Cyanides, Isocyanides, and Hydrides of Zn, Cd and Hg from Metal Atom and HCN Reactions: Matrix Infrared Spectra and Electronic Structure Calculations

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Zinc and cadmium atoms from laser ablation of the metals and mercury atoms ablated from a dental amalgam target react with HCN in excess argon during deposition at 5 K to form the MCN and MNC molecules and CN radicals. UV irradiation decreases the higher energy ZnNC isomer in favor of the lower energy ZnCN product. Cadmium and mercury atoms produce analogous MCN primary molecules. Laser ablation of metals also produces plume radiation which initiates H-atom detachment from HCN. The freed H atom can add to CN radical to produce the HNC isomer. The argon matrix also traps the higher energy but more intensely absorbing isocyanide molecules. Further reactions with H atoms generate HMCN and HMNC hydrides, which can be observed by virtue of their C–N stretches and intense M–H stretches. Computational modeling of IR spectra and relative energies guides the identification of reaction products by providing generally reliable frequency differences within the Zn, Cd and Hg family of products, and estimating isotopic shifts using 13C and 15N isotopic substitution for comparison with experimental data.

1. Introduction

For investigation of novel cyanide and isocyanide molecules with infrared (IR) spectroscopy, we have produced by laser ablation Group 2, 3, 4, 7, 8 and 13 metal atoms as well as the Ce, Nd, Th, and U atoms. We have observed their reactions with cyanogen and U with hydrogen cyanide, and Zn with oxygen and hydrogen mixtures.1–20 Cyanogen generally reacts with metal atoms to form dicyanides or even tetracyanides.10–19 Our work with the Group 2 metal atoms and HCN or with (CN)2 provided IR spectra for the MNC isocyanides with no obvious amount of the cyanide isomer MCN.18 The laser ablated Zn atom reaction with (CN)2 in excess argon during condensation at 5 K produced ZnNC with an intense N–C stretching frequency of 2074.6 cm⁻¹; the observed effects of 13C and 15N substitution were consistent with the isocyanide structure.18 The ZnNC isomer could also be produced by UV irradiation of Zn atoms and (CN)2 co-deposited in solid argon.18 The N–C stretch for the analogous CdNC molecule was observed at 2069.7 cm⁻¹ in experiments reacting Cd and (CN)2,20 and the analogous higher energy HgNC molecule is produced here from HCN and Hg ablated from a dental amalgam target.9 This novel use of a dental amalgam ablation target adds more to the known chemistry for mercury.

The Largo group has performed a theoretical investigation of the structure, bonding, and isomerization barriers for the first-row transition metal cyanides and isocyanides.21,22 Of particular interest to us, they have predicted a modest dissociation energy for ZnNC to Zn + CN (ca. 50 kcal/mol) and a low isomerization barrier for ZnNC to Zn-CN (ca 5 kcal/mol) which are supported by the trends reported here.

Infrared detection of the weakly absorbing cyanogen species is more challenging, which is why we used here the more reactive HCN precursor. A previous Microwave spectroscopic investigation revealed that ground state ZnCN was formed from a mixture of zinc vapor and cyanogen gas in a DC discharge, and provided its bond length values.23 Later work using Zn(CH3)2 as a source of zinc and hydrogen atoms with cyanogen in the discharge produced a related molecule, HZnCN, which was also identified from its microwave spectrum,24 and the hydrde absorptions here increase on annealing to allow diffusion and reactions of H atoms first trapped in this matrix isolation investigation. The use of HCN as the reagent provides H atoms to react with cyanides and isocyanides to form their monohydrides.

Reported calculations provide helpful information on experimentally known and hypothesized (H, M, C, N) systems. For example, vibrational frequencies, energies, and structures are obtained directly. Andrews and Cho18 explored the reaction of Zn with NCCN and employed a density functional (B3LYP)
method and a coupled-cluster expansion CCSD(T) method with a large Dunning basis (aug-cc-pVTZ) to describe isomers of general form (C,N)Zn\textsubscript{n} (C,N). They reported the impact of \textsuperscript{13}C isotopic substitutions on vibrational frequencies, and for the CCSD and DFT calculations, supplied the vibrational intensities. The latter calculation showed that the C–N stretch is much more intensely absorbing for Zn–N–C than for Zn–C–N, while ZnCN is more stable than ZnNC by 20.7 kJ/mol at CCSD(T) level.

Here follows the complementary investigation of Zn, Cd and Hg atom reactions with HCN, which are more productive than the reactions with NCCN studied earlier\textsuperscript{[18,20]} Additional information was obtained with the H\textsuperscript{13}CN reagent and the resulting isotopic shifts and isotopic frequency ratios became useful diagnostics. Our goal using the matrix isolation technique was to produce the zinc, cadmium, and mercury cyanides MCN and isocyanides MNC in these metal atom reactions with HCN, and to measure their argon matrix infrared spectra. As anticipated from previous work we observed a 5 fold-stronger signal for ZnNC produced from the HCN reagent than for the Zn reaction with cyanogen\textsuperscript{[18]}. We have found HCN to be more reactive than NCCN, which gives us a much better possibility for observing the ground state ZnCN isomer with the lower IR intensity for this essentially covalent ZnC=N bond vibration. In addition, the easy production of H atoms from HCN allows formation of the related HMCN and HMNC monohydride molecules. In the final analysis the use of two different CN bearing precursors to make the same new molecules provides a better understanding of their spectra and quite different bonding properties. This gives us the unique opportunity to investigate these new (Ar\textsubscript{n})\textsuperscript{+}CN\textsuperscript{+} complexes where \(n\) could be 1,2,3,4, for example.

**Experimental Methods**

Laser-ablated Zn and Cd and Hg atoms were reacted with the HCN and H\textsuperscript{13}CN molecules in argon host gas during their deposition at 5 K using a closed-cycle helium refrigerator (Sumitomo Heavy Industries, RDK-205D) inside of a self-made vacuum chamber, which has been described in detail previously\textsuperscript{[9]}. Hydrogen cyanides were synthesized by reacting KCN with dilute H\textsubscript{2}SO\textsubscript{4} and drying the HCN gas by trap to trap distillation. The H\textsuperscript{13}CN was made from K\textsuperscript{13}CN (99\%) in the same way. The 1064 nm fundamental of a Nd:YAG laser (10 Hz repetition rate with 10 ns pulse width), and a pulse energy up to 55 mJ) was focused onto different rotating metal targets. An amalgam previously in common use for dentistry served as targets for Hg. This amalgam, which was composed of about 50\% Hg and 50\% of the more noble Cu, Ag and Sn metals by weight, was pressed into a disc of about 13 mm diameter and 4 mm thick, and glued onto a sample holder for rotation (picture in supporting information, SI). This method produced ample Hg atoms for our work\textsuperscript{[9,17]} Infrared (IR) spectra were recorded on a FTIR vacuum spectrometer (Bruker Vertex 70v) equipped with a transfer optic and Mid-IR MCT detector (4000–450 cm\textsuperscript{-1}). Matrix samples were annealed at different temperatures and cooled back to 5 K, and selected samples were irradiated in the ultraviolet (UV) by a medium pressure mercury arc street lamp (175 W)\textsuperscript{[21–24]} Observed and calculated vibrational frequencies are reported here in cm\textsuperscript{-1} units.

**Computational Methods**

We conducted density functional, and coupled-cluster calculations primarily with Gaussian 16 software\textsuperscript{[25]} Frequencies, absorption intensities, bond lengths, and energies were obtained with the B3LYP hybrid density functional\textsuperscript{[27]} using the aug-cc-pVTZ basis set\textsuperscript{[20]} for C, N, and Zn and the aug-cc-pVTZ-pp\textsuperscript{[29]} basis for Cd and Hg. The structures obtained by geometry optimization were verified to be relative minima in energy by vibrational analysis. The calculated harmonic frequencies are reported here unscaled. Such DFT calculations predict vibrational frequencies 2 to 10 cm\textsuperscript{-1} higher than observed in the argon matrix for metal compounds\textsuperscript{[10]}. These comparisons between model and data suffer from (1) the accuracy of the theoretical method involved, (2) the usually higher value of harmonic (calculated) relative to anharmonic (observed) frequencies, and (3) the common matrix shift to lower frequencies owing to the generally weak dispersive interaction of the matrix host with the guest molecule.

Separate calculations were done with CCSD, to provide absorption intensities not available in the Gaussian CCSD(T) implementation. Zero-point energies derived from vibrational calculations are included in the energy reports. Additional calculations were done with the Molpro system\textsuperscript{[30]} to address an issue that came up in the modeling of the metal cyano systems.

