MARVEL analysis of the measured high-resolution spectra of ¹⁴NH

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Abstract

Rovibronic energy levels are determined for four low-lying electronic states ($X^{3}\Sigma^{-}$, $A^{3}\Pi$, $a^{1}\Delta$, and $c^{1}\Pi$) of the imidogen free radical (¹⁴NH) using the MARVEL (Measured Active Rotational-Vibrational Energy Levels) technique. Compilation of transitions from both laboratory measurements and solar spectra, found in 18 publications, yields a dataset of 3002 rovibronic transitions forming elements of a measured spectroscopic network (SN). At the end of the MARVEL procedure, the majority of the transitions form a single, self-consistent SN component of 2954 rovibronic transitions and 1058 energy levels, 542, 403, and 58 for the $X^{3}\Sigma^{-}$, $A^{3}\Pi$, and $c^{1}\Pi$ electronic states, respectively. The $a^{1}\Delta$ electronic state is characterized by 55 A-doublet levels, counting each level only once. Electronic structure computations show that unusually the CCSD(T) method does not accurately predict the $a^{1}\Delta$ excitation energy even at the complete basis set limit.

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Figure 1: Schematic representation of the three electronic band systems of NH analyzed in this work, with Hund's case a quantum numbers spin (Σ) and orbital (Λ) projections considered for each state separately; note that these quantum numbers are in general not good ones for NH. The total project is given by $\Omega = |\Lambda + \Sigma|$. The spin-forbidden intercombination band system is depicted using a dashed line.

1. Introduction

Imidogen, the NH free radical, is easily formed by flash photolysis of ammonia, often in excited electronic states [1, 2]. The spectral signature of NH is observed in a variety of plasmas, including laser-produced plasmas in air [3] and liquids [4], plasmas used for waste water treatment [5], and it also serves to measure ammonia concentration in plasmas [6]. Emissions can also be observed from NH trace species due to impurities in or seeding of fusion plasmas [7, 8]. NH emissions are also observable in flames [9, 10]. NH has also received considerable attention as an ultracold species that can be trapped using buffer gas cooling, magnetic trapping, or Zeeman relaxation [11–14] and has been the subject of detailed, state-resolved energy transfer experiments [15, 16]. Lasing action has also been observed involving particular rovibronic states of NH [17]. NH is an important astronomical species, whose $A^{3}\Pi-X^{3}\Sigma^{-}$ spectrum was first recorded in the laboratory by Eder in 1893 [18]. This electronic transition has been detected in the Sun [19, 20], in the comet Cunningham [21], in the interstellar medium by its absorption against various stars [22–25], and in the atmospheres of cool stars themselves [26]. The c¹\Pi–a¹ Δ band system was originally observed by Dieke and Blue in 1934 [27] (see Figure 1). Line lists are available which provide line strengths for rovibrational and rotational transitions within the X³ Σ^{-} ground state of ¹⁴NH [28, 29] and for the A³ $\Pi-X^{3}\Sigma^{-}$ band system [30].

In this paper we analyse all the available high-resolution spectra of NH recorded in the laboratory [31–54] and obtained from spectroscopic observations of the Sun [54–56]. The purpose of this work is to provide accurate empirical energy levels with dependable uncertainties for ¹⁴NH using the Measured Active Rotational-Vibrational Energy Levels (MARVEL) approach [57–60] (though the traditional name refers to rovibrational states, there is nothing in the original proposal of MARVEL [57] which prevents the use of rovibronic transitions in a MARVEL analysis). MARVEL energy levels have been used in the past to compute accurate, temperature-dependent ideal-gas thermodynamic data [61], to facilitate the empirical adjustment of potential energy surfaces [62], and to improve the accuracy of computed line lists [63, 64].

