


 Cite this: *RSC Adv.*, 2023, **13**, 33786

Delocalization quantitatively mapped for prototypic organic nitroanions as well as azidoform anions†

 Anatoly M. Belostotskii 

Delocalization of occupied orbitals impacts the chemical bonding in the simplest known pernitroanions $[(NO_2)_3C]^-$ (1) and $[(NO_2)_2N]^-$ (2) as well as other functionalized organic anions. By quantitatively mapping it onto molecular backbones of 1, 2, $[CH_2NO_2]^-$ (3), $[CH_3NNO_2]^-$ (4) and $[C(N_3)]^-$ (6) anions (all modeled by QM calculations), the Weinhold's NBO analysis refines their chemical structure, enabling to explain and even predict their essential chemical behaviour. In detail, the HOMO of 1 and 2 is associated with the central atom to the degree of 70.7% and 80.4%, respectively, while the HOMO localization on O atoms for 3 and 4 is 85.3% and 81.1%, respectively. Predomination of C-alkylation for 1 and that of O-alkylation for 3 in non-coordinating solvents thus becomes clear. The important news is that the easiness of homolytically disrupting the N–N bond in 2, a constituent of inexpensive powerful explosives, is because of the occupancy of the related σ^* -orbital increases with stretching this bond. The same is true for electrocyclic extrusion of NO_3^- from this molecule. This antibonding effect may be assumed to be the common cause of the proneness of aliphatic nitro compounds to decompose. Pyramidal anion 6 is a highly localized carbanion. Its isomer of molecular symmetry C_s has a unique chemical structure of its azido substituents: each of them is represented by one high-weight resonance structure, e.g., $N-N\equiv N$. The prediction is that the dinitrogen-eliminating decomposition of this isomer is more facile than of the isomer of C_3 symmetry. In summary, this study affords three novel particular insights into the chemical structure and reactivity of these anions: chemically telling delocalization-augmented molecular structures, a reasonable hypothesis of the common cause of thermally triggered instability of aliphatic nitro compounds, and discovered one-resonance structure azido groups.

 Received 2nd October 2023
 Accepted 9th November 2023

DOI: 10.1039/d3ra06708d

rsc.li/rsc-advances

Introduction

Nitroformate $[(NO_2)_3C]^-$ (1) and dinitroamide $[(NO_2)_2N]^-$ (2) are the only known anions whose molecular structure is one atom bearing nitro groups. Their salts are easily available high-energy compounds and the thoroughly selected ones (e.g., hydrazinium nitroformate, ammonium dinitroamide, and hydrazinium dinitroamide) are an inexpensive chief component of excellent explosives and propellants. In view of this important practical circumstance as well as the general interest in the structure of prototypic molecules, anions 1 and 2 have been the focus of many structural studies (mainly by XRD^{1,2} and quantum mechanics (QM) molecular modeling^{1a,2b,c,3a,4-7} as well as episodically by ¹³C, ¹⁴N, ¹⁵N NMR, ^{1c,3a} IR^{3a,4} and Raman spectroscopies^{3a,4}). Astonishingly, chemical bonding in these homoleptic anions has remained inadequately understood.

Department of Chemistry, Bar-Ilan University, Ramat-Gan, 5290002, Israel. E-mail: belost@biu.ac.il

† Electronic supplementary information (ESI) available: Cartesian coordinates, energies, frequencies for transition states, Wiberg indexes, orbital occupancies, some reaction energy profiles. See DOI: <https://doi.org/10.1039/d3ra06708d>

Their chemical structure is mainly considered as a significantly delocalized anion,^{2b,c,3,5,6a} a near-ideal planarity of the central elementary molecular fragment (C)NNN in 1 urges to accept this view. Among them, QM calculations undertaken for anion 2 using Bader's atoms-in-molecules approach are notable: they show that the π bonding is dissipated along the whole molecular backbone.^{2c,5} An extreme viewpoint is that anion 1 is a Y-aromatic system;^{6a} the number n of the Hückel's aromaticity criterion ($4n + 2$) is 0 in this case as it is for the archetypal Y-aromatic molecule of trimethylenemethane dication.^{6b} Unfortunately, the convenient magnetic shielding-based criterion of aromaticity (abbreviated to TSNMRS) is unable to prove or disprove this.^{6c} Contradicting the aromaticity hypothesis, early DFT calculations characterize anion 1 to be not sizably delocalized.⁷ Thus, while establishing the fact of delocalization of anions 1 and 2, the undertaken molecular modeling^{2b,c,3a,b,5-7} does not unambiguously conclude what the degree of this delocalization is.

Obviously, conclusions regarding the chemical bonding in anions 1 and 2, which were drawn from age-old QM calculations, require verification because these studies have used either insufficiently accurate calculation methods, or unbalanced basis



sets, or an unsuccessful combination of them. For instance, in modeling anion **2**, Pople's basis sets together with various post-HF calculation methods were sometimes used or tried.^{3a,4} This combination rather delivers an essentially corrupted molecular geometry if applied to conjugated systems.⁸

The description of anions **1** and **2** as a σ -bonded molecular backbone enforced by a framework of substantially delocalized π -bonding is not in line with their considerable chemical instability: as a rule, their salts are explosive substances sensitive to storing and handling conditions (except the cases where crystal packing stabilizes the salt^{16,3c}). Also, the central atom in anions **1** and **2**, though carrying substituents both of a potent $-I$ and $-M$ effect, shows an appreciable nucleophilicity. Tetrafluoroammonium salt NF_4SbF_6 oxidizes anion **2** in SO_2 at -64°C with forming fluorodinitroamine $\text{FN}(\text{NO}_2)_2$.⁹ This low-temperature fluorination does not surprise only if assuming a poor delocalization of the N-anion. On the other hand, an extremely high acidity of both parent C–H and N–H acid-trinitromethane ($\text{p}K_a = 0.17$) and dinitraminic acid ($\text{p}K_a = -5.0$) – is a rationale in favor of the hypothesis of a sheer anion delocalization.

Herein, homoleptic nitroanions **1** and **2** (Fig. 1) are computationally revisited with quantitatively mapping delocalization in their molecular backbones by means of the Weinhold's natural bond orbital (NBO) method¹⁰ that analyses chemical structure with 'translating' orbital consideration to that in chemical bond terms. Conformational dynamics of these anions is modeled as well. Their mononitro analogs, nitronate anion $[\text{CH}_2\text{NO}_2]^-$ (**3**), *N*-methyl nitramide anion $[\text{CH}_3\text{NNO}_2]^-$ (**4**), tris(dimethylboranyl)methane anion $[(\text{Me}_2\text{B})_3\text{C}]^-$ (**5**), and azidoform anion $[\text{CN}_3]^-$ (triazidomethane anion; **6**) are similarly approached. The chemistry of anions **1–6** is considered in the light of the calculated orbital delocalization.

Results and discussion

Studied molecules are modeled for vacuum conditions with using the Dunning basis set aug-cc-pVTZ or Karlsruhe basis set QZVP. The Møller–Plesset post-HF calculation methods are

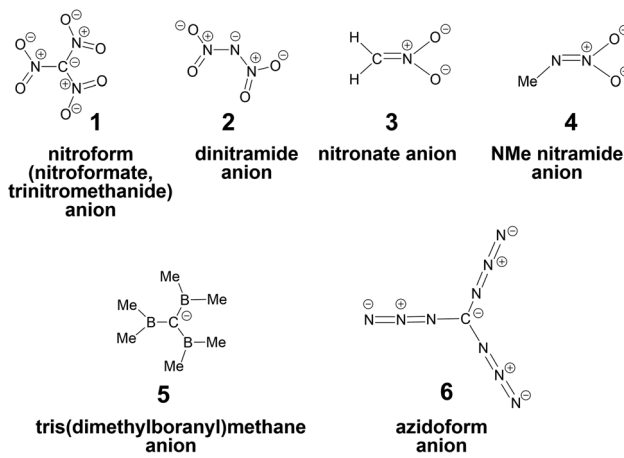


Fig. 1 Chemical structure of anions **1–6** shown in a plain notation of localized bonds. This formal view is very far from reflecting the chemical bonding in **1–6**.

exploited; that of the second-order perturbation –MP2– for anions **1**, **2**, **5** and that of the third-order perturbation –MP3– for anions **1** and **2**. As expected, it is possible to model molecular geometry of these anions within the limits of harmonic approximation. When examining applicability of the latter on the example of anion **2**, only insignificant differences are found to distinguish between the vibrationally averaged molecular geometry and the equilibrium geometry (e.g., the difference is 7×10^{-4} nm for the N–N bond length and 0.09° for the bond angle N–N–N; ESI†). MP2/aug-cc-pVTZ and B2PLYP/QZVP calculations provide a very similar molecular geometry for anion **6**, and this double-hybrid functional in combination with this quadruple-zeta basis set has been chosen for analyzing orbital delocalization in this anion.

