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STRUCTURE AND VIBRATIONAL ASSIGNMENT OF gauche-1,3-BUTADIENE

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ABSTRACT

The completely optimized structures and harmonic force fields of the s-trans(anti) and gauche-isomers of 1,3-butadiene have been computed at the ab initio Hartree—Fock level using the 6-31G basis set. The gauche dihedral angle was found to be 348° from the planar s-cis(syn)-configuration of the 1,3-butadiene molecule. Seven scale factors for correcting the theoretical force constants of the trans-isomer were calculated from the experimental frequencies of light and heavy trans-1,3-butadienes. The correction of the gaiche-butadiene force field was then carried out using these scale factors, and the vibrational problems were solved for gauche forms of C₄H₆. CD₂CHCHCD₂, C₂D₆. ¹³CH₂CHCH¹³CH₂, and ¹³CH₂CHCHCHC₂. The total assignment of the experimental vibrational frequencies of these isotopomera is given. The conclusion drawn is that the quantum mechanical geometry and the scaled quantum mechanical force field correctly simulate the structure and the experimental frequencies of the butadiene gauche-conformer

INTRODUCTION

During the last few years a series of studies has been completed in which many of the experimental vibrational frequencies of the second rotational isomeric form of 1,3-butadiene have been observed [1–4]. However, the suggested interpretations of the experimental spectrum were different in some cases [1–8]. The analysis of the "hot" band progression of the Raman active torsional overtone $2\nu_{13}$ of 1,3-butadiene allowed the construction of a potential energy curve of internal rotation around the central C–C bond [9]. This curve turned out to be applicable to the series of its isotopomers [10]. According to this experimental curve the second form of the C_4H_6 molecule is rotated about 23° from the planar s-cis-form(syn-form) [9, 10]. The most recent quantum mechanical calculations give an angle of internal rotation of about 38° from the planar s-cis-form [11–14]. The direct electron diffraction measurement does not allow the determination of the exact structure of the second form [15].

Calculations of the vibrational frequencies have been carried out for a planar s-cis-form of 1,3-butadiene [3-8], the force constants being transferred in some cases from the s-trans-form force field [3, 4, 8]. A calculation for the gauche-form (40° from the planar cis-configuration) has also been published; however, the same force field was used as for the s-cis-configuration [4] which is probably the reason why its results practically coincided with those on the cis-configuration calculation [4].

In this connection it is of interest to perform the quantum mechanical computation of the structure and the harmonic force field of the second rotational isomeric form of 1,3-butadiene and to predict the vibrational frequencies of this conformer. This in turn can be used to facilitate the assignment of the observed frequencies for this conformer.

METHOD OF COMPUTATION

The geometrical parameters and force constants of the trans- and gauche-isomers of 1,3-butadiene were computed by the ab initio Hartree—Fock method using the 6-31G basis set and gradient optimization [16] The completely optimized structural parameters of these two forms of 1,3-butadiene are given in Table 1.

At present, the combination of the theoretical and the experimental information was accepted as usual practice in the force field calculations of homologous series of molecules (see, for example, ref. 17). According to this technique the force fields are calculated for a standard (calculated ab initio and then experimentally corrected) geometry. Further the force constants are corrected by empirical scale factors. However, the main purpose of this work is to predict the vibrational frequencies of gauche-C₄H₆ and to assign those that have been observed experimentally. Hence the transition to a

TABLE 1

Completely optimized geometry of trans- and gauche-1,3-butadiene

Parameter	$trans$ - $C_{4}H_{6}$	gauche-C _c H _e
C=Ca	1 3276	1 3266
CC2	1 4645	1 4739
C-H _{trens-terminal} a	1 0726	1 0730
C—H _{cis-terminal} a	1 0748	1.0741
C-Ha	1.0767	1 0769
$\angle C = C - C_p$	124.29	125.76
∠C=C—H _{trans-terminal} o	121.79	121.51
LC=C-H _{CIS} terminal b	121.80	122.04
∠C=C—Hb	119 55	118 89
Dihedral angle ^b	0 0	34 78
Energy ^c	-154 864576	-154.859731

Angstrom u., ts bDegrees Atomic units.

standard geometry was not made and force fields were computed for the theoretical structures of the *trans*- and *gauche*-isomers. The force constants as obtained in cartesian coordinates were transformed into the internal vibrational coordinates in order to correct them using empirical scale factors. As internal coordinates, the local valence coordinates were chosen [18]. The coupling force constants are small in these coordinates, and it is easier to assign the calculated frequencies by their vibrational forms (normal-mode vectors, or eigenvectors). The definition of these coordinates for *trans*-1,3-butadiene is given in ref. 17.