The configuration of the ground electronic state of NH is $1\sigma^2 2\sigma^2 3\sigma^2 1\pi^2$. The doubly occupied π orbital gives rise to the X³ Σ^- ground electronic state and the lowlying metastable excited a ¹ Δ and b ¹ Σ^+ states. Rearrangement of the electrons to $1\sigma^2 2\sigma^2 3\sigma^1 1\pi^3$ leads to two further low-lying electronic states designated A ³ Π and c ¹ Π . *Ab initio* calculations show that these are indeed the dominant configurations of these five electronic states [65, 66]. Following some earlier studies [67, 68], Owono Owono *et al.* [65] provide a comprehensive set of *ab initio* potential energy curves (PEC) for the states considered here. The work of Owono Owono *et al.* [65] is complemented by a high-accuracy study of the X³ Σ^- ground-state PEC by Koput [69] and by other calculations [66]. This work started out by concentrating on the five lowest-lying electronic states of NH. However, although the metastable $b^{1}\Sigma^{+}$ state has been observed spectroscopically [1, 34], there are insufficient data to warrant a MARVEL study including this state's energy levels. Analysis of this state therefore awaits the results of further laboratory measurements. Figure 1 illustrates the electronic states and the electronic band systems of NH which we consider in this work. We also revisit the issue of obtaining accurate electronic excitation energies using standard quantum chemistry methods for this 8 electron system.

2. Methodological details

2.1. MARVEL

The MARVEL approach [57–60] enables a set of assigned experimental transition wavenumbers to be converted into empirical energy levels with associated uncertainties that are propagated from the input transitions to the output energy levels. This conversion relies on the construction of a spectroscopic network (SN) [70–73], which contains all measured and assigned inter-connected transitions. For a detailed description of the approach, algorithm, and code, we refer the reader to Ref. [59].

The MARVEL approach has been used to validate compiled experimental rovibronic transitions for the astronomically important species ${}^{48}\text{Ti}{}^{16}\text{O}$ [74], which proved to be an important step in providing a greatly improved line list for TiO [75]. Although the spectroscopy of TiO is somewhat more complicated, there are similarities between this study and the one presented in Ref. [74]. Other MARVEL studies on chemically or astronomically important molecules include those for ${}^{12}\text{C}_2$ [76], ${}^{12}\text{C}_2\text{H}_2$ [77], ${}^{14}\text{NH}_3$ [78, 79], three isotopologues of SO₂ [80], $\text{H}_2{}^{32}\text{S}$ [81], ${}^{90}\text{Zr}{}^{16}\text{O}$ [82], isotopologues of H_3^+ [83, 84], and a series of investigations on isotopologues of water [60, 85–90]. Note that the water studies provided the original motivation for developing the MARVEL procedure.

2.2. Quantum numbers

MARVEL requires that all transitions are assigned with a unique set of descriptors, usually quantum numbers, which are self-consistent across the entire dataset. Many studies considered here used Hund's case b quantum numbers for NH and these descriptors were adopted for this work. Each transition is therefore labeled by an electronic state label (X ${}^{3}\Sigma^{-}$, A ${}^{3}\Pi$, a ${}^{1}\Delta$, or c ${}^{1}\Pi$), a vibrational quantum number (v), the total angular momentum quantum number (J), the rotational angular momentum quantum number (N), and parity (e/f) [91]. For singlet states N is set equal to J.

Figure 1 gives a schematic representation of the rovibronic band systems considered in this work; it also shows the spin (Σ) and orbital (Λ) angular momentum projections considered for each state. In the Hund's case *b* representation employed here these quantum numbers are not actually used.

In constructing a unified set of quantum numbers the following changes were made. The label F adopted for components of the $X^{3}\Sigma^{-}$ state by Bernath and co-workers [35, 41, 48, 54, 92] were mapped to the Hund's case b quantum number N as follows:

$$N = \begin{cases} J - 1, & \text{if } F = 1 \text{ and } p = e \\ J, & \text{if } F = 2 \text{ and } p = f \\ J + 1, & \text{if } F = 3 \text{ and } p = e \end{cases}$$
(1)

The parity of a rovibronic level incident to a rovibronic transition must satisfy one of the following relations:

$$\Delta J = 1 : e \leftrightarrow e \text{ or } f \leftrightarrow f,$$

$$\Delta J = 0 : e \leftrightarrow f.$$
(2)

The interested reader should consult Figure 1 of Dixon [93] for an explanation how the various notations represent the transitions within the $A^{3}\Pi - X^{3}\Sigma^{-}$ band system.