Quantitative orbital delocalization is computed in the form of natural semi-localized molecular orbitals (NLMOs), where each NLMO is a linear combination of the parent natural bond orbital (NBO) and the delocalization tail (contributions of the parent NBO to other, vacant NBOs¹⁰). Notably, in spite of its apparently easy interpretability, this 'light construction', NLMO, is not a step-down theoretical simplification: it does not decrease the accuracy of the *ab initio* theory in representing the wave function.

Chemical bonding in anions **3** and **4**

Computational screening of the PES for **3** (including initial structures with a pyramidal C atom) shows that this molecule is planar as well as single-shaped in the ground state (Fig. 2), confirming less accurate early calculations.¹¹ NBO analysis of the chemical structure of anion **3** shows that there are two NLMOs in bonding the C and N atoms, both of two-electron

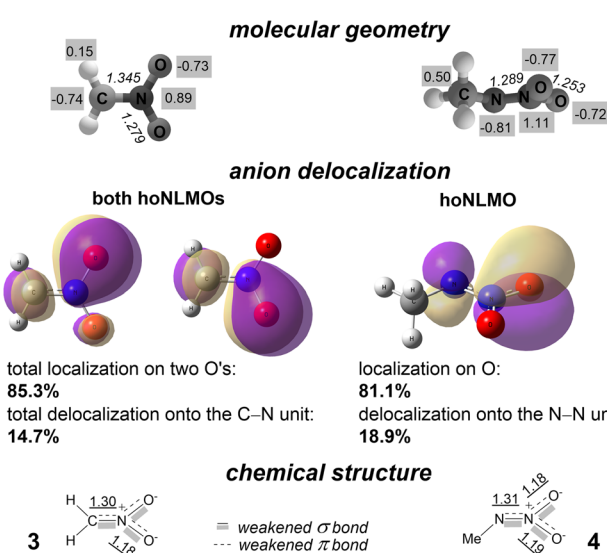


Fig. 2 Molecular geometries of anions **3** and **4** (numbers in italics display bond lengths, Å, while those with gray background show CHELPG atomic charges¹⁴), hoNLMOs with quantitatively mapped anion delocalization, and Lewis structures of anions with the bonding that reflects the weakening effect of partial σ^* and π^* antibonds. Underlined numbers show calculated bond orders (as Wiberg indexes).



occupancy near 100%, while there is no occupied NLMO solely related to the C atom. Thus, the carbon and nitrogen are both σ - and π -bonded (Lewis structure C=N), as usually accepted for this anion. This chemical structure does not point to the still considerable C-nucleophilicity¹² in S_N2 reactions in aprotic solvents of low polarity.

The chemical structure C=N does not accurately describe the bonding of the C and N atoms in 3. For each O atom, there is a remarkable, 7.3% delocalization tail of one NLMO occupied by a pair of p electrons. Thus, the O,O-anion delocalization is 14.7% (Fig. 2). These two NLMOs of the same energy are the highest occupied NLMOs (hoNLMOs) for 3. For each of these NLMOs, the delocalization tail is due to a partial donation of unshared electrons of these O's from the p orbital to the π^* orbital of the C=N unit; the related orbital interaction energy is large (109.7 kcal mol⁻¹ for each pair of the p and π^* NBOs). A weighty 0.58 occupancy of this π^* orbital means that the bonding of the C and N atoms is appreciably weakened vs. pure bonding C=N formally assigned earlier to this fragment.¹³ Also, the NBO analysis rejects the single bond structure N-O 'canonized' (ibid) for both diatomic N,O units of anion 3. There is a $n \rightarrow \pi^*$ vicinal donation that strengthens the bonding of the N and both O atoms by 'connecting' them by a partial π bond (in addition to the σ bond N-O; Fig. 2).

Hyperconjugation also affects the chemical bonding in 3. Each pair of p electrons of both O atoms, which does not participate in occupation of the π orbital of the C=N unit, partially occupies the σ^* orbital of the C-N (each related stabilization energy is 8.6 kcal mol⁻¹). In addition, one of these pairs of both oxygens is in part donated to the σ^* orbital of the geminal N-O unit (each related stabilization energy is 22.3 kcal mol⁻¹). Stabilizing the molecule as a whole, this delocalization of the O,O unshared valence electrons weakens the σ bonding both in C-N and N-O units. With including these delocalizations of lone electron pairs, the bonding of the C and N atoms in nitronate anion 3 appears the 'sum' σ bond + partial σ^* antibond + π bond + partial π^* antibond, and the bonding of the N and O atoms appears σ bond + partial σ^* antibond + partial π^* bond. Bond orders calculated for 3 confirm that the bonding in its C, N fragment is significantly weaker than that of the pure double bond C=N (σ bond + π bond), and the bonding in its N, O fragment is stronger than that in the single bond N-O.

Structural conclusions are similar for nitramide anion 4. There are two doubly occupied NLMOs associated with bonding the two nitrogen atoms; they both do not have a considerable delocalization tail. Thus, this anion is a diazenium-1,1-diolate (Fig. 2). The performed NBO analysis adds more detail to the familiar chemical structure [MeN = N(O)O]⁻ deduced from IR spectra and early *ab initio* calculations.¹⁵ There is a highly stabilizing $n \rightarrow \pi^*$ electron donation from the both O's to the unit N=N: the π^* orbital of this unit has a 0.65 occupancy while the occupancy of this donating n orbital of each O atom decreases to 1.65. As the related NLMOs show, delocalization of this partially donated lone electron pair is 18.9% for the *trans*-positioned oxygen and 15.6% for the *cis*-positioned oxygen. The other p electron pair of both O atoms also is not O-localized.

This pair is partially donated to the π^* orbital of the N=N fragment as well as to the σ^* orbital of the N-O fragment with the related stabilization energies of 13.4 and 14.8 kcal mol⁻¹, respectively. That is, the bonding which binds the N atoms in 4 is σ bond + partial σ^* antibond + π bond + partial π^* antibond, and the bonding which ties the atoms N and O of the nitro group is σ bond + partial σ^* antibond + partial π^* bond (Fig. 2).

Chemical bonding in anions 1 and 2

There are two conformers **1A** (molecular symmetry D_3) and **1B** (molecular symmetry C_{2v}) for 1; they interconvert with synchronously rotating the C substituents and changing interatomic distances C···N (Fig. 2 and 3).[‡] Transformation **1A** ⇌ **1B** is associated with rotating all three NO₂ groups in the conrotatory fashion, while degenerate transformation **1A** ⇌ **1A** is provided by rotating two NO₂ groups conrotatory and one in the opposite direction. Diastereomerization **1A** ⇌ **1B** appears a very fast intramolecular process: the calculated barrier $\Delta G_{AB}^\#$ is 1.8 kcal mol⁻¹ at 298 K (the barrier corresponds to transition state **TS_{AB}** in Fig. 3). Interconversion of degenerate (structurally identical) conformers **1A** occurs slower (calculated barrier $\Delta G_{AA}^\#$ = 6.8 kcal mol⁻¹; the related transition state is **TS_{AA}** in Fig. 3) while degenerate conformers **1B** do not directly interconvert. In total, stereodynamics of anion 1 occurs as interconversion of six degenerate structures **1A** and six degenerate structures **1B**.

The NLMO associated with the C lone electron pair is the hoNLMO for both conformers. It is significantly C-localized, though somewhat differently for them (Fig. 4). That is, in contrast to the mononitro analog 3, anion 1 is a carbanion with a 29.3% and 33%-delocalization for conformers **1A** and **1B**, respectively. Its essential C-localization indicates that this molecule cannot be assigned to Y-aromatic systems. A strong Y-aromaticity for 1 would require the delocalization tail to be of the 65–75% magnitude, *i.e.*, the contribution of each related molecular fragment (the carbon atom and three NO₂ groups) into this hoNLMO should be more or less equal.

Together with this, the delocalization tail of the hoNLMO both for **1A** and **1B** is not negligible. The occupancy of the parent C-localized NBO for conformers **1A** and **1B** is substantially less than 2 (1.42 and 1.32, respectively). This deficiency in the orbital occupancy mainly is because of partially accommodating these p electrons of the C atom in the antibonding π^*

[‡] Isomers **1A** and **1B** are a borderline case in classifying isomers either as stereoisomers or as chemical isomers. Traditional understanding of covalent chemical structure (a molecular framework of 2c-2e bonds) only implies full occupation of bonding orbitals. From this perspective, **1A** and **1B** are conformers (rapidly interconverting stereoisomers – a simplified form of a strict definition of conformer from monograph A. M. Belostotskii, Conformational Concept for Synthetic Chemist's Use, World Scientific Publishing, Singapore, 2015). However, a sizeable donation to a vacant π orbital actually means a partial π bond. Such a 2c-ne bond ($n < 1.8$) – a bond in the sense of a certain, definable chemical bonding – may be considered the structural determinant to the same extent as the canonical doubly occupied 2c-2e bond is. The molecule of **1A**, in contrast to **1B**, does not have partially occupied π orbitals for all three diatomic fragments C-N (Fig. 4). Thus, the number of bonds in **1A** and **1B** is different. From this viewpoint, these structures are chemical isomers.



