Trinitromethanide and Tricyanomethanide Salts Restricted to C, H, N, and O Atoms

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Technical Report

August 7, 1992

See Attached Report
Trinitromethanide and Tricyanomethanide Salts Restricted to C, H, N, and O Atoms

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ABSTRACT

Trinitromethane combined with oximes of cyclopentanone, cyclohexanone, and diphenylcyclopropenone and with melamine and two 1,3-dialkyl-2,4-dialkylimino-1,3-diazetidines to give simple trinitromethanide salt adducts. Tricyanomethane added to diphenylcyclopropenone oxime to give hydroxylamino-2,3-diphenylcyclopropenylum tricyanomethanide.

INTRODUCTION

Utilization of the energy rich trinitromethyl group in energetic molecules composed of C, H, N, and O atoms has been limited to a few ammonium trinitromethanide salts RNH₃⁺ −C(NO)₃ [1] and to molecules with covalent attachment of trinitromethyl to a carbon atom [2–5]. Examples of trinitromethyl attachment to an oxygen atom are unknown and examples of the rarely encountered covalent attachment to a nitrogen atom are found in tetrinitromethane and in arylazotrinitromethanes 1 in equilibrium mixtures with aryldiazonium trinitromethanides [6,7].

In a recent report on 2-nitropropan-2-ylbis-(p-methoxyphenyl)methylium trinitro-
tromethanide 2, a steric inhibition toward the C-C covalent isomer was proposed [8].

A predominance of ionic over covalent C-C bonding to the trinitromethyl group in an organic molecule was also shown in our elucidation of the ionic structure of guanidinium trinitromethanide 3 without detection of a C-C covalent isomer [9–11].

\[
\begin{align*}
\text{ArN} & \equiv \text{N} \quad \text{C(NO}_2\text{)}_3 \\
\text{ArN} & = \text{NC(NO}_2\text{)}_3 \\
(p-H_3\text{COC}_6\text{H}_4)_2\text{C} & + \text{C(CH}_3\text{)}_2\text{NO}_2 \\
\text{(H}_2\text{N})_3\text{C}^+ & -\text{C(NO}_2\text{)}_3
\end{align*}
\]

An interest in the discovery of additional molecules restricted to C, H, N, and O atoms with suppression of a covalent attachment of the trinitromethyl group to a carbon atom in favor of the dissociated isomeric trinitromethanide salt led to the present investigation. With recognition of the claim [2] that it does not yield isolable adducts with aldehydes and ketones other than formaldehyde, the addition of trinitromethane [HC(NO₂)₃, nitroform], pKₐ 0, [2] to ketones, ketoximes, and amidines was selected for initial investigations. In contrast with a scarceness of literature on the 1,2-addition of trinitromethane to carbonyl and azomethine linkages, Michael additions to give β-trinitromethyl derivatives is well established [2].

RESULTS AND DISCUSSION

Molecules structurally related to trinitromethylcyclopropene 4 and/or cyclopropenylum trinitromethanide 5 and their derivatives were sought for the energy to be derived from both trinitromethyl and the strained ring [12–14]. When an adduct from cyclopropen-
one and nitroform, obtained as an impure unstable yellow viscous oil tentatively identified as hydroxycyclopropenylium trinitromethanide 6 on the basis of its absorption at \( \lambda_{\text{max}} \) (ethanol) 350 nm (see below), decomposed on storage and warming, our attention shifted to similar adducts from ketoximes.

\[
\begin{align*}
\begin{array}{ccc}
\triangle C(NO_2)_3 & \Xbar & -C(NO_2)_3 \\
4 & 5 & 6 \ X = H \qquad & \ X = OH \\
11 & n = 4 & 12 \ n = 5
\end{array}
\end{align*}
\]