**2. Zinc Studies**

**2.1. Spectroscopic Results and Discussion (Zinc Systems)**

Figure 1 displays infrared spectra for zinc and hydrogen cyanide reaction products. Many vibrational bands are labeled; red numbers identify new data. Black numbers in the spectra identify absorption bands observed in our previous work using laser ablated zinc and the (CN\textsubscript{2})\textsuperscript{+} reagent\textsuperscript{[18]}. Spectra begin with (a) the deposition of 10 mbar of argon gas from the 1.1 L vacuum manifold containing 1% HCN\textsuperscript{[13]} in argon and continue through stages of processing scans (b-f).

**2.2. Non-Metal Systems**

Matrix spectra of HCN and its oligomers were first measured with prism, filter, and grating spectrometers\textsuperscript{[31]}. King and Nixon report signals for HCN in argon at 3303.3 (H–CN stretch), 2093 (C–N stretch) and 720.2 cm\textsuperscript{-1} (bend)\textsuperscript{[31]} The band at 2027.6 cm\textsuperscript{-1} was assigned to the HNC isomer by Milligan and Jacox, who produced the signal by UV photolysis of CH\textsubscript{3}N\textsubscript{2} in solid argon and then by the vacuum ultraviolet photolysis of HCN during deposition in excess Ar\textsubscript{2}\textsuperscript{[32]} A signal at 2046 (in an argon matrix using an older instrument) was attributed to the stretch for CN radical by Milligan and Jacox\textsuperscript{[33]}, our recent value using FTIR is 2044.4 cm\textsuperscript{-1}.

Modern FTIR instruments provide inherently more accurate values compared with the earlier numbers\textsuperscript{[31–33]} especially toward higher frequencies for example, we measured the C–H stretch of HCN at 3305.5 cm\textsuperscript{-1}, (cf. 3303.3 for King and Nixon\textsuperscript{[31]}). The H–C–N bending mode at 720.9 cm\textsuperscript{-1}, (cf. 720.9\textsuperscript{[13]}) and its overtone at 1426.4 cm\textsuperscript{-1} were found in our 0.2% (1/500) HCN sample in argon. Abbate and Moore observed the bands at
We consider that our values are accurate within about $0.2 \text{ cm}^{-1}$ but there are a number of other factors which contribute to uncertainty such as guest concentration in the matrix, band width, substrate window temperature, and the resulting rate of sample deposition. Another consideration is local matrix site splittings: linear molecules such as CNCN and OCO have antisymmetric fundamentals that are split by the matrix. These will be shown later in Supporting Information (SI), and the major matrix site splitting bands for CNCN are given in Table 1.

Figure 1 shows our spectrum of HCN in argon first then more of the same sample co-deposited with Nd-YAG fundamental laser ablated Zn. For spectrum a, the stronger sharp band on the right side at 2093.1 cm$^{-1}$ is due to isolated HCN,$^{[21,33]}$ and the strong broad band at 2103 cm$^{-1}$ arises from HCN dimers and higher polymers labeled (HCN)$_x$. The remaining spectra in Figure 1(b) through (f) illustrate absorptions for a sequence of zinc reaction products in the 1% HCN sample in argon. The spectrum (b) followed the co-deposition of 120 mbar more of argon with HCN from a constant 1.1 Liter volume, and laser ablated zinc using 50% of maximum laser energy for 2 h. Cyanogen provides the band at 2153.5 cm$^{-1}$ presumably from the dimerization of CN radicals. The sharp band at 2054.0 cm$^{-1}$ shown in spectrum (b) of Figure 1 corresponds to a feature found in our studies of transition metals, uranium, and thorium with cyanogen.$^{[10–14]}$ This is due to the CNCN photoisomer of cyanogen.$^{[18]}$ In the current work with HCN, it appeared at 2053.7 cm$^{-1}$. Such small differences are a part of the matrix isolation experiments. The sharp band at 2954.0 on (b) in Figure 1 is favored by metals with cyanogen, and it is assigned to the CNCN photoisomer of NCCN.$^{[18]}$ Both matrix sites at 2054.0 and 2057.7 are observed with the larger product yield in Figure 1. More examples of matrix site splittings are found in our ref. $^{[18]}$.

2.2.1. Appearance of ZnCN, ZnNC, HZnCN and HZnNC

The CN radical$^{[1,2,18]}$ presents a strong absorption at 2044.4 cm$^{-1}$ on Ar/Zn/HCN sample deposition (Figure 1, spectrum b). Upon annealing to 20 K (spectrum c) the CN signal decreases by about 90% while the 2153.5 cm$^{-1}$ (CN)$_2$ band increases only slightly because other, namely metal atom, reactions with CN, are more favorable than its dimerization. Weak bands are found in Figure 1 spectrum (b) at 2196.6 and 2193.2 cm$^{-1}$, in excellent agreement with the site splittings first reported for the dizinc molecule NCZnZnCN.$^{[18]}$ Important newly discovered (red-type)
bands lie at 2162.2 and 2184.7 cm\(^{-1}\) in Figures 1 and 2 where they are labeled with the above numbers and the latter is also labeled HZnCN since there is room in the spectrum for this. The 2184.7 band increases slightly on annealing to 20 K (shown in spectrum c) while the 2162.2 band decreases slightly: These bands are only 22.5 cm\(^{-1}\) apart, which suggests a common CN group, with stretching frequency affected slightly differently by the Zn- or HZn- substituents. The 2162.2 band assigned here to ZnCN, which readily adds H upon annealing to form HZnCN. The very strong sharp 2074.6 band due to ZnNC, which also provides more details for ZnCN, ZnNC, HZnCN and HZnNC. Table 1 includes HCN, HNC, and ZnNC. Table 1 provides more details for ZnCN, ZnNC, HZnCN and HZnNC. The isotope effect on the C-N stretch is larger for the cyano systems than the isocyano systems, suggesting that displacement of the labeled Carbon is greater in the normal mode for ZnCN than for ZnNC.

### Table 1. Frequencies [cm\(^{-1}\)] observed from reactions of Zn atoms with H\(_{12}\)CN and H\(_{13}\)CN during co-deposition in excess argon at 5 K. Bold type marks the major trapping site. Common absorptions for non-metal species (CN), CNCN, HNC and CN are listed first. Frequencies for zinc-bearing systems follow, which have counterparts in all the Group 12 metal experiments. "da" = decreased intensity upon annealing.

| Non-Metal Species A \(\text{H}^{12}\)CN | \(\text{H}^{13}\)CN | 12/13 freq. ratio, shift | assignment |
|--------------------------------------|------------------|--------------------------|------------|
| 2294.9                               | 2241.7           | 1.02373, 52.3            | sym CNCN mode |
| 2153.5                               | 2207.8           | 1.02655, 55.7            | (CN) \(_2\)  |
| 2138.4                               | 2091.5           | 1.02252, 46.9            | CO         |
| 2097.9, 2093.1, 2089.9                | 2063.9, 2060.1, 2056.8 | 1.01602, 33.0 | HCN matrix sites |
| 2100                                 | 2069             | 1.01498, 31              | (HCN) \(_2\) |
| 2054.2, 2053.7                       | 2017.7, 2109.4   | 1.01809, 34.5            | asym CNCN&site |
| 2044.4                               | 2001.9           | 1.02123, 42.5            | CN         |
| 2027.6, 2023.5 (da)                   | 1984.6, 1980.6   | 1.02167, 43, 42.9        | HNC and site variants |

Zinc-containing molecules B: cyano and isocyano species

| \(\text{H}^{12}\)N | \(\text{H}^{13}\)N | Isotopic \(\nu\) Ratio | Computed \(\nu\) | Assignment |
|------------------|------------------|------------------------|----------------|------------|
| 2184.7           | 2136.2           | 1.02270                | 2198           | HZnCN      |
| 2162.2           | 2114.9           | 1.02236                | 2177           | ZnCN       |
| 2097.9           | 2060.1           | 1.01830                | 2102           | HZnNC      |
| 2074.6           | 2034.0           | 1.01996                | 2083           | ZnNC\([18]\) |
| 2196.0           | 2151.3           | 1.02078                | 2077           | ZnNC\([18]\) |
| 2097.9           | 2058.3           | 1.01914                | 2030           | HZnNC\([18]\) |
| 2200.5           | 2153.9           | 1.02164                | 2044           | HZnNC\([18]\) |
| 2097.0           | 2057.5           | 1.01920                | 2048           | HZnNC\([18]\) |
| 1996.0           | 1956.0           | 1.00000                | 1972.1         | HZnNC      |
| 1972.1 look at 2023.9 | 1972.1, 2023.9 | 1.00000                | 2023.9         | HZnNC      |