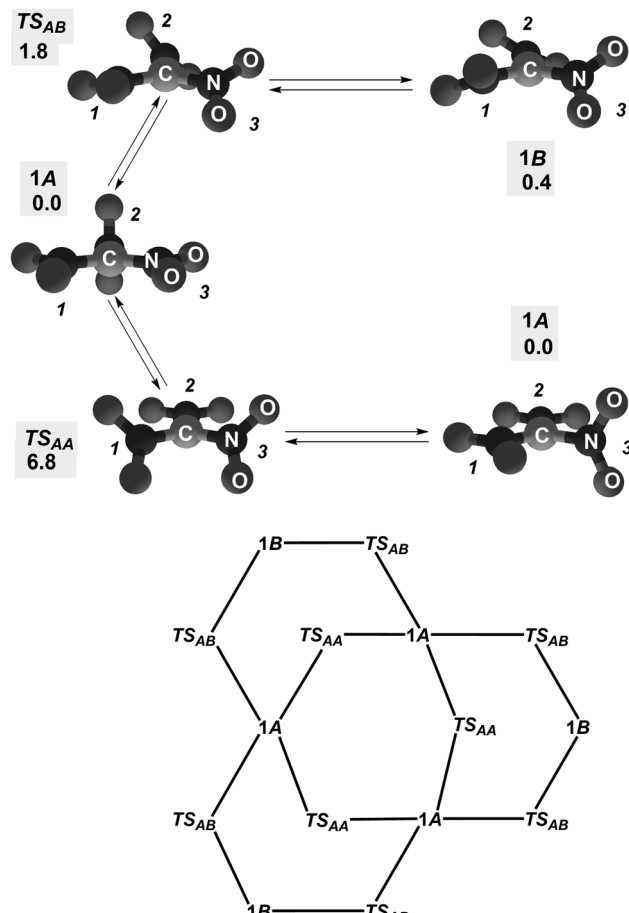


Fig. 3 Interconversion of conformers **1A** and **1B**. Top: interconversion in the pair **1A** and **1B** and interconversion in the pair of two degenerate conformers **1A**. Bold numbers with gray background indicate calculated relative free energies (kcal mol^{-1}) for these structures, and numbers in bold italics are formal labels of NO_2 groups. For clarity, the central atom and atoms of one NO_2 group are supplied with related chemical element symbols. Bottom: the representative graph of conformational pathways for anion **1** (interconversion of three degenerate conformers **1A** and three degenerate conformers **1B** of axial chirality R_a).

NBOs of the $\text{N}=\text{O}$ units, as the distinctively high values of the stabilization energy for these n donor and π^* acceptor NBOs show (more than $130 \text{ kcal mol}^{-1}$ for each of three nitro group in **1B** and $260 \text{ kcal mol}^{-1}$ for each of two coplanar nitro groups in **1A**). Thus, there is a partial π bonding in connecting the C atom with each of the three N atoms in **1B** while this π bonding connects the C atom with only two N atoms in **1A**. This orbital interaction also explains why C–N bond lengths both in **1A** and **1B** are different for their nitro groups differently oriented rotationally: the considered bond-shortening conjugation is maximal at the coplanar disposition of three atoms of the nitro group and the carbon atom (Fig. 4).

The hoNLMO delocalization tails for **1A** and **1B** expose that the carbanion delocalization is dissimilar for them: the delocalization onto one among their NO_2 groups is 1.1% and 11.0% for **1A** and **1B**, respectively (Fig. 4). In particular, this means that the conformers of ambident anion **1** possess somewhat

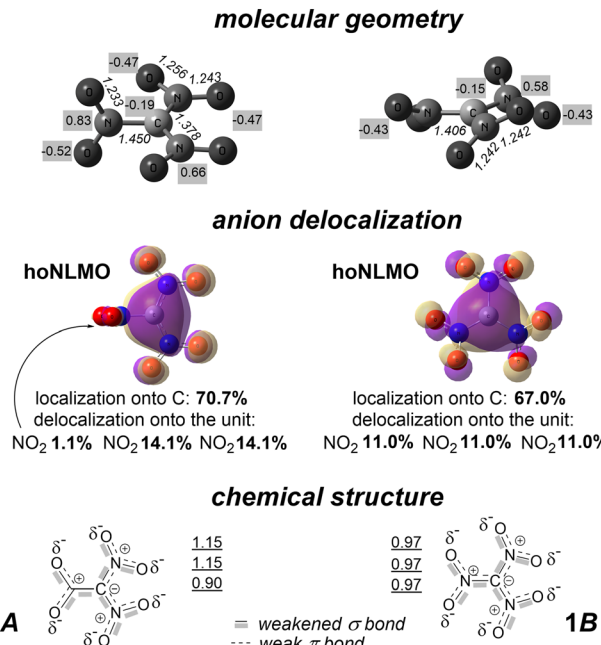


Fig. 4 Conformers **1A** and **1B**: calculated molecular geometry (numbers in italics point out interatomic distances, \AA ; numbers with gray background present CHELPG atomic charges¹⁴), the hoNLMO with quantitatively mapped carbanion delocalization, and Lewis structure with indicating the bond weakening due to partial occupancy of σ^* and π^* antibonding orbitals. Underlined numbers display calculated C–N bonding orders (as Wiberg indexes).

different C-nucleophilicity. Nevertheless, it would be rather impossible to discern this difference in experiments. Nucleophilic reactions of this anion are not controlled conformationally: the kinetic barrier of interconversion of **1A** and **1B** is very low (Fig. 3).

Unshared electron pairs of the oxygen atoms of **1** are not localized. There is a partial donation of one p electron pair of each O atom of each nitro group to the σ^* orbital of the corresponding C–N fragment [with energy of $16.0 \text{ kcal mol}^{-1}$ for the electron pair in **1B** and $15.8 \text{ kcal mol}^{-1}$ (in average) for the electron pair in **1A**]. Summarizing all these delocalizations propagated to this diatomic fragment, one can conclude that the bonding that ties the C and N atoms in **1** is σ bond + partial π bond + partial σ^* antibond for **1B** as well as for the two coplanar nitro groups of **1A**. While the aforementioned vicinal $n \rightarrow \pi^*$ conjugation reinforces this bonding, the $n \rightarrow \sigma^*$ hyperconjugation weakens it (see Fig. 4 for the calculated bond orders) together with strengthening the bonding between the N and O atoms. The bonding that ties the central C atom and the nitro group of **1A**, which is orthogonal to the other NO_2 groups, is σ bond + partial σ^* antibond.

Concerning the nitro groups of **1** themselves, the bonding in $\text{N}=\text{O}$ units is identical in each NO_2 group of conformer **1B**. Similarly, it is the same in the two coplanar NO_2 groups of conformer **1A**. In each of these nitro groups, there are both $\text{N}=\text{O}$ and $\text{N}–\text{O}$ units with oxygen lone electron pairs partially delocalized in the NO_2 fragment. For **1A**, the delocalization includes high energy $n \rightarrow \pi^*$ vicinal donation from the $\text{N}–\text{O}$

unit to the N=O unit (188.8 kcal mol⁻¹ for each from two coplanar NO₂ groups and 283.9 kcal mol⁻¹ for the NO₂ group orthogonal to them). Such a donation characterized by a 241.0 kcal mol⁻¹ energy for each NO₂ group of **1B**. In other words, there is an essential partial π bonding in each N–O unit of anion **1**.

Anion **2** is more localized on the central atom (80.4%; Fig. 5) than carbanion **1**. Nevertheless, it is not a highly localized N-anion. The 19.6% delocalization tail of the hoNLMO associated with one unshared electron pair of the central N encompasses both nitro groups, *i.e.*, this lone electron pair is involved by almost one fifth in forming the bonding which ‘attaches’ each NO₂ substituent to the central N atom.

The performed NBO analysis indicates that each NO₂ group has both the N=O and N–O unit (correspondingly, the N···O distances are different for them). The mentioned p electron pair of the central N atom is in part donated to the π^* orbital of the N=O unit of each nitro group, diminishing the occupation of the donating n NBO to 1.61. That is, the additional, delocalization-delivered bonding in the σ -bonded fragment N–N–N is a partial π bonding (see Fig. 5 for the calculated bond orders). Correspondingly, both interatomic distances N···N are decreased, and N···O distances in both units N=O are increased.

Similarly, for each nitro group, there is a high stabilization energy (more than 160 kcal mol⁻¹) donation of one p electron pair of the O atom of the N–O unit to the π^* orbital of the N=O unit of the same nitro group (the resulting occupation of this antibonding orbital is 0.54). In chemical bond terms, a partial π bond, in addition to the σ bond, ties the N and O atoms in both N–O units.

