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View Article Online **PAPER**



Cite this: Phys. Chem. Chem. Phys., 2021, 23, 22057

Received 7th June 2021, Accepted 15th September 2021

DOI: 10.1039/d1cp02537f

rsc.li/pccp

Rovibronic spectroscopy of PN from first principles*

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We report an ab initio study on the rovibronic spectroscopy of the closed-shell diatomic molecule phosphorous mononitride, PN. The study considers the nine lowest electronic states, X $^{1}\Sigma^{+}$, A $^{1}\Pi$, C $^{1}\Sigma^{-}$, D $^{1}\Delta$, E $^{1}\Sigma^{-}$, a $^{3}\Sigma^{+}$, b $^{3}\Pi$, d $^{3}\Delta$ and e $^{3}\Sigma^{-}$ using high level electronic structure theory and accurate nuclear motion calculations. The ab initio data cover 9 potential energy, 14 spin-orbit coupling, 7 electronic angular momentum coupling, 9 electric dipole moment and 8 transition dipole moment curves. The Duo nuclear motion program is used to solve the coupled nuclear motion Schrödinger equations for these nine electronic states and to simulate rovibronic absorption spectra of 31p14N for different temperatures, which are compared to available spectroscopic studies. Lifetimes for all states are calculated and compared to previous results from the literature. The calculated lifetime of the $A^1\Pi$ state shows good agreement with an experimental value from the literature, which is an important quality indicator for the ab initio A-X transition dipole moment.

1 Introduction

Phosphorus is considered to be one of the key elements as a source of life and replication on our planet, 1,2 with PN being one of the candidates in star and meteorite evolution to provide the necessary life building material. There have been multiple observations of PN in different media in space: hot dense molecular clouds, 3,4 energetic star forming regions, 5,6 cold cloud cores, 5,7 red giant stars⁸⁻¹⁰ and protoplanetary nebula.⁸ Regardless of high astrophysical and astrobilogical importance, phosphorous mononitride is one of the experimentally least-well studied diatomic molecules of its isoelectronic group (P2, SiO, N2, CS).

The molecule was reported for the first time by Curry et al. 11 using a spectroscopic study. Many other subsequent spectroscopic studies have taken place including photoelectron, 12 fluorescence, 13 matrix infrared, 14 microwave 15-17 and Fourier Transform Infra-Red (FTIR).18 Most of the high resolution spectroscopy experiments with PN concentrated on the electronic system A ${}^{1}\Pi$ -X ${}^{1}\Sigma^{+}$, ${}^{11,13,19-21}$ with E ${}^{1}\Sigma^{+}$ -X ${}^{1}\Sigma^{+}$ being confirmed afterwards, 22-24 and several valence and Rydberg states have been studied as well.25,26

Measured lifetimes can be important indicators of intensities and Einstein A coefficients. So far only one experimental work reports lifetime measurements for the A ¹Π state of PN molecule²⁷ using Hanle effect, with several *ab initio* works providing computed values for PN lifetimes.^{28,29}

The mass spectrometric experiment by Gingeric³⁰ reports PN's dissociation energy D_0 to be 6.35 \pm 0.22 eV, which is lower than experimental D_0 values by Huffman et al.³¹ and Uy et al.,³² 7.1 ± 0.05 eV and 7.57 ± 0.03 eV, respectively. In a combination of a high-level ab initio and microwave spectroscopy study, Cazzoli et al. 17 suggested a D_0 value of 6.27 eV. This is close both to the originally predicted value by Curry et al. 33 and to the experimental value of Gingeric.30

Several theoretical investigations of PN are available in the literature. The most recent was carried out by Qin et al. 29 who reported spectroscopic constants for the lowest five singlet $(X^{1}\Sigma^{+}, A^{1}\Pi, D^{1}\Delta, C^{1}\Sigma^{-}, 2^{1}\Pi)$, six triplet (including a $^{3}\Sigma^{+}$, b ${}^{3}\Pi$, d ${}^{3}\Delta$, e ${}^{3}\Sigma^{-}$) and two quintet electronic states of PN. In that paper all the states were studied at the internally contracted multi-reference configuration interaction (icMRCI) level of theory with Davidson correction (+Q). Similarly, Abbiche et al. 34 reported an *ab initio* study of seven states of PN, $X^{1}\Sigma^{+}$, $A^{1}\Pi$, $D^{1}\Delta$, C $^{1}\Sigma^{-}$, 2 $^{1}\Pi$, E $^{1}\Sigma^{+}$, 3 $^{1}\Sigma^{+}$, as well as of 10 triplet and 3 quintet electronic states. The main purpose of their study was to interpret perturbation and predissociation effects in the observed transitions.

In this work we present a comprehensive ab initio spectroscopic model for the nine lowest electronic states of phosphorous mononitride, $X^{1}\Sigma^{+}$, $A^{1}\Pi$, $C^{1}\Sigma^{-}$, $D^{1}\Delta$, $E^{1}\Sigma^{+}$, $a^{3}\Sigma^{+}$, $b^{3}\Pi$, $d^{3}\Delta$, e ${}^{3}\Sigma^{-}$, consisting of potential energy curves (PECs), transition dipole moments curves (TDMCs), spin-orbit coupling curves (SOCs) and angular momentum coupling curves (AMCs) using

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[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/ d1cp02537f

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the icMRCI + Q method and calculate the rovibronic energies and transition probabilities as an ab initio line list for PN. Producing such line lists for molecules of astrophysical significance is one of the main objectives of the ExoMol project.³⁵ These curves, with some simple adjustment of the minimum energies of the PECs, are used to solve the coupled nuclearmotion Schrödinger equation with the program Duo.³⁶ The spectroscopic model and ab initio curves are provided as part of the supplementary material. Our open source code Duo can be accessed via http://exomol.com/software/. The results of Duo calculation are then used to generate rovibronic spectra of PN and compare to systems previously reported in the literature.

