	THIS PAGE					(/
AD-A21	0 062	DOCUMENTATIO	DN PAGE			Form Approved OMB No. 0704-0188
NCLASSIFIED		TIC	1b. RESTRICTIVE			ALL SALA
SECURITY CLASSIFICATION	AUTHONE E	LECTE		VAVAILABILITY		
DECLASSIFICATION / DOWN	NGRADI	£€ 0 5 1989	DISIRIBU	TON UNLIMI.	LED	
PERFORMING ORGANIZATIC	ON REF TUMB		5. MONITORING	ORGANIZATION	REPORT NUN	MBER(S)
NAME OF PERFORMING O		66. OFFICE SYMBOL	7a. NAME OF N	IONITORING ORG	ANIZATION	
.J. Seiler Researc	h Laboratory	, (If applicable) FJSRL/NC				
ADDRESS (City, State, and	ZIP Code)		7b. ADDRESS (C	ity, State, and Zll	Code)	
SAF Academy olorado 80840-652	28					
NAME OF FUNDING/SPON ORGANIZATION AF OF	NSORING fice of	8b. OFFICE SYMBOL (If applicable)	9. PROCUREMEN	IT INSTRUMENT !	DENTIFICATIO	ON NUMBER
cientific Research ADDRESS (City, State, and		AFOSR	10 SOURCE OF	FUNDING NUMBE		
1dg 410			PROGRAM	PROJECT	TASK	WORK UNIT
olling AFB DC 203	32		ELEMENT NO	NO. 2303	NO F3	ACCESSION N
PERSONAL AUTHOR(S)		3	······			<u></u>
	OD. L.L.P. S	tewart and W R	Carpor			
.P. Nash, T.E.Nels	13b. TIME C	OVERED	14. DATE OF REP		n, Day) 15	PAGE COUNT
.P. Nash, T.E.Nels a TYPE OF REPORT burnal Article	13b. TIME C FROM				n, Day) 15	PAGE COUNT
.P. Nash, T.E.Nels Ba TYPE OF REPORT DURNAL Article	13b. TIME C FROM	OVERED	14. DATE OF REP		n, Day) 15	PAGE COUNT
P. Nash, T.E.Nels a type of report ournal Article SUPPLEMENTARY NOTATION	13b TIME C FROM	OVERED	14. DATE OF REP 88 Novemb	er		4
P. Nash, T.E.Nels a TYPE OF REPORT burnal Article SUPPLEMENTARY NOTATIO	13b TIME C FROM	OVERED TO	14. DATE OF REP 88 Novemb	er		4
P. Nash, T.E.Nels a TYPE OF REPORT purnal Article SUPPLEMENTARY NOTATION COSATI CO FIELD GROUP	ON ODES SUB-GROUP	OVERED TO TO	14. DATE OF REP 88 Novemb	er		4
P. Nash, T.E.Nels a TYPE OF REPORT burnal Article SUPPLEMENTARY NOTATION COSATI CO FIELD GROUP ABSTRACT (Continue on re the x-ray structure stermined at 130K. IDO-PM3 method and stermolecular hydro	ON ODES SUB-GROUP everse if necessary es of 2,4,6- The molecu are in agre ogen contact	OVERED TO 18. SIJBJECT TERMS and identify by block of trinitrotoluene lar structure an ement with expense s correlate with	14. DATE OF REP 88 Novemb (Continue on rever number) and 2,4,6-t nd vibration rimental res h the known	er se if necessary an rinitrotolu al spectra ults. The	ene-a-dig are calco short in	4 y block number) have been ulated by the tra~and
P. Nash, T.E.Nels a TYPE OF REPORT burnal Article SUPPLEMENTARY NOTATIO	IB TIME C FROMON ODES SUB-GROUP everse if necessary es of 2,4,6- The molecu are in agre ogen contact rinitrotolue	OVERED TO 18 SUBJECT TERMS and identify by block r trinitrotoluene lar structure and ement with expenses s correlate with ne.	14. DATE OF REP 88 Novemb (Continue on rever humber) and 2,4,6-t nd vibration rimental res h the known CALL	er se if necessary an rinitrotolu al spectra ults. The thermal and f f CURITY CLASSIFI ASSIFIED (Include Area Coc	ene-á-d jare calco short in ionizat:	4 y block number) have been ulated by the tra-and ion degradatic

0584 8539 89 \$3.00 + 0.00 1989 Pergamon Press plc

Molecular structure and vibrational analysis of 2,4,6-trinitrotoluene and 2,4,6-trinitrotoluene- α - d_3

C. P. NASH, T. E. NELSON, J. J. P. STEWART* and W. R. CARPER*

Department of Chemistry, University of California, Davis, CA 95616, U.S.A.; *Frank J. Seiler Research Laboratory, United States Air Force Academy, Colorado Springs, CO 80840, U.S.A.; and †Department of Chemistry, Wichita State University. Wichita, KS 67208, U.S.A.

