Title: A Simple Complex on the Verge of Breakdown: Isolation of the Elusive Cyanofomate Ion

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Abstract:

Why does cyanide not react destructively with the proximal iron centre at the active site of 1-aminocyclopropane-1-carboxylic acid (ACC) oxidase, an enzyme central to the biosynthesis of ethylene in plants? It has long been postulated that the cyanofomate anion, [NCCO2]−, forms and then decomposes to CO2 and CN− during this process. We have now isolated and crystallographically characterized this elusive anion as its tetraphenylphosphonium salt. Theoretical calculations show that cyanofomate has a very weak C-C bond and that is thermodynamically stable only in low dielectric media. Solution stability studies have substantiated the latter result. We propose that cyanofomate shuttles the potentially toxic cyanide away from the low dielectric active site of ACC oxidase, before breaking down in the higher dielectric medium of the cell.

Main Text:

Nature manifests a myriad of efficient strategies to manage and manipulate reactive small molecules, such as water, hydrogen peroxide, carbon dioxide and oxygen, in the sensitive environs of enzyme active sites. The cyanofomate ion, [NCCO2]−, a complex of carbon dioxide and cyanide, has been implicated in the biosynthesis of ethylene (1, 2) but because it has eluded detection in condensed media the details of its decomposition have not been elucidated. Ethylene is an important plant hormone that regulates such essential biological processes as fruit ripening (3, 4). Its production is catalyzed by the enzymes 1-aminocyclopropane-1-carboxylic acid (ACC) synthase, which generates ACC, and ACC oxidase, which converts ACC to ethylene, cyanide and carbon dioxide. A simplified mechanism of the reaction catalysed by ACC oxidase is provided in Fig. 1. Mechanistic and structural studies strongly support binding of ACC and oxygen to the iron centre located in the active site of ACC oxidase. Sequential single-electron transfer from the co-factor, ascorbate, with elimination of water generates an iron(IV)-oxo species, followed by ring-opening of bound ACC to eliminate ethylene and cyanofomate, which then decomposes to CN− and CO2. It is not definitively known whether this decomposition occurs...
at the active site or elsewhere (5). Bicarbonate has been identified as an essential mediator for proton transfer in the process (6-8).

Figure 1: Simplified reaction scheme for the generation of ethylene, cyanide and carbon dioxide from ACC, via cyanoformate, at the active site of ACC oxidase.

It remains unclear as to how the iron-containing active site of ACC oxidase is protected from the cyanide by-product, a known deactivating agent of iron-containing enzymes in general and Fe(III) in particular (9). Although the hydrogen cyanide is ultimately metabolised, primarily by β-cyanoalanine synthase to produce the amino acid β-cyanoalanine (10), coordination to the iron center of ACC oxidase would seem likely beforehand. A recent publication by Dilley et al. (11) revealed that ACC oxidase is activated at cyanide concentrations between 0.1 to 1 mM whereas above 1 mM the cyanide becomes inhibitory. These authors propose that cyanide is bound as a ligand to Fe(II) in ACC oxidase, which then assists ACC binding. It should be noted, vis-à-vis our later discussion, that in most studies cyanide levels are typically determined from the bulk plant material (12, 13) rather than intra- or extra-cellularly. How then are the higher levels of cyanide removed from the active site of ACC oxidase for safe metabolism? We believe that the generation of cyanoformate, a simple coordination complex of cyanide and CO₂, plays a vital role in this regard.

Surprisingly, much of the fundamental coordination chemistry of CO₂, particularly with regards to small ligands and their corresponding reactivity, has yet to be explored. For example, little is known of the reactions between carbon dioxide and halides or pseudo-halides (i.e., OH, CN, OCN, SCN, N₃,...), although this gap can be attributed, at least partially, to the instability of these donor-acceptor complexes. There are only two well-characterised families of halide/pseudo-halide complexes of CO₂, the more common being bicarbonates, species that form numerous materials stabilized through extensive hydrogen bonding (14). The only other known example is fluorocarbonate, [FCO₂]⁻. This deceptively simple anion, isoelectronic with [NO₃]⁻,
might superficially be predicted to be stable, but it has only recently been prepared, and the fluoride affinity of carbon dioxide was calculated to be $-111$ kJ mol$^{-1}$, which “appears to be the lower limit for a reaction” (15).