2 Computational details

Ab initio calculations

Using MOLPRO 2020, 37 ab initio calculations were performed for nine low-lying electronic states of PN. Apart from PECs, we

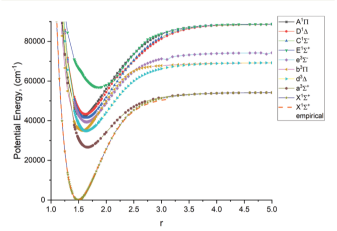


Fig. 1 Ab initio (icMRCI + Q/ECP10MWB) PECs of PN: The lowest 5 singlets and 4 lowest triplets

also computed (T)DMCs, SOCs and EAMCs. For the ab initio calculations we used the icMRCI method38 in conjunction with the effective-core-potential (ECP) method ECP10MWB (Stuttgart/ Cologne) for phosphorus³⁹ and aug-cc-pCV5Z⁴⁰ basis set for nitrogen. The initial complete active space self-consistent field (CASSCF) calculation over which the configuration interaction calculations were built was for the X $^{1}\Sigma^{+}$ state only. In conjunction with ECP for phosphorus, the active space was selected to be (6,2,2,0) with (1,0,0,0) closed orbitals. The state averaging set contained 96 states: 24, 24, 24 and 24 states of the singlets, triplets, pentets and septets. This level of theory will be referenced to as icMRCI + Q/ECP10MWB.

The calculations of the spin-orbit couplings were too difficult to perform for the icMRCI + Q/ECP10MWB level of theory, taking too long to complete and producing wrong data. We therefore decided to use a non-ECP level of theory for our spin-orbit calculations. To this end, we selected the ab initio level of theory similar to that used by Oin et al. 29 with an active space of (9,3,3,0). In this case the state averaging set consisted of 11 singlet configurations (4A₁, 2B₁, 2B₂ and 3A₂). The Douglass-Kroll correction was taken into account with or without core-valence correlation. These levels of theory will be referenced to as icMRCI/aug-cc-pV5Z(-DK) and icMRCI/aug-ccpWCV5Z(-DK), respectively, with or without DK.

Fig. 1-4 show all PECs, SOCs, EAMSc and (T)DMCs generated in this study. If the MOLPRO calculations at some geometries did not converge, they were interpolated or extrapolated from the surrounding points as part of the Duo calculations (see below). We used an adaptive ab initio grid consisting of 150 bond lengths ranging from 0.7 to 8 Å with more points around the equilibrium region. The grid points with the corresponding ab initio values (if converged) are included in the supplementary material and also shown in Fig. 1-4. The icMRCI/aug-cc-pV5Z(-DK) calculations have only converged up to about 3.5 Å due to a smaller number of configurations in the internally contacted reference set used.

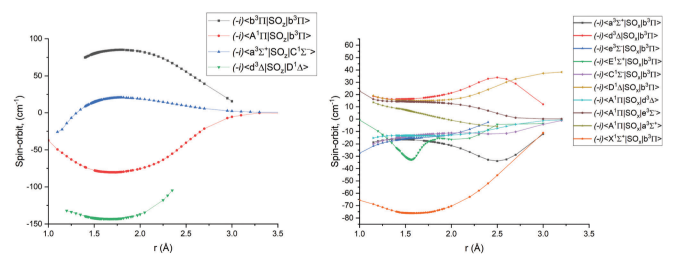


Fig. 2 Ab initio spin-orbit matrix elements (i|SO_x|j) for PN at the icMRCI/aug-cc-pV5Z-DK level of theory. The MOLPRO values of the magnetic quantum numbers $m_{\rm S}$ for the curves can be found in Table 2.

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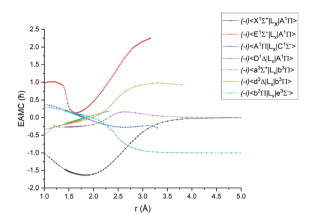
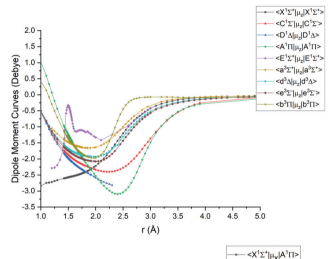


Fig. 3 Ab initio (icMRCI + Q/ECP10MWB) electronic angular momentum couplings for PN in the units of \hbar .



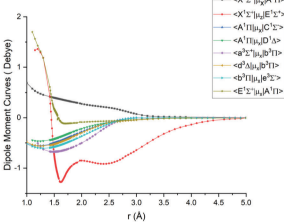


Fig. 4 *Ab initio* (icMRCI + Q/ECP10MWB) dipole moment curves for PN: diagonal (upper), off-diagonal (lower).

