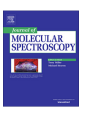
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# Microwave measurements of rotational transitions and nitrogen quadrupole coupling for 2-aminopyridine



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#### ABSTRACT

The high-resolution microwave rotational spectrum was measured for 2-aminopyridine in the 4.5–13.2 GHz range using a pulsed beam Fourier transform microwave spectrometer. The measured transitions for the parent isotopologue in the 0<sup>+</sup> vibrational state of the amino group inversion vibration were used to determine accurate rotational constants and quadrupole coupling constants for the two  $^{14}N$  atoms in the molecule. The molecular parameters determined for the 2-aminopyridine in 0<sup>+</sup> vibrational state have the following values: A = 5780.374(1) MHz, B = 2733.5017(3) MHz, C = 1857.6768(3) MHz,  $1.5\chi_{aa}$  ( $^{14}N_1$ ) = 3.5789(45) MHz,  $0.25(\chi_{bb} - \chi_{cc})$  ( $^{14}N_1$ ) = 1.5033(24)MHz,  $1.5\chi_{aa}$  ( $^{14}N_2$ ) = -0.0958(70) MHz, and  $0.25(\chi_{bb} - \chi_{cc})$  ( $^{14}N_2$ ) = -1.1615(21) MHz. The measured quadrupole coupling constants are in excellent agreement with those calculated using different computational methods and basis sets. It is noted that the calculation using the MP2 functional with cc-pvDz basis set closely reproduces the experimental values for quadrupole coupling constants for the two  $^{14}N$  atoms.

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#### 1. Introduction

2-aminopyridine is similar to the nucleic acid base adenine and has two potential hydrogen binding sites. The 2-aminopyridine-2-pyridone complex is an analog of the adenine-uracil base pair and was studied using laser spectroscopy by Leutwyler, et al. [1]

Electronic spectra for the 2-aminopyridine-ammonia complex were reported by Brutschy, et al. [2] Microwave spectra for 2-aminopyridine were measured in the 29–40 GHz range by Kydd and by Mills [3] using a Stark-cell, waveguide spectrometer. The molecule is nonplanar, with the hydrogen atoms of  $\mathrm{NH}_2$  out of the plane of the pyridine ring and is shown in Fig. 1.

Due to its feature as a derivatizing reagent, 2-aminopyridine has been utilized as a fluorescent label in various applications such as mass spectrometric analysis, oligosaccharide detection and chromatographic separation. [4–6] Similarly 2-aminopyridine and its derivatives have been found to be successful fluorescent chemosensors and are therefore good candidates for molecular sensing applications [7]. It is also effective in inhibiting the corrosions of steels and therefore 2-aminopyridine and its derivatives can serve as excellent anticorrosion agents for industrial usages. [8] Not only does 2-aminopyrdine perform its functions in chemical and industrial fields, it also emerges as a pharmacophore for

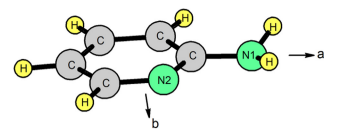
medical and pharmaceutical applications. A variety of drugs containing 2-aminopyridine available on the market have already been shown to be effective in treating infections caused by microorganisms, tumors, Alzheimer, diabetes and many other medication conditions. [9]

Another principal motivation for the structural investigation of 2-aminopyridine stems from its resemblance to tautomers 2-hyroxypyridine and 2-pyridone for which microwave measurements have been previously reported by our group. [10] In a similar way, 2-aminopyrdine can exhibit hydrogen-bonded interactions to form dimers with its tautomer and other derivatives, which can be compared to basic DNA base pairs interactions.

The microwave spectra and rotational constants for  $0^-$  and  $0^+$  vibrational states of 2-aminopyrdine and several of its deuterium-substituted isotopomers have been reported previously by Kydd and Mills. [3] The two inversion states arise from the NH<sub>2</sub> out of plane motion. Since those rotational spectrum measurements were collected in 1972, with the low resolution of the spectrometer quadrupole hyperfine structure was not resolved in that earlier work. It would also have been more difficult to resolve on the high-J transitions which were measured. Measurements of accurate quadrupole coupling constants can provide information on the electric field gradients and the electronic structure for 2-aminopyrdine. With pulsed-beam, Fourier-transform spectrometers, the quadrupole coupling splitting in 2-aminopyridne for low

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**Fig. 1.** Structure of 2-aminopyridine. N1 is the Nitrogen atom in the amino group, N2 is the Nitrogen in the ring (pyridine).

J transitions can be resolved, and the accuracies of rotational constants be significantly improved.