None of the studies considered here were able to resolve the Λ -doubling of the a¹ Δ state where each (v, J = N) state should appear with either e or f parity. We decided to treat these (v, J = N) combinations as a single state and to adopt a parity label of d (for degenerate) in each case. This simplification means that odd-numbered cycles become possible in the experimental SN of ¹⁴NH. In fact, while the great majority of the cycles within the minimum cycle basis [94] of the experimental SN of ¹⁴NH have cycles of length four, the remaining cycles are largely split equally between cycles with length three and five. An example is the cycle formed by the transitions 27 550.1873 [40] and 30 826.5245 cm⁻¹ [40] connecting the a ¹ Δ and c ¹ Π states with the same upper rovibronic level, whereby the two lower rovibronic states are connected by the measured transition 3276.3820 cm⁻¹ [39]. Validation of these odd-numbered cycles supports the assertion that Λ -doubling in the a ¹ Δ state is indeed small.

3. Compilation of experimental sources

¹⁴NH has a pronounced hyperfine structure due to both ¹⁴N (I = 1) and H $(I = \frac{1}{2})$ nuclear spins and most studies of its pure rotational spectrum resolve this structure. However, as none of the studies at higher wavenumbers resolve the hyperfine structure, it was decided not to consider hyperfine effects and treat the rotational spectra hyperfine unresolved. The hyperfine structure was therefore removed from sources 04LeBrWiSi [52], 82VaMeDy [37], and 04FlBrMaOd [51]. Studies that only consider hyperfine transitions within a given (v, J, N) state [38, 45] were excluded from our analysis at this stage. Averaging the hyperfine components, of course, leads to some loss of accuracy in the pure rotational levels for which it should be possible to obtain hyperfine-resolved energies. A number of older sources [1, 32, 33, 95] were found to provide less accurate data than available from more recent measurements. These sources were not considered further.

Our preliminary analysis also showed that including the transition data of 82RaSa [36] caused conflicts within the SN. In 82RaSa [36] there are 70 measured lines in the $c^{1}\Pi$ - $a^{1}\Delta$ band system in the range 27 105–27 697 cm⁻¹ with a claimed uncertainty of about 0.005 cm⁻¹. Nearly all these lines were remeasured as part of a much more

extended study by 86RaBe [40]. Removing six lines of 82RaSa [36] from the singlet subnetwork allowed the SN to form correctly once the average uncertainty had been increased to 0.2 cm^{-1} . We note that 82RaSa used a grating spectrograph and these measurements are essentially superseded by the more accurate Fourier transform spectroscopy (FTS) measurements of 86RaBe; these two datasets agree once the uncertainty for 82RaSa is increased. Remarkably, it was found necessary to remove only one further line, due to 86BrRaBe [41], during MARVEL's validation process.

