

CH3CN PSEUDO-LINELIST

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INTRODUCTION.

This document gives information on the CH3CN pseudo-linelist derived at JPL in May 2005. The linelist was created based on 29 laboratory spectra taken at the Pacific Northwest National Laboratory (PNNL) and provided by Steven Sharp. These are not the same as the CH3CN cross-sections on the HITRAN website, which have been mutilated by converting all negative values to zero. The measurements and the absorption cross sections (incl. assignments of major bands) are described by Rinsland et al. (2005).

The measurement conditions for each of these spectra are tabulated below. Each measurement used the same cell of 8.1576 m length. Each spectrum covers a region between 600 and 6500 cm⁻¹ with a resolution of 0.1125 cm⁻¹ and a spectral point spacing of 0.0603 cm⁻¹.

#	File	Temp	P_tot	P_ch3cn
1	"CH3CNA.D01"	298.7	750.5	0.137679
2	"CH3CNA.D02"	298.7	749.4	0.068739
3	"CH3CNA.D03"	298.7	749.0	0.206106
4	"CH3CNA.D04"	298.7	747.8	0.411552
5	"CH3CNA.D05"	298.7	747.8	0.274368
6	"CH3CNA.D06"	298.7	747.9	0.686012
7	"CH3CNA.D07"	298.7	748.1	0.548956
8	"CH3CNA.D08"	298.7	748.1	1.029293
9	"CH3CNA.D09"	298.7	748.2	1.235317
10	"CH3CNA.D10"	298.7	748.3	0.864837
11	"CH3CNA.D11"	298.7	748.6	0.343327
12	"CH3CNB.D01"	276.1	752.7	0.069041
13	"CH3CNB.D02"	276.1	752.7	0.096658
14	"CH3CNB.D03"	276.1	752.4	0.138028
15	"CH3CNB.D04"	276.1	752.3	0.207014
16	"CH3CNB.D05"	276.1	752.3	0.276019
17	"CH3CNB.D06"	276.1	752.2	0.413974
18	"CH3CNB.D07"	276.1	752.2	0.551965
19	"CH3CNB.D08"	276.1	752.0	0.689773
20	"CH3CNB.D09"	276.1	751.9	1.034521
21	"CH3CNC.D01"	324.1	750.0	0.068794
22	"CH3CNC.D02"	324.1	750.0	0.137588
23	"CH3CNC.D03"	324.1	749.9	0.275139
24	"CH3CNC.D04"	324.1	749.9	0.687846
25	"CH3CNC.D05"	324.1	750.1	0.206409
26	"CH3CNC.D06"	324.1	750.1	0.412818
27	"CH3CNC.D07"	324.1	750.2	0.550497
28	"CH3CNC.D08"	324.1	750.4	0.344152
29	"CH3CNC.D09"	324.1	750.6	1.032733

Temp - Temperature in K

P_tot - total pressure in torr

P_ch3cn - CH3CN partial pressure in torr

DESCRIPTION.

First, the cross-sections were converted back into transmittance spectra from knowledge of the cell length and gas concentrations. The resulting laboratory transmittance spectra were then simultaneously fitted (using the GFIT algorithm) by iteratively adjusting the strengths and ground-state energies of the pseudo-lines.

Due to the resolution of the laboratory spectra of 0.1125 cm⁻¹ a pseudo-line spacing of 0.05 cm⁻¹ was considered to be appropriate. Fitting was performed in the frequency regions around 900 cm⁻¹ (where the nu₄ band is located), around 1050 cm⁻¹ (where the nu₇ band is located), and around 1450 cm⁻¹ (for the nu₃, nu₆, and nu₇+nu₈ bands). These regions include the two bands with the strongest absorption features. A zero level offset of 0.2% has been assumed throughout, based on fits to spectra in which the absorption feature at 1463 cm⁻¹ was saturated. The result of the fitting process is a continuous pseudo-linelist containing 15601 lines between 870 and 1650 cm⁻¹.

CALCULATION OF S, E", and ABHW.

At each line frequency, an effective strength and ground-state energy was derived by simultaneous non-linear least squares fitting to the 29 spectra. Furthermore, the ABHW was calculated from the ground-state energy using the formula $ABHW = 0.04 * (E'' + 2000)/(E'' + 1000)$, giving a ABHW of 0.08 cm⁻¹/atm for E''->0 and a ABHW of 0.04 cm⁻¹/atm for E''->∞. This formulation seemed to be the most appropriate to fit the feature at 1042 cm⁻¹, which is the narrowest feature in the considered frequency region. These widths are smaller than those measured by Drouin (2003) in the microwave region, but we found that using larger widths produced significantly poorer fits to the sharp spectral features. As part of the fitting, the strengths and ground-state energies were both constrained to be positive.

PARTITION FUNCTION.

The rotational partition function for CH₃CN was assumed to be (296/T)^{1.5}. The vibrational partition function was calculated in the way it had been done for the ATMOS experiment, as described e. g. by Norton and Rinsland (1991). The following vibrational frequencies and degeneracies were assumed:

freq.	2954	2267	1385	920	3009	1448	1041	362
deg.	1	1	1	1	2	2	2	2

ACCURACY.

To estimate how well the pseudo-linelist represents the PNNL spectra, test retrievals were performed in which the laboratory spectra were fitted using the pseudo-linelist. The retrieved scale factors for the CH₃CN abundances in the different spectra are tabulated below.

#	Scale factors retrieved in freq. region		
	900 cm ⁻¹	1050 cm ⁻¹	1450 cm ⁻¹
1	1.0073	0.9984	1.0065
2	1.0180	0.9993	1.0071
3	1.0043	0.9938	1.0013
4	1.0089	0.9964	1.0032
5	1.0073	0.9973	1.0037
6	1.0058	0.9953	0.9998

7	1.0018	0.9925	0.9975
8	1.0013	0.9928	0.9949
9	1.0001	0.9930	0.9946
10	0.9968	0.9880	0.9906
11	1.0033	0.9905	0.9969
12	0.9847	0.9733	0.9820
13	0.9993	0.9935	1.0019
14	1.0013	1.0003	1.0079
15	0.9885	0.9830	0.9908
16	0.9985	0.9963	1.0032
17	1.0040	1.0015	1.0083
18	1.0034	1.0004	1.0068
19	0.9930	0.9908	0.9972
20	1.0016	1.0006	1.0064
21	0.9968	0.9974	1.0086
22	1.0020	0.9944	1.0052
23	0.9987	0.9928	1.0045
24	0.9999	0.9952	1.0048
25	0.9987	0.9934	1.0054
26	0.9918	0.9891	1.0000
27	0.9993	0.9928	1.0035
28	1.0087	1.0009	1.0119
29	0.9958	0.9940	1.0013

mean	1.00072	0.99403	1.00158
stddev	0.00655	0.00588	0.00655

The pseudolines correctly represent the PNNL spectra to within 0.7% of the given CH₃CN amount in all bands analyzed. The main exception to this is spectrum 12 which appears to contain ~2% less CH₃CN than advertised.

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