In the present study, we have resolved the hyperfine splitting of low-J transitions due to nuclear quadrupole coupling of the two nitrogen atoms ( $^{14}$ N, I = 1) to the overall rotation. Measurements were made on the 2-aminopydine molecule in the 4.5–13.2 MHz range using a pulsed-beam Fourier transform spectrometer. A series of theoretical computations were done to determine the quadrupole coupling strengths of the nitrogen atoms. The comparisons between experimental data and calculated values can serve as an excellent benchmark for the application of computational chemistry in molecules similar to 2-aminopyridine.

#### 1.1. Calculations

The rotational constants and quadrupole coupling constants for two <sup>14</sup>N atoms in 2-aminopyridine 0<sup>+</sup> vibrational state were calculated using DFT (Minnesota -M11 and B3LYP) and ab initio (Moller-Plesset second-order perturbation theory(MP2)) methods. M11 with Def2QZVPP basis set, B3LYP with cc-pVQZ, and MP2 with cc-pVDZ basis set and MP2 with cc-pVQZ basis set calculations were performed on the high performance computing system (HPC) at University of Arizona. The software used for these calculations is Gaussian 16. [11–12] The condition "output = pickett" was included in the Gaussian script files in order to yield microwave parameters in the Gaussian output files. The results of the calculations are compared with experimental values in Table 1. The excellent agreement of the MP2 calculations, column 3 suggests that the structure is accurately represented by these calculations so the atom Cartesian coordinates are given in Table 2.

#### 2. Experimental

Rotational transitions were measured for 2-aminopyridine in the  $4.5-13.5\,$  MHz range at University of Arizona using the

**Table 2**Calculated Cartesian coordinates in the a,b,c principal axis system from the MP2(cc-pVO7) calculation. in Å.

Atom	a	b	С
С	1.030002	-1.202934	0.000524
C	-0.895261	0.033715	-0.004740
N	-2.278280	0.036694	0.053188
Н	1.468944	-2.192166	0.000878
Н	-2.731313	0.826674	-0.371975
Н	-2.679278	-0.847594	-0.212993
N	-0.307432	-1.163937	-0.007782
C	1.834095	-0.074849	0.001544
C	-0.169288	1.231371	-0.000290
C	1.211426	1.170263	-0.002006
Н	-0.685775	2.180849	0.010455
Н	1.794020	2.080535	-0.001592
Н	2.909041	-0.168914	0.003503

Flygare-Balle type pusled-beam Fourier transform microwave spectrometer that was previously described by our group. [13-14] 2-aminopydine (98%) was purchased from Sigma-Aldrich and used without any further purification. In order to successfully carry out the gas phase microwave spectral measurement of 2aminopyridine, a sublimation test was done to determine the temperature at which it sublimes at as such information is not provided in the study reported by Kydd and Mills. A glass sample cell containing 2-aminopyridine was heated to around 85 °C in order to produce the few torr of vapor pressure necessary for microwave measurements. One end of the glass cell was connected to the pulsed valve of the microwave spectrometer while the other end was connected to a reservoir holding the carrier gas. The carrier gas system uses neon as carrier gas and the backing pressure was maintained at around 0.7 atm during the measurement. The pressure inside the spectrometer chamber was kept at  $10^{-6}$  to  $10^{-7}$  Torr powered by a diffusion pump system in order to cool molecules through supersonic expansion and populate them at their ground vibrational and rotational states. Other adjustments that include time delay, power, attenuator and cavity mode, were made to optimize molecular signal as well as signal to noise ratio. Under these conditions, the molecules were pulsed into the microwave cavity at 2 Hz through a General Valve pulsed valve and a FID with 5/1 S/N could be observed for a-type  $2_{0212}$ - $1_{0101}$  (quantum numbers J,  $K_{a}\text{, }K_{c}\text{, }F_{1}$  and F) transition at 9019.6016 MHz in 25 pulsed beam cycles. Some typical experimental spectra showing the resolved hyperfine splittings are shown in Fig. 2.

The rotational constants from the study by Kydd and Mills<sup>3</sup> as well as the calculated quadrupole coupling constants from Moller-Plesset second-order perturbation theory (MP2) functional with cc-pvDz basis were entered into the SPCAT program in Pickett's program suite [15] to predict the rotational transitions with

Table 1
The experimental and calculated molecular parameters for 2-aminopyridine in MHz.

