

# NOVEL MODIFICATIONS OF ELEMENTAL NITROGEN AND THEIR MOLECULAR STRUCTURES – A QUANTUM-CHEMICAL CALCULATION

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Using the quantum-chemical calculation methods QCISD and G3, the possibility of the existence of nitrogen molecules with the composition  $N_4$ ,  $N_6$ ,  $N_8$  and  $N_{10}$  has been discussed. On the basis of the data obtained, the conclusion about possibility of existence of three novel polymorphic modifications of elemental nitrogen with an even number of atoms in molecules, namely  $N_4$  with rectangular and regular tetrahedron shapes, and  $N_6$  in a form remotely resembling an "open book", has been made. The values of bond lengths, valence and torsion angles, and oscillations frequencies in each of the above-mentioned forms of elemental nitrogen have been presented.

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atoms.  $^{10\text{-}12}$  In this connection, in the this article the possibility of the existence of polyatomic molecules of nitrogen with an even number of atoms, namely nitrogen molecules of  $N_4$ ,  $N_6$ ,  $N_8$  and  $N_{10}$  compositions will be discussed.

### INTRODUCTION

Molecule of elemental nitrogen is known to consist of two atoms with a triple bond between them.<sup>1</sup> The length of this bond is only 109.5 pm, and, owing to this, the N<sub>2</sub> molecule (dinitrogen) is characterized by a very high dissociation energy (941.64 kJ mol<sup>-1</sup>) and by very less chemical activity.

Three crystalline modifications of  $N_2$  are known. In the temperature range 36.61-63.29 K,  $\beta$ - $N_2$  phase having a hexagonal dense packing, a space  $P6_3$  mmc<sup>-2</sup> group, lattice parameters a=3.93 Å and c=6.50 Å, exists. At a temperature lower than 36.61 K, a stable  $\alpha$ - $N_2$  phase having cubic lattice, a space  $P2_13$  group and a period of a=5.660 Å, occurs. At a pressure of more than 3500 atmospheres and a temperature below 83.0 K, a hexagonal phase of dinitrogen, namely  $\gamma$ - $N_2$ , is formed.

There are some theoretical indications that other nitrogen oligomers and polymers may be possible. These nitrogen modifications may have potential applications as materials with a very high energy density and as powerful propellants or explosives.1 For most neutral polynitrogens are not expected to have a large barrier towards decomposition, and that the few exceptions would be even more challenging to synthesize than the tetranitrogen N<sub>4</sub> which is an analogue of tetrahedrane C<sub>4</sub>, and has potential as a high-performance energetic material.<sup>2</sup> Nevertheless, cationic and anionic polynitrogens, namely cations of triazenium  $(N_3^+)$ , tetrazenium  $(N_4^+)$ , pentazenium  $(N_5^+)$ , and azide-anione (N<sub>3</sub><sup>-</sup>), pentazolide-anione (cyclic aromatic N<sub>5</sub><sup>-</sup>) have been characterized.<sup>1,3-9</sup> However, all these nitrogen compounds have a positive or negative charge. Up to now, however, there is no information in the literature on the existence of other neutral simple substances consisting solely of nitrogen

### CALCULATION METHOD

The calculation of molecular structures of polyatomic nitrogen molecules of  $N_4$ ,  $N_6$ ,  $N_8$  and  $N_{10}$  compositions was carried out using the QCISD(T)/TZVP method, as described in detail earlier,  $^{13}$  in combination with the Gaussian09 software package.  $^{14}$  The initial structures of the  $N_4$ ,  $N_6$ ,  $N_8$  and  $N_{10}$  molecules for carrying out quantum-chemical calculations are shown in figure 1.

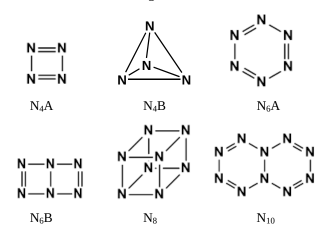


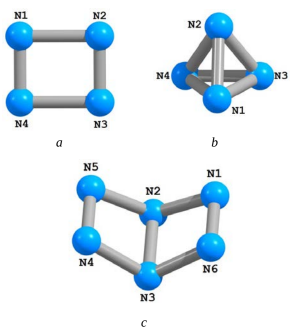
Figure 1. The assumed initial structures of nitrogen molecules.