Simple adducts from nitroform and ketoximes 7, 9 were readily obtained as cyclopentanone and cyclohexanone oxime trinitromethanides 8a, 10a, eq(1), isolated as viscous yellow oils. In each the trinitromethanide anion was shown to be present by UV absorption: 8a (ethanol), \( \lambda_{\text{max}} \) 350 nm, \( \log \epsilon \) 4.20; 10a (ethanol), \( \lambda_{\text{max}} \) 350 nm, \( \log \epsilon \) 4.30; for comparison potassium trinitromethanide gave absorption for the anionic chromophore at \( \lambda_{\text{max}} \) (ethanol) 350 nm, \( \log \epsilon \) 4.31 (lit [15] \( \lambda_{\text{max}} \) (water) 350 nm, \( \log \epsilon \) 4.16). Insofar as covalent trinitromethane derivatives including nitroform, were transparent in this spectral region the absence (or minimal presence) of the covalent isomers 8b, 10b was revealed by both the absorption and the molar absorption coefficient values. Facile thermal and hydrolytic dissociations of the adducts 8a, 10a regenerated nitroform and the oximes 7, 9. The absence of Beckmann rearrangements of oximes 7, 9 to lactams 11, 12 attested to noninteraction between the oxime hydroxyl function and nitroform or the trinitromethanide anion.

\[
\begin{align*}
\begin{array}{ccc}
(\text{CH}_2)_n \ C = \text{NOH} + \text{HC(NO}_2)_3 & \Leftrightarrow & (\text{CH}_2)_n \ C = \text{N}^+\text{HOH} \ -C(NO_2)_3 \\
7 \ n = 4 & & 8a \ n = 4 \\
9 \ n = 5 & & 10a \ n = 5 \\
(\text{CH}_2)_n \ C(\text{NHOH})C(NO_2)_3 & \Leftrightarrow & (\text{CH}_2)_n \ C(NO_2)_3 \\
8b \ n = 4 & & 10b \ n = 5 \text{ eq (1)}
\end{array}
\end{align*}
\]
Somewhat more stable adducts from both nitroform and cyanoform [HC(CN)₃, tricyanomethane], pKₐ -5, [2] with the oxime of diphenylcyclopropenone were readily obtained as amorphous yellow solids identified as hydroxylaminodiphenylcyclopropenylum trinitromethanide 13 and tricyanomethanide 14 on the basis of elemental and spectroscopic analysis. The hydroxylaminodiphenylcyclopropenylum cation was detected by (a) absorption at λ_max 310 nm, log ε 4.07, for compound 13 (ethanol) and at λ_max 312 nm, log ε 4.13, for compound 14 (ethanol) in close agreement with absorption at λ_max 312 nm, log ε 4.12, for the known hydroxylaminodiphenylcyclopropenylum chloride 15 (ethanol) and (b) IR absorption at ν 1908 for compound 13 and ν 1912 for compound 14 in agreement with absorption at ν 1920 reported for the salt 15 [16]. The trinitromethanide anion was detected in compound 13 (ethanol) by absorption at λ_max 350 nm, log ε 4.15 (compare values above for potassium trinitromethanide). The tricyanomethanide anion was detected in compound 14 by IR absorption at ν 2160 [17]. Further confirmation of the structure for the salt 13 was found in its EI-MS fragmentation pattern, eq. (2) (see Experimental).

\[
\begin{align*}
&\text{13 } X = \text{C(NO}_2\text{)}_3 \\
&\text{14 } X = \text{C(CN)}_3 \\
&\text{15 } X = \text{Cl} \\
&\text{16 } X = \text{(C}_2\text{H}_5\text{)}_2\text{N}, \ Y = \text{C}_6\text{H}_5, \ Z = \text{C(NO}_2\text{)}_3 \\
&\text{17 } X = \text{Y = (C}_2\text{H}_5\text{)}_2\text{N}, \ Z = \text{C(NO}_2\text{)}_3 \\
&\text{18 } X = \text{(C}_2\text{H}_5\text{)}_2\text{N}, \ Y = \text{C}_6\text{H}_5, \ Z = \text{BF}_4 \\
&\text{19 } X = \text{Y = (C}_2\text{H}_5\text{)}_2\text{N}, \ Z = \text{ClO}_4
\end{align*}
\]
An attempt to obtain the trinitromethane salt 13 from the corresponding chloride 15 [16] in an ion exchange reaction with potassium trinitromethanide was unsuccessful. Similar attempts to obtain diethylaminodiphenylcyclopropenylium and trisdiethylaminocyclopropenylium trinitromethanides 16, 17 by treating the respective tetrafluoroborate 18 [18] and perchlorate 19 [19] with potassium trinitromethanide were also unsuccessful.