The scaling procedure has been previously described [5, 19]. It is a congruent transformation of the force constant matrix. The values of the scale factors were obtained by minimizing the weighted mean-square deviation between the calculated and the experimental fundamental frequencies for the trans-form of light and heavy 1,3-butadiene using a program written by Dr. G. Pongor The experimental vibrational frequencies and their assignment for the trans-conformers of the C₄H₆ and C₄D₆ molecules are well known since 14 isotopomers of this molecule have been investigated [3, 20, 21]. The values of the scale factors obtained are given in Table 2. As a starting approximation, the scale factors used were those calculated for the trans-form of 1,3-butadiene in ref. 17 (see the last column in Table 2).

Correction of the theoretical force constant matrix of gauche-1,3-butadiene was performed using the seven scale factors calculated for the transconformer. The empirically corrected force constants obtained for the transand gauche-forms of butadiene are given in Tables 3 and 4. The vibrational problems were solved with these force fields for the trans-isomers of C_4H_6 and C_4D_6 (Table 5) and the gauche-isomers of C_4H_6 , $C_4H_2D_4$, C_4D_6 , $1,4^{-13}C_2-C_4H_6$, and $1^{-13}C-C_4H_6$ (Table 6). The assignment of the experimental frequencies of the gauche-conformer was carried out using data on the potential energy distribution taking into account the isotopic shifts

TABLE 2
Optimized scale factors

Scale factor	Basis set		
	6-31G (this work)	4 31G [22]	4-21G [17]
C—C stretch	0.8514	C.887—0.909	0.9215
C=C stretch	0 7573	0 869-0.872	0 8657
C-H stretch	0.8301	C 881-0.890	0.8677
C=C-C, C=C-H in-plane	0.7944	C.793-0.816	0 8025
C—C torsion	3.9979		1.095
C-H, =CH, wag	0.6706	C.718-0.746	0 7217
C=C twist	0 7986	C 793—0 816	0 7837

TABLE 3

Scaled force constant matrix of trans-1,3-butadiene^a

	1	2	3	4	5	6	7	8
1	5 087							
2	0 299	8.521						
3	0 29°	-0 069	8.521					
4	-0 00∮	0 046	0.007	5.209				
5	-0.004	0 007	0.046	0.002	5.209			
6	0.004	0.064	-0 002	0.026	0.001	5.138		
7	0 004	-0.002	0 054	0.001	0.026	C 3	5.138	
S	0.054	0.068	0.019	0.015	-0.001	00	0.011	5.048
9	0 054	0 019	0 068	-0.001	0.015	0.011	0.0	0.0
10	0 211	9 168	0.965	0 047	0.013	-0.059	-0.002	-0 150
11	0 211	0 065	0.168	0 013	0 047	-0 002	-0.059	-0 048
12	0.180	-0 184	-0.011	0.009	-0.003	-0 027	0 011	-0 002
13	0.180	-0.011	-0.184	-0 003	0 009	0 011	-0.027	0.022
14	-0 005	-0.197	-0 001	0.058	0.0	0.056	0.001	-0 007
15	-0 005	-0.001	-0.197	00	0.058	0 001	0.056	~ે 00€
16	0.04%	0.008	0 009	0 097	0.002	-0.105	0.003	-0.035
17	0 041	0 009	0 008	0.002	0 097	0.003	-0.105	C 003
	9	10	11	12	13	14	15	16
	5.048			_				
10	0 048	1.055						
11	-0.150	0.153	1 055					
12	0.022	9.005	-0 04(,	0.525				
13	-0.002	-0 046	0 005	0 037	0.525			
14	-0.006	-0 012	-0.009	0.015	-0.001	0.447		
15	-0 007	-0 009	-0 012	-0 001	0.015	00	0 447	
16	-0 003	0.083	0 011	0 039	0.0	0 002	-0.001	0 527
17	-0.035	ი_ი11	0 083	e o	0 039	- 0 001	0 002	0 003
	17	₹8	19	20	21	22	23	24
17	0 5 2 7							
18		0.023						
19		0 003	0 246					
20		0.003	-0 009	0.246				
21		0.0	0 024	-0 003	0.232			
22		0 0	-0 003	0 024	0 0	0.232		
20		0.005	0.0	-0 013	0 0	-0.004	0 126	
24		0.005	-0 013	0.0	─ 0 004	0.0	-0 004	0 120