The choice of these initial structures was determined by two factors. First, the valence possibilities of the nitrogen atom (which capable to bind with one, two or three neighboring atoms by means of three chemical bonds according to the exchange mechanism) and secondly, with the greatest typicality of these structures compared with other structures with a corresponding number of atoms. In this connection, the regular octahedron in the case of  $N_6$ , the

hexagonal bipyramid and the dodecahedron in the case of N<sub>8</sub>, the two-capped cube or the two-capped dodecahedron in the case of N<sub>10</sub> were not included in the number of initial structures. The correspondence of the found stationary points to energy minima was proved in all cases by the calculation of the second derivatives of energy with respect to the atom coordinates All equilibrium structures corresponding to the minima on the potential energy surfaces has only positive frequencies. The values of the standard thermodynamic characteristics of the nitrogencontaining compounds under examination were calculated using the G3 method described in earlier in detail.<sup>15</sup> All quantum-chemical calculations were carried out in the Joint Supercomputer Center, Kazan Branch of RAS - Branch of Federal Scientific Center "Research Institute for System Studies of the Russian Academy (http://www.jscc.ru).

## RESULTS AND DISCUSSION

According to the results of our calculations, only three of the six structures of polyatomic nitrogen molecules mentioned above are stable, namely the  $N_4A$  structure in the form of a rectangle,  $N_4B$  in the form of a regular tetrahedron, and the  $N_6B$  structure in the form of an "open book". All these modifications have been shown in figure 2.



**Figure 2.** Molecular structures of three theoretically possible polymorphic modifications of elemental nitrogen.

The geometric parameters of these above-mentioned stable structures (bond lengths, valence and torsion angles) have been presented in the Table 1. As can be seen from these data, in the polyatomic nitrogen molecules under examination, the nitrogen–nitrogen bond lengths are much larger than those in the  $N_2$  molecule; it is quite natural, because the theoretically expected multiplicity of bonds in them should be lesser than one in a dinitrogen molecule. It should be noted in this connection that for the  $N_2$  molecule itself, according to the QCISD(T)/TZVP method, the bond length (N1N2) is 110.3 pm that is very close to the

experimental value of (109.5-110.0) pm.\(^1\) It is interesting that, according to our calculations, for a "planar" version of the tetranitrogen molecule  $N_4A$ , not a square or orthorhombic structure is realized, as might be expected, but a rectangular structure, with rather considerably different "longitudinal" and "transverse" bond lengths (154.6 and 127.1 pm, respectively) (Figure 2). At the same time, for the "tetrahedral" variant  $N_4B$ , the structure of the regular tetrahedron, where all the lengths of nitrogennitrogen bonds are exactly the same and have an intermediate values between the nitrogen- nitrogen bond lengths in the structure of  $N_4A$ .

**Table 1.** Parameters of the molecular structure of four- and six-atom nitrogen molecules.

uom muogen me		la and a NI A				
Molecule N <sub>4</sub> A						
N-N bond leng	ths, pm	Valence angles, d	leg			
(N1N2)	154.6	(N1N2N3)	90.0			
(N2N3)	127.1	(N2N3N4)	90.0			
(N3N4)	154.6	(N3N4N1)	90.0			
(N4N1)	127.1	(N4N1N2)	90.0			
Torsion (dihedral) angles, deg						
(N1N2N3N4)	0.0	(N2N3N4N1)	0.0			
(N3N4N1N2)	0.0	(N4N1N2N3)	0.0			
Molecule N <sub>4</sub> (B)						
N-N bond leng	ths, pm	Valence angles, deg				
(N1N2)	146.7	(N1N2N3)	60.0			
(N1N3)	146.7	(N1N2N4)	60.0			
(N1N4)	146.7	(N1N3N2)	60.0			
(N2N3)	146.7	(N1N3N4)	60.0			
(N2N4)	146.7	(N1N4N2)	60.0			
(N3N4)	146.7	(N1N4N3)	60.0			
Torsion (dihedral) angles, deg						
(N1N2N3N4)	70.6	(N2N3N4N1)	-70.5			
(N3N4N1N2)	-70.6	(N4N1N2N3)	-70.5			
Molecule N <sub>6</sub> (B)						
N-N bond lengths, pm		Valence angles, deg				
(N1N2)	147.8	(N1N2N5)	109.1			
(N2N3)	153.0	(N2N5N4)	95.3			
(N3N6)	147.7	(N5N4N3)	95.3			
(N6N1)	125.8	(N4N3N6)	109.1			
(N2N5)	147.7	(N3N6N1)	95.3			
(N5N4)	125.8	(N6N1N2)	95.3			
(N4N3)	147.7	(N2N3N4)	84.7			
		(N2N3N6)	84.7			
		(N3N2N1)	84.7			
		(N3N2N5)	84.7			
Torsion (dihedral) angles, deg						
(N1N2N5N4)	-82.5	(N6N3N4N5)	82.5			
(N1N6N3N4)	-82.6	(N6N1N2N5)	82.5			
(N1N2N3N6)	0.0	(N5N2N3N4)	0.0			
(N2N3N6N1)	0.0	(N2N3N4N5)	0.0			
(N3N6N1N2)	0.0	(N3N4N5N2)	0.0			
(N6N1N2N3)	0.0	(N4N5N2N3)	0.0			
(N5N2N3N6)	-109.8	(N1N2N3N4)	109.8			