Zinc-containing molecules B: continued hydrides

| H | D | Isotopic \(\nu\) Ratio | Computed \(\nu\) | Assignment |
|---|---|------------------------|----------------|------------|
| 2162.2 | 2131.7 | 1.01431 | 2177 | ZnCN |
| 2074.6 | 2041.0 | 1.01646 | 2083 | ZnNC\([18]\) |

Zinc-containing molecules B: continued hydrides

| \(\text{HN}^\text{14}\) | \(\text{HN}^\text{15}\) | Isotopic \(\nu\) Ratio | Computed \(\nu\) | Assignment |
|-----------------|-----------------|------------------------|----------------|------------|
| (1870.2)        | (1870.2)        | (1870.2)                | 2177           | ZnH\(_2\)\([6]\) |
| (1493.9)        | (1493.9)        | (1493.9)                | 2083           | ZnH\(_2\)\([6]\) |
| 1889.4          | 1371.6          | 1.37752                 |                | ZnH\(_2\)\([6]\) |

addition of H to the ZnCN cyanide and ZnNC isocyanide isomers above to make the new HZnCN and HZnNC hydrides, which are assigned here. This radical-radical combination is highly exothermic (Energetics are in Table S1).

### 2.2.2. Isotope Effects

Figure 2 compares vibrational features for the normal and \(^{15}\)C labeled species including HCN, HNC, and ZnNC. Table 1 provides more details for ZnCN, ZnNC, HZnCN and HZnNC. The isotope effect on the C-N stretch is larger for the cyano systems than the isocyano systems, suggesting that displacement of the labeled Carbon is greater in the normal mode for ZnCN than for ZnNC.

The weak bands at 2054.0 and 2057.7 in Figure 1 are matrix site splittings for CNCN and they are more intense in experiments with the cyanogen precursor, which is isoelectronic with CNCN.\([18]\) Space limitations may determine what can be used for labels in a particular figure. Two matrix site bands are observed for NCZnZnCN in Figure 1 at 2193.2 and 21966. More of these bands were observed at higher HCN concentration in Figure 1 than at lower HCN concentration in Figure 2.

Our previous work using a statistical mixture of cyanogen with \(^{14}\text{NCN}^{15}\text{N}\) provided the sharp above 2074.6 cm\(^{-1}\) band for ZnNC and a similar band at 2041.0 which exhibited the same...
increase on annealing and decrease on photolysis as the first band. The $^{14}$N/$^{15}$N isotopic frequency ratio 1.01646 is less than the $^{12}$C/$^{13}$C isotopic frequency ratio 1.01986 owing to its lower reduced mass. Thus the strong 2074.6 band is due to the isocyanide ZnNC and the 33.6 cm$^{-1}$ lower band at 2041.0 is the right amount for the Zn$^{15}$NC shift and 14/15 ratio 1.01646. Another band in the statistical sample spectrum at 2131.7 was 90.7 cm$^{-1}$ higher than the 2041.0 band for Zn$^{15}$NC, which is appropriate for its cyanide isomer ZnC$^{15}$N as assigned. Figure 2 also contains the new band at 2184.7 cm$^{-1}$ in the neighborhood near the ZnC–N stretch, 22.5 cm$^{-1}$ higher than the new 2162.2 band assigned above to ZnCN. The 2184.7 cm$^{-1}$ band has its $^{13}$C counterpart at 2136.2 cm$^{-1}$, just below the CO band, which gives a healthy 48.5 cm$^{-1}$ shift and a 1.02270 $^{12}$C/$^{13}$C frequency ratio. This shift and ratio closely match the 47.3 cm$^{-1}$ $^{13}$C shift and ratio 1.02236 associated with the 2162.2 cm$^{-1}$ band now assigned to ZnCN. This indicates that the 2184.7 cm$^{-1}$ and 2136.2 cm$^{-1}$ bands are associated with a ZnC–N functional group. HZnCN is the clear candidate to explain the 22.5 shift for the 2184.7 band. The sharp 2097.8 band is due to HZnNC owing to the large yield of the ZnNC radical for favorable reaction with H atoms.

Figure 3 shows that the two lower frequency monohydride absorptions clearly increase on annealing which promotes H atom reactions, but the more strongly absorbing HNC band at 2074.6 cm$^{-1}$ declines in intensity on photolysis while the weaker absorption at 2162.2 cm$^{-1}$ increases upon photolysis. The key here is the lower carbon $^{12}$C/$^{13}$C frequency ratio, i.e. (2074.6/2034.22) = 1.01986, is diagnostic of the isocyno stretch, to be compared with the higher ratio for the 2162.2 band, (2162.2/2114.9) = 1.02236 characteristic of the cyano stretch. The 2184.7 band is just above HCN, which is observed in Figure 2 using lower reagent concentration, and this band is due to HZnCN. Figure 3 also features two additional new IR bands in the Zn–H stretching region at 1962.2 and 1972.1 both above ZnH$^2$ at 1870.2 cm$^{-1}$ (1889.43, gas phase). These are associated with an annealing step, which allows trapped H atoms to diffuse and react with open shell molecules in the matrix. The assignment of these modes as HZn stretches is assured by their zero carbon-13 shifts. The same frequency bands are observed in scans a,b,c, also with HCN, as in d,e,f. The experimental frequency separation between the H–Zn and C–N mode at 2184.7 and the ZnC–N mode at 2162.2 cm$^{-1}$ (22.5 cm$^{-1}$) is similar to the frequency separation between the calculated HZnN–C mode at 2097.8 cm$^{-1}$ and the ZnN–C mode at 2074.6 (that is, 23.2 cm$^{-1}$). These two isocyano absorptions decrease on UV photolysis as the corresponding lower energy cyano signals increase. This parallel allows us to confirm the assignment of both the absorptions at 2184.7 and 1962.2 cm$^{-1}$ to HZnCN’s cyano C–N isomer and H–Zn motions and the two bands at 2097.8 and 1972.1 cm$^{-1}$ to HZnNC’s isocyno Zn–N–C and H–Zn stretches.
Table 1 collects the numerical values of the observed C–N frequencies, the isotopic frequency ratios of $^{12}$C and $^{13}$C species, and assignments for the observed frequencies. Cyanogen and its isomer CNCN, CO, CN, HCN, and HNC have been identified. Frequencies have been assigned to Zn cyanide and isocyanide and to the hydrogen addition hydrides as well. Data for NCZnZnCN and ZnH$_2$ from previous work are also included.\(^{[18]}\) The HNC and HCN isomers are related by photolysis.\(^{[26]}\) The cyano stretch for HZnCN is 22.5 cm$^{-1}$ higher than the cyano stretch in ZnCN; the isocyano stretch for HZnNC lies 23.2 cm$^{-1}$ higher than the isocyano stretch in ZnNC. This suggests that the formation of the hydride affects the (C–N) groups in small and similar ways. The observed blue shifts for the hydrides are slightly higher than the CCSD computed values for the cyano, 6.2 and isocyano 14.4 cm$^{-1}$. The $^{12}$C/$^{13}$C isotopic frequency ratios for Zn-bearing species (also shown in Table 1) fall into two sets, (A) greater than 1.021 for cyano systems and (B) smaller than 1.021 for isocyano species. This generalization applies to systems with one or two Zn atoms and two (iso)cyano fragments, according to previous work.\(^{[6,18]}\) We observed that the band at 1972.1 cm$^{-1}$, assigned to the H–ZnCN mode for HZnCN, lies 101.9 cm$^{-1}$ higher than the Zn-asymmetric stretch for ZnH$_2$ at 1870.2 (in solid argon). More 12/13 frequency ratios are in Table 1 below.

The $^{15}$N isotopic data are from statistical NC=C$^{15}$N. We confine discussion to the strongest bands of ZnNC and ZnCN.

Figure 3. The infrared spectra between 2060 and 1800 cm$^{-1}$ for the reaction products from laser ablated Zn co-deposited with 0.2 % HCN in argon at 5 K. Spectra (a–e) are observed after the following steps: (a) deposition for 120 min, (b) annealing to 20 K and cooling back to 5 K, (c) full Hg arc irradiation for 20 min, (d) annealing to 30 K, and (e) annealing to 35 K.