Together with this, hyperconjugation $n \rightarrow \sigma^*$ slightly weakens the σ bond framework in anion **2**. Both oxygens of each NO₂ group partially donate one lone electron pair to the σ^* orbital of the related N–N fragment (the orbital interaction energy is \sim 16.7 kcal mol⁻¹ for each of the N–O oxygens and 22.8 kcal mol⁻¹ for each of the N=O oxygens). The occupancy of this antibonding orbital of both N–N fragments is further increased to 0.08 due to the partial $n \rightarrow \sigma^*$ donation (with energy of $2 \times$ 12.7 kcal mol⁻¹) of the unshared N electron pair which does not participate in occupying the π^* orbital of the N=O units. For both N–O units, one O atom partially donates (with energy of 20.9 kcal mol⁻¹) to the σ^* orbital of the N=O fragment of this NO₂ group, while the O atom of the N=O fragment similarly donates a p electron pair to the σ^* orbital of the geminal N–O unit with stabilization energy of 22.8 kcal mol⁻¹. When describing these delocalizations of lone electron pairs in terms of chemical bonding, chemical structure of anion **2** appears somewhat different from that considered^{12,13} earlier. The bonding of the N atoms is σ bond + partial π bond + partial σ^* antibond. The bonding in the N=O units is σ bond + π bond + partial σ^* antibond + partial π^* antibond, while the bonding in the N–O units is σ bond + partial σ^* bond + partial π^* antibond. Clearly, there are two opposite tendencies in bonding atoms in units N–N and N–O: the bond weakening due to the $n \rightarrow \sigma^*$ hyperconjugation and its strengthening due to the $n \rightarrow \pi^*$ conjugation. The calculated bond orders for these diatomic units (Fig. 5) reflect the final balance of these tendencies.

molecular geometry and conformational dynamics

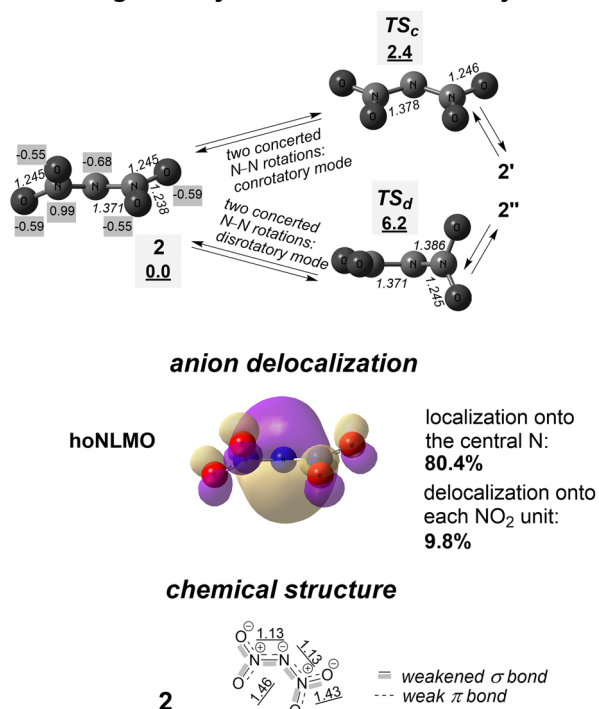


Fig. 5 Anion **2** and interconversion of degenerate conformers **2**, **2'** and **2''** through transition states **TS_c** and **TS_d**. Shown are calculated molecular geometry (numbers in italics point out interatomic distances, Å; numbers with gray background present CHELPG atomic charges¹⁴), relative free energies (underlined numbers with gray background; kcal mol⁻¹), the hoNLMO with quantitatively mapped N–anion delocalization, and Lewis structure with indicating bond weakening due to partial σ^* and π^* antibonds. Underlined numbers show calculated bond orders (as Wiberg indexes).

Chemical bonding in some other methanide anions

Delocalization of tri(dimethylboranyl) methanide anion **5** (Fig. 6) is the single orbital interaction significantly stabilizing the structure; others are of essentially lower energy. It is a σ occupied \rightarrow σ vacant orbital interaction that takes place in diatomic fragments of neighboring proper atoms, another type of formal electron donation.

This orbital interaction is highly expedient energetically in borane-stabilized C anions,¹⁶ where such 1,2-positioned occupied (for C) and vacant (for B) p orbitals are separated by a modest energy gap and their C_n symmetry axes are coplanar or near-coplanar. Both conformers **5A** and **5B** have the geometrical motif of conformers **1A** and **1B**, and their donor and acceptor p orbitals are almost coplanar for **5B** and coplanar for two C substituents of **5A**. Nevertheless, anion **5** is not Y-aromatic. The delocalization of this anion comprises the four-atom fragment C(B₃), and the anion is localized there onto the C atom by 67.0% in one conformer and by 67.3% in the other (Fig. 6), similarly to anion localization in **1**. The example of anions **1** and **5** that have different channels of strong electron donation indicates that



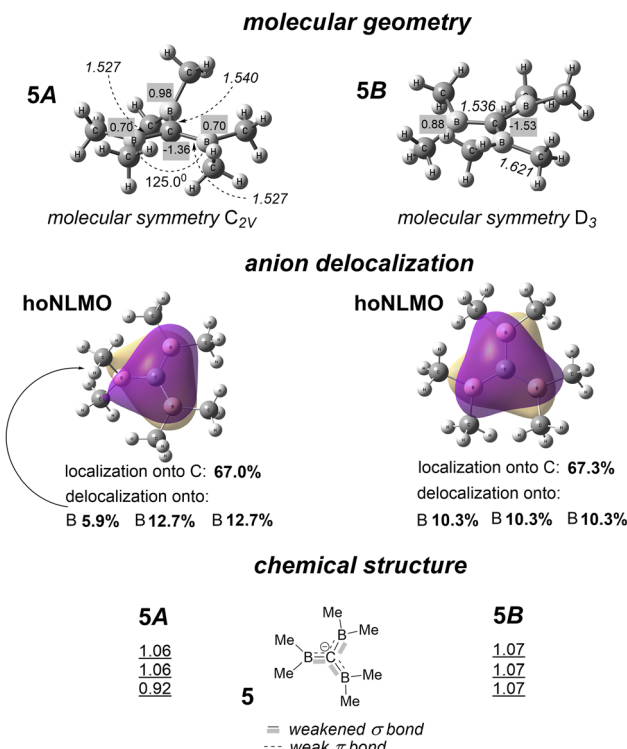


Fig. 6 Conformers 5A and 5B (designations are the same as they are in Fig. 4). Calculated bond orders are shown for diatomic C–B units.

carbanions bearing three substituents of the same chemical structure have a certain, carbon-specific and substituent-invariant, limit in delocalization magnitude ($\sim 30\%$ of delocalization). That is, Y-aromaticity is not inherent in triply substituted methanide anions.

Azidoform anion 6, one of the simplest geminal azide molecules,¹⁹ is unknown. It is difficult to *a priori* judge whether, *e.g.*, salt $\text{Li}^+ \cdot 6$ could be prepared in a short way, for instance, from the parent, very unstable azidoform¹⁷ with using neopentyl lithium (a non-nucleophilic organolithium reagent) or from triazidocarbenium tetrafluoroborate¹⁸ if reducing this salt by Li/NH_3 or lithium naphthalene. Nevertheless, QM modeling of this methanide anion is undertaken here in order to reveal what is the degree of its delocalization; the starting hypothesis is that the $n-\pi^*$ conjugation similar to that in vinyl azides²⁰ stabilizes this energy-rich molecule.