The structure of  $N_6B$ , as it should be expected, is non-coplanar. There are three different kinds of nitrogen-nitrogen bonds having different lengths in it. Two of them

are relatively short (125.8 pm) and correspond to the double N=N bond; the other five in own length correspond to a single N-N bond, but one of them forming sui generis "binding" of this "open book" itself, is noticeably longer than the other four, 153.0 and 147.7 pm, respectively.

**Table 2.** The oscillation frequencies in the N<sub>4</sub>A, N<sub>4</sub>B and N<sub>6</sub>B.

Oscillation Assignment of oscillation frequency				
frequency, cm <sup>-1</sup>	Assignment of oscination frequency			
	Molecule N <sub>4</sub> (A)			
402	Wagging			
493	Stretching (asym.) with participation of			
	atoms (N1N2) and (N3N4)			
904	Stretching (sym.) with participation of atoms			
	(N1N2) and (N3N4)			
998	Scissoring			
1296	Stretching (asym.) with participation of			
1504	atoms (N1N4) and (N2N3)			
1526	Stretching ( <i>sym.</i> ) with participation of atoms			
	(N1N4) and (N2N3)  Molecule N <sub>4</sub> (B)			
* *				
695 695	Scissoring with change of all valence angles Scissoring with participation of atoms			
073	(N1N4) and (N2N3), concerning			
	(N2N4) bond			
915	Stretching (asym.) with change of (N1N4),			
	(N1N3) and (N2N3) bond lengths			
915	Stretching with change of (N1N3), (N1N4),			
	(N2N3) and (N2N4) bond lengths			
917	Scissoring with change of angles (N3N1N4)			
	and (N3N2N4)			
1288	Stretching (sym.) with participation of all			
	participation of all N atoms			
	Molecule N <sub>6</sub> (B)			
383	Conjugation of two rocking oscillations of			
4.45	(N4N3N6) and (N1N2N5)			
447	Scissoring with participation of atom			
468	groupings (N3N4N6) and (N1N2N5) Scissoring with participation of atom			
400	grouping (N3N4N6)			
578	Stretching (sym.) with change of bond			
	lengths in pairs (N2N5), (N1N2) and			
	(N4N3), (N3N6)			
716	Stretching (asym.) with change of bond			
	lengths in pairs (N2N5), (N1N2) and			
	(N4N3), (N3N6)			
749	Stretching (sym.) with change of bond			
	lengths in pairs(N1N2), (N2N5), and			
0.41	(N4N3), (N3N6), and (N2N3), too			
841	Scissoring with participation of atom			
917	groupings (N1N2N5) and (N3N4N6)			
917	Rocking with participation of atom groupings (N1N2N5) and (N3N4N6)			
946	Scissoring with change of bond length			
710	(N2N3)			
1030	Conjugation of several scissoring			
	oscillations			
1447	Stretching (asym.) with change of bond			
	lengths (N1N6) and (N4N5)			
1510	Stretching (sym.) with change of bond			
	lengths (N1N6) and (N4N5)			

As a result of the above differences, both sides of the "cover" of this "open book" are in fact not rectangles, as might be expected at first glance, but are identical isosceles trapezoids. The sum of the valence angles (N3N6N1) + (N6N1N2) + (N1N2N3) + (N2N3N6) and (N4N3N2) + (N3N2N5) + (N2N5N4) + (N5N4N3) are the same and are equal to exactly 360.0°. What is remarkable, the values of the valence angles (N1N2N5) and (N4N3N6) are much closer to  $90^\circ$  than to  $180^\circ$ , so the degree of non-coplanarity of the molecular structure of  $N_6B$  should be considered as very significant.

Oscillation frequencies of polynitrogen molecules  $N_4A$ ,  $N_4B$  and  $N_6B$  are depicted in Table 2. According to theoretical expectations, 6 oscillations must be active in the IR spectrum in the case of  $N_4A$  and  $N_4B$ , and 12, in the case of  $N_6B$ . Our calculated values match the theoretical expectations.(Table 2). As may be seen from these data, the sets of vibrational frequencies in polynitrogen molecules are quite different among themselves. At the same time, what is interesting, in  $N_4B$ , there are two pairs of oscillations of different nature with the almost identical in frequency, 695 and 915 cm<sup>-1</sup> whereas in  $N_4B$  and  $N_6B$ , nothing like this occurs.