(Received 18 October 1988; in final form 23 November 1988; accepted 25 November 1988)

Abstract The X-ray structures of 2,4,6-trinitrotoluene and 2,4,6-trinitrotoluene- α - d_3 have been determined at 130 K. The molecular structure and vibrational spectra are calculated by the MNDO-PM3 method, and are in agreement with experimental results. The short intra- and intermolecular hydrogen contacts correlate with the known thermal and ionization degradation pathways of 2,4,6-trinitrotoluene.

INTRODUCTION

In earlier studies [1-3], the vibrational spectra of 2,4.6-trinitrotoluene (TNT) in solid, liquid, and vapor phases have been compared with vibration frequencies calculated by using semi-empirical quantum mechanical models. The structures predicted theoretically were in reasonable agreement with that found in a room temperature X-ray diffraction study [4] for the average of the two independent molecules (designated forms A and B both here and in Ref. [4]) in the unit cell. The experimental structures, however, showed unreasonably large variations in both the C-C (1.364-1.408 Å) and C-N (1.443-1.496 Å) bond distances, which prompted the low temperature (130 K) redetermination of the crystal structure which we now report. We have also determined the structure of TNT- α - d_3 at 130 K to obtain additional verification on the short inter- and intramolecular hydrogen contacts (potential hydrogen bonds) that were reported in the previous study [4]. Finally, we have recalculated both the predicted molecular structure and the predicted gas-phase vibration spectrum using the MNDO-PM3 [5, 6] method.

EXPERIMENTAL

The synthesis of 2,4,6-trinitrotoluene and 2,4,6-trinitrotoluene- α - d_3 have been described elsewhere [7]. Crystals of each compound suitable for X-ray analysis were cut from larger ones grown by the slow evaporation of solutions of the compounds in *n*-propanol (TNT) or *n*-propanol-OD (TNT- d_3). Data were collected at 130 K on a low temperature-accessorized Syntex P2₁ diffractometer.

THEORETICAL

The calculations were accomplished with the MOPAC program [8] using the MNDO-PM3 method [5, 6]. The MNDO-PM3 method uses optimized parameters that represent a considerable im-

‡Author to whom correspondence should be addressed.

provement over those parameters used with AM1 [9] and other semi-empirical methods. Typically, use of these parameters results in a significant decrease in heats of formation errors [6]. All ground state geometries were optimized using the BFGS function minimization procedure [10–13]. The Hessian matrix for the normal coordinate analysis was calculated as described previously [1].

RESULTS AND DISCUSSION

The heavy atom bond distances and bond angles obtained from the low temperature X-ray structures of 2,4,6-trinitrotoluene and 2,4,6-trinitrotoluene-x- d_3 , and the corresponding dimensions of TNT calculated by the MNDO-PM3 method are given in Tables 1 and 2. The estimated standard deviations of the heavy-atom dimensions are: angles 0.4 (TNT) or 0.4–0.5 (TNT- d_3); C-C bonds 0.006–0.007 Å (TNT) or 0.007 Å (TNT) or 0.006 Å (TNT) or 0.006–0.007 Å (TNT- d_3).

In general, the bond distances found in the present low temperature studies are in better agreement with those calculated by the MNDO-PM3 method than are the corresponding distances obtained in the previous room temperature structure determination [4]. The ring C-C bond distances span a much narrower range in the low temperature structures and the C-N bond distances are also closely similar.

The dihedral angles between the ring planes and the nitro groups differ by only a few degrees from those found in the earlier study, as shown in Table 3. The oxygen atoms on the o-nitro groups that are nearest the methyl group lie on the same side of the ring plane. In the calculated structure (gas phase) the 4-nitro group is almost coplanar with the phenyl ring, stabilized by hydrogen interactions as shown in Fig. 1, while those in the 2- and 6-positions are significantly twisted out of this plane.

