

Coordination Chemistry of Proton. In Situ Synthesis and X-Ray Structural Analysis of 2-Hydroxyethanaminium Picrate, $\text{HOCH}_2\text{CH}_2\text{NH}_3^+ \cdot \text{C}_6\text{H}_2\text{N}_3\text{O}_7^-$

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2-Hydroxyethanaminium picrate, $\text{HOCH}_2\text{CH}_2\text{NH}_3^+ \cdot \text{C}_6\text{H}_2\text{N}_3\text{O}_7^-$, is prepared through a metathetical reaction of $\text{Mg}(\text{Pic})_2$ and $\text{HOCH}_2\text{CH}_2\text{NH}_2$ (2-aminoethanol, MEA) in ethanol through in situ generation of HPic. Orange yellow crystals ($\text{MEAH}^+\text{Pic}^-$) (mp 163 °C) of the "Salt" are produced along with the colorless $\text{Mg}(\text{CH}_2\text{NH}_2\text{CH}_2\text{O})_2$, even when MEA is less than 1:1 with respect to $\text{Mg}(\text{Pic})_2$. The crystals are monoclinic, space group $\text{P}2_1/\text{c}$, $a=11.797(4)$, $b=14.288(3)$, $c=7.124(2)$ Å, $\beta=97.63(2)^\circ$, $Z=4$, $D_{\text{cal}}=1.62$ g cm⁻³. Structural analysis from the 1617 reflection data ($R=0.059$, $R_w=0.056$) reveals that the picric acid proton is formally transferred to the MEA nitrogen (N–H, 0.968 Å) and is merely bonded to the phenoxide of Pic ($\text{H}\cdots\text{O}^-$, 1.951 Å) and weakly also to an oxygen of *o*-nitro group of the anion ($\text{H}\cdots\text{ONO}$, 2.379 Å). The transferred proton becomes an integral part of the cation MEAH^+ and completes the fourth covalent bond of the sp^3 N of MEA, and does not show any special affinity for the anionic oxygen of Pic; the N–H $\cdots\text{O}^-$ (142.8°) and N–H $\cdots\text{ONO}$ (141.6°) angles are closely comparable.

During in vitro chemical studies of alkali (M^+) and alkaline earth (M^{2+}) cations (general abbr. M^{2+}), which are being executed to throw light on the in vitro functions^{1,2} of Na^+ , K^+ , Mg^{2+} , and Ca^{2+} , we frequently use a foreign proton donor, HL, in the ML_x -ligand reaction mixtures, where L is an organic charge neutralizer and HL its parent acid. The aim is usually to stabilize L through homoconjugation, formation of $\text{L}\cdots(\text{HL})_n$, so that M^{2+} -ligand interaction is favored to enable the synthesis of the ML_x -ligand. Such studies with 1,10-phenanthroline,³ 2,2'-bipyridine,⁴ *N,N,N',N'*-tetramethylethylenediamine (TMED),⁵ and *N*-donor ligands in general, led frequently to the synthesis of HL-ligand by products which have constantly been of the 1:1 stoichiometry. In the meantime we developed interest in the study of interaction chemistry of the proton so that the trends in the interaction chemistry of small alkali and alkaline earth cations, Li^+ and Mg^{2+} , are understood more confidently and hence the H^+ -complexes such as the title compound.

For the ML_x -MEA systems, wherein MEA denotes 2-aminoethanol and L is 2-nitrophenolate (Onp), 2,4-dinitrophenolate (Dnp), or 2,4,6-trinitrophenolate (picrate; Pic), each ML_x -MEA reaction gave rise to a metathetical reaction (in the order $\text{M}^+ < \text{M}^{2+}$, $\text{Ca} < \text{Sr} < \text{Ba} < \text{Mg}$, $\text{Onp} < \text{Dnp} < \text{Pic}$) so that $\text{M}(\text{CH}_2\text{NH}_2\text{CH}_2\text{O})_2$ salt plus HL-MEA were obtained even when amount of MEA in the reaction was not more than the equivalent of ML_x .⁶ For $\text{Mg}(\text{Pic})_2$, as also evident from the foregoing efficiency orders, metathetical reaction and hence the production of HPic-MEA (instead of the Mg^{2+} -complex) was only noted. Because of our interest in the coordination chemistry of the proton, we undertook X-ray analysis of HPic-MEA which in

fact proved to be $\text{MEAH}^+ \text{Pic}^-$.