We began our investigation into the possibility of isolating cyanoformate with a preliminary computational study. Table S6 of the supplementary material includes a full list of the calculated energies for the association reaction of CO$_2$ and CN$^-$. At all of the levels of theory employed the formation of cyanoformate was found to be thermodynamically favoured in vacuo. However, the binding energy of CO$_2$ was even smaller than that of the aforementioned fluorocarbonate ion. With the computations indicating that a thermodynamically stable, albeit very fragile, species could exist, we set out to try and isolate a derivative. We had recently shown that higher order zincates could be stabilized and isolated from ion rich media using tetraphenylphosphonium halide salts (16). Using a similar approach, we exposed a concentrated solution of [PPh$_4$]CN to an atmosphere of CO$_2$, and colorless crystals immediately precipitated. The solvent was decanted and the crystals were isolated and characterized using crystallographic and spectroscopic techniques. Solid state NMR of both singly-labelled [PPh$_4$][NC$_{13}$CO$_2$] and doubly-labelled [PPh$_4$][NC$_{13}$C$_{13}$CO$_2$] isotopologues confirmed formation of the complex with $\delta$(CO$_2$) = 144.4 ppm, consistent with the chemical shift observed for the fluorocarbonate ion, ca. 142 ppm (15). The bonding between the cyano and carbonyl groups was also confirmed by direct observation of one bond coupling ($^{1}J_{13C-13C}$ = 61(±4) Hz) between the $^{13}$C-labelled carboxyl and $^{13}$C-labelled cyano fragments, based on the resolved $^{13}$CO$_2$ signal in the spectrum of the latter compound (see Fig. S19).

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**Figure 2**: Solid state structure of [PPh$_4$][NC$_{13}$CO$_2$]. Thermal ellipsoids are drawn at the 50% probability level. Unique atoms of the cation and all atoms of the anion are labelled. The anion comprises only three unique atoms, carbon and oxygen atoms with site occupancies of 0.50 and nitrogen with a site occupancy of 0.25. The remainder of the anion is generated by symmetry. To the right of the arrow the anion has been deconvoluted into its two overlapping components, both of which have 50% occupancy in the crystal structure.
Figure 2 shows the structure of [PPh₄][NCCO₂] as determined by single crystal x-ray diffraction (see Figs. S32 and S33, Tables S11-S17 and the accompanying supplementary discussion for further details). Isolation of the fragile anion in the solid state was likely aided by the crystal lattice stabilization of the salt. The C-C bond in the anion (1.480(9) Å) is equal to that reported by Drück et al. (17) in the structure of cyanoformamide, NCC(O)NH₂. The cyanide C-N bond length (1.056(12) Å) proved to be significantly shorter than that in cyanoformamide (1.141(8) Å). Investigation of bond lengths reported in the Cambridge Structural Database (CSD) (18) showed that the observed cyanide bond length is reasonable, if relatively short, suggesting a presence of a strong CN bond with only a weak C-C interaction to the CO₂ portion of the adduct. The strengthening of the CN interaction in the cyanide ligand upon coordination to electron deficient sites has previously been discussed in the literature (19).

The packing of the ions in the structure of [PPh₄][NCCO₂] in the tetragonal space group I-4, is similar to that observed in many other simple [PPh₄]X salts, where X = Br, I, SCN, OCN, N(CN)₂, etc. (20-24). Viewed down the Z axis, the cations and anions form discrete stacked columns. Because of the large size of the cations, the anions can be thought of as occupying the channels formed between the stacked columns of cations. There are few inter-ion interactions in the structure and those that are present are generally long and presumably weak. Only the interaction C(2)-H(2)...O(1)' is characterised by a separation of less than 2.5 Å, H(2)...O(1)' = 2.37 Å. The lack of strong interactions between the ions likely results in the relatively free rotation of the anion about the S₄ rotation axis in the cavity produced by the parallel stacking of the cations down the Z axis.

The theoretically predicted metrical parameters of [NCCO₂]⁻ show very little variation with respect to the employed method and are in good overall agreement with the results from the x-ray structural study, although some small systematic deviations do exist; numeric data are presented in Table S4. Heterolytic splitting of the C-C bond in the anion and subsequent analysis of the bonding energy contributions (electrostatic, orbital interactions and Pauli repulsion) using the Ziegler-Rauk-Morokuma energy decomposition procedure (25-27) show that the linear cyanide and bent carbon dioxide fragments interact predominately (ca. 90%) via a σ-type donor-acceptor interaction (28). Furthermore, the results show that the fragile nature of the adduct arises not so much because of the intrinsic characteristics of the C-C bond but because of the significant energy release (ca. 200 kJ mol⁻¹) associated with linearization of the CO₂ fragment (29). The morphologies of the key frontier orbitals as well as the results from the analysis of the electron localisation function of cyanoformate are both consistent with the view of cyanide acting as a two electron donor to carbon dioxide (see Figs. S29 and S30 and the accompanying supplementary discussion for full details). Hence, the C-C interaction in [NCCO₂]⁻ is similar to the bonding in adducts of CO₂ with N-heterocyclic carbenes (30-33), though, to our knowledge, the latter systems have not been subjected to a thorough theoretical treatment.