2.2 Duo calculations

We use the program Duo^{36,43} to solve the coupled Schrödinger equation for 9 lowest electronic states of PN. Duo is a variational program capable of solving rovibronic problems for a general (open-shell) diatomic molecule with an arbitrary number of

couplings, see, for example, ref. 44–47. All *ab initio* couplings between these 9 states are taken into account as described below. The goal of this paper is to build a comprehensive *ab initio* spectroscopic model for this electronic system of PN based on the icMRCI + Q/ECP10MWB and icMRCI/aug-cc-pV5Z-DK *ab initio* curves. We therefore do not attempt a systematic refinement of the *ab initio* curves by fitting to the experiment, which will be the subject of future work. In order to facilitate the comparison with the experimental data, we, however, perform some shifts of the equilibrium energy $T_{\rm e}$ and bond length $r_{\rm e}$ values, as described in further detail below.

In Duo calculations, the coupled Schrödinger equations are solved on an equidistant grid of points, in our case 501, with bond lengths r_i ranging from r = 0.85 to 5 Å using the sinc DVR method. Our *ab initio* curves are represented by sparser and less extended grids (see above). For the bond length values r_i overlapping with the *ab initio* ranges, the *ab initio* curves were projected onto the denser Duo grid using the cubic spline interpolation.

The following functional forms were used for extrapolation outside the original *ab initio* range:³⁶

$$f_{PEC}^{short}(r) = A + B/r,$$

$$f_{TDMC}^{short}(r) = Ar + Br^{2},$$

$$f_{other}^{short}(r) = A + Br,$$
(1)

for short range and

$$f_{\text{PEC}}^{\text{long}}(r) = A + B/r^{6}$$

$$f_{\text{EAMC}}^{\text{long}}(r) = A + Br,$$

$$f_{\text{other}}^{\text{long}}(r) = A/r^{2} + B/r^{3}$$
(2)

for long range, where A and B are stitching parameters.

The vibrational basis set was taken as eigensolutions of the nine uncoupled 1D problems for each PEC. The corresponding basis set constructed from 9 \times 501 eigenfunctions was then contracted to include about 60–80 vibrational functions from each state, enough to fill the PEC up to the corresponding dissociation energies. This gives a total of 640 states. These vibrational basis functions were then combined with the spherical harmonics for the rotational and electronic spin basis set functions. All calculations were performed for 31 P 14 N using atomic masses.

3 Results and discussion

3.1 Results of ab initio calculations

The lowest 9 singlet and triplet PECs (X $^1\Sigma^+$, A $^1\Pi$, C $^1\Sigma^-$, D $^1\Delta$, E $^1\Sigma^+$, a $^3\Sigma^+$, b $^3\Pi$, d $^3\Delta$, e $^3\Sigma^-$) calculated at the icMRCI + Q/ECP10MWB level and are shown in Fig. 1. Table 1 presents spectroscopic constants for these states estimated using the corresponding (spin–orbit-free) PECs and compares to previous studies. 17,19,29,33,34,41,42 These values agree well for the states X $^1\Sigma^+$, C $^1\Sigma^-$ D $^1\Delta$, E $^1\Sigma^+$, a $^3\Sigma^+$, e $^3\Sigma^-$, b $^3\Pi$, d $^3\Delta$ as estimated using the corresponding (spin–orbit-free) PECs. For the E $^1\Sigma^+$

Table 1 Comparison of spectroscopic constants taken from previous works and calculated from our *ab initio* curves (icMRCI + Q/ECP10MWB): Dissociation energy $D_{\rm e}$ in cm⁻¹ (rounded to 3 s.f.), electronic equilibrium energy $T_{\rm e}$ in cm⁻¹, equilibrium bond length $r_{\rm e}$ in Å, harmonic constant ω in cm⁻¹, rotational constant $B_{\rm e}$ in cm⁻¹