The results are unexpected. The molecular shape of two most stable, near-energy isomers 6A and 6B of 6 \ddagger is a trigonal pyramid, pointing out that this anion is delocalized insignificantly. In addition, C–N bonds in anion 6 (Fig. 7) are slightly longer than the C–N bond in methyl azide,²¹ but appreciably longer than that in π , π -conjugated vinyl azides.²⁰

NBO analysis firmly confirms the above guess, demonstrating that there is no such a vicinal interaction of the 2e-occupied p orbital of the carbon and the vacant π^* orbital of

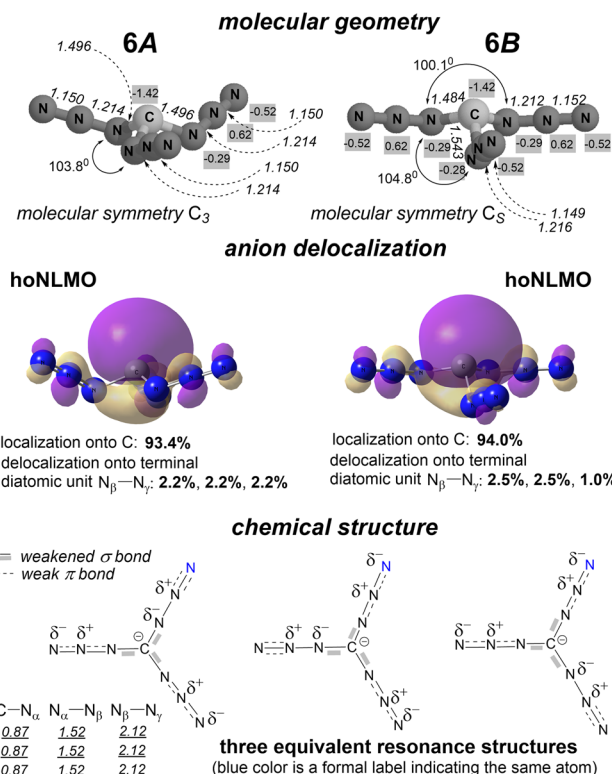


Fig. 7 Isomers 6A and 6B: calculated molecular geometry (numbers in italics point out interatomic distances, Å; arc arrows depict CN bond angles; numbers with gray background present CHELPG atomic charges¹⁴), the hoNLMO with quantitatively mapped carbanion delocalization, equivalent resonance structures of isomers of 6 with indicating the bond weakening due to a partial occupation of π^* antibonding orbitals, and calculated bond orders (as Wiberg indexes) for isomer 6A (underlined numbers; see ESI† for 6B). The chemical structure of isomer 6A is represented by three displayed resonance structures together, while one of these equivalent structures represents the chemical structure of isomer 6B.

the $\text{N}_\alpha-\text{N}_\beta$ fragment. Instead, there is localization of the p orbital of the C lone electron pair on this atom by 93.4 and 94.0% for 6A and 6B, respectively (Fig. 7). The small delocalization tail of 6.6 and 6.0%, respectively, is due to a partial electron donation from this occupied orbital of the C atom to the π^* orbital of each terminal N–N unit (a long range $n \rightarrow \pi^*$ electron donation).

For anion 6, NBO analysis endows two azido substituents with the σ and π framework $\text{N}=\text{N}=\text{N}$ of chemical bonding, and one azido substituent with that of $\text{N}=\text{N}\equiv\text{N}$ (for simplicity, formal charges are omitted here). Excluding a 5% delocalization of each C–N bond (see below), both σ and π bonds are largely localized in both isomers 6A and 6B. Essential delocalizations (i)–(iii) are only associated with orbitals of unshared electron pairs of nitrogen atoms. (i) The p orbital of one unshared electron pair of the N_γ atom in each of two $\text{N}=\text{N}=\text{N}$ substituents both in 6A and 6B interacts with the vacant π^* antibonding orbital of the related $\text{N}_\alpha=\text{N}_\beta$ fragment with decreasing its own localization to 63.9% and populating this antibonding orbital to a significant value of 0.74. The other 2e-occupied p orbital of

§ 6A and 6B are considered as isomers, assuming that a barrier of the height more than 1.5 kcal mol⁻¹ separates these molecular structures in the potential energy surface.

this nitrogen remains N-localized by 98.5% both for **6A** and **6B**. (ii) The p orbital of the unshared electron pair of the N_{α} atom both in $N=N=N$ and $N-N\equiv N$ substituents both in **6A** and **6B** interacts with the antibonding π^* orbital of the corresponding terminal unit $N_{\beta}\equiv N_{\gamma}$ with delocalization of 13.0% (for both $N=N=N$ substituents) and 11.8% (for the $N-N\equiv N$ substituent), respectively. This formal electron donation, together with the $\sigma \rightarrow \pi^*$ donation from the σ orbital of each C–N unit as well as the above mentioned small delocalization tail of the p orbital of the C atom, leads to a 0.32 occupancy of this antibonding orbital for **6A** and 0.32 (for two symmetrically positioned azido substituents) as well as 0.34 occupancy (for the third azido substituent) for **6B** thus loosening the bonding in each of these diatomic units. The $\sigma \rightarrow \pi^*$ donation weakens the σ bond in all C–N units, and the calculated bond orders (Fig. 7) independently 'detect' this weakening. (iii) The second lone electron pair of the N_{α} atom of the $N-N\equiv N$ substituent both of **6A** and **6B** is copiously donated to the second antibonding π^* orbital of the related terminal unit $N_{\beta}\equiv N_{\gamma}$ with populating it by 0.58 and 0.54, respectively, and delocalizing the parent non-bonding orbital by 27.2% for both conformers. Calculated bond orders well reflect the total decrease of the strength of chemical bonding in diatomic units N–N resulted from weakening their π bonds (Fig. 7 and ESI†).

As mentioned above, NBO analysis supplies the chemical bonding framework $N=N=N$ to two azido substituents and chemical bonding $N-N\equiv N$ to one azido substituent for both structures **6A** and **6B** (Fig. 7 shows their Lewis chemical structures with taking into account the delocalization of p orbitals of nitrogen lone pairs). However, all three C substituents in isomer **6A** of C_3 symmetry have the identical molecular geometry (see Fig. 7 for bond lengths), and, hence, the same chemical bonding framework. Neither azido substituent in **6A** therefore has the $N-N\equiv N$ or $N=N=N$ chemical structure. Three equivalent resonance structures together (Fig. 7; those also indicate the π bond weakening due to orbital delocalizations) represent the chemical structure of **6A**, *i.e.*, each azido substituent of this isomer is represented by two-thirds by chemical connectivity $N=N=N$ and by one-third by connectivity $N-N\equiv N$, both with a weakened π bonding.

Concerning isomer **6B** of molecular symmetry C_S , interatomic distances in diatomic units $C-N_{\alpha}$, $N_{\alpha}-N_{\beta}$ and $N_{\beta}-N_{\gamma}$ in two azido substituents, which are mirrorly positioned in the molecule, are identical. The bond lengths of the same diatomic units $C-N_{\alpha}$, $N_{\alpha}-N_{\beta}$ and $N_{\beta}-N_{\gamma}$ in the azido substituent, which lies in the symmetry plane of the molecule, differ from the corresponding ones in the above mentioned symmetrically positioned azido substituents (Fig. 7), indicating that the chemical bonding framework in this substituent is distinct from chemical bonding in other two C substituents in this isomer. This azido substituent is supplied by the NBO analysis with chemical bonding $N-N\equiv N$; two other substituents have that of $N=N=N$. Thus, the chemical structure of this isomer is represented by one structure from three equivalent ones shown in Fig. 7. This means that the chemical structure of isomer **6B** is more certain than the structure of ordinary alkyl azides. While four classical resonance structures together represent the azido

group (see, *e.g.*, ref. 22), each azido substituent of **6B** is actually represented by one predominantly highly weighted resonance structure. The clear 'off-resonance' chemical structure of the triazide fragment of **6B** renders the azide chemistry of this isomer to be more predictable than that of **6A** (see below).

Chemical structures of **6A** and **6B** are explicitly not identical; in isomerism terms, **6A** and **6B** are chemical isomers and not conformers (rapidly interconverting stereoisomers), in contrast to the border case of **1A** and **1B** explained in distinguishing between chemical isomer-stereoisomer.‡

Interestingly, the deduced chemical structure of isomer **6A** (Fig. 7) is related to a diverse group of molecular structures which cannot be adequately filed with exploiting standard formats of computer-readable representation of chemical structure ubiquitously used in cheminformatics (*e.g.*, SMILES, InChI, MDL Mol, *etc.*). This non-Lewis structures thus is technically problematic for including it into datasets focused on applying computerized AI algorithms to molecular or crystal-level design of high-energy materials (exemplified by datasets used in dozens of related recent studies²³). It is an open question whether other geminal polyazides, for instance, tetraazidomethane,²⁴ fall under such a 'hard nut' category.

Explainable chemistry of anions **1–4** and chemical predictions for anion **6** (examples). Quantitative mapping of delocalization in anions **1–4**, **6** assists in understanding and predicting their chemical reactivity. For instance, in reactions, which involve nitro groups of methanide anion **1**, the reactivity of these groups NO_2 of distinct rotational orientation in conformer **1A** should be somewhat dissimilar. Namely, the nitro group of **1A**, which does not essentially delocalize the unshared electron pair of the carbon (Fig. 4), should be more prone to rupturing the C–N bond than two other NO_2 groups. Indeed, DFT modeling of decomposition pathways of molecules of $K^+ \cdot \mathbf{1}$ in both conformations²⁵ shows that the lower barrier of disruption of the C–N bond is for the conformer of geometrical motif of conformer **1A**. NO_2 groups in molecules of crystalline $NH_4^+ \cdot \mathbf{4}$ have different N–N rotational orientations.²⁶ According to this QM modeling, the NO_2 group which is near orthogonal to the NNN plane of the molecule, *i.e.*, minimally participates in delocalization of the anion, is kinetically most prone to loss its chemical bonding with the central nitrogen atom, similarly to the above described case of anion **1**.