Parameter	Experimental	3. MP2(cc-pVQZ)	4. B3LYP(cc-pVQZ)	5. M11(Def2QZVPP)	6. MP2(cc-pVDZ)	7. Kydd and Mills <sup>3</sup>
A	5780.374(1)	5827	5834	5863	5699	5780.34
В	2733.5017(3)	2745	2744	2757	2699	2733.57
C	1857.6768(3)	1869	1868	1877	1836	1857.66
1.5 χ <sub>aa</sub> (N1)*	3.5789(45)	3.455	3.657	4.042	3.119	
$0.25(\chi_{bb} - \chi_{cc})(N1)$	1.5033(24)	1.470	1.662	1.797	1.431	
$1.5\chi_{aa} (N2)^*$	-0.0958(70)	-0.380	-0.187	-0.219	-0.121	
$0.25(\chi_{bb}-\chi_{cc})(N2)$	-1.1615(21)	-1.304	-1.293	-1.427	-1.131	
χ <sub>aa</sub> (N1)	2.3859(30)	2.303	2.438	2.695	2.079	
$\chi_{bb}(N1)$	1.814(47)	1.788	2.105	2.247	1.822	
χ <sub>cc</sub> (N1)	-4.120(47)	-4.092	-4.543	-4.941	-3.902	
χ <sub>aa</sub> (N2)	-0.0639(47)	-0.253	-0.126	-0.146	-0.081	
$\chi_{bb}(N2)$	-2.291(42)	-2.481	-2.524	-2.781	-2.222	
χ cc (N2)	2.355(42)	2.735	2.648	2.927	2.302	

<sup>\*</sup> N1: Nitrogen in amino group, N2: Nitrogen in the ring (pyridine).

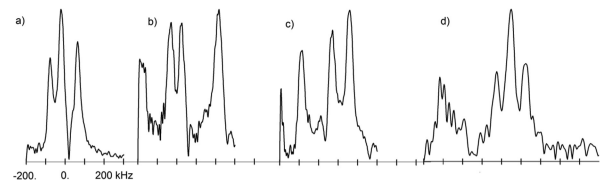


Fig. 2. Spectra showing the resolved <sup>14</sup>N hyperfine structure, a) 5973 MHz, b) 6781 MHz, c) 7638 MHz and d) 9019 MHz. The relative frequency scale is shown under a). Tick marks on the frequency scale are separated by 0.1 MHz.

**Table 3** Measured microwave transitions and the deviations from the calculated values for  $0^+$  vibrational state of 2-aminopyridine. All values are shown in MHz. Measured Transitions  $(v_{obs})$  (MHz) and deviations of the best-fit calculated frequencies  $(v_{o-c})$  (MHz) The standard deviation for the fit is  $\sigma$  = 0.0062 MHz.

(MHz) The standard deviation for the fit is $\sigma$ = 0.0062 MHz.			
Quantum number	assignments	2-Aminopyridin	e
J' Ka' Kc' F1' F'	J" Ka" Kc" F1" F"	$v_{ m obs}$	$v_{o-c}$
10101	00011	4589.9780	-0.0075
10101	00012	4589.9910	0.0055
10122	00011	4591.0410	-0.0069
10122	00012	4591.0560	0.0081
10123	00012	4591.0700	0.0076
10110	00011	4591.7570	-0.0019
10112	00012	4591.7700	-0.0035
10112	00011	4591.7800	0.0065
10111	00012	4591.7910	0.0075
1 1 1 0 1	00010	7636.9204	-0.0045
1 1 1 2 2	00012	7637.4668	-0.0016
11110	00011	7637.9414	0.0104
1 1 1 2 3	00012	7638.0786	0.0041
11112	00011	7638.5313	-0.0003
11111	00012	7639.3188	-0.0124
20212	10112	9017.8291	0.0038
20233	10123	9018.9316	-0.0036
20233	10122	9018.9629	0.0131
20234	10123	9019.1250	0.0053
20222	10111	9019.2676	-0.0095
20223	10112	9019.3711	-0.0051
20212	10101	9019.6016	-0.0117
20223	10123	9020.0872	-0.0071
20210	11111	5970.4204	-0.0076
20232	11111	5970.9219	0.0058
20212	11112	5971.0635	0.0019
20222	11111	5971.7422	0.0006
20221	11111	5971.8960	0.0015
20234	11123	5972.1045	-0.0031
20232	11121	5972.1821	-0.0019
20233	11122	5972.5352	0.0060
20223	11112	5972.6172	0.0046
20212	11101	5972.6704	-0.0034
20222	11121	5973.0190	0.0095
2 0 2 2 3 2 0 2 2 1	1 1 1 2 3 1 1 1 1 0	5973.0684 5973.2778	-0.0068
20221	11110		-0.0048
20222	11122	5973.5972 5973.7372	0.0050 -0.0055
31232	30332	6778.6372	0.0005
31232	30332	6778.9922	0.0003
31234	30334	6779.0757	-0.0031
31252	30333	6779.2778	-0.0031 -0.0177
31234	30334	6779.3262	-0.0006
31243	30333	6779.6660	0.0055
31233	30333	6779.7627	-0.0004
31233	30334	6779.9316	-0.0022
31233	30332	6780.0825	0.0068
31243	30343	6780.2275	0.0004
31232	30343	6780.3657	-0.0029
31232	30345	6780.5920	-0.0023
31234	30343	6780.7285	0.0051
31231	50511	0700.7203	0.0051