Infrared spectra of the $X^{3}\Sigma^{-}$ state of ¹⁴NH are also available from two distinct space-borne observations of solar spectra. The ATMOS [96] experiment, part of the Atmospheric Laboratory for Applications and Science (ATLAS) Space Shuttle program, provides solar spectra at a resolution of about 0.01 cm^{-1} . The Atmospheric Chemistry Experiment (ACE) [97] is a Canadian satellite mission which performs infrared solar occultation observations of the Earth's atmosphere producing detailed solar spectra in the infrared as a byproduct [98]. While the resolution of these spectra is lower than can be obtained in the laboratory, typically 0.01 cm^{-1} or lower, the high temperature of the Sun means that they probe rotational states not easily accessible in laboratory experiments. 91GeSaGrFa [56] used ATMOS to study pure rotational transitions of NH in the ground and first vibrationally excited states spanning levels with N from 20 to 42. 90GrLaSaVa [55] provides ATMOS R-branch spectra of the vibrational fundamental and the first hot band. However, these spectra do not resolve the fine structure splittings and there is only a single transition, the fundamental R(N'' = 27) line, at 3389.147(10) cm⁻¹, which was not available from much higher resolution laboratory studies. As a result, we decided not to include transitions from 90GrLaSaVa [55] in our final compilation. Conversely, 10RaBe [54] analyses ACE spectra in the region of the vibrational fundamental where ACE gives much higher signal-to-noise ratio than ATMOS. 10RaBe also present laboratory spectra, so we divided their results into two segments and label them 10RaBe and 10RaBe_S2 to distinguish these independent subsets of data, with the latter source segment containing the less accurately measured

transitions.

Table 1: Data source segments and some of their characteristics for the $^{14}\rm NH$ molecule^a

Segment tag	Band system	Range	A/V	ESU	ASU	MSU
97KlTaWi [47]	$X{}^3\Sigma^ X{}^3\Sigma^-$	31.570 - 33.356	3/3	3.00e-06	3.00e-06	3.00e-06
04FlBrMaOd [51]	$X{}^3\Sigma^ X{}^3\Sigma^-$	65.213 - 162.67	12/12	5.00e-06	5.00e-06	5.00e-06
07RoBrFlZi [53]	$X{}^3\Sigma^ X{}^3\Sigma^-$	28.977 - 156.21	34/34	1.00e-05	1.00e-05	1.00e-05
82VaMeDy [37]	$X{}^3\Sigma^ X{}^3\Sigma^-$	32.505 - 33.356	2/2	2.00e-05	7.54e-05	1.19e-04
86LeEvBr [42]	$X{}^3\Sigma^ X{}^3\Sigma^-$	98.495 - 98.495	1/1	2.00e-05	2.00e-05	2.00e-05
04LeBrWiSi [52]	$X{}^3\Sigma^ X{}^3\Sigma^-$	32.505 - 66.169	4/4	1.00e-04	1.84e-04	3.99e-04
86BoBrChGu [44]	$X{}^3\Sigma^ X{}^3\Sigma^-$	2310.1 - 3456.8	310/310	5.00e-04	7.82e-04	3.83e-03
82BeAm [35]	$X{}^3\Sigma^ X{}^3\Sigma^-$	2950.0 - 3293.1	29/29	1.00e-03	1.64e-03	5.28e-03
85HaAdKaCu [39]	$a{}^1\Delta - a{}^1\Delta$	3177.9 - 3537.2	35/35	1.00e-03	3.86e-03	3.88e-02
86BrRaBe [41]	$A{}^3\Pi-X{}^3\Sigma^-$	26289 - 33080	1239/1238	1.00e-03	3.48e-03	4.11e-02
91GeSaGrFa [56]	$X-X{}^3\Sigma^-$	622.45 - 941.28	103/103	1.00e-03	4.16e-03	2.80e-02
99RaBeHi [48]	$X{}^3\Sigma^ X{}^3\Sigma^-$	2151.9 - 3459.0	500/500	1.00e-03	2.34e-03	2.63e-02
10RaBe [54]	$X{}^3\Sigma^ X{}^3\Sigma^-$	2151.8 - 2530.2	36/36	1.00e-03	1.00e-03	1.00e-03
$10RaBe_{S2} [54]$	$X{}^3\Sigma^ X{}^3\Sigma^-$	2662.1 - 3459.0	266/266	5.00e-02	5.00e-02	5.00e-02
86UbMeTeDy [43]	$c{}^{1}\Pi - a{}^{1}\Delta$	30059 - 30742	38/38	2.00e-03	5.57e-03	1.17e-02
03VaSaMoJo [50]	$A{}^{3}\Pi-a{}^{1}\Delta$	17002 - 17193	52/52	2.00e-03	1.10e-02	2.74e-02
86RaBe [40]	$c{}^{1}\Pi - a{}^{1}\Delta$	26653 - 32938	250/250	3.00e-03	4.57e-03	3.85e-02
82RaSa [36]	$c{}^{1}\Pi - a{}^{1}\Delta$	27106 - 27697	70/64	5.00e-03	2.64e-02	6.98e-02
90HaMi [46]	$c{}^{1}\Pi - a{}^{1}\Delta$	21608 - 24525	18/18	2.00e-02	2.00e-02	2.00e-02