State		D_{e}	$T_{ m e}$	$r_{\rm e}$	ω_{e}	$B_{\rm e}$
$X^1\Sigma^+$	This work	53100.00	0.00	1.49	1303.23	0.78
	Expt. ³³	50812.91	0.00	1.4869	1337.24	0.78549
	Expt.17	51938.86	0.00		1336.992	0.78648
	Calc ²⁹	51454.07	0.00	1.4918	1339.61	0.78549
	Calc. ⁴¹	50645.15	0.00	1.4948	1333.84	0.7823
	Calc. ³⁴	49683.73	0.00	1.4977	1328.21	0.77872
$A^1\Pi$	This work	44900.00	42143.66	1.56	1041.07	0.71
	Expt. ³³	41134.26	39688.52	1.5424	1103.09	0.73071
	Expt. ¹⁹		39805.90		1103	0.731
	Calc ²⁹	40995.53	40032.55	1.5476	1104.81	0.7306
	Calc. ⁴¹	41331.86		1.5501	1100.24	0.727505
	Calc.34	40624.26	40610.00	1.556	1078.42	0.7213
$D^1\Delta$	This work	43100.07	43287.03	1.63	1016.44	0.66
	Calc. ²⁹	39993.49	42106.86	1.6073	1018.98	0.67724
	Calc. ⁴²	33713.96	41618.19	1.622	967.9	0.664
	Calc. ³⁴	39448.89	41835.96	1.614	1010.647	0.671
	Calc. ⁴¹	41555.28		1.6196	1002.269	0.665056
$C^1\Sigma^-$	This work	45300.00	41417.18	1.63	998.65	0.66
	Calc. ²⁹	42766.02	39298.72	1.6211	977.5	0.6639
	Calc. ⁴²	37827.38	37504.77	1.617	1108.8	0.669
	Calc. ³⁴	41852.09	39505.02	1.627	973.927	0.6597
$E^1\Sigma^+$	This work	30200.00	56793.82	1.86	702.48	0.50
	Calc.34	26188.80	55103.80	1.889	743.74	0.4893
$a^3\Sigma^+$	This work	26600.62	26632.70	1.66	917.51	0.63
	Calc. ²⁹	25249.32	26065.66	1.6481	941.61	0.64253
	Calc. ⁴²	19599.26	24761.21	1.669	787.4	0.628
	Calc. ³⁴	23785.28	25817.80	1.655	913.765	0.6375
$e^3\Sigma^-$	This work	32670.00	39370.41	1.64	974.15	0.65
	Calc. ²⁹	32081.01	37951.02	1.624	981.37	0.66149
	Calc. ⁴²	37827.39	37504.77	1.617	1108.8	0.669
	Calc. ³⁴	30665.19	37956.44	1.632	982.255	0.6556
$d^3\Delta$	This work	33300.00	34860.41	1.65	960.01	0.65
	Calc. ²⁹	29679.37	32975.56	1.6324	949.99	0.65376
	Calc. ⁴²	23551.38	32504.13	1.666	770.3	0.629
	Calc. ³⁴	28471.36	33125.18	1.64	959.263	0.6495
$b^3\Pi$	This work	33400.00	35146.16	1.56	1070.81	0.71
	Calc. ²⁹	29197.77	33671.79	1.5449	1113.24	0.73301
	Calc. ⁴²	21696.31	34359.20	1.558	1124.3	0.72
	Calc. ³⁴	27672.87	33843.01	1.555	1102.494	0.7224

state, however we note that there are very few studies of that state, so further computational investigation is potentially required. For the A $^1\Pi$ state, the $T_{\rm e}$ value (the equilibrium electronic energy relative to the minimum value of ${\rm X}^1\Sigma^+$) is by about 2300 cm $^{-1}$ larger than previous experimental and *ab initio* values, which in turn affects the $D_{\rm e}$ value. We consider this to be an effect of the ECP method used in the *ab initio* calculations and we note that there is a similar magnitude shift in $T_{\rm e}$ of all other states.

The *ab initio* SOCs, EAMCs and (T)DMCs are shown in Fig. 2–4 respectively, with their phases indicated and the reference distance for phases taken as 1.5 Å. For all these curves the post-processing of *ab initio* calculations included (i) inter- and extrapolations for the missing points using neighboring geometries and (ii) phase mapping using a procedure similar to the one described by Patrascu *et al.*⁴⁴ Some of the curves also cover only part of the range, which can be seen with the TDMCs $\langle X^{1}\Sigma^{+}|\mu_{z}|E^{1}\Sigma^{+}\rangle$ in Fig. 4. The rest of the points portrayed incoherent behaviour and hence were

Table 2 MOLPRO magnetic quantum numbers m_S values for the *ab initio* spin-orbit matrix elements displayed in Fig. 2

SOC	Bra $m_{\rm S}$	Ket $m_{\rm S}$
$\langle A^1 \Pi_y SO_x d^3 \Delta_{xy} \rangle$	0	1
$\langle A^1 \Pi_y SO_x e^3 \Sigma^2 \rangle$	0	1
$\langle a^3 \Sigma^{\dagger} SO_x b^3 \Pi_y \rangle$	0	1
$\langle a^3 \Sigma^+ SO_x A^1 \Pi_y \rangle$	1	0
$\langle d^3 \Delta_{xy} SO_x b^3 \Pi_y \rangle$	0	1
$\langle e^3 \Sigma^- SO_x b^3 \Pi_x \rangle$	0	1
$\langle X^1 \Sigma^+ SO_x b^3 \Pi_y \rangle$	0	1
$\langle E^1 \Sigma^+ SO_x b^3 \Pi_y \rangle$	0	1
$\langle C^1 \Sigma^- SO_x b^3 \Pi_y \rangle$	0	1
$\langle D^1 \Delta_z SO_x b^3 \Pi_x \rangle$	0	1
$\langle b^3 \Pi_x SO_z b^3 \Pi_x \rangle$	1	1
$\langle A^1\Pi_x SO_z b^3\Pi_x \rangle$	0	0
$\langle a^3 \Sigma^+ SO_z C^1 \Sigma^- \rangle$	0	0
$\langle d^3 \Delta_z SO_z D^1 \Delta_z \rangle$	0	0

dropped in the post-processing. When calculating spectra for curves with dropped points, the lines were extrapolated using cubic splines as part of the Duo calculations.