²For definition and order of internal coordinates see ref. 17

DISCUSSION

The calculated geometrical parameters of the two molecular forms corroborate the changes which are expected when passing from a trans to a gauche-conformer. An accentuation in the alternation of the carbon—carbon bonds

TABLE 4
Scaled force constant matrix of gauche-1 3-butadiene²

	1	2	3	4	5	6	7	8
1	4.968						~	
2	0.277	8.543						
3	0 277	-0.061	8 543					
4	-0.003	0 048	0.002	5.190				
5	-0.003	-0 002	0.048	0 001	5 190			
6	-0.008	0.061	0.613	0.026	00	5 166		
7	-0.008	0.013	0 061	0.0	0.026	0 009	5.166	
8	0.068	0.077	0.003	0 015	0.005	-0.001	0.0	5.034
9	0.068	0 003	0 077	0.005	0.015	0.0	-0 001	0 015
10	0 266	0.151	-0.022	0.050	-0 008	− 0 077	-0.011	 0.150
11	6ر،0.2	-0 022	0.151	-0 008	0 050	-0 011	0.077	0.041
12	0.186	-0.191	0 024	0 009	0 007	-0.027	0.002	-0 003
13	0 186	0.024	⊸ 0 191	0 007	0.009	0 902	-0 027	00
14	-0 006	-0.199	⊸ 0 007	0.059	0.001	0.058	0.001	-0.037
15	-0 006	-0 007	-0 199	0 001	0.059	-0 001	0 058	- 0.001
16	0.029	0 010	-0 003	0 095	0.002	-0 098	0.003	-0.037
17	0.029	-0 003	0 010	0 002	0 095	0.003	0.098	0 008
18	0 009	0 018	0.018	-0 001	-0 001	0 001	0.001	− 0 012
19	-0 014	0.0	0 004	0 003	0.0	0.001	0 001	—ე.008
20	-0.014	0 004	00	0.0	0 003	0 001	0 001	00
21	-0 013	0 003	0 003	0 002	0.001	-0.007	00	0,0
22	-0 013	0.003	0 003	0 001	0 002	0 0	-0.007	0.0
23	0.028	-0.001	0.003	0.002	0.0	0.004	0.001	-0.001
24	0 028	0.003	-0 001	0 0	0.002	0.001	0.004	- 0.004
	9	10	11	12	13	14	15	16
9	5 034			·				
10	0.041	1.091						
11	 0 150	0.016	1.091					
12	0.0	0.008	0 062	0.531				
13	-0.003	0.062	0.008	-0.009	0.531			
14	-0.001	-0.012	-0 004	0.015	0 003	0 446		
15	-0 007	-0.004	-0.012	-0 003	0 015	0 003	0.446	
16	0.008	0 067	-0.021	0 040	0.009	0.0	0.002	0 529
17	-0 037	− 0 021	0 067	0.009	0.040	0.092	80	0.005
18	-0.012	0 035	0.035	0.001	0.001	-0 004	-C.004	-0.007
19	00	-0.016	0.006	0.004	0.011	0 003	0,0	0 004
20	-0.008	0.006	-0.016	-0 011	0 004	0.0	0 003	0.005
21	0.0	-0.009	0.001	0.001	 0 004	0,001	0.002	0 004
22	00	0 001	-0 009	- ₿ 004	0 001	0 002	C 001	0.002
23	-0.004	0 014	0 039	0,001	-0.008	0.0	-0 003	-0 002
24	-0.001	0.039	0 014	-0.008	0 001	 0 003	0.0	-0.001

TABLE 4 (continued)

	17	18	19	20	21	22	23	24
 17	0 529							
18	0.007	0.017						
19	0 005	− 0 001	0.243					
20	0 004	-0.001	0.008	0.243				
21	0.002	0.0	0 026	C.004	0.235			
22	0.004	0.0	0.004	0.026	0.0	0 235		
23	-0 001	0.003	-0 002	0.008	0.0	0 003	0.128	
24	-0 002	0 003	0.008	-0.002	0.003	0.0	0.004	0.1

^aFor definition and order of internal coordinates see ref. 17.

is observed with decreasing length of the C=C bond and increasing length of the C=C bond (see Table 1). Furthermore, the value of the C=C—C angle increases noticeably, resulting in some decrease of the C=C—H angle. These changes are possibly connected with disturbances of the conjugation in the skew configuration.

