no means a single reference state, which requires additional contributions from higher excitations to accurately predict its thermodynamic properties. As the pentuple contributions are quite computationally demanding, as noted earlier, the contribution from the pentuples (*i.e.*, as in  $s$ -ccCA) was not calculated, but its value may still be significant, particularly considering the size of the quadruple contribution. For the NH fragment,  $f$ -ccCA and HEAT render similar  $D_0$  values, with a difference between  $f$ -ccCA and HEAT energies of 1.03 kcal mol<sup>-1</sup>.

For LuNH, three dissociation pathways were probed (routes (1)–(3)). The first route entailed breaking the N–H bond from LuN (LuN + H). The second route was to break the NH fragment from lutetium (Lu + NH) and the third route was a total atomization scheme, *i.e.*, Lu + N + H. For the first route,  $f$ -ccCA and HEAT predictions do not agree energetically, the difference between methods is 4.09 kcal mol<sup>-1</sup>. It is interesting to note that the quadruples contribution for the  $\lambda$ -CCSDT(Q) from HEAT is  $-2.91$  kcal mol<sup>-1</sup>, to break the LuN–H bond.

**Table 4** Dissociation routes for LuH, LuN, and LuNH. All  $D_0$  are in kcal mol<sup>-1</sup>. The LuNH atomization is determined directly (Lu + N + H), and by the  $D_0$  of LuN *via* route (1)

	$f$ -ccCA $D_0$	HEAT $D_0$
LuH	72.96	71.67
LuN	83.95	78.50
NH	77.50	78.47
LuNH routes		
(1) LuNH $\rightarrow$ LuN + H	106.96	111.05
(2) Lu + NH	112.86	111.76
(3) LuNH $\rightarrow$ Lu + N + H	190.36	190.23
Dissociation energy <i>via</i> route 1 + LuN $D_0$	196.81	189.55



However, the core–valence correlation calculated from HEAT is only  $0.01 \text{ kcal mol}^{-1}$  versus  $1.36 \text{ kcal mol}^{-1}$  for *f*-ccCA, which accounts for a part of the overall energetic differences predicted by the methods. The full triples correction at the triple- $\zeta$  level renders  $8.74 \text{ kcal mol}^{-1}$ . This value is the primary difference between the *f*-ccCA and HEAT energies. The isolated fragment requires  $77.50$  and  $78.47 \text{ kcal mol}^{-1}$  for *f*-ccCA and HEAT, respectively, to break the N–H bond. However, when complexed with lutetium, this bond becomes much stronger, *i.e.* breaking the N–H bond within the complex requires  $35.36$  and  $32.58 \text{ kcal mol}^{-1}$ , as predicted by *f*-ccCA and HEAT respectively.

For route (2), breaking the NH fragment from lutetium (Lu + NH), HEAT predicts the  $D_0$  to be  $111.76 \text{ kcal mol}^{-1}$ , while *f*-ccCA predicts  $112.86 \text{ kcal mol}^{-1}$ . When comparing the energy contributions in Tables S8 and S9 (ESI<sup>†</sup>), the core valence energetic contributions are quite different. For *f*-ccCA, the contribution is positive,  $1.36 \text{ kcal mol}^{-1}$ , but for HEAT, it is negative,  $-0.21 \text{ kcal mol}^{-1}$ . For this route, the quadruples contribution is quite small ( $0.41 \text{ kcal mol}^{-1}$ ) compared to the contributions for LuN or LuN + H. The full triples correction is much smaller ( $-0.68 \text{ kcal mol}^{-1}$ ) than for route (1). For both composite methods, it is easier to break the LuN–H bond, than the Lu–NH bond. However, for *f*-ccCA, the difference between routes is  $6.1 \text{ kcal mol}^{-1}$ , while for HEAT, the difference is only  $0.71 \text{ kcal mol}^{-1}$ . The higher order coupled–cluster correlations (full triples and higher) play a crucial role in obtaining full energetic contributions, for highly multireference ground states such as LuN.