3.2 The A ${}^{1}\Pi$ -X ${}^{1}\Sigma^{+}$ band using different levels of *ab initio* theory

As the A $^{1}\Pi$ –X $^{1}\Sigma^{+}$ band is one of the most important spectroscopic systems of PN, here we compare the X $^{1}\Sigma^{+}$ and A $^{1}\Pi$ PECs and the A $^{1}\Pi$ –X $^{1}\Sigma^{+}$ TDMC computed using different levels of theory: icMRCI + Q/ECP10MWB, icMRCI/aug-cc-pV5Z-DK and icMRCI/aug-cc-pwCV5Z-DK, see Fig. 5 and 6. As part of the spectroscopic model for PN, even a slight change of these curves can have a profound effect on the simulated spectra or lifetimes

In Fig. 6, we present a comparison of TDMCs of the $A^1\Pi$ and $X^1\Sigma^+$ states calculated at different levels of theory and results previously calculated by Qin *et al.*²⁹ The TDMC calculated with the ECP10MWB stands out the most with larger values of the dipole moment around the equilibrium (*i.e.* in the spectroscopically relevant region) and the rest of the ECP-free methods giving curves which lie close to each other. We show below that

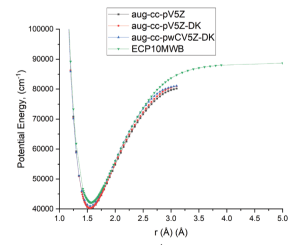


Fig. 5 Comparison of PECs for the A $^1\Pi$ state of PN at different levels of theory.

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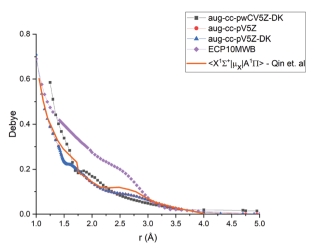


Fig. 6 Comparison of transition dipole moment between A $^{1}\Pi$ and X $^{1}\Sigma^{+}$ states of PN at the different levels of theory (ECP10MWB, icMRCI/aug-ccpV5Z-DK, icMRCI/aug-cc-pWCV5Z-DK) and previously reported results.²⁹

the ECP10MWB DMC has improved the A-X lifetime of PN, the only known experimental evidence of the transition probability of this system.

The aug-cc-pWCV5Z-DK and previous results by Qin et al.²⁹ show smaller values of TDMC which will lead to weaker intensities and longer lifetimes.

3.3 Results of Duo calculations

In this study we work directly with the ab initio data in the grid representation without representing the curves analytically, with exception of the X $^1\Sigma^+$ state for which we use an empirical potential energy function from Yorke et al.49 Using their PEC, we essentially reproduce the X $^{1}\Sigma^{+}$ states ro-vibrational energies of the YYLT line list.

For Duo calculations, we selected the following set of curves: the icMRCI + Q/ECP10MWB PECs shown in Fig. 1 for all but the $X^{1}\Sigma^{+}$ state. The comparison between the $X^{1}\Sigma^{+}$ ab initio PEC from this work and the empirical PEC from 49 can be seen in Fig. 7. Similarly for (T)DMCs all curves are ab initio, see Fig. 4, apart from the ground state dipole moment $\langle X^{1}\Sigma^{+}|\mu_{z}|X^{1}\Sigma^{+}\rangle$,

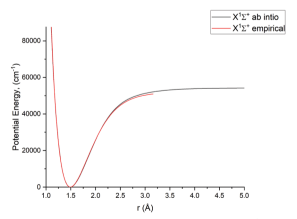


Fig. 7 Comparison between a pseudopotential *ab initio* X $^1\Sigma^+$ PEC and previously calculated empirical PEC⁴⁹ of PN.

which was also taken from Yorke et al.49 The comparison between two DMCs is shown in Fig. 8. For SOCs, TDMCs and EAMCs we use ab initio obtained curves shown in Fig. 2-4.

The rovibronic wave functions of PN were computed with Duo and then used in conjunction with the ab initio (T)DMCs to produce Einstein A coefficients for all rovibronic transitions between states considered in this work, for the wavenumber range from 0 to 88 000 cm⁻¹ and I < 270. In these calculations, the lower states were capped by the energy threshold 60 000 cm⁻¹, which is close to the lowest dissociation threshold, while the upper state threshold was set to 88 000 cm⁻¹ (highest asymptotic channel of our nine electronic state system). The line list contains 233 418 251 transitions between 390 368 rovibronic states.

These Einstein A coefficients, organised as per the ExoMol format⁵⁰ in a line list, were then used to calculate lifetimes and spectra. The aforementioned format uses a two file system to represent relevant spectroscopic information, with the energies and specific state quantum numbers included into the States file (.states) and Einstein coefficients appearing in the Transitions file linking different states (.trans). The states and trans files produced by Duo are used in conjunction with Exocross⁵¹ to produce spectra and lifetimes (see below).

3.4 Partition function

The partition function of 31P14N computed using our ab initio line list is shown in Fig. 9, which is compared to that recently reported by Barklem and Collet.⁵² Since ¹⁴N has a nuclear spin degeneracy of 3 and 31P has nuclear spin degeneracy of 2, we have multiplied Barklem and Collet's partition function by a factor of six to compensate for the different conventions used; we follow ExoMol and HITRAN53 and include the full nuclear spin in our partition functions. The differences in higher temperatures can be attributed to incompleteness in the model used by Barklem and Collet.52

3.5 Spectral comparisons

Using the ab initio 31P14N line list, spectral simulations were performed with our code ExoCross.51 ExoCross is an open

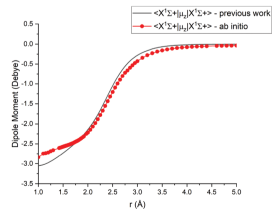


Fig. 8 Comparison between absolute values of the ab initio $X^1\Sigma^+$ DMCs from this work (icMRCI + Q/ECP10MWB) and by 49 (MRCI + Q/aug-ccpCV6Z).