The present study confirms the existence of the various short intra- and intermolecular hydrogen contacts (hydrogen bonds?) identified in the earlier



585

	TN	Г	$TNT-d_3$		ŤNT		TNT-d3
Distance*	Obs.	Calc.	Obs.	Angle	Obs.	Caic.	Obs.
C1-C2	1.509	1.487	1.495	C1-C2-C3	123.7	120.7	123.7
C2-C3	1.403	1.408	1.394	C2-C3-C4	124.3	121.3	124.5
C3-C4	1.390	1.395	1.396	C3-C4-C5	117.4	119.9	116.7
C4-C5	1.378	1.397	1.366	C4-C5-C6	122.7	119.8	123.3
C5-C6	1.389	1.396	1.372	C5-C6-C7	116.4	119.9	116.7
C6C7	1.382	1.396	1.379	C6-C7-C2	125.7	121.3	125.2
C7-C2	1.399	1.405	1.394	C7-C2-C3	113.6	117.7	113.4
C3-N1	1.475	1.516	1.470	C2-C3-N1	119.6	121.8	119.9
C5-N2	1.475	1.510	1.483	C4-C3 N1	116.1	116.9	115.6
C7-N3	1.475	1.517	1.485	C2 C7-N3	118.9	121.8	118.6
(N-O)	1.225(6)	1.211	1.218(9)	C6 C7 N3	115.6	116.9	116.1
				C4-C5-N2	119.0	119.1	118.0
				C6-C5-N2	118.3	120.1	118.7
				(C-N-O)are	117.5(6)	118.5	117.5 (9
				(O-N O)	125.0(3)	123.0	124.9 (3

Table 1. Bond distances (Å) and bond angles (degrees) for form A TNT and TNT- d_3

*Atom designations as in Ref. [4].

Table 2. Bond distances (Å) and bond angles (degrees) for form B TNT and TNT- d_3

	TN	т	TNT-d,		TNT	~	TNT-d3
Distance*	Obs.	Calc.	Obs.	Angle	Obs.	Calc.	Obs.
 С8-С9	1.494	1.487	1.492	C8-C9-C10	124.4	120.7	124.9
C9-C10	1.404	1.408	1.397	C9-C10-C11	124.7	121.3	125.1
C10-C11	1.380	1.395	1.379	C10-C11-C12	117.4	119.9	117.0
C11-C12	1.381	1.397	1.390	C11-C12-C13	122.8	119.8	123.0
C12-C13	1.379	1.396	1.376	C12-C13-C14	116.1	119.9	115.9
C13-C14	1.375	1.396	1.386	C13-C14-C9	126.4	121.3	126.1
C14-C9	1.400	1.405	1.403	C14-C9-C10	112.6	117.7	112.9
C10-N4	1.482	1.516	1.488	C9-C10-N4	119.5	121.8	119.5
C12-N5	1.467	1.510	1.464	C11-C10-N4	115.8	116.9	115.5
C14-N6	1.486	1.517	1.474	C9-C14-N6	117.8	121.8	118.0
(N-O)ave	1.228(4)	1.211	1.221(3)	C14-C12-N5	115.8	116.9	115.8
				C11C12N5	119.8	119.1	118.3
				C13-C12-N5	118.4	120.1	118.6
				$(C-N-O)_{ave}$	117.4 (5)	118.5	117.3(5)
				(O-N-O)	125.2(4)	123.0	125.4 (8

*Atom designation as in Ref. [4].

Table 3.	Dihedral angles (degrees) between nitro groups and	
	ne of the ring in 2,4,6-trinitrotoluene isotopomers	

Molecular	Nitro group					
form	2	4	6			
A (Ref. [4])	51	24	43			
A (this work)	52.0	22.3	43.4			
A-d, (this work)	51.8	23.0	42.3			
B (Ref. [4])	60	30	45			
B (this work)	58.7	33.0	39.6			
B-d, (this work)	58.9	33.0	39.5			
Calc. (this work)	55.1	9.3	76.2			



Fig. 1. Calculated molecular structure of 2,4,6-trinitrotoluene. The atom designation is that of TNT structure A in Ref. [4], and distances are in Å.

work. We observe $O \cdots H$ hydrogen contacts between two A molecules (05 H2) and an A and a B molecule (04 H10) that approximate distances of 2.5 Å. Only the first of these involves the kind of methylhydrogen-to-nitro-oxygen interaction that has been implicated in the thermal degradation of TNT. Of greater significance is the complete lack of hydrogen bonding involving the *para*-nitro groups. Such

586

absences are consistent with the *para*-nitro group being unreactive in the thermal or ionization breakdown of TNT [14-16].