Another example is S_N2 reactions of ambident anions **3** and **1** with primary alkyl halides. If very roughly assessing the anion delocalization, the mononitro anion **3** is an O-anion, while the pernitro anion **1** is a C-anion (Fig. 1 and 3). Trivial alkylation of mononitronate anions crucially depends on counterion and solvent, *i.e.*, factors which determine the strength of the heteroatom-cation coordination in solution and thus interfere with the nucleophilicity of the naked anion. Nevertheless, the common experimental conclusion is that the inherent reactivity of nitronate anions is that of the O-anion.²⁷ In contrast, C-alkylation absolutely predominates in similar reactions of trinitromethanide anion **1**.²⁸ The quantitatively mapped anion delocalization well explains both experimental observations.

The 'prominent' anion **2** is the energy-storing core of superior low-cost explosives forged in the 70 s–80 s and the oxidizing



constituent in some modern propellants.²⁹ Both the electrocyclic extrusion of the nitrate anion and homolytically initiated release of gaseous nitrogen oxides in decomposition of this nitroanion are characterized by relatively low kinetic barriers.³⁰ For these barriers, the undertaken calculations provide free energy values $\Delta G^\#$ of 43.0 and 36.3 kcal mol⁻¹, respectively (ESI);† they are satisfactory close to the barrier values calculated earlier.³⁰ The crucial factor of this easiness of molecules of 2 to fall apart is exposed if including orbital delocalization into consideration. As appears, the occupancy of the σ^* antibonding orbital in unit N-N of anion 2 archly increases when stretching this diatomic fragment. The same is true for the σ^* orbital of unit N-O being stretched, too. Homolytic decomposition of anion 2 starts from dissociation of one N-N bond (Fig. 8). The increase in occupancy from 0.08 (see above for the hyperconjugation in this anion) to 0.62 in the transition state on the route to the intermediate product $\text{N}_2\text{O}\cdot\text{N}_2\text{O}_4^-$ (ESI);† softens this bond utterly. The occupancy further increases to 0.69 when reaching this supramolecule in the reaction pathway.

The above mentioned electrocyclic deintegration of the molecule of 2 requires two bonds, the N-N and N-O, to be concertedly broken (Fig. 8 and ESI†). Thus, the kinetic stability of 2 could be expected to be high while, in fact, it is relatively low, as noticed above. The revealed NO₂ group-associated feature of the increase of the occupancy of the σ^* orbital under stretching the related σ bond plays a significant role in decomposition of anion 2 also according to this molecular mechanism. In the related transition state (Fig. 8), the σ^* orbital occupancy for both bonds being disrupted, namely the N-N bond and the N-O bond, increases to 0.77 and 0.55 (from 0.09 for the parent anion 2), respectively. Not surprisingly, decomposition temperatures for molten salts cation 2 are lower by \sim 100 °C and more than those for molten salts cation 4, where the anion has a stronger bonding of a partially double bond character in the N₂N unit (Fig. 2).

Both different mechanisms of decomposition of nitroanion 2 are associated with a smooth increase of the occupancy of the antibonding σ^* orbital in parallel with stretching the related σ bond. One can reasonably ask whether this molecular feature – the coupling of elongation of a σ bond with a peculiar decrease of the strength of this bond (originated from elevating the σ^* orbital occupancy) – is the common cause of proneness of nitro compounds to decompose under thermal impact.

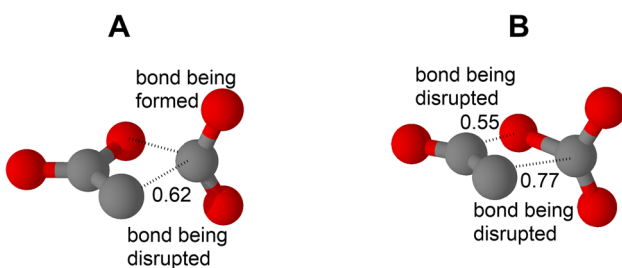


Fig. 8 (A) The located transition state in homolytic decomposition of anion 2 (TS_{low} in ESI†). (B) The located transition state of the concerted electrocyclic decomposition of this anion (ESI†). Numbers show the occupancy of the antibonding σ^* orbitals for both A and B.

Concerning triazido methanide 6, the established high localization of the anion on the central atom permits to formally separate the chemistry of 6 in that of the C-anion, *i.e.*, a trivial chemistry of carbanions, and that of the azido substituents, *i.e.*, a difficult-to-predict chemistry of geminal polyazides.^{24,31} However, this anion localization has a clear consequence: the σ bonding framework in the central four-atom fragment (C)NNN of this anion is not reinforced by an additional (*i.e.*, π) bonding, in sharp contrast to the case of nitroanions 1 and 2. Moreover, the elucidated chemical structure of 6 (Fig. 7) reflects that C–N bonds are weakened in this fragment.

In regard to carbanion chemistry, if a salt cation 6 is synthetically available and possesses a considerable stability at moderately low temperatures, one may foresee that one-electron or two-electron oxidation of 6 with producing the corresponding radical (N₃)₃C[·] or cation (N₃)₃C⁺,¹⁸ respectively, is quite feasible: the carbanion is only insignificantly delocalized. One can also expect an appreciable C-nucleophilicity for this anion.

The chemical structure of isomer 6B (Fig. 7) prompts that the thermally induced nitrene-generating decomposition of the azido group³² is facilitated for this molecule. Indeed, the single-resonance-structure chemical bonding N–N≡N in one of its azido groups (the absence of appreciably weighted resonance structures which disallow N₂ extrusion without rearranging chemical bonding and changing interatomic distances C···N_z, N_z···N_β and N_β···N_γ in this substituent; Fig. 6) clearly points out this chemical lability of 6B to be higher than that of 6A, the isomer without azido substituents of a ‘pure’ N–N≡N resonance structure.

This conclusion can be extended to claiming that isomer 6B is kinetically more labile than any other ordinary alkyl azide. Probably, an isomer of the parent azidoform also has an ‘off-resonance’ structure N–N≡N in its triazide fragment (similar to that in 6B) or the chemical structure with a predominated weight of resonance structure N–N≡N of one of its azido substituents. This reasonable assumption well explains explosive degradation of azidoform in solution observed at the room temperature.^{17a} Homolytic decomposition of 6 and, thus, azidoform with dissociating the C–N bond and yielding azide radical³³ could be scarcely assessed to be a dark reaction possible to occur at this temperature although the σ bonding framework has a diminished strength in this anion as stated above.

The ‘pure’ dipolar N[–]–N⁺≡N chemical structure of one azido substituent of 6B also indicates an increased reactivity of this isomer in [3 + 2] cycloadditions. Indeed, isomerization of this ‘off-resonance’ structure into reactive 1,3-dipole N[–]–N=N⁺ is associated with the energy cost of only changing one interatomic distance N_β···N_γ.

Conclusions

The undertaken QM modelling followed by NBO analysis finds planar nitroanions 1 and 2 to be delocalized to the 33% and 20% extent, respectively. Consequently, the conclusion, which finalizes the long-term discussion on aromaticity suggested for



anion **1**, is that it is not a Y-aromatic system. The delocalization-prone planar anion **5** is delocalized to the same 33% extent, and, thus, the numerical value of one-third probably is the upper limit of delocalization of homoleptic three-substituted methanide anions. Azidoform anion **6** appears to be pyramidal and is almost entirely C-localized. One from its two lowest energy isomers has a unique chemical structure: one azido substituent possesses the chemical bonding of a 'pure' (predominantly highly weighted) resonance structure $\text{N}-\text{N}\equiv\text{N}$, while two others have the chemical bonding of single resonance structure $\text{N}=\text{N}=\text{N}$ highly weighted to the same extent. An important result of these calculations is that the essential chemistry of the studied anions becomes understandable or even predictable when quantitatively mapping orbital delocalization in the molecular backbone. For instance, the outcome of nucleophilic reactions of ambident nitroanions **1** and **3** is clear in the light of the mapped anion delocalization.

A remarkable example of this 'through-the-delocalization-prism' understanding of reactivity is the revealed cause of the easiness of the high-energy anion **2** to initiate its decomposition both *via* N–N bond homolysis and electrocyclic extrusion of nitrate anion. Delocalization of one 2e-occupied p orbital of both O's of the departing NO_2 group onto the antibonding σ^* orbital of this bond significantly increases in parallel with stretching this diatomic unit N–N. This finding leads to reasonably conclude that this cause of the proneness of a molecule to start disintegration – the raising increase in the occupancy of the σ^* orbital related to the being dissolved σ bond is probably common for aliphatic nitro compounds.