Although tetrachlorocyclopropene 20 reacted with a mixture of cyanamide and triethylamine at −30°C to give the zwitterionic product 21 [20] a replacement of cyanamide with nitramine [21] failed to give the zwitterion 22 and led instead to an intratable mixture, a result also obtained in reactions between tetrachlorocyclopropene 20 and either ethyl carbamate or hydrazine. An independent treatment with triethylamine brought about a dissociation of nitramine [22].

An extension of the investigation of guanidinium trinitromethanide 3 [9] led to
the discovery of similar amidinium trinitromethanides. Melamine and nitroform readily gave a 1:1 yellow solid ionic adduct identified as melaminium trinitromethanide 23 by elemental analysis and absorption at $\lambda_{\text{max}}$ (ethanol), 350 nm, log $\varepsilon$, 4.15. Although di- and triprotonation of melamine were known [23] attempts to obtain 1:2 and 1:3 adducts between melamine and nitroform were unsuccessful. In similar manner unstable ionic yellow viscous oil adducts (1:2) 26 (R = isopropyl) and 27 (R = cyclohexyl), $\lambda_{\text{max}}$ (ethanol) 350 nm, log $\varepsilon$ 4.15 and 4.24 respectively, were obtained from the 1,3-dialkyl-2,4-dialkylimino-1,3-diazetidines 24, 25 and nitroform.

$$\begin{align*}
24 & \quad R = (\text{CH}_3)_2\text{CH} \\
25 & \quad R = c-C_6\text{H}_{11} \\
26 & \quad R = (\text{CH}_3)_2\text{CH} \\
27 & \quad R = c-C_6\text{H}_{11}
\end{align*}$$

The thermal instability of all trinitromethanide salts reported here precludes their practical application as energetic materials.

**EXPERIMENTAL**

Melting and decomposition points were determined on a Mel-Temp II apparatus and are uncorrected. Elemental analyses were obtained from Midwest Microlab, Indianapolis, IN. Instruments included: Perkin Elmer 1600 series FT-IR, HP5985 (70 eV) GC/MS, Cary 17 UV Spectrometer. The preparation of potassium trinitromethanide from tetranitromethane [24], its conversion to nitroform [24], and reactions of nitro-
form were carried out in a hood. Potassium trinitromethanide was stored at 0°C; it should be handled as an unstable compound capable of decomposition on extended storage or warming [4]. Diphenylcyclopropenone was commercially available; its oxime was prepared as reported [16]. Melamine was commercially available. 1,3-Diisopropyl-2,4-bis(isopropylimino)-1,3-diazetidine 24 and 1,3-dicyclohexyl-2,4-bis(cyclohexylimino)-1,3-diazetidine 25 were prepared as reported [25]. Solvents were removed by rotary evaporation under reduced pressure. Analyses for nitrogen content were generally unsatisfactory and not reproducible. Similar results from attempted analysis for elemental nitrogen content in trinitromethyl derivatives were reported [4].

Nitroform

A suspension of potassium trinitromethanide (0.19 g, 1.0 mmol) and anhydrous ether (15 ml) in a 3-necked RB flask equipped with a calcium chloride guard tube and a nitrogen inlet was held at 0–5°C. A solution of hydrogen chloride in ether was added in drops with stirring until the yellow color disappeared. Nitrogen was bubbled through the suspension for 15 min to remove hydrogen chloride. The colorless nitroform solution at 0–5°C was transferred to another 3-necked flask by filtration.