Tables 4 and 5 contain the calculated [5, 6] and observed [1, 3] vibration frequencies of TNT and TNT- d_3 . The symmetry designations [1] are based on the nominal $C_{2\nu}$ symmetry of the calculated structure, with the principal axis in the z-direction and the ydirection orthogonal to the plane of the ring. The A_2 and B_1 vibrations are out-of-plane; A_2 vibrations are i.r. inactive if the molecule has genuine $C_{2\nu}$ symmetry, but such is not the case in the present instance and at least some of the A_2 -species vibrations may become observable. In correlating calculated vibrational frequencies to those observed, emphasis was placed on absorption intensities and isotopic substitution [3]. In the majority of cases, calculated absorption frequencies with large transition dipoles were identified with the more intense absorption bands. The major improvement in the calculations occurs in the NO₂ v_{sym} and v_{asym} stretching frequencies, where the error compared to previous calculations [3] decreases from 34% to 22%. The C-H stretching frequencies in the 2800-3100 cm⁻¹ range are reproduced with an average error of 1.3% vs 3.0% using the AM1 method. The remaining frequencies over the range of 400-1800 cm⁻¹ are reproduced with an average error

Table 4. Observed and calculated frequencies (cm⁻¹) for 2,4,6-trinitrotoluene

v K Br	• Vapour	V _{calc}	Symmetry	Assignment+	Transition dipole	_
465		444	B_1, B_2	11, 14	0.10	~
467		480	Β,	12	0.28	
565		533	\boldsymbol{B}_1	12	0.34	
579		558	B ,	15	0.21	
639		634	A_2	5, 12	1.40	
664		669	A_1	17	2.37	
664		673	A_1	17	2.10	
704, 720	705, 721	743	A_2	17	0.78	1
		745	Α,	15	0.06	l l
735	730	749	B ₁	15	0.56	CORN
793	792	769	B ₂	15	1.15	INSPECTED /
		805	B	15	0.13	
-	868	849	A'	12	1.21	\bigcirc
909, 940	907, 938	939	<i>B</i> ,	16	1.32	
		971	A 2	13	0.11	ccesion For
		990	B ₁	6	0.10	iccesion For
		995	A	16	1.00	TIC COLOR
1026	1028, 1067	1032	B ₁	12, 13	0.30	ITIS CRA&I
1086	1075	1049	\vec{B}_2	4,6	0.75 C	TIC TAB
1171		1182	.A ₁	10		Inannounced
1208	1217, 1230	1247	B ₂	10, 12	012	
		1271	A_1^{-2}	12	0.05	ustification
		1334	B ₂	9	0.19	
		1360	A_1	4	0.16	
1356	1352	1581	A ₁	19	5,90	ly
1356	1352	1589	B,	19)istribution /
1356	1352	1604	A ₁	19	0.80	(Stabullar)
1000		1376	B ₂	4	0.09	A
1406	1420	1414	A,	9, 16	0.62	Availability Codes
1437, 1466	1437, 1457	1449	- A ₁	7, 11	0.28	
		1551	B ₂	9		Avail and for
1541	1508	1918		18	3.51	Special Special
1541	1539	1921	B,	18	7.58	
1541	1539	1925	B ,	18	7.26	
1603, 1619	1617	1614	A ₁	8,9	1.12	-120
1005, 1017	1734	1770	B ₂	9	1 19	
		1790	A_1	9	2.67	
2955	2901	2983	B_2	í	0.50	
3015	2971	2991		i	0.42	
3058	2991	3057	A ₁	2	0.42	
3087	3085	3070	B ₂	$\frac{1}{2}$	0.18	
3087 3096			A1	3		
0000	3096	3176	A1	3	0.30	

*Data from Ref. [1, 3].

†Assignments of vibrational frequencies: 1. Ring C-H stretch; 2. Methyl C-H asym. stretch; 3. Methyl C-H sym. stretch; 4. Methyl H-C-H asym. bend; 5. Methyl H-C-H sym. bend; 6. Methyl rock; 7. Methyl umbrella; 8. C-C stretch: methyl to ring; 9 C-C ring stretch: 10 King H-C-C in plane bend; 11. Methyl ring CCC in-plane bend; 12. Ring CCC bend; 13. Out-of-plane bend of ring H; 14. Out-of-plane bend of methyl C; 15. C-N bend; 16. C-N stretch; 17. C-N-O bend; 18. NO₂ asym. stretch; 19. NO₂ sym. stretch.