Our use of the optimized geometry, rather than a standard or experimental geometry, gave a set of force constant scaling factors somewhat different from those of other authors (Table 2) for the trans-form. This is most noticeable for the scale factors of the stretching force constants, since they depend strongly on the adopted geometry. This effect was greatest in the case of the scale factor for stretching the C=C bond since the deviation of the calculated value of this bond length from the standard geometry was the largest. Deviation from the scale factors given in ref. 17 is caused by the use of a different basis set in the present work; in addition a reversal of the assignment for the ν_{10} and ν_{22} frequencies of trans-C₄D₆ was accepted (see below). The procedure of scaling was also slightly different from that used in Ref. 22. Nevertheless, the set of scale factors obtained as a whole is quite close to that obtained in ref. 17, and thus the structures of the force constant matrices for the transform are similar in the present work and in ref. 17 (Table 3).

Comparing the force fields of the trans- and gauche-conformers, it is interesting to note the appearance of large non-diagonal elements in the force constant matrix of the gauche-form which are situated at the "intersection" of in-plane and out-of-plane coordinates. This clearly demonstrates that using the force constants of the trans-form to calculate the vibrational frequencies of the gauche-form is not in general justified.

Frequencies of the trans-form, calculated with the empirically corrected force field, are in good agreement with the experimental results (Table 5). It should be noted that there is some disagreement as to the assignment of the ν_{10} and ν_{23} bands of trans- C_4D_6 . In refs. 3, 4, and 17 it was suggested that the assignment of bands at 768.8 cm⁻¹ and 735.6 cm⁻¹ be interchanged in comparison with the earlier suggested assignment [20]. However, experimental study of the IR spectra of the trans- C_4D_6 molecule in the crystalline phase

TABLE 5

Experimental and calculated fundamental vibrational frequencies of trans forms of 1,3 butadiene and d.-1,3-butadiene

Assignment	trans	CH ₃ = (JH-CH=C	11,					trans-(בית:	vans-CD, =CD—CD=CD,	D,				
	Sym,	a	Exp [20]	Calc	Sym.	-	Exp. [20]	Calc,	Sym,	a	Exp. [8, 20]	Calc.	Sym,	<u> </u>	Exp [3, 20]	Calc.
ν(CH ₂)str.	<u>ש</u>	1	3101	3110	ρ'n	17	3102	3110	a _n	1	2341	2316	$b_{\mathbf{u}}$	17	2336	2916
$\nu(C-II)$ str.	•	~	3014	3035	•	18	3020	3036	:	~	2262	2251	!	18	2266	2251
v(CH,)atr		c.	3014	3021		19	3010^{a}	3027		~	2205	2205		18	2215	2206
"(C *C)str.		4	1643	1668		20	1599	1591		7	1583	1604		20	1523	1497
δ(CH ₂)sc.		ಶ	1442	1466		21	1385	1388		⋍	1048	1050		21	1042	1053
p(C −II)r.		9	1291"	1290		22	12962	1304		9	919	926		22	1009	1005
$\nu(C-C)str$		-	1205	1214		l	l	I			1136	1178		١	İ	i
$\rho(\mathrm{CH}_{\mathbf{i}})_{\mathbf{i}}$		8	890	878		53	9066	900 1		20	739	730		23	7688	749
$\delta(C=C-C)b$.		0	613	203		24	301	202		6	440	439		73 4	253 6	246
$\lambda(C-H)w$	a _n	10	1013.2	1021	$b_{\mathbf{k}}$	14	190	976	ď	10	7356	768	$\rho_{\mathbf{g}}$	14	796	196
$x(CH_2)w$		11	9078	915	ì	15	110	911	i	Π	7.18,5	716	ı.	16	702	687
7(CH ₂)tw.		12	524 6	512		16	753	739		12	381	374		16	603	598
r(C—C)tor.		13	163	163		l	ļ	ł		13	140	141		ı	i	i

⁴Corrected on Penni resonance Values expressed in cm⁻¹

TABLE 6

Experimental and enleulated fundamental vibrational frequencies of gauche-forms of 1,3-butadiene, 1,1,4,4, d,·1,3 butadiene, d,·1,3 butadiene, and 1.¹3C-1,3 butadiene, and 1.¹3C-1,3 butadiene, and 1.¹3C-1,3 butadiene, and 1.¹3C-1,3 butadiene, and 1.³C-1,3 butadiene, and