Hydroxycyclopropenylium trinitromethanide 6

To a solution of nitroform (0.45 g, 3.0 mmol) in ether (25 ml), cyclopropenone [26] 0.16 g, 3.0 mmol, was added and stirred for 2 h at 0–5°C. Removal of ether left the salt 10 as a yellow viscous oil, (0.15 g, 75%). IR (neat): v 3028, 2984, 2900, 1590, 1305, 1097, 1073, 941, 837, 777, 626; UV (C<sub>2</sub>H<sub>5</sub>OH) λ<sub>max</sub> 350 nm, log ε 4.14. Instability precluded its further purification.
**Cyclopentanone oxime trinitromethanide 8a**

To a solution of nitroform (0.15 g, 1.0 mmol) in ether (15 ml) at 0–5°C, cyclopentanone oxime (0.1 g, 1.0 mmol) was added and stirred for 2 h. Evaporation of the solvent left the salt 8a as a viscous yellow liquid 0.22 g (88%); IR (neat): v 3441, 2890, 1592, 1487, 1404, 1272, 1158, 1066, 940, 787; UV (C₆H₅OH) λ_max 350 nm, log ε 4.20. Attempts at further purification were unsuccessful.

**Cyclohexanone oxime trinitromethanide 10a**

Treatment of cyclohexanone oxime (0.11 g, 1.0 mmol) with a solution of nitroform (0.15 g, 1.0 mmol) in ether (15 ml) at 0–5°C for 2 h gave the salt 7 as a yellow viscous liquid 0.24 g, (90%) after solvent was removed under vacuum at 20 °C. IR (neat): v 3421, 2945, 1591, 1545, 1484, 1282, 1158, 1061, 941, 787; UV (C₆H₅OH): λ_max 350 nm, log ε 4.30. Anal. calcd for C₇H₁₂N₄O₇: C, 31.82; H, 4.55. Found: C, 32.55; H, 4.57.

Water (20 ml) was added to the salt 10a (2.0 g, 7.5 mmol) and the mixture was stirred for 20 h at room temperature. It was extracted with dichloromethane (3 x 20 ml) and the combined extracts were washed with water (1 x 10 ml) and dried over anhydrous magnesium sulfate. Evaporation left cyclohexanone oxime (0.11 g, 13%) which melted at 89°C (lit [27] 87–88°C). A similar hydrolysis of the salt 8a gave cyclopentanone oxime.

**Hydroxylamino-2,3-diphenylcyclopropenylium trinitromethanide 13**

To a solution of nitroform (0.15 g, 1.0 mmol) in ether (15 ml), 2,3-diphenylcyclopropenone oxime (0.22 g, 1.0 mmol) was added at 0–5°C and stirred for 6 h. After iso-
lation, washing with cold pentane, and drying under a vacuum the salt 13 was obtained as a yellow solid, 0.32 g (85%), mp 75–76°C (dec); IR (KBr): ν 3221, 2976, 2814, 1908, 1597, 1542, 1277, 787, 734; UV (C<sub>6</sub>H<sub>6</sub>O): \( \lambda_{\text{max}} \) 350 nm, log ε 4.15. Anal. calcd for C<sub>16</sub>H<sub>2</sub>N<sub>4</sub>O: C, 51.61; H, 3.23. Found: C, 51.23; H, 3.36. EI-MS, \( m/z \) (%): 178 (88), 105 (96), 77 (100), 51 (77), 50 (44), 46 (42), 43 (56). Diphenylacetylene, mw 178, was known to give the fragments \( m/z \) 77, 51, and 50 [28].