Calculation methodology

Geometry optimization and frequency calculations were performed at the MP2/aug-cc-pVTZ level (anions **1**, **2**, and **5**), and MP3/aug-cc-pVTZ level (anions **3** and **4**) for the gas phase with using harmonic approximation; both MP2/aug-cc-pVTZ and B2PLYP/QZVP calculations were applied to modeling of anion **6** (Gaussian16 calculation package, Revision A.03).³⁴ Anharmonic vibrational frequency calculations were undertaken for anion **2** and both transition states **TS_c**, and **TS_d**. NBO program implemented in Gaussian16 was exploited for NBO analysis with setting the program option Highly Delocalized Structure. A convenient narrow-specialized computer program Chemcraft³⁵ was used for generating files of the Gaussian's cub format from the Gaussian-produced files of the fchk (formatted checkpoint) format. The Gaussian's graphical interface program GaussianView6 provided visualization of calculated orbitals.

Energy minimization was performed with exploiting the Berny algorithm implemented in the used Gaussian16 version. The GEDIIS algorithm embedded in Gaussian was employed when optimizing the geometry of stable (energy-minimum) structures, while the Rational Function Optimization was the algorithm option involved in optimizing the geometry of transition states.

Transition states **TS_{AB}**, **TS_{AA}**, **TS_c**, and **TS_d** were located *via* generating rotamers by sequential on-the-screen 10°-step rotation of one substituent relative to the other one

(GaussianView) and optimizing the geometry of the each resulting rotamer. Transition state **TS_h** of anion **2** were located by randomly varying interatomic distances $\text{N}_1\cdots\text{N}_7$ (ESI, page S34†) in the range of 1.8 to 1.9 Å, $\text{N}_2\cdots\text{O}_6$ in the range of 1.6 to 1.7 Å, and $\text{N}_1\cdots\text{O}_6$ in the range of 1.7 to 1.8 Å, all with increment of 0.1 Å, optimizing the geometry of the each generated structure under 'freezing' interatomic distances for the above indicated atom pairs and re-optimizing the resulting geometry with removing geometry constraints. Transition state **TS_{low}** was located in the similar way with varying interatomic distances $\text{N}_1\cdots\text{N}_7$ (ESI, page S36†) in the range of 1.9 to 2.0 Å, $\text{N}_2\cdots\text{O}_6$ in the range of 1.3 to 1.4 Å, and $\text{N}_1\cdots\text{O}_6$ in the range of 1.7 to 1.8 Å. The IRC procedure (a Gaussian utility) was applied to establishing the relationship of each located transition state with the corresponding energy-minimum structures.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The governmental KAMEA program coordinated by the Council for Higher Education of Israel has provided the financial ground for this work. The author is always very thankful to a group of civic-minded professors of Israeli universities headed by Prof. Benjamin Fein who, in their times, volunteered to develop and advance this nonsymbolic arrangement in the Israel Academy of Sciences and Humanities and succeeded in their high initiative.

References

- (a) M. Göbel and T. M. Klapötke, Potassium-, Ammonium-, Hydrazinium-, Guanidinium-, Aminoguanidinium-, Diaminoguanidinium-, Triaminoguanidinium- and Melaminiumnitroformate – Synthesis, Characterization and Energetic Properties, *Z. Anorg. Allg. Chem.*, 2007, **633**, 1006–1017; (b) J. C. Bryan, M. N. Burnet and A. A. Gakh, Tetrabutylammonium and Caesium Salts of Trinitromethane, *Acta Crystallogr., Sect. C*, 1998, **54**, 1229–1233; (c) A. A. Gakh, J. C. Bryan, M. N. Burnett and P. V. Bonnesen, Synthesis and structural analysis of some trinitromethanide salts, *J. Mol. Struct.*, 2000, **520**, 221–228; (d) L. Yang, J. Zhang, T. Zhang, J. Zhang and Y. Cui, Crystal structures, thermal decompositions and sensitivity properties of $[\text{Cu}(\text{ethylenediamine})_2(\text{nitroformate})_2]$ and $[\text{Cd}(\text{ethylene-diamine})_3](\text{nitroformate})_2$, *J. Hazard. Mater.*, 2009, **164**, 962–967; (e) A. F. Baxter, A. Martin, K. O. Christe and R. Haiges, Formamidinium Nitroformate: An Insensitive RDX Alternative, *J. Am. Chem. Soc.*, 2018, **140**, 15089–15098.
- (a) R. D. Gilardi, J. Flippen-Anderson, C. George and R. J. Butcher, A New Class of Flexible Energetic Salts: The Crystal Structures of the Ammonium, Lithium, Potassium, and Cesium Salts of Dinitramide, *J. Am. Chem. Soc.*, 1997, **119**, 9411–9416; (b) D. C. Sorescu and D. L. Thompson,

Classical and Quantum Mechanical Studies of Crystalline Ammonium Dinitramide, *J. Phys. Chem. B*, 1999, **103**, 6774–6782; (c) E. A. Zhurova, A. Martin and A. A. Pinkerton, Chemical Bonding in Biguanidinium Dinitramide and Biguanidinium Bis-Dinitramide from Experimental X-ray Diffraction Data, *J. Am. Chem. Soc.*, 2002, **124**, 8741–8750; (d) S. Venkatachalam, G. Santhosh and K. N. Ninan, An Overview on the Synthetic Routes and Properties of Ammonium Dinitramide (ADN) and other Dinitramide Salts, *Propellants, Explos., Pyrotech.*, 2004, **29**, 178–187.

3 (a) V. A. Shlyapochnikov, M. A. Tafipolsky, I. V. Tokmakov, E. S. Baskir, O. V. Anikina, Yu. A. Strelenko, O. A. Luk'yanova and V. A. Tartakovskiy, On the structure and spectra of dinitramide salts, *J. Mol. Struct.*, 2001, **559**, 147–166; (b) J. C. Bottaro, P. E. Penwell and R. J. Schmitt, 1,1,3,3-Tetraoxo-1,2,3-triazapropene Anion, a New Oxy Anion of Nitrogen: The Dinitramide Anion and Its Salts, *J. Am. Chem. Soc.*, 1997, **119**, 9405–9410; (c) L. He, G.-H. Tao, D. A. Parrish and J. M. Shreeve, Impact insensitive dinitromethane salts, *Chem. Commun.*, 2013, **49**, 10329–10331.

4 (a) K. O. Christe, W. W. Wilson, M. A. Petrie, H. H. Michels, J. C. Bottaro and R. Gilardi, The Dinitramide Anion, $\text{N}(\text{NO}_2)_2^-$, *Inorg. Chem.*, 1996, **35**, 5068–5071; (b) H. Michels and J. A. Montgomery, Jr., On the Structure and Thermochemistry of Hydrogen Dinitramide, *J. Phys. Chem.*, 1993, **97**, 6602–6606.

5 J. P. Ritchie, E. A. Zhurova, A. Martin and A. A. Pinkerton, Dinitramide Ion: Robust Molecular Charge Topology Accompanies an Enhanced Dipole Moment in Its Ammonium Salt, *J. Phys. Chem. B*, 2003, **107**, 14576–14589.

6 (a) J. Cioslowski, S. T. Mixon and E. D. Fleischmann, Electronic structures of trifluoro-, tricyano-, and trinitromethane and their conjugate bases, *J. Am. Chem. Soc.*, 1991, **113**, 4751–4755; (b) A. Dworkin, R. Naumann, C. Seigfried and J. M. Karty, Y-Aromaticity: Stable than the Butadienyl Dication?, *J. Org. Chem.*, 2005, **70**, 7605–7616; (c) E. Kleinpeter and A. Koch, Y-aromaticity – existing: yes or no? An answer given on the magnetic criterion (TSNMRs), *Tetrahedron*, 2016, **72**, 1675–1685.

7 H. Brand, J. F. Liebman, A. Schulz, P. Mayer and A. Villinger, Nonlinear, Resonance-Stabilized Pseudohalides: From Alkali Methanides to Ionic Liquids of Methanides, *Eur. J. Inorg. Chem.*, 2006, 4294–4308.

8 Energy minimization performed for conformers of anion **1** at the MP2/6-311+G(d,p) level (preliminary calculations in this work) delivers the molecular geometry with an appreciably pyramidal central carbon instead of that with a planar carbon (corresponding to the experimentally established geometry). This 'deplanarization' problem probably is due to the intramolecular basis set incompleteness error revealed when modeling planar aromatic systems: D. Moran, A. C. Simmonett, F. E. Leach, W. D. Allen, P. v. R. Schleyer and H. F. Schaefer, Popular Theoretical Methods Predict Benzene and Arenes To Be Nonplanar, *J. Am. Chem. Soc.*, 2006, **128**, 9342–9343.

9 K. O. Christe, W. W. Wilson, G. Bélanger-Chabot, R. Haiges, J. A. Boatz, M. Rahm, G. K. S. Prakash, T. Saal and M. Hopfinger, *Angew. Chem., Int. Ed.*, 2015, **54**, 1316–1320.