^a Tags denote experimental data-source segments employed during this study (the identifier associated with the first segment of a data source, ' $_S1$ ', is not written out explicitly). The column 'Range' indicates the range (in cm⁻¹) corresponding to validated wavenumber entries within the experimental linelist. 'A/V' is an ordered pair, where A and V are the number of assigned and validated transitions related to a given source segment, respectively, obtained at the end of the MARVEL analysis. ESU, ASU, and MSU designate the estimated, the average, and the maximum segment uncertainties in cm⁻¹, respectively. Rows of this table are arranged in the order of the ESUs with the restriction that the segments of the same data source should be listed one after the other.

There are a number of studies of the intercombination band system which link the singlet and triplet subnetworks [49, 50, 99]. 99RiGe [49] used stimulated emission pumping of the strongly forbidden a ${}^{1}\Delta - X {}^{3}\Sigma^{-}$ band system to study the singlet-triplet splitting. However, the data of 99RiGe [49] only have an accuracy of 0.1 cm⁻¹ or worse and the source does not actually provide the primary transition data, so this source was not considered further. 03VaSaMoJo [50] used optical pumping to study a ${}^{1}\Delta - A {}^{3}\Pi$ transitions with an accuracy of 0.03 cm⁻¹ or better. 03VaSaMoJo [50] also used the F



Figure 2: The experimental spectroscopic network of the ¹⁴NH molecule showing the different electronic states considered during this work with different color, red: $X^{3}\Sigma^{-}$, green: $A^{3}\Pi$, blue: $a^{1}\Delta$, and black: $a^{1}\Delta$. The relatively large number of bridges characterizing the spectroscopic network of ¹⁴NH is evident from this figure.

labelling convention; therefore, their assignments were transformed as described above. Lower resolution experimental studies of the $a^{1}\Delta - X^{3}\Sigma^{-}$ emission spectra by 84RoSt [99] do not provide any transition wavenumbers and were not considered further.

4. Results and Discussion

Table 1 presents a summary of the transitions data used in the final MARVEL run of this study. The latest version of the MARVEL code (intMARVEL [60]), with some minor modifications, was used to obtain the energy levels and the refined transitions. A total of 3002 assigned transitions from 18 distinct data sources were included in our final MARVEL analysis. Of these transitions and energy levels, 2954 and 1058, respectively, are contained within the principal component of the experimental SN of ¹⁴NH linking levels in all four electronic states of interest. 48 transitions and 62 energy levels could not be linked to the principal component, they are part of floating components, but

they are retained in the dataset as they may be linked to the principal component when new experimental data become available.

After the reassignments and other manipulations considered in the last section, only 7 transitions had to be removed from the dataset considered by MARVEL as not consistent with the other transitions. These transitions are retained in the final transition list but are given as negative wavenumber entries.

The principal component of the experimental SN of ¹⁴NH assembled during this study links together 1058 energy levels (see Figure 2). Of these energy levels, 44, 818, 17, 179 have resistances P^+ , P^- , S, and U, respectively (definitions of the labels can be found in the ReadMe.txt file of the Supplementary Material). Energy levels with resistance P^+ are indeed dependable, while the others may change when even more accurate lines will be included in the database of experimental transitions. Although it is likely that even the non-P⁺ energy levels are correct, especially those of P⁻, they cannot be regarded as fully verified by the present analysis.