2.3. Estimate of Abundances for Zinc Cyanide and Isocyanide

We measured absorbances A for the 2074.6 cm$^{-1}$ ZnNC and 2162.2 cm$^{-1}$ ZnCN bands in 0.2% and 1.0% concentration experiments (Figures 2 and 1). In the deposit with 1% reagent we find the band absorbance measured is 0.1344 for the 2074.6 cm$^{-1}$ band while the value for the 2162.2 cm$^{-1}$ band is 0.02364. We can obtain an estimate of the relative abundance of cyano and isocyano species from the computed absorption intensities (see next section). Normalize by dividing the absorbances by the CCSD computed intensities for these two modes: 276 and 26 km/mol, respectively, we find a measure of abundance estimated for the 2074.6 isocyano band to be $4.9 \times 10^{-4}$ and for the 2162.2 cm$^{-1}$ cyano band to be $9.1 \times 10^{-4}$. These numbers show that our matrix traps 2× more of the more stable ZnCN cyanide isomer than the 17.3 kJ/mol (CCSD) higher energy ZnNC isocyanide. The same treatment can be applied to the hydrides HZnNC and HZnCN, whose 2097.8 and 2184.7 cm$^{-1}$ bands are 5/3 in relative absorbance in
the 0.2% experiment. (We could not measure the 2097.8 cm\(^{-1}\) band in the 1% HCN experiment owing to interference from HCN polymers). The relative populations are roughly 6:1 in favor of the more stable HZnCN. Only the extremely high intrinsic infrared intensities for the isocyano systems makes their presence prominent in the infrared spectra.

### 2.4. Computational Modeling (Zinc Systems)

One of the most demanding, and for predominantly single-determinant states one of the most reliable electronic structure methods is the coupled cluster expansion. Usually the CCSD(T) form, expressed in an extended basis set such as aug-cc-pVTZ approaches chemical accuracy in energy terms and estimates harmonic vibrational frequencies within at most a few tens of wave numbers. Computational modeling provides energetics and structures. We defer discussion of these properties to a later comparison of all three metal systems. Here we focus on the computed vibrational frequencies including isotope effects for the Zinc bearing molecules. We report both CCSD(T) and CCSD results since absorption intensities are available in the Gaussian 16 implementation for CCSD, and CCSD provides a comparison point for the CCSD(T) vibrational frequencies.

Vibrational frequencies for HCN and H\(^{14}\)C reaction products with of Zinc using the CCSD and CCSD(T) methods.

Table 2 contains computed vibrational frequencies for isotopic variants of Zinc systems. The computed \(^{12}\)C/\(^{13}\)C frequency ratios clearly distinguish cyano from isocyano systems, and are consistent with the experimental ratios. Specifically, the observed Zn\(^{12}\)CN/Zn\(^{13}\)CN frequency ratio (2162.2/2114.9) is 1.0224; the ratios calculated from the CCSD and CCSD(T) frequencies are 1.02271 and 1.02313, in close agreement. The consistency confirms assignment of the new observed 2162 cm\(^{-1}\) band to the ZnC–N mode. The theoretical (harmonic) frequency ratios are slightly larger than the experimental values, which are due to their anharmonicity.

The naturally occurring abundance of \(^{12}\)N in \(^{14}\)N–\(^{15}\)N gave ZnC\(^{12}\)N and Zn\(^{15}\)NC absorptions at 2131.7 and 2041.0 cm\(^{-1}\) as shown in Table 1. The ratio of CN stretching frequencies for ZnC\(^{12}\)N/Zn\(^{15}\)NC isotopic substitution is 1.01457 according to CCSD calculations, 1.01511 by CCSD(T). The observed \(^{14}\)N/\(^{15}\)N frequency ratio for ZnNC is 1.01996 while the computed values are 1.02003 and 1.01992 for CCSD and CCSD(T) respectively. The observed \(^{14}\)N/\(^{15}\)N frequency ratio for ZnNC is 1.01680; CCSD and CCSD(T) values are 1.01680 and 1.01689, respectively. Comparison with the ZnCN ratios suggests that there is more N motion and less C movement in the isocyanide than in the cyanide. A more thorough discussion of this statement is covered in Supplementary Information. The isotopic frequency (harmonic) frequency ratios are slightly larger than the experimental values, which are due to their anharmonicity.

The parallel between (C–N) stretching isotopic frequency ratios in the Zn(C, N) and HZn(C, N) species is clear. For HZnCN, the observed \(^{12}\)C/\(^{13}\)C ratio, 1.02270, matches CCSD and CCSD(T) values, 1.02293 and 1.02301, respectively. The corresponding observed \(^{12}\)C/\(^{13}\)C value for HZnNC, 1.01830, is to be compared

### Table 2

**Coupled cluster calculations for sigma stretching modes in Zinc systems using G-16.**

| 2A: CCSD | v1 σ (C, N) (Intensity) | Isotopic Ratios\(^{\text{[a]}}\) | v2 σ (H)Zn-(C, N) (Intensity) | v4 σ H–Zn (Intensity) | v1–v4 |
|----------|------------------------|-----------------------------|-----------------------------|------------------------|-------|
| ZnCN     | 2260.3 (26)            | 1.02236                     | 418.8 (58)                  | 413.7 (57)             |       |
| Zn\(^{13}\)CN | 2210.1 (23)        | 1.01457                     | 412.8 (56)                  | 412.8 (56)             |       |
| Zn\(^{15}\)CN | 2227.6 (28)        | 1.01293                     | 462.3 (38)                  | 462.3 (38)             |       |
| ZnNC     | 2148.2 (276)          | 1.02003                     | 464.2 (78)                  | 464.2 (78)             |       |
| Zn\(^{15}\)NC | 2106.0 (276)       | 1.01680                     | 465.4 (79)                  | 465.4 (79)             |       |
| Zn\(^{15}\)NC | 2112.7 (258)       | 2008.0 (85)                  | 258.5                       |                        |       |
| HZnCN    | 2266.5 (25)           | 467.8 (39)                  | 2008.0 (85)                  | 258.5                  |       |
| DZnCN    | 2266.5 (25)           | 466.7 (39)                  | 2008.0 (85)                  | 258.5                  |       |
| HZn\(^{15}\)CN | 2215.7 (23)      | 2008.0 (85)                  | 258.5                       |                        |       |
| HZn\(^{15}\)CN | 2234.4 (27)       | 461.0 (38)                  | 2008.0 (84)                  | 258.5                  |       |
| HZn\(^{15}\)NC | 2162.6 (264)      | 512.2 (56)                  | 2027.4 (62)                  | 135.2                  |       |
| DZn\(^{15}\)NC | 2162.6 (261)      | 519.7 (55)                  | 2027.4 (62)                  | 135.2                  |       |
| HZn\(^{15}\)NC | 2120.9 (265)      | 513.3 (54)                  | 2027.4 (59)                  | 135.2                  |       |
| HZn\(^{15}\)NC | 2126.3 (249)      | 515.0 (55)                  | 2027.4 (61)                  | 135.2                  |       |
| 2B: CCSD(T) | v1 σ Isotopic Ratios\(^{[b]}\) | v2 σ (H)Zn-(C, N) (Intensity) | v4 σ H–Zn (Intensity) | v1–v4 |
| Zn\(^{12}\)CN | 2277.6            | 1.02313                     | 411.4                       | 411.4                  |       |
| Zn\(^{13}\)CN | 2226.1            | 1.01511                     | 410.5                       | 410.5                  |       |
| Zn\(^{15}\)CN | 2243.7            | 1.01992                     | 459.7                       | 459.7                  |       |
| Zn\(^{13}\)NC | 2083.6            | 1.01689                     | 460.0                       | 460.0                  |       |
| Zn\(^{15}\)NC | 2049.0            | 2010.8                      | 185.2                       |                        |       |
| HZn\(^{12}\)CN | 2196.0            | 467.3                       | 185.2                       |                        |       |
| HZn\(^{13}\)CN | 2196.0            | 466.3                       | 185.2                       |                        |       |
| HZn\(^{12}\)NC | 2146.6            | 462.0                       | 185.2                       |                        |       |
| HZn\(^{13}\)NC | 2165.1            | 460.7                       | 185.2                       |                        |       |
| HZn\(^{15}\)NC | 2099.3            | 519.1                       | 2028.2                      | 71.1                   |       |
| DZn\(^{15}\)NC | 2099.3            | 517.7                       | 2028.2                      | 71.1                   |       |
| HZn\(^{15}\)NC | 2059.1            | 511.2                       | 2028.2                      | 71.1                   |       |
| HZn\(^{15}\)NC | 2063.9            | 513.0                       | 2028.2                      | 71.1                   |       |

\(^{[a]}\) \(^{12}\)C/\(^{13}\)C or \(^{14}\)N/\(^{15}\)N, \(^{[b]}\) \(^{12}\)C/\(^{13}\)C or \(^{14}\)N/\(^{15}\)N.
that CCSD(T) estimates the isocyano stretch very accurately in ZnCN systems the difference is 21 cm
stretches (no intensities are produced). The difference between H
computed an almost pure Zn
structural formula.

frequencies in HZnNC, (2099.3/2059.1
computed with 1.01966 and 1.01952 computed values. The CCSD(T)
values for the isocyano systems suggest that the N atom
displacement is greater in the isocyano systems. A parallel
argument indicates that the C atom displacement is greater in
the cyano systems than in the isocyano systems.