Experimental

Synthesis of HPic-MEA: $\text{Mg}(\text{Pic})_2$ (0.1 mmol) was dissolved with MEA (0.2 mmol) in 5 ml ethanol. Colorless $\text{Mg}(\text{CH}_2\text{NH}_2\text{CH}_2\text{O})_2$ salt, produced through the metathetical reaction, was filtered after ca. 30 min. The filtrate was subjected to crystallization of the title compound through slow evaporation at ca. 25 °C. Orange yellow single crystals (mp 163 °C) were filtered as produced (6–8 h). The title product can also be synthesized by a direct 1:1 reaction of HPic and MEA in ethanol but it is usually not as nicely crystalline. Elemental analysis of HPic-MEA: Found: C, 33.45; H, 3.38; N, 19.14%; Calcd: C, 33.10; H, 3.44; N, 19.31%.

X-Ray Structural Analysis: Cell parameters were determined by a least square fitting of the settings of 15 reflections ($\pm hkl$) measured on a Syntx diffractometer. Intensity data were collected in the θ - 2θ -mode ($2\theta \leq 50^\circ$) with graphite-monochromatized $\text{Mo K}\alpha$ radiation. The structure was solved by direct methods (MULTAN-82)⁷ and refined by full-matrix least squares to $R=0.059$, $R_w=[\sum(w|F_o|-|F_c|)^2/\sum w|F_o|^2]^{1/2}=0.056$ for 1617 ($2\theta \leq 50^\circ$) independent reflections with $F_o \geq 1.5 \sigma(F_o)$ [221 parameters, maximal shift esd=0.24]. Weights were given by the expression $w=k[\sigma^2(F_o)+0.0002 F_o^2]^{-1}$. Hydrogen atoms were located in a difference synthesis and their positions refined freely together with individual isotropic temperature factors in the last cycles of refinement. The non-hydrogen atoms were refined anisotropically. A final difference synthesis was effectively contourless (maximum 0.36 e Å⁻³).

Results

Molecular Structure: Structure of the molecule is depicted in the labelled Fig. 1 and a projection of the unit cell content perpendicular to [001] in Fig. 2. Final coordinates for the nonhydrogen and hydrogen atoms with equivalent isotropic temperature factors are shown in Table 1.⁸ $\text{C}_8\text{H}_{10}\text{O}_8\text{N}$ is monoclinic, space

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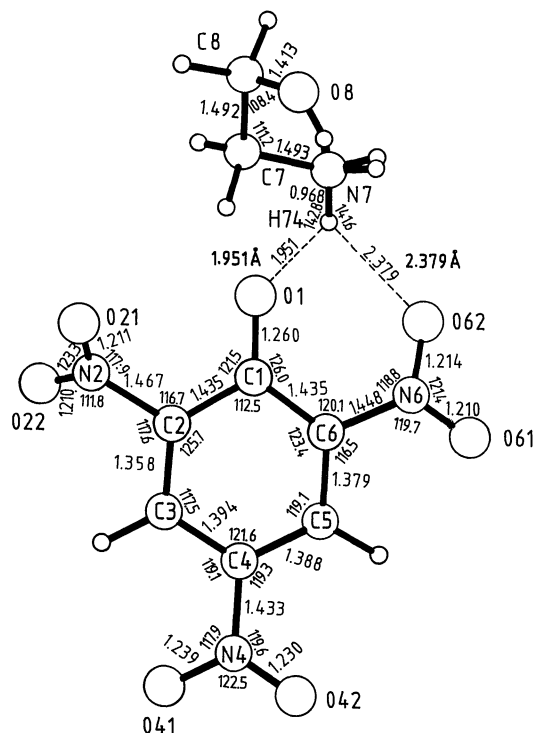


Fig. 1. Labelled diagram of the molecular structure of 2-hydroxyethanaminium picrate, $\text{HOCH}_2\text{CH}_2\text{-NH}_3^+\text{C}_6\text{H}_2\text{N}_3\text{O}_7^-$ with their bond distances and angles.