Due to the lack of truly high-accuracy first-principles energies for ¹⁴NH, we used spectroscopic constants available in the literature to confirm the empirical (MARVEL) rovibrational energy levels derived during this study. For the $X^{3}\Sigma^{-}$ state, we used the PGOPHER [100] file of 15BrBeWe [29] to compute the rovibrational energy levels. Figure 3 shows the differences between the MARVEL levels and this earlier work. As seen in Figure 3, there are only few energy levels where the difference is larger than 0.05 cm^{-1} so the agreement is considered to be very good.

For the energy levels of the a¹ Δ state, we used the molecular constants of 86RaBe [40], in the cases of v = 0 and 1, and 90HaMi [46], for v = 2 and 3. Prior to 1974 [101], no intercombination transitions could be measured; therefore, the energy separation of the singlet and triplet states of NH had to be estimated by indirect experimental data or by *ab initio* calculations. This is the reason why in 1963 McBride [102] set the value of $T_{\rm e}({\rm a}^{1}\Delta)$ to 14 922 cm⁻¹ but in 1979, in their famous book, Huber and Herzberg [103] published 12 566 cm⁻¹ for this value, and 10 years later Gurvich [104]

IN11.									
frozen-core	$\Delta E_{\rm e}({\rm UHF})$	δ [MP2]	δ [CCSD]	δ [CCSD(T)]	δ [CCSDT]	δ [CCSDTQ]	$\delta[\text{CCSDTQP}]$	$\delta[\text{CCSDTQPH}]$	$\Delta E_{\rm e}[{\rm FCI}]$
aug-cc-pwCVDZ	23300.4	-3981.3	-2229.6	-1079.7	-1787.2	-148.6	-4.3	-0.0	14069.9
aug-cc-pwCVTZ	23222.7	-4838.1	-2043.0	-1078.9	-2002.3	-229.9	[-4.3]	[-0.0]	[11947.3]
aug-cc-pwCVQZ	23236.1	-5168.2	-1914.7	-1074.6	-2034.0	-247.7	[-4.3]	[-0.0]	$[11\ 735.8]$
aug-cc-pwCV5Z	23240.9	-5316.0	-1836.4	-1074.9	-2039.4	-	[-4.3]	[-0.0]	[11665.0]
CBS	${\bf 23241.7(20)}$	-5471.1(500)	-1754.2(250)	-1075.3(10)	-2045.1(40)	-266.5(250)]	[-4.3(20)]	[-0.0(10)]	${\bf 12625.2}({\bf 750})$
all-electron	$\Delta E_{\rm e}({\rm UHF})$	δ [MP2]	δ [CCSD]	δ [CCSD(T)]	δ [CCSDT]	δ [CCSDTQ]	$\delta[\text{CCSDTQP}]$		$\Delta E_{\rm e}$
aug-cc-pwCVDZ	23300.4	-3995.9	-2133.8	-1093.6	-1846.8	-159.8	-4.9		14065.6
aug-cc-pwCVTZ	23222.7	-4823.6	-1924.2	-1089.4	-2100.5	-254.3	[-4.9]		
aug-cc-pwCVQZ	23236.1	-5149.6	-1794.2	-1083.1	-2136.0	-276.0	[-4.9]		
aug-cc-pwCV5Z	23240.9	-5296.1	-1716.1	-1082.6	_	-	[-4.9]		
CBS	23 241 7(20)	-5449 9(500)	-1634 1(250)	-1082.0(20)	-2168.0(40)	-298 7(250)	[-4.9(20)]		126041(750)

Table 2: Focal-point-analysis table of the $T_{\rm e}$ excitation energy between the $X^{3}\Sigma^{-}$ and $a^{1}\Delta$ states of 14 NH.