Ionic carboxylates exhibit two characteristic C-O stretching bands in their IR spectra. The higher energy asymmetric band (νasCO₂) typically appears as a very strong and broad band, whereas the lower energy band (νsCO₂) is usually less intense. The spectra collected on samples of [PPh₄][NCCO₂], Figs. S3-S5, exhibit both of these expected signals. Consistent with the theoretical predictions for the IR and Raman bands (Tables S8-S10), the cyano stretch (νCN) is not observed in the infrared spectrum, but is clearly visible in the Raman (2196 cm⁻¹, Fig. S7). This frequency, when compared to that of other cyano groups, such as in acetonitrile (2253 cm⁻¹),
Fig. S6) and [PPh₄]CN (2069 cm⁻¹, Fig. S10), supports the notion of a strong CN interaction in [NCCO₂]. We also prepared ¹³C-labelled samples to confirm our assignments and this data is included in the supplementary material.

Figure 3: Electrostatic potentials (atomic units) superimposed on the total electron density isosurface (0.01 a.u.) of (A) CN⁻ and (B) [NCCO₂]⁻ at the ωB97X-D/aug-cc-pVTZ level of theory.

To visualize the effect of cyanide coordinating to CO₂, we have superimposed maps of the electrostatic potential on the total electron density of the reactant and product ions (Fig. 3). These plots aid in the assessment of reactivity, in particular prediction of potential sites of attack for nucleophiles and electrophiles. In CN⁻, the surface around the carbon atom is electron poor compared to the nitrogen center, but both atoms have electron rich areas directly on the bond axis, consistent with the two possible coordination modes for this anion. The electrostatic potential of [NCCO₂]⁻ shows that it is now the oxygen centers that are electron rich and therefore the most favored sites for electrophilic interactions. This suggests that hydrogen bond donors and acidic metals will be attracted more favourably to the oxygen atoms, rather than to the cyanide portion of the [NCCO₂]⁻ anion. The delicate interplay of nucleophilic and electrophilic interactions is very important in supporting the many biological reactions involving CO₂ (34) and the chemistry of the cyanoformate anion is no exception in this regard.

We have also calculated the thermodynamics for the formation of cyanoformate as a function of various solvents at different density functional levels of theory (see Figs. 4 and S31, Table S7 and the accompanying supplementary discussion for full details). The results suggest that [NCCO₂]⁻ is thermodynamically stable only in non-polar solvents (toluene); in high dielectric media the anion becomes thermodynamically unstable with respect to fragmentation to CN⁻ and CO₂ (Fig. 4). However, the anion is kinetically stabilized, albeit not by much, in all media as breaking the C-C bond has an activation barrier of approximately 40 kJ mol⁻¹ at these levels of theory. Although many enzymatic processes occur in bulk water, the local dielectric constant at the active site of an enzyme is typically low (7, 35), in the range of 4-20, as compared to 78 for water (36, 37). In the cell, at the low dielectric active site of the ACC oxidase enzyme, the cyanide ions could remain complexed with CO₂ in the form of [NCCO₂]⁻, protecting the iron...
centre from attack. However, as the cyanoformate anion migrates out to the bulk aqueous solution of the cytoplasm, the thermodynamics are modified, and the complex decomposes to the observed products, namely hydrogen cyanide and carbon dioxide.

Figure 4: Calculated reaction profile for the formation of \([\text{NCCO}_2^-]\) from \(\text{CN}^-\) and \(\text{CO}_2\) at the \(\omega\text{B97X-D/aug-cc-pVTZ}\) level with different solvent models.