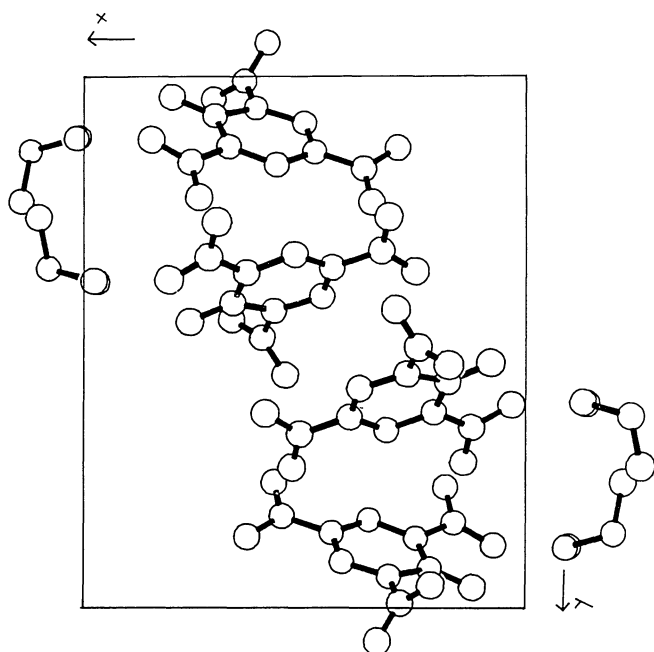


Fig. 2. Projection of the unit cell contents perpendicular to [001].

group $\text{P2}_1/\text{c}$ with $a=11.797(4)$, $b=14.288(3)$, $c=7.124(2)$ Å, $\beta=97.63(2)^\circ$, $Z=4$, $D_{\text{calc}}=1.62 \text{ g cm}^{-3}$.

Figure 1 shows that the HPic proton is rather formally transferred to the MEA nitrogen, $\text{N}_7(\text{N-H}$,

Table 1. Atom Coordinates with Isotropic Temperature Factors ($\text{\AA}^2$)

Atom	x/a	y/b	z/c	U_{eq}
O ₁	0.1659(2)	0.5387(1)	1.0825(3)	0.040(1)
C ₁	0.1659(2)	0.5743(2)	1.0537(4)	0.030(1)
C ₂	0.3626(2)	0.5596(2)	1.1825(4)	0.032(1)
N ₂	0.3507(2)	0.5091(2)	1.3581(4)	0.044(2)
O ₂₁	0.2898(2)	0.5430(2)	1.4641(4)	0.086(2)
O ₂₂	0.4052(3)	0.4380(2)	1.3934(4)	0.078(2)
C ₃	0.4692(2)	0.5876(2)	1.1547(4)	0.033(2)
C ₄	0.4791(2)	0.6417(2)	0.9948(4)	0.030(1)
N ₄	0.5900(2)	0.6732(2)	0.9616(3)	0.036(1)
O ₄₁	0.6747(2)	0.6344(2)	1.0500(3)	0.045(1)
O ₄₂	0.5980(2)	0.7362(2)	0.8466(3)	0.052(1)
C ₅	0.3841(2)	0.6659(2)	0.8677(4)	0.032(2)
C ₆	0.2782(2)	0.6321(2)	0.7598(4)	0.032(2)
N ₆	0.1823(2)	0.6608(2)	0.7598(4)	0.042(2)
O ₆₁	0.1923(2)	0.7268(2)	0.6568(4)	0.070(2)
O ₆₂	0.0928(2)	0.6183(2)	0.7533(4)	0.086(2)
N ₇	-0.0485(2)	0.6159(2)	1.1060(4)	0.038(1)
C ₇	-0.0241(2)	0.6138(2)	1.3170(5)	0.043(2)
C ₈	-0.1275(3)	0.6399(3)	1.4055(5)	0.047(2)
O ₈	-0.1524(2)	0.7351(2)	1.3655(3)	0.031(1)
H ₃	0.542(2)	0.570(2)	1.251(3)	0.047(8)
H ₅	0.386(3)	0.706(2)	0.745(3)	0.058(9)
H ₇₁	0.000(3)	0.545(1)	1.355(5)	0.070(11)
H ₇₂	0.044(2)	0.662(2)	1.357(5)	0.066(11)
H ₇₃	-0.096(2)	0.664(2)	1.052(4)	0.053(10)
H ₇₄	0.020(2)	0.605(3)	1.048(5)	0.094(14)
H ₇₅	-0.086(3)	0.675(2)	1.056(6)	0.088(14)
H ₈₁	-0.108(3)	0.633(2)	1.552(3)	0.066(11)
H ₈₂	-0.198(2)	0.597(2)	1.357(4)	0.041(8)
H ₈₃	-0.115(3)	0.773(3)	1.304(6)	0.058(8)