10 F. Weinhold and C. Landis, *Valence and Bonding*, Cambridge University Press, New-York, 2005.

11 W.-K. Li, An *ab initio* Molecular Orbital Study of the Nitromethyl Anion, *Croat. Chem. Acta*, 1990, **63**, 127–134.

12 (a) P. G. Gildner, A. A. S. Gitterer, D. Cui and D. A. Watson, Benzylation of Nitroalkanes Using Copper-Catalyzed Thermal Redox Catalysis: Toward the Facile C-Alkylation of Nitroalkanes, *J. Am. Chem. Soc.*, 2012, **134**, 9942–9945; (b) K. W. Shimkin and D. A. Watson, Recent developments in copper-catalyzed radical alkylations, *Beilstein J. Org. Chem.*, 2015, **11**, 2278–2288.

13 (a) K. E. Edgecombe and R. J. Boyd, Molecular orbital treatment of substituent effects. I. Structures of some carbon acids and their conjugate bases, *Can. J. Chem.*, 1983, **61**, 45–49; (b) V. A. Shlyapochnikov, M. A. Tafipolsky, I. V. Tokmakov and G. I. Oleneva, Structure and vibrational spectra of the salts of mononitroalkanes, *J. Mol. Struct.*, 2000, **520**, 19–27; (c) R. B. Metz, D. R. Cyr and D. M. Neumark, Study of the 2B1 and 2A2 States of CH_2NO_2 via Ultraviolet Photoelectron Spectroscopy of the CH_2NO_2^- Anion, *J. Phys. Chem.*, 1991, **95**, 2900–2907.

14 C. M. Breneman and K. B. Wiberg, Determining Atom-centered Monopoles from Molecular Electrostatic Potentials. The Need for High Sampling Density in Formamide Conformational Analysis, *J. Comput. Chem.*, 1990, **11**, 361–373.

15 (a) N. Jonathan, The Infrared Spectra and Structures of O,N -Dimethyl Nitramide and the Silver and Ammonium Salts of Methyl Nitramide, *J. Mol. Spectrosc.*, 1960, **5**, 101; (b) V. G. Avakyan, V. A. Shlyapochnikov and A. Chekrygin, Structure of Nitramine Salts, *Bull. Acad. Sci. USSR, Div. Chem. Sci.*, 1972, **21**, 902–904.

16 R. J. Maza, E. Fernández and J. J. Carbó, Mapping the Electronic Structure and the Reactivity Trends for Stabilized α -Boryl Carbanions, *Chem. – Eur. J.*, 2021, **27**, 12352–12361.

17 (a) A. Hassner, M. Stern, H. E. Gottlieb and F. Frolov, Utility of a Polymeric Azide Reagent in the Formation of Di- and Triazidomethane. Their NMR Spectra and the X-ray Structure of Derived Triazoles, *J. Org. Chem.*, 1990, **55**, 2304–2306; (b) Y.-H. Joo, *Doctoral dissertation*, Chemnitz University of Technology, 2007.

18 M. A. Petrie, J. A. Sheehy, J. A. Boatz, G. Rasul, G. K. S. Prakash, G. A. Olah and K. O. Christe, Novel High-Energy Density Materials. Synthesis and Characterization of Triazidocarbenium Dinitramide, -Perchlorate, and -Tetrafluoroborate, *J. Am. Chem. Soc.*, 1997, **119**, 8802–8808.

19 (a) A. P. Häring and S. F. Kirsch, Synthesis and Chemistry of Organic Geminal Di- and Triazides, *Molecules*, 2015, **20**, 20042–22006; (b) B. D. Tsyrenova, P. S. Lempert and V. G. Nenajdenko, Di- and polyazides. Synthesis, chemical transformations and practical applications, *Russ. Chem. Rev.*, 2023, **92**, 1–48.



20 D. J. R. Duarte, M. S. Miranda and J. C. G. E. da Silva, Computational Study on the Vinyl Azide Decomposition, *J. Phys. Chem. A*, 2014, **118**, 5038–5045.

21 (a) B. J. Costa Cabral and M. L. Costa, Density functional study of molecular properties of hydrazoic acid and methyl azide, *J. Mol. Struct.*, 1995, **343**, 31–41; (b) R. L. Livingston and C. X. R. Rao, An Electron Diffraction Investigation Of The Molecular Structure Of Methyl Azide, *J. Am. Chem. Soc.*, 1960, **64**, 756–759.

22 S. Bräse, C. Gil, K. Knepper and V. Zimmermann, Organic Azides: An Exploding Diversity of a Unique Class of Compounds, *Angew. Chem., Int. Ed.*, 2005, **44**, 5188–5240.

23 X. Huang, C. Li, K. Tan, Y. Wen, F. Guo, M. Li, Y. Huang, C. Q. Sun, M. Gozin and L. Zhang, Applying machine learning to balance performance and stability of high energy density materials, *iScience*, 2021, **24**, 102240.

24 K. Banert, Y.-H. Joo, T. Rüffer, B. Walfort and H. Lang, The Exciting Chemistry of Tetraazidomethane, *Angew. Chem., Int. Ed.*, 2007, **46**, 1168–1171.

25 F.-L. Zhang, H.-S. Huang and X.-H. Xie, Theoretical Studies on the Thermal Decomposition Mechanism of Potassium Nitroformate, *Chin. J. Struct. Chem.*, 2016, **35**, 514–520.

26 M. Rahm and T. Brinck, On the Anomalous Decomposition and Reactivity of Ammonium and Potassium Dinitramide, *J. Phys. Chem. A*, 2010, **114**, 2845–2854.

27 (a) K. W. Shimkin and D. A. Watson, Recent developments in copper-catalyzed radical alkylations of electron-rich π -systems, *Beilstein J. Org. Chem.*, 2015, **11**, 2278–2288; (b) N. Ono, in *Organic Nitro Chemistry Series*, ed. H. Feuer, Wiley-VCH, New York, NY, 2001, pp. 126–128.

28 (a) G. S. Hammond, W. D. Emmons, C. O. Parker, B. M. Graybill, J. H. Waters and M. F. Hawthorne, Silver Salt Alkylation Reactions And The Preparation of Polynitroalkanes, *Tetrahedron*, 1963, **19**, 177–195; (b) S. A. Shevelev, L. I. Dalinger, V. M. Vinogradov and A. A. Fainzilberg, Alkylation of Alkali Metal Salts of Trinitromethane Under Interphase Catalysis Conditions In Solid Phase–Liquid Systems, *Bull. Acad. Sci. USSR, Div. Chem. Sci.*, 1990, **39**, 1647–1650.

29 (a) F.-Y. Chen, C.-L. Xuan, Q.-Q. Lu, L. Xiao, J.-Q. Yang, Y.-B. Hu, G.-P. Zhang, Y.-L. Wang, F.-Q. Zhao, G.-Z. Hao and W. Jiang, A review on the high energy oxidizer ammonium dinitramide: Its synthesis, thermal decomposition, hygroscopicity, and application in energetic materials, *Def. Technol.*, 2023, **19**, 163–195; (b) S. G. Zlotin, I. L. Dalinger, N. N. Makhova and V. A. Tartakovsky, Nitro compounds as the core structures of promising energetic materials and versatile reagents for organic synthesis, *Russ. Chem. Rev.*, 2020, **89**, 1–54; (c) X. Zhang, Y. Liu, F. Wang and X. Gong, A Theoretical Study on the Structure, Intramolecular Interactions, and Detonation Performance of Hydrazinium Dinitramide, *Chem.-Asian J.*, 2014, **9**, 229–236.

30 (a) M. Rahm and T. Brinck, Kinetic Stability and Propellant Performance of Green Energetic Materials, *Chem. – Eur. J.*, 2010, **16**, 6590–6600; (b) S. Alavi and D. L. Thompson, Decomposition pathways of dinitramic acid and the dinitramide ion, *J. Chem. Phys.*, 2003, **19**, 232–240.

31 (a) A. P. Häring and S. F. Kirsch, Synthesis and Chemistry of Organic Geminal Di- and Triazides, *Molecules*, 2015, **20**, 20042–20062; (b) B. D. Tsyrenova, P. S. Lempert and V. G. Nenajdenko, Di- and polyazides. Synthesis, chemical transformations and practical applications, *Russ. Chem. Rev.*, 2023, **92**, RCR5066.

32 J. Soto, M. Algarra and D. Peláez, Nitrene formation is the first step of the thermal and photochemical decomposition reactions of organic azides, *Phys. Chem. Chem. Phys.*, 2022, **24**, 5109–5115.

33 M. Shee and N. D. P. Singh, Chemical versatility of azide radical: journey from a transient species to synthetic accessibility in organic transformations, *Chem. Soc. Rev.*, 2022, **51**, 2255–2312.

34 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, and D. J. Fox, *Gaussian 16, Revision A.03*, Gaussian, Inc., Wallingford CT, 2016.

35 G. A. Zhurko, *Chemcraft*, Ivanovo, 2005.