**Table 3.** Calculated standard thermodynamic parameters of formation of  $N_4A$ ,  $N_4B$  and  $N_6B$  molecules from dinitrogen.

Compound	$\Delta_{\mathrm{f}}H^{0},$	$S^0$ ,	$\Delta$ f $G^0$ ,
	kJ mol <sup>-1</sup>	J mol <sup>-1</sup> K <sup>-1</sup>	kJ mol <sup>-1</sup>
N <sub>4</sub> A	771.6	248.6	810.8
N <sub>4</sub> B	771.0	230.9	815.6
N <sub>6</sub> B	1009.5	278.2	1096.7

The values of standard thermodynamic characteristics, namely  $\Delta_t H^0$  (kJ mol<sup>-1</sup>),  $S^0$  (J mol<sup>-1</sup> K<sup>-1</sup>) and  $\Delta_t G^0$  (kJ mol<sup>-1</sup>), of the formation of N<sub>4</sub>A, N<sub>4</sub>B and N<sub>6</sub>B were calculated with using of G3 method and are presented in the Table 3. As follows from the data given in it, they all have positive values. In this connection, this fact attracts its attention that the values of these standard parameters for compounds N<sub>4</sub>A and N<sub>4</sub>B despite the very significant difference in their molecular structures, are nevertheless extremely close to each other. It is easy may be shown with using of these data and well-known values of  $\Delta_t H^0$  and  $S^0$  for dinitrogen (0 kJ mol<sup>-1</sup> and 191.6 J mol<sup>-1</sup> K<sup>-1</sup>, respectively, <sup>16</sup> they cannot be formed directly from N<sub>2</sub> within the framework of the isobar process by general eqns. (1-3)

$$2N_2(gas) \to N_4A (gas) \tag{1}$$

$$2N_2(gas) \to N_4B (gas) \tag{2}$$

$$3N_2(gas) \rightarrow N_6(gas)$$
 (3)

**Table 5.** Calculated standard thermodynamic parameters of formation of N<sub>4</sub>A, N<sub>4</sub>B and N<sub>6</sub>B molecules from atomic nitrogen.

Reaction	$\Delta H^0$ , kJ	S <sup>0</sup> , J K <sup>-1</sup>
4N (gas) → N <sub>4</sub> A (gas)	-1119.6	-364.6
$4N (gas) \rightarrow N_4B (gas)$	-1120.2	-382.3
$6N (gas) \rightarrow N_6 (gas)$	-1827.3	-641.6

The point is that for each of these processes, the value of the standard enthalpy of the reaction  $(\Delta H^0_{298})$  is positive, and, on the contrary, the standard entropy ( $S^{0}_{298}$ ) is negative (Table 4), and in accordance with the classical Gibbs-Helmholtz equation  $\Delta G^0 = \Delta H^0 - T \Delta S^0$ , their synthesis directly from the "ordinary" dinitrogen N2 is thermodynamically forbidden process. However, from the thermodynamic point of view, the formation of N<sub>4</sub>A, N<sub>4</sub>B and N<sub>6</sub>B from atomic nitrogen according to schemes (4-6) is perfectly permissible and, nevertheless, as can be shown using the values of  $\Delta_f H^0$ and S<sup>0</sup> for atomic nitrogen (472.8 kJ mol<sup>-1</sup> and 153.3 J mol<sup>-1</sup>  $K^{-1}$ , respectively), <sup>16</sup> for each of reactions (4-6)  $\Delta H^0$  as well as  $\Delta S^0$  are negative (Table 5) so that in a certain temperature range each of them are thermodynamically resolved. It should be noted in this connection that according to our calculation using the G3 method, the standard enthalpy of the  $N_2(gas) \rightarrow 2N$  (gas) process, in fact, the energy of N=N bond in the dinitrogen molecule is 936.60 kJ mol<sup>-1</sup>, which is in very good agreement with the experimental value of 941.64 kJ mol<sup>-1</sup>. How to synthesize these novel polymorphic modifications of nitrogen is another problem, the discussion of which is beyond the scope of this paper, and further researches are needed for it.

$$4N (gas) \rightarrow N_4(A) (gas) \tag{4}$$

$$4N (gas) \rightarrow N_4(B) (gas) \tag{5}$$

$$6N (gas) \rightarrow N_6 (gas) \tag{6}$$

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# **CONFLICT OF INTEREST**

The authors declare that they have no conflict of interest, financial or otherwise.

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