^{*a*} The symbol δ denotes the increment in the relative energy ($\Delta E_{\rm e}$) with respect to the preceding level of theory in the hierarchy HF \rightarrow MP2 \rightarrow CCSD \rightarrow CCSD(T) \rightarrow CCSDT \rightarrow CCSDTQ \rightarrow CCSDTQP \rightarrow CCSDTQPH (\equiv FCI in the case of the frozen-core approximation). CBS = complete basis set. The basis set extrapolations are based on the cardinal number X of the aug-cc-pCVXZ Gaussian basis-set family, for electron correlation increments they follow the formula X^{-3} and employ the largest two X values available. Uncertainties are given in parentheses. All energy values are given in cm⁻¹.

used a value of 12 577.1 cm⁻¹ for T_e when calculating the thermochemical functions of the ¹⁴NH molecule. The large shift in the T_0 value of the a ¹ Δ state over time can be explained by the pronounced multireference character of this state, due partly to the fact that it dissociates to an excited state of the N atom. The considerable difficulties the gold standard CCSD(T) level has in providing a correct estimate of $T_e(a^{1}\Delta)$ can be seen from the frozen-core and all-electron focal-point-analysis-type [105, 106] entries of Table 2. The CCSD to CCSDT increment is huge and CCSD(T) can recover only less than half of this increment, a rather unusual situation. Though the CCSDTQ increment is not particularly large, it is unusually basis-set dependent, showing the necessity of a multireference treatment for obtaining the excitation energy of the a¹ Δ state. The increments above CCSDTQ are very small, as usual and expected. Overall, the excitation energy is only about half of its unrestricted Hartree–Fock (UHF) estimate. One can estimate $T_e(a^{1}\Delta)$ starting from the all-electron CBS CCSDTQ value



Figure 3: Differences between the empirical MARVEL energy levels of this study and the earlier literature results of 15BrBeWe [29] related to the $X^{3}\Sigma^{-}$ electronic state.

of Table 2, 12604(75) cm⁻¹, and adding to this the higher-order coupled-cluster estimates, -5(2) cm⁻¹, as well as the relativistic correction (computed at the all-electron aug-cc-pwCVQZ CCSD(T) level), -2 cm⁻¹. Adding the zero-point vibrational energy (ZPVE) correction, +27(5) cm⁻¹, obtained again at the aug-cc-pwCVQZ CCSD(T) level, yields our final $T_e(T_0)$ estimates of 12597(12624) cm⁻¹, with a conservative 2σ uncertainty estimate of ± 80 cm⁻¹. These excitation energies support the best previous experimental estimates mentioned above.

The first high-accuracy measurement of the singlet-triplet splitting, between the $a^{1}\Delta$ (v = 0, J = N = 2) and $X^{3}\Sigma^{-}$ (v = 0, J = 1, N = 0) states, was obtained by 86RaBe [40], the value is 12 688.39(10) cm⁻¹. This energy difference was given as 12 687.8(1) cm⁻¹ in 99RiGe [49] and probably the best value is given by 03VaSaMoJo [50], it is 12 688.622(4) cm⁻¹. The MARVEL value for this splitting is 12 688.612(2) cm⁻¹. Since the $T_{\rm e}$ values are not accurate enough to confirm the MARVEL levels and



Figure 4: Differences between the empirical MARVEL energy levels of this study and the earlier literature results of 86RaBe [40] and 90HaMi [46] related to the $a^1\Delta$ electronic state.



Figure 5: Differences between the empirical MARVEL energy levels of this study and the earlier literature results of 10RaBe [54] related to the $A^{3}\Pi$ electronic state.



Figure 6: Differences between the empirical MARVEL energy levels of this study and the earlier literature results of 86RaBe [40] related to the $c^{1}\Pi$ electronic state.

there are no high-accuracy values for T_v in the literature, we fitted a simple Hamiltonian, taken from 86RaBe [40], for the degenerate energy levels of the a ¹ Δ state. We obtain the following values for $T_e + T_v$: 12655.761, 15838.558, 18875.218, and 21768.236 cm⁻¹ for v = 0, 1, 2, and 3, respectively. Using these $T_e + T_v$ values and the rotational constants of 86RaBe [40] and 90HaMi [46], we could check the accuracy of the MARVEL energy levels. Figure 4 shows the differences between the MARVEL levels and the effective Hamiltonian results. It can be seen that for the v = 0 and 1 vibrational states the agreement is nearly perfect, but in the cases of higher vibrational states the differences usually are larger than 0.03 cm⁻¹.