0.968 Å) which infact becomes the integral part of the base molecule yielding $+\text{H}_3\text{N}\cdot\text{CH}_2\text{CH}_2\text{OH}$ while completing the fourth tetrahedral valency of sp^3 hybridized N_7 atom. The C_7 and C_8 atoms of MEA are approximately sp^3 with $\text{N}_7\text{-C}_7\text{-C}_8$ (111.2°) and $\text{C}_7\text{-C}_8\text{-O}_8$ (108.4°) angles being tetrahedral values. None of the N_7 -protons undergoes intramolecular bonding with the MEA oxygen, O_8 . However, the H_{74} atom bonds with the Pic phenoxide atom, O_1 (1.951 Å) and rather weakly also with an *ortho*-nitrooxygen atom, O_{62} (2.379 Å). The $\text{N}_7\text{-H}_{74}\cdots\text{O}_1$ (142.8°) and $\text{N}_7\text{-H}_{74}\cdots\text{O}_{62}$ (141.6°) angles are, however, comparable.

The Pic moiety displays a detectable quinonoid structure wherein $\text{C}_1\text{-O}_1$ (1.260 Å) and $\text{C}_4\text{-N}_4$ (1.433 Å) bonds are short enough to show a double-bond character; the bonds adjoining them, viz. $\text{C}_1\text{-C}_2$ (1.435 Å) and $\text{C}_1\text{-C}_6$ (1.435 Å), and $\text{N}_4\text{-O}_{41}$ (1.239 Å) and $\text{N}_4\text{-O}_{42}$ (1.230 Å) are longer than the analogous bonds in the Pic moiety.

Discussion

Chemical Significance: Direct reaction of organic acids with organic nitrogenous bases have led to the recognition of detectable charge transfer towards the acid proton⁹ to a formal proton transfer on the base nitrogen.¹⁰ HPic and HDnp have been noted to form

H⁺-transfer complexes with aromatic diamines and been characterized spectrophotometrically.¹¹ X-Ray structural analysis of the product HPic-serotonin·H₂O¹² has revealed a transfer of the proton to the base (and a charge-transfer stacking of Pic moiety with the protonated base). We report herein the first such product, MEAH⁺-Pic⁻, obtained in situ employing a M(Pic)₂-MEA (M²⁺=Mg²⁺) reaction in a protic medium (ethanol); initial Mg²⁺L₂⁻-MEA complexation leads to Mg²⁺(L⁻)₂ pair loosening as well as "cationization" of those MEA protons which belongs to the NH₂ and OH functions. This is adequate enough for Pic⁻ to eliminate H⁺ from the complexed molecule of MEA; HPic so produced forms HPic-MEA with unreacted MEA.

A direct reaction of MEA with HOnp, HDnp and HPic leads to the synthesis of complexes with progressively increased melting points (115, 136, 163 °C, respectively). The present X-ray structural results indicate that the H⁺ charge transfer with MEA in this order, ultimately ends up in formal H⁺-transfer. A progressively increased melting point of the complexes in this order is, therefore, an index of accordingly an enhanced ionicity of the system. The salt MEAH⁺-Pic⁻, in a sense, is a H⁺-complex wherein Pic is the charge neutralizer and MEA a monodentate ligand. The H⁺...N(MEA) is, however, strong to the extent that interatomic penetration takes place yielding a formal H-N (MEA) covalent bond. The lengths of the H₇₄-O₁⁻ and H₇₄-O₆₂ bonds may be different (for structural and conformational reasons) but the proton, because of its hydrogen like nature, does not show any preference for any of the two oxygens; the N₇-H₇₄-O₁ (142.8°) and N₇-H₇₄-O₆₂ (141.6°) angles are practically equal. The concerned proton has not only shown an indifference to the MEA-hydroxyl oxygen (MEA moiety being not intramolecularly bonded) but also an exclusive preference for the neutral nitrogen N₇, over the anionic phenoxide, O₁. Also, the M(Pic)₂-MEA reactions become favored⁶ in the orders Cs to Li and Ba to Mg, so that the most efficient metathetical reaction for Mg(Pic)₂, and hence the most highly facilitated generation of HPic, is attributed to

the most strong Mg²⁺-N(MEA) interaction. Both the points collectively suggest that the hard M²⁺ ions are not only O-philic as usually described¹³⁻¹⁵ but also N-philic and that the preference of the M²⁺ (M⁺ or M²⁺) for the N-sites increases with the charge density (Charge/radius ratio) of the cation, unlike noted for O-sites.

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