For route (3), for the total atomization approach, both HEAT and *f*-ccCA result in a very similar  $D_0$  ( $\sim 190 \text{ kcal mol}^{-1}$ ). However, if the dissociation energy is calculated *via* route (1), *i.e.*, summing the atomization of LuN to the value obtained through route (1) ( $D_0 \text{ LuNH} \rightarrow \text{LuN} + \text{H} + D_0 \text{ LuN}$ ), *f*-ccCA predicts the  $D_0$  to be  $196.81 \text{ kcal mol}^{-1}$ , while HEAT results in an energy of  $189.55 \text{ kcal mol}^{-1}$ , which is near to the total atomization energy predicted *via* route (3).

The total atomization energies calculated in route (3) are the common ground when comparing the different dissociation energies.

## 4 Conclusions

In this project, for the first time (to our knowledge), the elucidation of the electronic structure of multiple building blocks of a lutetium nitride doped network have been described. The PECs of LuH, LuN, and LuNH are described in detail including an accounting for spin–orbit effects. The LuH molecule ground state is a well-separated ground ( $X^1\Sigma^+$ ) and it is formed by pairing two electrons on the  $1s$  orbital of hydrogen and  $6s$  of lutetium. For lutetium hydride, after constructing the Breit–Pauli Hamiltonian, severe mixing of the  $\Omega = 1$  states of the  $1^3\Pi$ ,  $1^3\Delta$ ,  $1^3\Sigma^+$  was observed, which consequently shifted the first excited state of this complex; now a  $1^3\Pi_0^-$  followed by  $1^3\Delta_1$ . In fact, the three components of the  $1^3\Delta$  state are separated by  $2138.8 \text{ cm}^{-1}$  at a spin–orbit level.

Lutetium nitride is a more complex molecule from the electronic structure standpoint. The ground state is constructed by three different components, which makes single reference methodologies breakdown. The ground state of LuN is also a  $1^1\Sigma^+$ , which originates from a crossing with a potential ionic channel. The excited states of this molecule are not in the Frank–Condon region, however, herein are still characterized by MRCI + Q, and at spin–orbit level. For LuN, there is a lot less mixing when comparing the composition tables of this molecule versus LuH, and there are multiple crossings.

For LuNH, the ground and two first excited states were investigated. The ground state is a  $1^2\Sigma^+$ , which also comes from a higher dissociation channel. It is observed that the linear molecule, with the nitrogen as the center atom is the most stable structure, followed by the bent (N–Lu–H) and linear (N–Lu–H).

An in-depth analysis of the different thermochemical routes for bond breaking of the following complexes: LuH, LuN and LuNH bonds was performed with *f*-ccCA and HEAT schemes. Both *f*-ccCA and HEAT energetics agree for LuH and NH, but are  $\sim 5.3 \text{ kcal mol}^{-1}$  different for LuN. The HEAT composite scheme considers quadruple excitation contributions through  $\lambda$ -CCSDT(Q), in this case at a double- $\zeta$  level that show the importance of these higher-level contributions. In addition, the full triples contribution is quite important, especially for LuN. The HEAT and *f*-ccCA composite schemes show that LuNH  $\rightarrow$  LuN + H bond requires the least energy to break (route (1)). However, the full triple and  $\lambda$ -CCSDT(Q) energetic contributions are crucial for accurate energetics profiles when breaking the LuN bond. In addition, both methods agree in the energy predictions using the total atomization scheme LuNH  $\rightarrow$  Lu + N + H at  $190.36$  and  $190.23 \text{ kcal mol}^{-1}$  for *f*-ccCA and HEAT, respectively.

To the best of our knowledge this is the first *ab initio* study of the different fragments of the nitrogen-doped lutetium hydride network. Herein, the foundations of bond formation of the different molecules that compose the LuNH<sub>x</sub> lattice, along with bond lengths, harmonic and anharmonic frequencies, and accurate thermodynamic dissociation energies of LuH, LuN and LuNH are provided. This investigation will provide experimentalists with detailed information of the building blocks of a nitrogen-doped lutetium hydride network, enabling a full understanding of its electronic structure.

## Conflicts of interest

There are no conflicts to declare.

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