It is interesting to examine the computed frequencies
associated with HZnCN and HZnNC with those of ZnCN and ZnNC. The isocyano and Zn–H stretching modes for HZnNC are
computed with CCSD as 2162.6 cm\(^{-1}\) (intensity 264 km/mol) and 2027.4 (intensity 62) while the cyano and Zn–H stretches are 2266.5 and 2008.0 cm\(^{-1}\) (Table 2A, below). The N–C stretch in HZnNC is computed by CCSD to be more strongly absorbing than the CN stretch by 264/62 = 4.3 times. Frequencies obtained using CCSD(T) are 2099.3 and 2028.2 cm\(^{-1}\) for the HZnN–C and H–Zn stretching modes, respectively. CCSD(T) calculations on HZnCN yield values of 2196.0 for C–N and 2010.8 for H–Zn
stretches (no intensities are produced). The difference between
computed cyano and isocyano stretches (96.7 cm\(^{-1}\)) is consistent
with the widely observed 100 cm\(^{-1}\) differences. For the (H)
Zn NC systems as described by Molpro, the NC stretching
frequency is about 19 cm\(^{-1}\) higher for the hydride, while for (H)
Zn NC systems the difference is 21 cm\(^{-1}\). Gaussian values do not
follow this pattern; see discussion below.

The 12–13 shift for the observed cyano stretch 2162.2–
2114.9 is 47.3 cm\(^{-1}\), and it is less, 40.6, for the isocyano. The observed 14/15 ratio for ZnNC is 1.01651 and the shift is
33.6 cm\(^{-1}\) while the computed shift for ZnCN is 32.9 cm\(^{-1}\) and the ratio is 1.01466. Thus the N moves more in the isocyano
than in the cyano, which is obvious from the Zn–N–C structural
formula.

The \(^{12}\text{C}/^{13}\text{C}\) ratio from CCSD(T) for isocyano stretching
frequencies in HZnNC, (2099.3/2059.1 = 1.01952), is characteristic
of an isocyano stretch. The CCSD(T) band at 2028.2 cm\(^{-1}\) is
unaffected by \(^{13}\text{C}\) isotopic substitution, which tells us that it is
an almost pure Zn–H stretching mode. This is borne out by the computed \(^{1}\text{H}/^{2}\text{H}\) isotopic ratio, (2028.2/1446.7) = 1.40195. Note
that CCSD(T) estimates the isocyano stretch very accurately
in ZnNC, (2075 cm\(^{-1}\) observed, 2083 cm\(^{-1}\) calculated) but is less
faithful to the hydride molecule’s H–Zn stretch (2098 cm\(^{-1}\) for
this ionic bond, 2074 cm\(^{-1}\))

Comparisons among computed stretching frequencies are
possible. The significance of the perturbative triples (PT) in the
CCSD(T) correction is minor for the Zn–H stretch, and for the
cyano- and isocyano hydrides; it adds +2.8 cm\(^{-1}\) and
+0.8 cm\(^{-1}\) to the cyano and isocyano H–Zn modes respectively.
These modes, at 2010.8 and 2028.2 cm\(^{-1}\), are almost entirely
local, and lie well to the blue of the asymmetric stretch for
ZnH\(_2\) at 1870.2 cm\(^{-1}\). These ionic bonds are more difficult to model.
The cyanide and isocynide Zn–C and Zn–N stretches in the
hydrides are virtually unaffected by the PT as well.

However the PT correction softens the cyano and isocyano
stretches in the hydrogen compounds, by −70.5 and
−63.3 cm\(^{-1}\). For the Zn(CN) radicals, the PT correction softens
the isocyano stretch by about that same amount (−64.6 cm\(^{-1}\)),
but stiffens the cyano stretch slightly (+17.3 cm\(^{-1}\)). This
anomaly, suggesting a flaw in the PT for the ZnCN, will be
encountered in the CdCN and HgCN radical analogs as well.

3. Cadmium Studies

3.1. Spectroscopic Results and Discussion (Cadmium Systems)

Figure 4 displays infrared spectra for cadmium atoms con-
densed in argon with 0.2 % HCN. Familiar nonmetal species
including CO, CN, HCN, HNC, cyanogen, and its CNCN isomer,
are identified explicitly. Upon first deposition (spectrum a)
features appear at 2259.2, 2207.0, 2177.7, and 2069.7 cm\(^{-1}\).
These signals persist through first annealing (spectrum b). Upon
first annealing a new band emerges at 2196.7 cm\(^{-1}\), and upon
annealing to 20 K. (spectrum c) another band appears at
2140.9 cm\(^{-1}\).

3.1.1. CdCN and CdNC

The spectra of cadmium cyanides and isocyanides are slightly
lower in frequency than those observed for the zinc cyanides
and isocyanides. The isocyano stretching mode for CdNC is
observed at 2069.7 cm\(^{-1}\) in the cyanogen reaction.[19] The band
at 2140.9 cm\(^{-1}\) is a candidate for the cyanano stretch for CdCN,
which is 21.3 lower than the analogous mode in behavior for
ZnCN, and the CdNC product is only 4.9 cm\(^{-1}\) lower than its
zinc analog.

These and other comparisons suggest that Cd acts like a
heavier isotope of Zn. Although the Cd product bands are
weaker, photolysis decreases the higher energy CdCN isomer in
favor of the lower energy CdCN isomer.

3.1.2. HCdCN and HCdNC

By analogy to observations in the Zn system, we expect to find
the products of H radical-CdCN combination. In Figure 4 we see
new bands at 2177.7 and 2140.9 cm\(^{-1}\) separated by just
36.8 cm\(^{-1}\). In analogy to the Zinc case, these modes can
reasonably be associated with the cyano stretches of the
HCdCN and CdCN molecules. The bands observed at 1884.3
and 1864.8 cm\(^{-1}\) (see Figure 6) are candidates for Cd–H
stretches in HCdNC and HCdCN. In this case the lower observed
band at 2140.9 goes with the lower computed frequency at
2167 and the lower terminal Cd–H frequency at 1864.8 cm\(^{-1}\).

3.1.3. Isotopic Effects

The strong isocyano absorption at 2069.7 cm\(^{-1}\) has a \(^{13}\text{C}\) counterpart at 2028.2 cm\(^{-1}\) (Figure 5, Table 3). Its \(^{12}\text{C}/^{13}\text{C}\)
isotopic frequency ratio, 1.02046, is only slightly higher than the
1.01996 value found for the zinc counterpart. We may infer a
larger amplitude motion for C against the heavier metal in
Cd–C–N. The $^{12}$C/$^{13}$C frequency ratio for the 2177.7 cm$^{-1}$ band is 1.02249, which suggests the cyano isomer HCdC–N and another C–N band. Figure 5(b) red shows a substantial 75% decrease in the CN radical and an increase in HCN and cyanogen NCCN on annealing to 15 K, which also happens to the carbon-13 analogs.

The signal at 2140.9 cm$^{-1}$ shown in Figure 5 spectrum c obtained after annealing to 20 K, can be attributed to CdCN by indirect argument. The signal for the cyan stretch for Cd$^{15}$CN is obscured by the intense absorption of HCN, but (1) the proximity of the 2140.9 cm$^{-1}$ band to the 2177.7 cm$^{-1}$ band attributable to the HCd$^{15}$CN cyano stretch and (2) HCd$^{13}$CN’s isotopic counterpart at 2129.8 cm$^{-1}$ with ratio 1.02249, together support the CdCN assignment.