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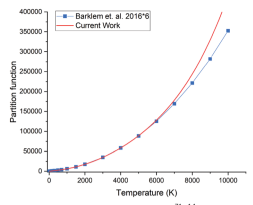


Fig. 9 Comparison of partition functions for ³¹P¹⁴N: ab initio this work and the values of Barklem and Collet. 52

source Fortran 2003 code, see http://exomol.com/software/ or https://github.com/exomol, whose primary use is to produce spectra of molecules at different temperatures and pressures in the form of cross sections using molecular line lists as input. Amongst other features, ExoCross can generate spectra for nonlocal thermal equilibrium conditions characterized with different vibrational and rotational temperatures, lifetimes, Landé g-factors, partition and cooling functions.

An overview of the PN absorption spectra in the form of cross sections at the temperature T = 2000 K is illustrated in Fig. 10. Here, a Gaussian line profile with a half-width-at-halfmaximum (HWHM) of 1 cm⁻¹ was used. This figure shows contributions from each electronic band originating from the ground electronic state. The only band systems that have so far been characterized experimentally are X $^1\Sigma^+$ -X $^1\Sigma^+$ A $^1\Pi$ -X $^1\Sigma^+$, E ${}^{1}\Sigma^{+}$ -X ${}^{1}\Sigma^{+}$. Here we mainly concentrate on discussing these band systems, due to the lack of experimental detection for other bands calculated in this work.

3.5.1 $X^{1}\Sigma^{+}$ band. The pure rotational X state transitions represent the main source of the PN observations in interstellar-medium and stellar spectra.3-10 Apart from astrophysical observations, the band has been analysed in multiple

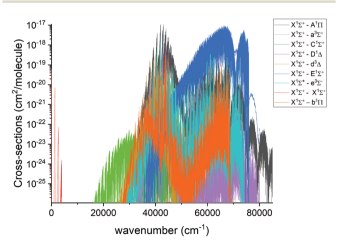


Fig. 10 Overview of the calculated absorption spectra of PN from our model at T = 2000 K. A Gaussian line profile of HWHM = 1 cm⁻¹ is used.

lab experiments as well. 14-16,18 An overview of the calculated absorption band of PN at 2000 K is shown in Fig. 11.

3.5.2 A ${}^{1}\Pi$ -X ${}^{1}\Sigma^{+}$ band. The visible A-X band system was first observed by Curry et al.33 and there have been several subsequent laboratory observations. 13,19,21,54 compare our results to the experimental spectra; an overview spectrum for the A ${}^{1}\Pi$ -X ${}^{1}\Sigma^{+}$ band system at 2000 K is given in Fig. 12.

Fig. 13 compares a Duo generated spectrum of the A ${}^{1}\Pi$ -X ${}^{1}\Sigma^{+}$ (2,0) band to a part of the spectrum from recent experimental observations by Le Floch *et al.*²¹ Even at such a high resolution the main trends of the spectrum are reproduced. However, we have to note a shift of -2217 cm^{-1} and some differences in intensities. This will be corrected with a further improvement of our model by fitting the A ¹Π PEC to the experimental data.

Fig. 14 illustrates a simulated emission A ${}^{1}\Pi$ -X ${}^{1}\Sigma^{+}$ band system at 250 nm, which is compared to the chemiluminescent experimental spectrum of Saraswathy and Krishnamurty,54 where we used the rotational temperature setting as 2000 K. However, when trying to adjust this spectrum, it proved to be very sensitive to the $T_{\rm e}$ and $r_{\rm e}$ values for the A $^{1}\Pi$ state. Therefore, for the purpose of this illustration we shifted the

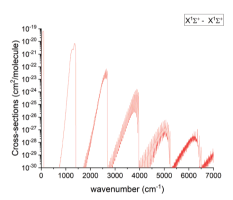


Fig. 11 Overview of the calculated X $^1\Sigma^+$ state rotation-vibration absorption spectrum of PN at T = 2000 K with a Gaussian line profile of $HWHM = 1 cm^{-1}$

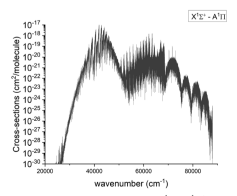


Fig. 12 Calculated absorption spectra of the A $^{1}\Pi$ – X $^{1}\Sigma$ + system at 2000 K with a Gaussian profile of HWHM = 1 cm^{-1} .

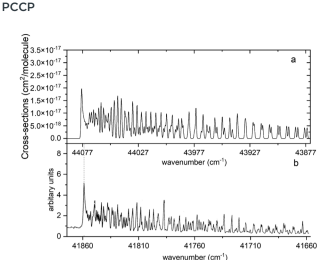


Fig. 13 Comparison of the A ${}^{1}\Pi$ –X ${}^{1}\Sigma^{+}$ (2,0) simulated absorption band (panel a) with that recorded by Le Floch et al.²¹ (panel b). The spectrum was simulated at T = 1173.15 K with a Gaussian profile of HWHM = 0.5 cm⁻¹ and is offset by 2217 cm⁻¹ so that the features align.