	JE A	сп,снспсн,	ICH,	CD,CHCHCD,	HCD,	co,cococo,	ocp,	13,HDE1	*CH,CHCH''CII,	чононочо.	сисн,
	}	Exp. [3, 4]	Calc,	Exp.	Calc,	Exp [3, 4]	Cale,	Exp.	Cale	Екр. [3]	Cale,
1 v(CH.)str. a		3103	3112	2337	2317	2335	2317	3068	3099	3100	3110
2 v(C-H)str		3030	3041	2013	3036	2256	2253	1	3038	ł	3040
3 v(CH,)str		3014	3026	2228	2219	2222	2207	3008	3027	3010	3030
4 v(C=C)str		1633	1635	1592	1590	1552	1582	1612	1614	1626	1632
6 8(CH ₁)3c.		1426	1448	. 384	1090	1	1009	1424	144.3	1424	1447
6 n(C-H)r		1	1318	1305	1311	I	838	1	1312	1	1315
7 ρ(CH ₂)r.		1	1057	I	723	708	723	1	1053	l	1055
8 x(C-II)w		983	987	927	921	J	776	ļ	986	982	986
9 x(C)1,)w,		916^{6}	918	726	744	712	703	907b	600	913	918
10 v(C-C)str.		ı	865	944	696	1166	1159	I	867	I	961
11, (C-C)tw.		727	729	694	592	587	588	731	728	731	729
12 5 (C≖C—C)b.		ļ	271	ł	234	J	232	1	265	j	268
13 r(C-C)tor.		ı	155	1	136	1	122	i	154	1	155
14 "(CII,) str b		3103	3109	2337	2314	2335	2316	I	3096	1	3098
15 "(C -II)atr		3039	3033	3025	3019	2266	2246	I	3023	i	3024
16 v(CH,)str		3011	3016	2228	2210	2222	2203	١	3016	i	3016
17 v(C=C)atr		1612	1629	1	1577	1528	1531	l	1608	i	1611
18 5 (CH,)5c		1403	1493	103T	1040	1056	1.068	1401	1422	1401	1422
19 p(C-H)r.		1	1286	1286	1292	877	875	ļ	1278	I	1281
$20 \ \rho(\mathrm{CH_2})_\mathrm{r}$		108/	1094	I	946	1	1009	1077	1064	1033	1089
$21 \ (C-II)w$		966	1002	014	913	763	761	906	1002	966	1002
$22 \chi(\mathrm{CH_3}) \mathrm{w}$		914	918	727	731	715	709	200	600	906	606
23 6 (C=C—C)b.		596	604	488	505	492	497	596	603	697	604
24 r(C-C)tw		468	461	373	358	360	346	467	450	467	451

Values expressed in cm⁻¹, ^aAll the vibrations belong to the σ symmetry type ^bValues as estimated from $\nu_{\rm p}$ i $\nu_{\rm p_2}$ (1829 cm⁻¹ and 1814 cm⁻¹, respectively) [3]

The first the state of the stat

allows, by analogy with C_4H_6 , assignment of the band at 735.6 cm⁻¹ to the out-of-plane vibration ν_{10} of the C-D bond. Provided that the structure of the C_4H_6 and the C_4D_6 crystals are the same, this assignment is corroborated by the value and the direction of shift, the value of splitting, and the polarization of components of the split band in the IR spectrum when passing from the gas phase to the crystal for the frequency pairs of 907.8—718.5 cm⁻¹ and 1013 2—735.6 cm⁻¹ (Table 5). The spectral region under consideration and a detailed discussion are given in ref. 23.

The calculation of the vibrational frequencies of trans- C_4H_6 showed that the ν_{19} frequency should be reassigned. Indeed, in the IR spectrum of C_4H_6 in condensed phase there are five bands instead of three, i.e. 2972, 2998, 3023, 3044, and 3084 cm⁻¹ [24]. Those at 2998 cm⁻¹ and 3023 cm⁻¹ are weak and are possibly combination bands. On going from the vapour to the solid state, shifts of 10–20 cm⁻¹ to the low frequency are observed. Such a low frequency of the strong band at 2972 cm⁻¹ (2985 cm⁻¹ in vapour) could be explained by a shift due to a Fermi resonance with a combination band (1643 + 1385 = 3028, $a_g \times b_u$) and by a phase shift Obviously, the value of the vibrational frequency ν_{19} in the condensed phase should be equal to (2972 + 3023)/2 = 2997 cm⁻¹. Then in the gas phase it should be situated approximately at 3010 cm⁻¹ (Table 5) and indeed, this vibrational frequency in the molecules CH₂CHCDCD₂ and trans-CHDCHCHCH₂ is observed at 2997 cm⁻¹ and 3003 cm⁻¹, respectively [20]. The vibrational frequency ν_6 in the C_4H_6 molecule is also corrected by Fermi resonance [20] (Table 5).