Table 3. Frequencies [cm$^{-1}$] observed from reactions of Cd atoms with H$^{12}$CN and H$^{13}$CN in excess argon at 5 K. Bold type marks the major trapping site. Harmonic frequencies computed by Molpro CCSD(T)/aug-cc-pVTZ guided the assignment of structures and vibrational modes. [4]

| Cadmium $^{15}$C | $^{13}$C | Isotopic Ratio | Computed | Species |
|-----------------|---------|---------------|----------|---------|
| 2177.7 (ia)     | 2129.8 (ia) | 1.02249       | 2190     | HCDNC   |
| 2140.9          | Under (H$^{13}$CN)$_x$ | – | 2167 | CdCN    |
| 2088 sh         | Under (H$^{13}$CN)$_x$ | – | 2094 | HCDNC   |
| 2069.7, 2066.2  | 2028.2, 2023.8 | 1.02046       | 2077     | CdNC, site variants |
| 1884.3 (da)     | 1884.3 (da) | 1925 |
| 1864.8 (ia)     | 1864.8 (da) | 1908 | HCDCN Cd–H stretch |
| 1754.2          | (1260.3) | (1.39348) | CdH$_2$ and CdD$_2$ |
| (1756.2)        | (1339.8) | (1.37459) | in Ar with a H$_2$/D$_2$ mix $^4$ |
| (1339.4)        | (974.4)  | (1.38585) | CdH and CdD |
| [1771.53] $^{[3]}$ | [1278.3] $^{[3]}$ | [35] |

[a] Band behavior: ia = increase on annealing; da = decrease on annealing; dp = decrease on photolysis; sh = shoulder. Frequency values in curly brackets were observed using other methods.
3.2. Computational Modeling (Cadmium Systems)

3.2.1. Vibrational Frequencies for Isotopic Variants of Cadmium Systems

Table 4 contains computed vibrational frequencies for isotopic variants of the cadmium systems. As we found for the zinc systems, computed $^{12}\text{C}/^{13}\text{C}$ frequency ratios clearly distinguish cyano from isocyano systems. The ratios calculated from the CCSD and CCSD(T) frequencies for CdCN are 1.02255 and 1.02252, in close agreement with the zinc analogs. Unfortunately, the Cd$^{12}$CN/Cd$^{13}$CN frequency ratio cannot be measured because the Cd$^{13}$CN frequency is obscured by the H$^{12}$CN band present as a 1% impurity in the H$^{13}$CN sample. The observed $^{12}\text{C}/^{13}\text{C}$ frequency ratio for CdNC is 1.02046 while the computed values are 1.02026 and 1.02023 using CCSD and CCSD(T), respectively.

The ratios of CN stretching frequencies for CdC$^{14}$N/CdC$^{15}$N isotopic substitution are 1.01472 according to CCSD calculation, 1.01478 by CCSD(T). The $^{14}\text{N}/^{15}\text{N}$ frequency ratio for CdNC is 1.01664 according to CCSD and 1.01668 by CCSD(T). Comparison of the CdCN ratios suggests that there is more N motion and less C movement in the isocyanide than in the cyanide.

The parallel between (C and N) stretching frequency isotope ratios in the Cd(C,N) and HCd(C,N) species is plain. For HCdCN, the CCSD and CCSD(T) values for the $^{12}\text{C}/^{13}\text{C}$ frequency ratios are 1.02280 and 1.02284 respectively, very close to the ratios for CdCN. The corresponding computed $^{12}\text{C}/^{13}\text{C}$ values for HCdNC are 1.01993 and 1.01983. The $^{14}\text{N}/^{15}\text{N}$ frequency ratios for HCdNC are 1.01690 according to CCSD and 1.01696 by CCSD(T), in close agreement with corresponding values for CdNC.

Calculations further show that the new band observed at 1864.8 cm$^{-1}$ is due to the Cd–H stretching mode of HCdCN (computed at 1907.5 cm$^{-1}$), and the higher new band in this region at 1884.3 cm$^{-1}$ is due to the Cd–H stretching mode for the HCdNC isomer (computed at 1949.6 cm$^{-1}$). CCSD and CCSD(T) calculations agree closely in $^{1}\text{H}/^{2}\text{H}$ isotopic ratio for the CdH stretch (1.40742 and 1.49733 respectively), but both overestimate the H–Cd stretch value considerably.

3.3. Discussion of (H)Cd–CN, Cd–H and (C–N) stretching frequencies

As in the HZn(C,N) systems, the significance of the perturbative triples (PT) in the CCSD(T) correction is modest for the H–Cd cyano ($\sim 3.5$ cm$^{-1}$) and isocyano (24.4 cm$^{-1}$) stretches. The PT corrections are $-9.2$ cm$^{-1}$ and $-4.5$ cm$^{-1}$ for the Cd–C and Cd–N stretches in the two isomers. However the PT correction softens the cyano and isocyno stretches in the hydrogen compounds, by $-70.8$ cm$^{-1}$ and $-61.3$ cm$^{-1}$ respectively. For the Cd(C,N) radicals, the PT correction softens the isocyno...
stretch by a like amount (−66.6 cm⁻¹), but has little effect on the cyano stretch (−1.8 cm⁻¹). This anomaly, seen in more striking form in the Zn system, will be encountered in the Hg systems as well.

There are some other persistent features of the stretching frequency differences between cyano and isocyano systems. Adding H stiffens the (H)Cd–(C=N) frequencies slightly, and the C=N frequencies for both radicals and hydrogenated compounds are higher for the cyano systems than for the isocyano systems, by a like amount. Once again the PT term for the CdCN system is anomalous. The H–Cd stretch is stiffer for the isocyano system, and there is a noticeable effect of the PT correction.

4. Mercury Studies

4.1. Spectroscopic Results and Discussion (Mercury Systems)

We generated Hg atoms by laser ablation of a 50% Hg dental amalgam target, a new procedure developed in our labs. A picture of this target is shown in supporting Information P-1. The Volatile Hg is produced preferentially, leaving behind the other more refractory components of the amalgam. Co-deposition at 5 K with 0.2% HCN and argon followed by further processing produced the spectra displayed in Figure 7.

4.1.1. Appearance of HgCN, HgNC, HHgCN and HHgNC

Besides the signals associated with familiar non-metallic species, the first deposition (spectrum a) produced bands at 2032.7 and 2120.4 cm⁻¹ as well as 2150.8 cm⁻¹ and 2066.2 cm⁻¹. The first annealing produced a new absorption (spectrum b, red) at 1933.0 cm⁻¹ and had little effect on the other bands: All bands were enhanced on photolysis (spectrum c) except the 1933 cm⁻¹ absorption, which was erased (spectrum c, blue). This feature will be considered after more experiments and calculations are done on possibilities for the 1933 band.

We will be following analogies set forth in the above Zn and Cd studies, which lead us to anticipate an intense signal from the HgNC isocyano stretch labeled in the spectra of the Figure 7 deposited sample. The strong band at 2032.7 cm⁻¹ (37 cm⁻¹ below CdNC), is a good candidate. The feature at 2120.4 cm⁻¹ meets our expectation for an HgCN stretch at ca. 100 cm⁻¹ higher frequency and less intensity than the isocyano stretch. Even so we expect that there are more of the weaker absorbing more stable HgCN molecules in our matrix: Band intensity is central to this work.
4.1.2. Isotopic Effects

These proposals are borne out by the presence of transitions at 1992.5 cm\(^{-1}\) and 2074.1 cm\(^{-1}\) in the spectra produced with H\(^{13}\)CN (Figure 8). These are linked to primary product absorptions for HgNC and HgCN, respectively to be assigned here. The isotopic ratio in the first case is 1.02018 (Table 5), characteristic of the isocyano stretch, and in the second case is 1.02232, close to the value for the cyano stretch established here in Zn and Cd systems. These \(^{12}\)C/\(^{13}\)C red shifts by 46.3 and 40.2 cm\(^{-1}\) to lower frequencies follow the profiles for Cd and Zn (compare Tables 1 and 2 for Zn, 3 and 4 for Cd above).