A ${}^{1}\Pi$ PEC to match experimental values of $T_{\rm e}$ and $r_{\rm e}$. There is also a difference in relative magnitudes, which is attributed to the difference in vibrational population distribution of the experiment and our LTE calculations, as there are non-LTE effects which contribute to the spectrum. Even though the PEC was adjusted to the experimental values previously reported by Ghosh et al. 19 to reproduce this spectrum, there is still a difference of 0.6 nm between ab initio spectrum and experiment, which we aim to resolve with further improvements of our spectroscopic model via empirical refinements.

Fig. 15 shows a simulated non-LTE absorption A ${}^{1}\Pi$ -X ${}^{1}\Sigma^{+}$ spectrum of PN to compare to the experiment of Moeller and Silvers. 13 Most of the major features are again shifted by \sim 13.5 nm. We attribute the relative difference in strength of

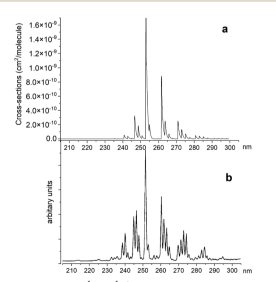


Fig. 14 Comparison of A ${}^{1}\Pi$ – X ${}^{1}\Sigma^{+}$ simulated emission spectra (panel a) with that recorded by Saraswathy and Krishnamurty.⁵⁴ (panel b). The spectra was simulated at 2000 K rotational temperature with a Gaussian profile and HWHM = 100 cm⁻¹

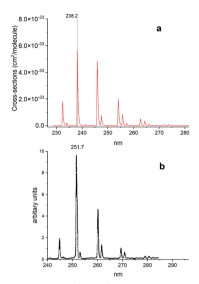


Fig. 15 Comparison of an A ${}^{1}\Pi$ -X ${}^{1}\Sigma^{+}$ simulated emission spectrum (panel a) with that recorded by Moeller and Silvers. 13 (panel b). The number at the top indicates the centre of the peak. The spectrum was simulated at $T_{\rm rot}$ = 500 K (rotational) and $T_{\rm vib}$ = 1200 K (vibrational) temperatures with a Gaussian profile of HWHM = 100 cm^{-1}

peaks between theory and experiment to difficulty in reproducing the non-LTE behaviour of the experiment, due to the lack of necessary information in the original paper.

3.6 Lifetimes

In this work, lifetimes of PN are calculated using ExoCross,⁵¹ based on the states and transitions files generated by Duo. ExoCross calculates lifetimes as follows:

$$\tau_i = \frac{1}{\sum_{j < i} A_{ij}} \tag{3}$$

where τ_i is radiative lifetime, A_{ii} is Einsteins A coefficients, and i and j stand for upper and lower states, respectively. While the lifetimes reported in other works cited below are for vibrational levels and the above formula is for an individual state; however we have found no strong J-dependency for the PN lifetimes. This means that I = 0 does indeed give a good approximation for the vibrational state lifetime. The lack of this dependency can be seen in Fig. 16. The methodology used is described in detail by Tennyson et al.55 Fig. 16 shows structures for different vibrational levels, which speaks to the perturbed nature of the A ${}^{1}\Pi$ state.

The lifetimes of $^{31}P^{14}N$ in the A $^{1}\Pi$ state for (v = 0) were measured by Moeller et al.,27 and also were calculated by Qin et al.²⁹ and de Brouckere et al.²⁸ While Moeller et al.²⁷ report a lifetime of 227 \pm 70 ns for ν' = 0 of A $^{1}\Pi$ our value is only slightly higher at 341 ns, which is closer than previous theoretical calculations of 695.4 ns²⁹ and 742.4 ns²⁸ by a factor of 2. Table 3 provides a more detailed comparison between our calculated lifetimes and those of Qin et al.29 We assume that the major differences in the A ${}^{1}\Pi$ lifetimes and previous calculation to the difference in the transition dipole moment

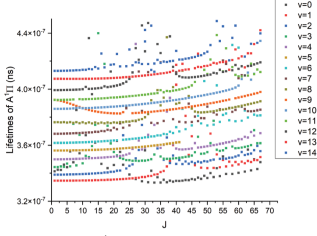


Fig. 16 Lifetimes of A $^{1}\Pi$ state vs. J. The top row of points is v=14 whereas the bottom is v=0.

between X $^1\Sigma^+$ and A $^1\Pi$ comparison of which can be seen in Fig. 6. In order to check that, we have extracted the A $^1\Pi$ TDMC data from the original source 29 and rerun the lifetime calculation, then getting a comparable result of 677 ns for $\nu'=0$. For the b $^3\Pi$ state, lifetimes seem to be in agreement with Qin *et al.* 29 for all but $\nu=0$, but same cannot be said for the D $^1\Delta$ and e $^3\Sigma^-$ states. We attribute the bulk of differences in lifetimes for these states to the differences in TDMCs.