Table 3 (continued)

Quantum number assignments		2-Aminopyridin	e
J' Ka' Kc' F1' F'	J" Ka" Kc" F1" F"	$v_{ m obs}$	$v_{o-c}$
3 1 2 2 3	30323	6781.1636	0.0028
3 1 2 4 4	30345	6781.2817	0.0117
3 1 2 3 3	30323	6781.5679	-0.0081
3 1 2 4 4	3 0 3 4 4	6781.7036	-0.0027
3 1 2 2 2	30322	6782.2310	-0.0002
30322	20212	13154.7710	0.0067
3 0 3 4 4	20233	13154.8660	-0.0024
3 0 3 4 5	20234	13155.0580	0.0049
3 0 3 4 3	20232	13155.0820	-0.0050
3 0 3 4 3	20233	13155.3200	-0.0045
30334	20223	13155.3980	0.0127
30321	20211	13155.4330	-0.0001
30332	20221	13155.4550	0.0041
30332	20222	13155.6100	0.0062

quadrupole splitting in the 4.5–13.5 MHz range. The observed transitions were assigned quantum numbers J,  $K_a$ ,  $K_c$ ,  $F_1$  and F, which follow the scheme of angular momentum coupling  $I_{N1} + J = F_1$  and  $F_1 + I_{N2} = F$ , based on the predictions given by SPCAT. These assigned transitions were used in the SPFIT program contained in Pickett's program suite to fit new and improved rotational constants as well as quadrupole coupling constants and previous rotational constants. [3]

#### 3. Results

Searching for molecular signals from 2-aminopyridine in 0<sup>+</sup> vibrational state was direct and efficient since the rotational constants from previous study help us pinpoint the frequency range for rotational transitions with low J's and quadrupole splitting. Rotational transitions for 2-aminopyridine in 0<sup>-</sup> state were also predicted and searched for but were not found. The supersonic expansion nozzle of the spectrometer probably cooled the vibrational distribution too much to have sufficient population in the 0<sup>-</sup> vibrational state. A total of 66 a-type and b-type transitions were observed for 2-aminopyridne and used in the SPFIT. These rotational transitions are listed in Table 3 and consist exclusively of 0–1, 1–2 and 2–3 transitions as quadrupole coupling tend to blend and collapse for high-J transitions. The best-fit experimental molecular constants are presented in Table 1 along with theoretically calculated parameters and previous rotational constants. [3]

# 4. Discussion

For the  $0^+$  vibrational state 2-aminopyridine, the experimentally measured rotational constants are in excellent agreement with values from previous microwave measurements by Kydd

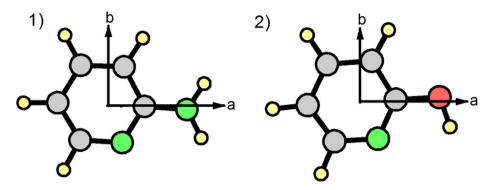


Fig. 3. Structures for 2-aminopyridine and 2-hydroxypyridine showing the a, b, c principal axis systems.

**Table 4**Comparison of <sup>14</sup>N quadrupole coupling values for 2-aminopyridne and 2-hydroxypyridne. [10] The N2 nitrogen is in the ring.

	2-aminopyridne	2-hydroxypyridne
χ <sub>aa</sub> (N2)	-0.0639(47)	-0.076(11)
$\chi_{bb}(N2)$	-2.291(42)	-2.2828(64)
$\chi_{cc}$ (N2)	2.355(42)	2.3588(64)

and Mills. [3] Numerical values for rotational constants A, B and C all agree within 1% accuracy.

There were no previous measurements of nuclear quadrupole coupling constants for the two <sup>14</sup>N atoms. The present study measured low-J rotational transitions and was able to resolve the quadrupole coupling splitting within those transitions using pulsed-beam Fourier transform microwave spectrometer. The <sup>14</sup>N atoms in 2-aminopyridine have very different quadrupole coupling strengths. By comparing experimentally measured quadrupole coupling constants with theoretically calculated values, it can be seen that calculations based on MP2 (column 3.) and the B3LYP functional (column 4.) yielded better agreements with experimental measurements than M11 functional.