### Table 4. Calculations for cadmium cyanide and isocyanide and their hydride isotopic frequencies [cm\(^{-1}\)] and intensities [km/mol]. From Gaussian calculations with the aug-cc-pVTZ-pp basis.

| Species  | \(\nu_1\) (\(\sigma(N-N)\)) (Intensity) | \(^{12}\)C/\(^{13}\)C Isotopic Ratio | \(\nu_2\) (\(^{14}\)N/\(^{15}\)N) | \(\nu_4\) (Intensity) | \(\nu_1-\nu_4\) |
|----------|-----------------------------------|-----------------------------------|-----------------|-----------------|-----------------|
| CdCN     | 2253.7 (21)                       | cal                               | 369.2 (56)      |                  |                  |
| Cd\(^{13}\)CN | 2204.0 (17)                      | 1.02255                           | 364.0 (54)      |                  |                  |
| Cd\(^{15}\)N | 2221.0 (22)                      | 1.01472                           | 363.3 (54)      |                  |                  |
| CdNC     | 2144.7 (258)                      | 1.02026                           | 390.8 (75)      |                  |                  |
| CdN\(^{13}\)C | 2102.1 (258)                     |                                   | 391.6 (76)      |                  |                  |
| Cd\(^{15}\)NC | 2109.6 (241)                     | 1.01664                           | 391.6 (76)      |                  |                  |
| HgCdCN   | 2261.2 (18)                       |                                   | 411.9 (38)      | 1911.0 (98)      | 351.2           |
| DgCdCN   | 2261.2 (18)                       |                                   | 411.5 (37)      | 1357.8 (52)      | 234.7           |
| Hg\(^{13}\)CN | 2210.8 (16)                      | 1.02280                           | 406.1 (37)      | 1911.0 (98)      | 234.7           |
| Hg\(^{15}\)CN | 2228.8 (19)                      | 1.01454                           | 405.1 (37)      | 1911.0 (98)      | 234.7           |
| HgCdNC   | 2160.0 (249)                      |                                   | 450.0 (55)      | 1925.3 (70)      | 234.7           |
| DgCdNC   | 2160.0 (247)                      |                                   | 449.5 (55)      | 1368.3 (40)      | 234.7           |
| HgCd\(^{13}\)N | 2171.8 (249)                     | 1.01993                           | 442.5 (53)      | 1925.3 (67)      | 234.7           |
| HgCd\(^{15}\)N | 2124.1 (234)                     | 1.01690                           | 443.7 (54)      | 1925.3 (70)      | 234.7           |

### Table 5. Observed frequencies and isotopic ratios for species produced by codeposition of Hg and HCN with argon (A) and reported frequencies for hydride HgH\(_2\) (B).\(^{[f]}\)

| MERCURY SYSTEMS A: cyanides and isocyanides \(^{13}\)C/\(^{15}\)N | Isotopic Ratio | Species |
|---------------------------------------------------------------|----------------|---------|
| 2150.8 May be hidden by \((\(^{13}\)HCN)\)_x           | –              | HgH\(_2\)CN |
| 2120.4 2074.1                                              | 1.02232        | HgCN 8HNC   |
| (2066.2) 2114.9                                            | 102236         | (HgH\(_2\)N–C) |
| 2072.6                                                      |                 | HN–C        |
| 2032.7 (da) 1992.5 (da)                                      | 1.02018        | HgN–C       |
| 1933.0 (ia, dp) 1892.4 (ia, dp)                              | 1.02145        | Work in progress |

| MERCURY SYSTEMS B: hydrides \(^{12}\)O/\(^{18}\)O | Species |
|-----------------------------------------------|---------|
| 11912.8 (gas)                                | HgH, asym str (a) |
| 1895.3 (Argon mat.)                         | HgH, asym str (b) |
| 1900.4: 1910.3 (e)                          |                 |
| 1209 (H/D)                                  | 899.8, 909.7   | 1.34363    | Hg–H str (c) |
| Tab5 cont                                   | 896.1         | 1.34348    | Hg–H str (d) |
| 1203.2 (gas)                                 |             |           | Hg–H str    |
| 1205                                        |             |           |             |

\(^{[a]}\) Ref. [37]. \(^{[b]}\) Ref. [38]. \(^{[c]}\) Ref. [8]. \(^{[d]}\) Ref. [39]. \(^{[e]}\) Ref. [40]. \(^{[f]}\) Band behavior: (ia = increase on annealing, da = decrease on annealing), dp = decrease on photolysis.
The information that leads to assignment of infrared bands for HgCN, HgNC, HHgCN and HHgNC is presented here in Table 5.

We anticipate signals for CN stretches in HHgCN and HHgNC that increase on annealing: Our Molpro calculation with CCSD(T) (Table 6) predicts the HHgCN cyano stretching mode to lie 33 cm$^{-1}$ higher than the HgCN's CN stretch. The weak band observed at 2066.2 is 33 cm$^{-1}$ higher than the HgNC band at 2032.7 cm$^{-1}$: It does not grow substantially on annealing, but it is in the region where it could be covered by reforming HCN. Thus the feature at 2066.2 cm$^{-1}$ may be attributed to HHgNC, but we cannot be as confident in such an assignment as we are with the isocyanide isomer. On the other hand the 2150.8 band does grow on first annealing and photolysis and it is probably due to HHgNC. This cyanide at 2150.8 and isocyanide at 2066.2 differ by 84.6 cm$^{-1}$ which is almost the ca. 100 cm$^{-1}$ expected difference.

4.2. Computational Modeling (Mercury Systems)

4.2.1. Vibrational Frequencies for Isotopic Variants of the Hg systems

CCSD/aug-cc-pVTZ/aug-cc-pVTZ-pp and CCSD(T)/aug-cc-pVTZ/aug-cc-pVTZ-pp calculations of selected frequencies are collected in Table 6. Following the pattern for Zn and Cd, the computed cyanide stretching frequencies in HgNC and HHgNC are about 100 cm$^{-1}$ higher than the corresponding values in the isocyanides. Likewise the $^{12}$C/$^{13}$C isotopic ratios are larger for the cyanides than for the isocyanides. The Hg–H stretches are placed at 2190.5 by CCSD and 2175.7 by CCSD(T) for HHgCN and at 2255.6 by CCSD and 2243.9 cm$^{-1}$ by CCSD(T) for HHgNC. They are identified by calculated $^2$H/$^3$H isotopic ratios 1.40986 and 1.41072. However these computed frequency values are higher than those assigned to Hg–H stretches in the experimental spectra (Table 7).

4.2.2. Discussion of (H)Hg–CN, Hg–H and (C, N) stretching frequencies

The significance of the perturbative triples (PT) correction in CCSD(T) is minor in the Hg–H stretches, $-14.8$ cm$^{-1}$ for the

| Table 6. Calculations on HgH$_2$ |
|----------------------------------|
| Model (Harmonic) | Sym. Stretch | Asym. Stretch | Hg–H distance |
|------------------|-------------|---------------|--------------|
| CCSD/aug…pp      | 2110.6      | 1986.9 obs. 1895 | 1.6361       |
| CCSD(T)/aug…pp   | 2098.6      | 1979.0         | 1.6366       |

Figure 7. Infrared spectra of the reaction products of laser ablated Hg from a dental amalgam target with 0.2 % HCN in argon during condensation at 5 K. The sample was exposed to ablation plume Hg resonance radiation during the deposition process. The bottom spectrum (a) grey followed co-deposition of the reagents for 120 min spectrum, (b) red recorded after annealing to 20 K and cooling back to 5 K. Spectrum (c) blue taken after full mercury arc photolysis for 20 min. Spectrum (d) green followed annealing to 30 K and spectrum (e) purple followed annealing to 35 K and cooling back to 5 K.
cyano isomer and $-11.7 \text{ cm}^{-1}$ for the isocyano isomer. The shift is only $-2.8 \text{ cm}^{-1}$ for the Hg to C (cyano) stretch and $-4.2 \text{ cm}^{-1}$ for the Hg to N (isocyano) stretch.

However the PT correction softens (red-shifts) the cyano and isocyano stretches in the hydrides. For the cyanide softening is $-70.9 \text{ cm}^{-1}$, while for the isocyanide the shift is $-79.6 \text{ cm}^{-1}$. For the Hg(C,N) radicals, the PT correction softens the isocyano stretch by $-67.9 \text{ cm}^{-1}$, but has little effect on the cyano stretch, the shift being $+1.4 \text{ cm}^{-1}$. However these computed values are far higher than those observed here. This anomaly is seen throughout the metals. The most immediately relevant impact is on the frequency assignment of the CN stretches in the MCN species, which is discussed below. There are some other persistent features of the stretching frequency differences between cyano and isocyano systems. Adding H increases the (H)Hg to cyano or isocyano stretching frequencies slightly, and the frequencies for both radicals and hydrogenated compounds are higher for the cyano systems than for the isocyano systems, by a like amount. The Hg–H stretching frequency is higher for the isocyano system.

5. Comparisons within the Group 12 Family

A good way to celebrate the chemistry of the Group 12 metals is to make comparisons within this family of metals. In the following sections we describe bonding in the Zn, Cd, and Hg cyanides and isocyanides and their mono hydrides, and draw some generalizations on the relative energies, the geometries, and the vibrational spectra of compounds of Zn, Cd, and Hg.