Lifetimes for other singlet and triplet states are reported in Table 4. From there we can see that the radiative lifetimes are in milliseconds for C $^1\Sigma^-$, d $^3\Delta$ and a $^3\Sigma^+$, and nanoseconds for E $^1\Sigma^+$. Long lifetimes for the C $^1\Sigma^-$, d $^3\Delta$ and a $^3\Sigma^+$ states are an indication of low probability of transition from these states, leading us to believe that these states would be very difficult to observe in an experiment. This is confirmed with current experimental evidence for PN, as C $^1\Sigma^-$, D $^1\Delta$, a $^3\Sigma^+$, d $^3\Delta$, e $^3\Sigma^-$, which have only been observed indirectly through perturbation with the state A $^1\Pi$ by Le Floch *et al.* 21 The b $^3\Pi$,

Table 3 Comparison of lifetimes from our current work (A) and Qin et al. 29 (B). An experimental lifetime 227 \pm 70 ns was reported by Moeller et al. 27 for ν' = 0 of A $^{1}\Pi$

	A ¹Π/ns		D ¹ Δ/μs		b ³ Π/μs		$e^{-3}\Sigma^{-}/\mu s$	
ν'	A	В	A	В	A	В	A	В
0	341.3	659.4	936.03	3872	18.76	49.2	128.51	206.9
1	334.6	674.3	301.51	1186	15.38	27.81	52.43	98.46
2	338.9	660.8	120.96	615.2	13.07	19.5	73.08	59.45
3	344.3	646.5	110.39	378.3	11.43	15.05	61.85	40.75
4	349.8	642.2	88.39	246.8	10.22	12.4	53.48	30.41
5	356.2	643	69.23	181.8	10.06	10.62	47.30	23.68
6	361.6	632.1	56.85	144.3	8.55	9.364	42.60	19.26
7	368.1	610.6	47.37	113	7.94	8.356	38.88	16.16
8	376.3	607.4	40.67	9.37	7.45	7.458	35.84	13.93
9	392.4	598.6	35.52	9.85	7.04	6.881	33.40	12.2
10	399.3	583.8	31.58	10.31	6.70	6.232	31.33	10.77
11	407.3	581.8	28.33	11.32	6.42	5.592	29.60	9.517
12	413.0	566.8	25.69	13.06	6.19	5.283	28.07	8.146
13	420.7	565.3	23.34	17.18	5.99	4.819	26.78	7.142
14	431.3	553.1	21.66	22.57	5.82	4.303	25.67	6.192

Table 4 Calculated lifetimes for the C $^1\Sigma^-$, a $^3\Sigma^+$ and d $^3\Delta$ E $^1\Sigma^+$ states with ν' up to 14. A full list of lifetimes is included in the Exomol states file

ν'	C $^1\Sigma^-/ms$	a $^3\Sigma^+/s$	$d^{3}\Delta/ms$	$E^{\ 1}\Sigma^{\scriptscriptstyle +}/ns$
0	65.02	2.07×10^{6}	4.19	58.70
1	53.72	9.58	9.81	52.43
2	19.31	4.54	5.86	48.71
3	8.21	2.85	2.77	44.25
4	4.35	2.02	1.44	39.88
5	2.68	1.50	0.64	37.44
6	1.83	1.18	0.03	34.03
7	1.33	0.97	0.39	30.09
8	1.02	0.81	0.19	28.67
9	0.81	0.69	0.35	27.89
10	0.66	0.06	0.31	27.76
11	0.56	0.41	0.27	27.97
12	0.48	0.33	0.23	28.23
13	0.42	0.09	0.17	28.71
14	0.37	0.04	0.05	29.77

d $^3\Delta$, e $^3\Sigma^-$ states have been also previously identified only through perturbation with $A^1\Pi$ state by Saraswathy and Krishnamurty. 20

4 Conclusion

In this work, a comprehensive ab initio spectroscopic model for the nine lowest electronic states of PN is presented. A full set of potential energy, (transition) dipole moment, spin-orbit coupling, and electronic angular momenta coupling curves for these 9 electronic states was produced ab initio using the icMRCI + Q/ECP10MWB and icMRCI/aug-cc-pV5Z(-DK) methods. These curves were then processed via the Duo program to solve the fully-coupled nuclear-motion Schrödinger equation. Many of the results show satisfactory agreement with previous computational works, but there are certain differences in lifetimes and predicted spectra which indicate that further investigation backed by experimental data is needed to obtain a reliable spectroscopic model. In the next work we aim to tackle such differences, by producing an accurate, empirical line list for ³¹P¹⁴N for use in astrophysical spectroscopy of distant stars and exoplanets. In order to achieve this, the ab initio curves will be refined by fitting to the experimental data collected and processed via MARVEL methodology. 56 Additionally non-adiabatic coupling effects, which are most important for the heavily perturbed $A^{1}\Pi$ state, will be included for finer accuracy. With PN being detected in multiple different interstellar media and phosphorous being a key element to life as we know it, PN is becoming a more important spectroscopic molecule. This work should provide an improved data quality for spectroscopic searches of PN in different astronomical environments, potentially leading to the detection of PN at ultraviolet wavelengths.

The supplementary materials in this paper contain our final spectroscopic model for PN in a form of a Duo input file, an ExoMol States file including lifetimes and a partition function file. This Duo file is ready to be directly used with the Duo program, see http://exomol.com/software/ allowing our results to be reproduced directly.

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Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors thank Alec Owens for helpful discussions during this project. This work was supported by UK STFC under grant ST/R000476/1. This work made use of the STFC DiRAC HPC facility supported by BIS National E-infrastructure capital grant ST/J005673/1 and STFC grants ST/H008586/1 and ST/K00333X/1. We thank the European Research Council (ERC) under the European Unions Horizon 2020 research and innovation programme through Advance Grant number 883830. We also want to thank Moscow Witte University for sponsoring the fellowship enabling this research. This work is additionally supported by Khalifa University of Science and Technology under Award No. CIRA-2019-054. Khalifa University High power computer was used for the completion of this work.

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