We can compare the quadrupole coupling tensors for 2-aminopyridine and 2-hydroxypyridine since the principal axes relative the ring nitrogen atom are nearly coincident. The principal axis systems for 2-aminopyridine and 2-hydroxypyridine are shown in Fig. 3.

Values for the ring nitrogen (N2) quadrupole parameters are given in Table 4. The microwave spectrum and quadrupole coupling for 2-hydroxypyridine were reported by Tanjaroon et al. [10] That paper contained a detailed analysis of the experimental and calculated quadrupole coupling strengths. We note that all values are very similar, indicating a very similar electron charge distribution for the pyridine ring.

### 5. Conclusions

Low-J rotational transitions were measured and quadrupole splitting were resolved in the 4.5–13.2 GHz range for 0<sup>+</sup> vibrational state of 2-aminopyridine using a pulsed beam Fourier transform microwave spectrometer. A total of 66 a-type and b-type rotational transitions were assigned and used to determine rotational constants and quadrupole coupling constants for <sup>14</sup>N. The measured quadrupole coupling constants were compared to those calculated by different computational methods and basis sets. Calculation using MP2 and B3LYP are in better agreement with the experimentally determined values for quadrupole coupling constants than calculation using M11 functional.

## **Declaration of Competing Interest**

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

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#### References

- A. Müller, F. Talbot, S. Leutwyler, J. Am. Chem. Soc. 124 (48) (2002) 14486– 14494.
- [2] Wu. Ronghu, Sascha Vaupel, Petr Nachtigall, Bernhard Brutschy, J. Phys. Chem. A 108 (2004) 3338–3343.
- [3] R.A. Kydd, I.M. Mills, J. Mol. Spectrosc. 42 (1972) 320-326.
- [4] Y.K. Takegawa, S. Ito, S. Yoshioka, A. Sano, K. Yoshinari, K. Kobayashi, H. Nakagawa, K. Monde, S.-I. Nishimura, Anal. Chem. 76 (24) (2004) 7294–7303.
- [5] N. Kamekawa, K. Hayama, S. Nakamura-Tsuruta, A. Kuno, J. Hirabayashi, J. Biochem. 140 (3) (2006) 337–347.
- [6] Y. Araki, A. Andoh, Y. Fujiyama, K. Hata, J. Makino, T. Okuno, F. Nakanura, T. Bamba, J. Chromatogr. B, Biomed. Sci. Appl. 753 (2) (2001) 209–215.
- [7] R.R. Koner, S. Sinha, S. Kumar, C.K. Nandi, S. Ghoshl, Tetrahedron Lett. 53 (18) (2012) 2302–2307.
- [8] M. Abdallah, E.A. Helal, A.S. Fouda, Corros. Sci. 48 (7) (2006) 1639–1654.
- [9] M. Marinescu, Int. J. Pharm Bio Sci. 8 (2) (2017) 338–355.
- [10] C. Tanjaroon, R. Subramanian, C. Karunatilaka, S.G. Kukolich, J. Phys. Chem. A 108 (44) (2004) 9531–9539.
- [11] Arizona High-Performance computing resources. https://public.confluence. arizona.edu/display/UAHPC/Compute+Resources.
- [12] M.J. Frisch, G.W. Trucks, H.B. Schlegel, G.E. Scuseria, M.A. Robb, J.R. Cheeseman, G. Scalmani, V. Barone, G.A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A.V. Marenich, J. Bloino, B.G. Janesko, R. Gomperts, B. Mennucci, H.P. Hratchian, J.V. Ortiz, A.F. Izmaylov, J.L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V.G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J.A. Montgomery, Jr., J.E. Peralta, F. Ogliaro, M.J. Bearpark, J.J. Heyd, E.N. Brothers, K. N. Kudin, V.N. Staroverov, T.A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A.P. Rendell, J.C. Burant, S.S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J.W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J.B. Foresman, D.J. Fox, GAUSSIAN 16, Revision A.O.3, Gaussian, Inc., Wallingford, CT, 2016.
- [13] R.E. Bumgarner, S.G. Kukolich, J. Chem. Phys. 86 (1987) 1083–1089, https://doi.org/10.1063/1.452248.
- [14] B.S. Tackett, C. Karunatilaka, A.M. Daly, S.G. Kukolich, Organometallics 26 (2007) 2070–2076, https://doi.org/10.1021/om061027.
- [15] H.M. Pickett, J. Mol. Spectrosc. 148 (1991) 371–377. http://spec.jpl.nasa. gov/ftp/pub/calpgm/spinv.html.