**Hydroxylamino-2,3-diphenylcyclopropenylum tricyanomethanide 14**

Diphenylcyclopropenone oxime (0.11 g, 0.5 mmol) was added to a moist ether solution of cyanoform [29] obtained without isolation from potassium tricyanomethanide (0.07 g, 0.5 mmol) [30]. The mixture was stirred for 1 h at room temperature, filtered, washed several times with cold water, and dried in vacuum to give the salt 14 as a pale yellow solid, 0.1 g (65%), mp 132°C (dec); IR (KBr): ν 3086, 2852, 2160 (-C(CN)<sub>3</sub>, [17]), 1912 (hydroxylaminocyclopropenylum cation, [16]), 1597, 1569, 1448, 1496, 1083, 946, 763, 681, and 567. Anal. calcd for C<sub>19</sub>H<sub>12</sub>N<sub>4</sub>O: C, 73.08; H, 3.85. Found: C, 72.97; H, 4.05.

**Melaminium trinitromethanide 23**

Melamine (0.13 g, 1.0 mmol) was added to a stirred solution of nitroform (0.15 g, 1.0 mmol) in ether at 0–5°C. After stirring for 2 h, a yellow solid was isolated, washed with cold pentane, and dried in vacuum to give the salt 23, 0.22 g (78%), mp 155°C (dec); IR (KBr): ν 3124, 1683, 1489, 1421, 1275, 1177, 793, 733; UV (C<sub>6</sub>H<sub>6</sub>O): \( \lambda_{\text{max}} \) 350 nm, log ε 4.15. Anal. calcd for C<sub>6</sub>H<sub>7</sub>N<sub>6</sub>O: C, 17.33; H, 2.53; N, 45.49. Found: C, 17.80; H, 2.65; N, 45.10.
I,3-Dialkyl-2,4-bis(alkylimino)-1,3-diazetidinium bistrinitromethanides 26, 27

To a solution of nitroform (0.30 g, 2.0 mmol) in ether (20 ml), 1,3-diisopropyl-2,4-bis-(isopropylimino)-1,3-diazetidine 24 (0.26 g, 1.0 mmol) was added at 0–5°C and the mixture was stirred for 6 h. Evaporation of the solvent afforded the salt 26 as a yellow viscous oil, 0.5 g (90%); IR (neat): ν 1676, 1542, 1288, 742; UV (C₂H₅OH): λ_max 350 nm, log ε 4.15. A similar treatment of 1,3-dicyclohexyl-2,4-bis(cyclohexylimino)-1,3-diazetidine 25 (0.41 g, 1.0 mmol) with nitroform (0.30 g, 2.0 mmol) gave the salt 27 as a yellow viscous oil, 0.6 g (85%); IR (neat): ν 1651, 1540, 1262, 733; UV (C₂H₅OH): λ_max 350 nm, log ε 4.24. Instability precluded further purification.

ACKNOWLEDGMENT

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FOOTNOTES AND REFERENCES

1964, 10642c) reported tetraaminomethylmethane as its tetrakistrinitromethanide salt, C(CH₂NH₃⁺)₄(C(NO₂)₃)₄; (c) K. Torssell *Arkiv För Kemi*, 23, 1965, 537 reported anilinium trinitromethanide; and (d) J. H. Boyer and A. M. Krishnan prepared *tert*-butylammonium trinitromethanide, mp 155–156°C (dec), elemental analyses for C, H, and N in agreement with theoretical values, from iodotrinitromethane and *tert*-butylamine (unpublished results).


[10] Guanidine nitroform, (H₂N)₂C=NH•HC(NO₂)₃, was reported in the *Encyclopedia of Explosives and Related Items*, Vol 6, ed. B. T. Federoff and O. E. Sheffield,
Picatinny Arsenal, Dover, New Jersey, 1974, p G151, and Vol 8, ed. S. M. Kaye, 1978, p M79. A preparative procedure was not reported and the compound was inadequately characterized by m.p. data, later corrected, [9] and by elemental analysis for nitrogen only. These reports were omitted in Chemical Abstracts.


[13] In a personal communication P. C. Redfern and P. Politzer described a theoretical derivation for specific impulses (I_\text{s}) [14] with values for structures 4 and 5 surpassing the standard value for HMX \( \text{[CH}_2\text{N(NO}_2\text{)]}_4 \), by 9 and 15 percent respectively.