For checking the MARVEL energy levels of the A³ Π state, we used the term values of 10RaBe [54]. The result of the comparison of the present empirical and the effective Hamiltonian energies can be seen in Figure 5. Only two MARVEL energy levels could mqtSbe reproduces twithin 0.03c emainties of the reference data are calculated as the square

ES	v	J	N	p	$E(MARVEL)/cm^{-1}$	$E(86BoBrChGu [44])/cm^{-1}$
${\rm X}{}^3\Sigma^-$	0	1	0	e	0	0
${\rm X}{}^3\Sigma^-$	1	1	0	e	3125.57241(50)	3125.57291(16)
${\rm X}{}^3\Sigma^-$	2	1	0	e	6094.87429(71)	6094.87502(20)
${\rm X}{}^3\Sigma^-$	3	1	0	e	8907.59729(87)	8907.59833(25)
${\rm X}{}^3\Sigma^-$	4	1	0	e	11562.3129(13)	11562.31434(42)

Table 3: Vibrational energy levels of ¹⁴NH.^a

root of the squared sum of the uncertainties found in Table 1 of Ref. [44].

We used the spectroscopic constants of 86RaBe [40] and our T_v values for the c¹ Π state to compute the energy levels of the c¹ Π state. Figure 6 shows the result of this comparison: the agreement is almost perfect.

The list of MARVEL energy levels and the corresponding experimental transitions are placed in supplementary data.

Table 3 compares the vibrational band origins obtained during the MARVEL analysis with available experimental results. It is striking that direct vibrational band origin information is available only for the ground electronic state of ¹⁴NH and only for v = 1-4 even though some experimental rovibrational data are available for v up to 6 in the $X^{3}\Sigma^{-}$ state, up to 3 in the $a^{1}\Delta$ state, up to 2 in the $A^{3}\Pi$ state, and for v = 0 and 1 in the $c^{1}\Pi$ state. It is also worth noting that the uncertainties obtained during the MARVEL analysis are slightly larger than the 1σ values reported in 86BoBrChGu [44]. Nevertheless, the agreement between MARVEL and 86BoBrChGu is excellent.

5. Conclusions

A MARVEL analysis of the observed and assigned rovibronic transitions has been performed for the parent imidogen radical, ¹⁴NH. After making a number of minor adjustments, a set of 1058 empirical rovibronic energy levels related to the principal component of the SN, containing 2954 validated transitions, are obtained for this radical spanning four low-lying electronic states. If desired, these energies provide suitable input for an effective Hamiltonian fit.

There are some high-resolution spectroscopic data available on higher-lying electronic states of NH including the $d^{1}\Sigma^{+}$ state [34, 107] and various Rydberg states [108–110]. However, at this stage there are insufficient data available for any of these states to make their inclusion in a MARVEL analysis productive. The ¹⁴NH data produced here will be included in the newly formed MARVEL project database [111] and can be actively updated should new high-resolution assigned spectra become available.

Finally, we note that a significant part of this work was performed by pupils from Preston Manor School in north-west London, as part of the ORBYTS (Original Research By Young Twinkle Students) project. Several other MARVEL studies have been completed as part of the ORBYTS project, including those on ${}^{48}\text{Ti}{}^{16}\text{O}$ [74], acetylene [77], H₂S [81], ${}^{90}\text{Zr}{}^{16}\text{O}$ [82], and methane [112]. Sousa-Silva *et al.* [113] discusses our experiences of working with high-school students on high-level research projects. High-school students in Hungary were also involved in a MARVEL analysis of the highresolution spectra of ${}^{16}\text{O}_2$ [114].

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