	obs				Transition	
KBr	Vapour	Vcale	Symmetry	Assignment†	dipole	
457, 471		465	B ₁	12	0.30	
563		527	В,	12	0.29	
573		554	B ₂	15	0.20	
625		623	A,	5, 15	1.28	
677		666	B ₂	15	1.65	
677		672	B_{i}^{-1}	15	2.35	
718	718	694	A_2, B_2	4	0.41	
733	731	730	A2	17	0.68	
_	749	743	A,	17	0.86	
758	758	749	A ₂	15	0.65	
775	778	769	B ₁	15	0.84	
826	826	804	A_1	15	0.84	
826	826	819	B ₂	15	0.79	
872, 909	907	894	B ₁	12	0.19	
938	938	971	$\overline{A_2}$	13	0.13	
938	938	982	B_2	16	2.51	
938	938	989	A_1	16	2.49	
030		1009	B ,	13	0.23	
030		1017	В,	13	0.31	
030		1025	B ₁	4	0.35	
057, 1095	1057, 1067	1070	A,	7	0.09	
179, 1198		1186	A_1	10	0.40	
219, 1264	1254	1245	B ₂	10	0.14	
302		1307	A_1	8	0.17	
		1333	B ₂	9	0.18	
354	1352	1581	.4	19	5.89	
354	1352	1589	B_2	19	5.91	
354	1352	1604	A_1	19	0.85	
410		1418	A_1	9, 16	1.06	
539	1508	1918	A_2	18	3.50	
541	1539	1921	$\vec{B_2}$	18	7.61	
541	1539	1925	в,	18	7.23	
599		1550	B_2	9	1.46	
619		1613	A,	8,9	1.13	
		1768	Β,	9	1.21	
		1790	A_1	19	2.67	
	2275	2258	B		0.38	
	2325	2273	\vec{B}_2	2 2 3	0.22	
	2325	2293	A_1^{-2}	3	0.26	
058	2907	2983	B ₂	ì	0.50	
096, 3087	2971	2991	A ₁	i	0.42	

Table 5. Observed and calculated frequencies (cm⁻¹) for 2,4,6-trinitrotoluene- α -d₃

*Data from Refs [1, 3].

†Assignments of vibrational frequencies. See Table 4.

of less than 21 cm⁻¹, down slightly from the 24 cm⁻¹ with the AM1 method [3].*

Acknowledgements - The authors wish to thank H. HOPE, in whose laboratory the X-ray work was done. W. R. C. wishes to acknowledge the NRC for fellowships and the generous support of the Frank J. Seiler Research Laboratory.

REFERENCES

- [1] J. J. P. STEWART, S. R. BOSCO and W. R. CARPER, Spectrochim. Acta 42A, 13 (1986).
- [2] W. R. CARPER, S. R. BOSCO and J. J. P. STEWART, Spectrochim. Acta 42A, 461 (1986).

*Tables of positional and thermal parameters and structure factors are available from the Directory of the Cambridge Crystallographic Data Centre, University Chemical Laboratory, Lensfield Road, Cambridge CB2 1EW, U.K. The request should be accompanied by the full literature citations for this report.

- [3] W. R. CARPER and J. P. STEWART, Spectrochim. Acta 43A, 1249 (1987).
- [4] W. R. CARPER, L. P. DAVIS and M. W. EXTINE, J. phys. Chem. 86, 459 (1982).
- [5] J. J. P. STEWART, J. comp. Chem. 10, 209 (1989).
- [6] J. J. P. STEWART, J. comp. Chem. 10, 221 (1989).
- [7] R. C. DOREY and W. R. CARPER, J. Chem. Eng. Data 29, 93 (1984).
- [8] J. J. P. STEWART, MOPAC, Quantum Chemistry Prog. Exch. No. 455 (1983).
- [9] M. J. S. DEWAR, E. G. ZOEBISCH, E. F. HEALY and J. J. P. STEWART, J. Am. chem. Soc. 107, 3902 (1985).
- [10] C. G. BROYDEN, J. Inst. Math. App. 6, 222 (1970).
- [11] R. FLETCHER, Comp. J. 13, 317 (1980).
- [12] D. GOLDFARB, Math. Comput. 24, 23 (1970).
- [13] D. F. SHANNO, Math. Comput. 24, 647 (1970).
- [14] T. M. MCKINNEY, L. F. WARREN, I. B. GOLDBERG and
- J. T. SWANSON, J. phys. Chem. 90, 1008 (1986). [15] J. T. SWANSON, L. P. DAVIS, R. C. DOPEY and W. R.
- CARPER, Magn. Reson. Chem. 24, 762 (1986).
- [16] W. R. CARPER, R. C. DOREY, K. B. TOMER and F. W. CROW, Org. Mass Spectrom. 19, 623 (1984).