The pivotal role assigned to the cyanoformate anion, based on its calculated (in)stability in solutions of different dielectric constant, was bolstered by carrying out further decomposition experiments (see the supplementary discussion for full details). Because of the insolubility of \([\text{PPh}_4][\text{NCCO}_2]\) in many traditional solvents, these studies were performed on tetrabutylammonium cyanoformate, \([\text{Bu}_4\text{N}][\text{NCCO}_2]\), generated \textit{in situ} from the cyanide precursor, in the ionic liquid (IL) trihexyl(tetradecyl)phosphonium bis(trifluoromethylsulfonyl)imide, \([\text{P}_6\text{6614}][\text{TFSI}]\). An aliquot of the \([\text{Bu}_4\text{N}][\text{NCCO}_2]/\text{IL}\) solution was added to the scrupulously dried solvent being studied and the disappearance of the \(\nu\text{CO}_2\) peak of cyanoformate (1689 cm\(^{-1}\)) monitored using real time Fourier transform (FT) IR spectroscopy. Half-lives were determined from linear regressions of plots of \(\ln(P/P_0)\), the ratio of peak height (P) to initial peak height (P\(_0\)), versus time (t). Solution half-lives for cyanoformate of 110 minutes, 55 minutes and 17 minutes were calculated for toluene, tetrahydrofuran and acetonitrile, respectively. As predicted by theory, the stability of the cyanoformate anion in solution rapidly decreased with increasing dielectric constant of the solution.

Although large scale experimental studies of the hydrolysis of cyanoformate were not feasible because of the possible release of HCN, the theoretical behaviour of the cyanoformate anion in aqueous environments was investigated in more detail by performing geometry optimizations (\(\omega\text{B97X-D/aug-cc-pVTZ}\) level) that included explicit H\(_2\)O molecules in addition to the bulk solvent (see the supplementary discussion for full details). In all cases, the calculations minimized to hydrogen bonded structures (O···H and/or N···H) with no indication of spontaneous hydrolysis. Although a concerted hydrolysis of cyanoformate in water thus seems unlikely, a sequential pathway can be envisioned. As shown computationally, cyanoformate readily
dissociates into \( \text{CN}^- \) and \( \text{CO}_2 \) in high dielectric media and the cyanide anion is an excellent nucleophile that can easily abstract a proton from water (\( pK_a \) of HCN = 9.24). The hydroxide ions generated can, in turn, attack the electrophilic carbon of \( \text{CO}_2 \) to yield the bicarbonate anion. Thus, in an aqueous environment containing dissolved \( \text{CO}_2 \), the addition of cyanide will lead to the formation of bicarbonate. This is a possible explanation for the observed activation of ACC oxidase by cyanide (11), since bicarbonate is an important participant in the catalytic cycle of the enzyme (6-8).

Our experimental work has clearly shown that cyanoformate is readily converted to bicarbonate (and HCN) in the presence of even a trace of water. Early attempts to isolate the cyanoformate salt often gave crystals of tetr phenylphosphonium bicarbonate, whose structure has been confirmed by crystallographic analysis. In our \( ^{13}\text{C} \) NMR study, the presence of a peak at 162.3 ppm is certainly attributable to bicarbonate, as the chemical shift correlates well with that reported for other such salts (38). That the bicarbonate originates from the added \( \text{CO}_2 \) is supported by the large size of this peak relative to others in the same spectrum, as \( ^{13}\text{CO}_2 \) was used in the original synthesis. We also probed the decomposition of the synthesized cyanoformate salts in a protic solvent by monitoring the \( \nu_{\text{as CO}_2} \) stretch of the cyanoformate anion (1689 cm\(^{-1}\)) with real time FTIR as 1-propanol was added to a dilute [Bu\(_4\text{N}\)][O\(_2\text{CCN}\)]/IL solution. This peak rapidly diminished during the addition while concomitantly a peak corresponding to bicarbonate appeared at 1652 cm\(^{-1}\) (39). Over the course of only two minutes, \( \nu_{\text{as CO}_2} \) for cyanoformate disappeared completely, while the bicarbonate peak reached a steady absorbance. All of the observed experimental results are entirely consistent with the computationally based proposal of a stepwise process for the decomposition and hydrolysis of the cyanoformate anion.

The stability of cyanoformate, a simple anion of \( \text{CO}_2 \) coordinated to \( \text{CN}^- \), has been shown to be dependent on the dielectric constant of its local environment. We propose that this property allows it to shuttle toxic \( \text{CN}^- \) away from the low dielectric active site of the enzyme ACC oxidase before its decomposition in higher dielectric media. In broader terms, the ability to manipulate solution stability of otherwise unstable or transient species, through changes to the dielectric constant, should find other important applications.

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