The assignment of the vibrational frequencies of gauche- C_4H_6 (Table 6) coincides in general with the results given in Refs. 3 and 4 and differs in the interpretation of frequencies from Ref. 8 for vibrations ν_{12} , ν_{23} , and ν_{24} . In Ref. 8 the experimental bands at 472 cm^{-1} and 475 cm^{-1} [2] are assigned to the ν_{12} and ν_{23} vibrations. Later, only one band ascribed to ν_{24} (Table 6) was observed in this region of the C_4H_6 spectrum [3, 4]. The 596 cm⁻¹ band interpreted as ν_{24} in Ref. 8 was reassigned to ν_{23} [3, 4].

Table 6 also contains the experimental and the calculated vibrational frequencies for the gauche-forms of the $C_4H_2D_4$ and the C_4D_5 molecules. In the case of the $C_4H_2D_4$ molecule the present calculation allows the reassignment of the vibrational frequencies ν_5 and ν_{10} in companion with the assignment in Ref. 4 The frequency of the =CD₂ group scissoring is very characteristic and should not depend on the rotation angle around the C—C bond. For the trans-conformers of the $C_4H_2D_4$ and the C_4D_6 molecules this frequency is about 1050 cm⁻¹ [20] which corroborates the assignment of the 1084 cm⁻¹ band to ν_5 . The experimental band at 927 cm⁻¹ corresponds very well to the 60 cm⁻¹ shift of the calculated vibrational frequency ν_8 when passing from C_4H_6 to $C_4H_2D_4$.

For the C_4D_6 molecule the present calculations confirmed the assignment of the experimental band at 877 cm⁻¹ to ν_{19} [3] in contradiction to its interpretation as ν_6 [4].

The most interesting fact in the assignment of the vibrational frequencies of the gauche-conformers of $C_4H_2D_4$ and C_4D_6 is the inverse isotopic shift [25] of the C—C bond stretching frequency ν_{10} (Table 6). A similar inverse isotopic shift was observed for the stretching frequencies ν_{10} and ν_{33} of the C—C bond in the trans-cis-trans-1,3,5-hexatriene molecule on going to its 2,3,4,5-tetradeuteroanalogues [26]. It should be noticed that in the transform substitution of the H atoms by D atoms shifted the stretching frequency of the C—C bond very little (ν_7 in Table 5), while in the gauche-form it resulted in an increase of this frequency to 300 cm⁻¹ (ν_{10} in Table 6).

The calculated values of the vibrational forms (normal-riode vectors or eigenvectors), distribution of potential energy [27, 25] and the isotopic shifts of the experimental vibrational frequencies confirmed the adopted assignments for all molecules studied

Table 6 gives the results of the vibrational frequency calculation of the gauche-forms of $1^{-13}C_1$,3-butadiene and $1,4^{-13}C_2$ -1,3-butadiene. The isotopic shifts for these molecules are small and do not give any essential information in addition to the data already existing for C_4D_6 and $C_4H_2D_4$. However, the spectra of these molecules give further confirmation for the experimental detection of the bands belonging to the second rotational isomeric form of 1,3-butadiene.

CONCLUSIONS

The calculation of the vibrational frequencies of the second rotational isomeric form of 1,3-butadiene showed that all its observed frequencies may be assigned to vibrations of the gauche-form with a rotation angle of about 34.8° from the planar s-cis-configuration. An additional experimental argument for the realization of the gauche-form is the manifestation in the IR spectrum of the bands of the ν_8 , ν_9 , and ν_{11} vibrations. In the case of an s-cisconfiguration these must belong to the a_2 symmetry type and be forbidden in the IR spectrum

Good agreement between the calculated and experimental vibrational frequencies of the gauche-form and the difference of its force field from that of the trans-form demonstrates the possibility of using the same set of scale factors for several structural isomers and shows in general the inaccuracy of transferring force constants from one rotational isomer to another.

The calculation performed is an additional example of the solution of the structural problem by means of the gradient technique the application of which was discussed in detail in Ref. 29.

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