5.1. Bonding in (H)HgNC

The Zn, Cd, and Hg cyanides and isocyanides are qualitatively similar in bonding. Starting with the MOs for $N_2$, we find symmetry-breaking polarization produces clear correlates in the CN anion to the central sigma bond, two pi bonds, and two lone pairs of $N_2$. As a metal cation with one odd $ns$ electron approaches along the molecular axis, a sigma bond forms to that metal. The ground state of the triatomic is $^{1}\Sigma^+^{(23)}$.

The qualitative nature of bonding in the Zn, Cd, and Hg hydrides is also very similar for all species. Figure 9 shows the specific case of (H)HgNC. The odd electron on HgNC found on the Hg atom (mainly the 6s AO) pairs with the odd electron of H atom as they approach one another along the molecular axis.
Table 7. Calculations for stretching modes of mercury cyanide and isocyanide and their hydride isotopic frequencies [cm\(^{-1}\)] and intensities [km/mol].

| Species          | ν1  | σ (Intensity) | Isotopic Fr ratios\(^a\) | ν2  | σ (Intensity) | ν3  | σ (Intensity) |
|------------------|-----|---------------|--------------------------|-----|---------------|-----|---------------|
| HgCN             | 2265.1 (11) | 351.1 (30)    |                          |     |               |     |               |
| Hg\(^{13}\)CN   | 2215.1 (9)  | 345.6 (29)    |                          |     |               |     |               |
| HgC\(^{15}\)N   | 2232.2 (12) | 345.0 (29)    |                          |     |               |     |               |
| HgNC             | 2140.0 (252) | 376.8 (42)    |                          |     |               |     |               |
| HgN\(^{13}\)C   | 2097.5 (250) | 370.1 (40)    |                          |     |               |     |               |
| Hg\(^{15}\)NC   | 2104.8 (238) | 370.9 (41)    |                          |     |               |     |               |
| HgCN             | 2278.2 (9)  | 432.5 (35)    |                          |     | 2190.5 (93)   |     |               |
| DHCN             | 2277.3 (16) | 432.5 (34)    |                          |     | 1553.7 (45)   |     |               |
| HHg\(^{13}\)CN  | 2228.1 (2)  | 426.0 (34)    |                          |     | 2189.3 (93)   |     |               |
| HHgC\(^{15}\)N  | 2246.8 (6)  | 424.8 (33)    |                          |     | 2189.8 (97)   |     |               |
| HHgNC            | 2167.7 (294) | 463.8 (57)    |                          |     | 2255.6 (32)   |     |               |
| DHNC             | 2168.3 (275) | 463.7 (57)    |                          |     | 1598.9 (28)   |     |               |
| HHgN\(^{13}\)C  | 2125.9 (288) | 455.3 (54)    |                          |     | 2255.4 (37)   |     |               |
| HHg\(^{15}\)NC  | 2131.4 (272) | 456.8 (55)    |                          |     | 2255.4 (38)   |     |               |

CCSD(T)/aug-cc-pVTZ/aug-cc-pVTZ-pp

| Species          | ν1  | Isotopic Fr Ratios\(^a\) | v2  | σ | v4  |
|------------------|-----|--------------------------|-----|---|-----|
| HgCN             | 2266.5 |                           | 346.8 |   |     |
| Hg\(^{13}\)CN   | 2216.6 | 1.02251                  | 341.5 |   |     |
| HgC\(^{15}\)N   | 2233.5 | 1.01478                  | 340.8 |   |     |
| HgNC             | 2072.1 |                           | 366.1 |   |     |
| HgN\(^{13}\)C   | 2031.1 | 1.02019                  | 359.6 |   |     |
| Hg\(^{15}\)NC   | 2044.0 | 1.01375                  | 387.3 |   |     |
| HHgCN            | 2207.3 |                           | 429.7 |   | 2175.7 |
| DHgCN            | 2203.5 |                           | 429.7 |   | 1545.2 |
| HHg\(^{13}\)CN  | 2182.9 | 1.01118                  | 423.3 |   | 2150.3 |
| HHgC\(^{15}\)N  | 2187.3 | 1.00914                  | 422.0 |   | 2164.8 |
| HHgNC            | 2100.9 |                           | 459.7 |   | 2243.9 |
| DHgNC            | 2101.4 |                           | 459.6 |   | 1590.7 |
| HHgN\(^{13}\)C  | 2060.6 | 1.01956                  | 451.3 |   | 2243.8 |
| HHg\(^{15}\)NC  | 2065.5 | 1.01714                  | 452.8 |   | 2243.8 |

\(^a\) (\(^{12}\)C/\(^{13}\)C) or (\(^{14}\)N/\(^{15}\)N).

Figure 9. Partial bonding diagram for HgNC and HHgNC. In HgNC the singly occupied MO is primarily the Hg sigma 6 s AO. The Hg delta lone pairs are isolated (d\(_d\), d\(_\delta\)) AOs, while the Hg (d\(_d\), d\(_\delta\)) pi lone pairs mix with carbon's (p\(_p\), p\(_\pi\)) AOs. These lone pairs lie lower than the Hg–N \(\sigma\) bond and the NC \(\pi\) bonds. In the formation of HHgNC, the new sigma H–Hg bond is a mixture of the HgNC singly occupied 6 s AO and the H 1s AO. The pairing of odd electrons makes the process strongly exothermic and leads to closed shell systems.
The HOMO of the hydride is primarily of the C lone pair, with some destabilizing admixture of the sigma bond.

5.2. Relative Energies of Cyano and Isocyano Systems

The data collected below in Table 8 and pictured in Figure 10 show that the cyano species are consistently more stable than isocyano species, with the difference increasing down Group 12. CCSD and CCSD(T) estimates (kJ/mol) are consistent within about 5 kJ/mol.

Table 8. Relative stability of cyanides and isocyanides: E(isocyano)−E(cyano) [kJ/mol]. Energies obtained with CCSD(T)/aug-cc-pVTZ (Zn) and aug-cc-pVTZ-pp (Cd and Hg).

| Species     | ΔE (Isocyano-Cyano) | ZnNC−ZnCN | HZnNC−HZnCN |
|-------------|---------------------|-----------|-------------|
| CCSD        | 17.3                | 19.2      |             |
| CCSD(T)     | 19.7                | 21.9      |             |
| CdNC−CdCN  | HCdNC−HCdCN         |           |             |
| CCSD        | 27.4                | 30.9      |             |
| CCSD(T)     | 29.3                | 33.0      |             |
| HgNC−HgCN  | HHgNC−HHgCN         |           |             |
| CCSD        | 54.6                | 48.4      |             |
| CCSD(T)     | 55.6                | 50.6      |             |

Figure 10 shows bond lengths and natural charges for the triatomic Zn, Cd, and Hg cyanides and isocyanides and for the tetra-atomic hydride cyanides. Parameters were obtained with CCSD(T)/aug-cc-pVTZ (Zn) and aug-cc-pVTZ-pp (Cd and Hg). Energies are reported in both CCSD and CCSD(T), which agree closely, within 3 kJ/mol.

5.3. Geometry from CCSD and CCSD(T)

In every case the (H)M−C−N metal to cyano carbon bond for cyano is longer than the (H)M−NC metal to nitrogen bond for isocyano systems. This may be a consequence of the smaller atomic radius for N compared with C, or greater inhomogeneity in charge in the isocyano structure compared with the cyano structure. Note, an accompanying shortening of the cyano bond relative to the isocyno bond is observed. Alternatively, this shortening may be a consequence of increasing electron density along the internuclear axis as the M−H bond is formed from the unpaired electrons on H and M. Adding H to form HM CN and HMNC has almost no effect on the C−N and N−C distances, but a shortening effect on the MC and MN bonds.

Microwave spectroscopic investigations of a mixtures of zinc vapor and cyanogen gas in a DC discharge observed the same two molecules ZnCN$^{23}$ and HZnCN$^{24}$ that we find with matrix isolation IR and laser ablated Zn atoms reacting with HCN in excess argon.

5.4. Trends in IR Spectra

Several features of the observed and computed IR spectra can be recognized.

1. Greater intensity of isocyanate stretch relative to cyano stretch: Isocyanides show consistently (almost tenfold) higher isocyanate stretching absorption intensities (CCSD computed) over the cyanides. The high intensities for the isocyanate stretching modes are due to the more substantially nonuniform polar charge distribution in the (H)MNC N to C bond relative to that of the C to N bond in cyanides (see natural charges in Figure 10).

2. Isotopic frequency ratios distinguish cyano from isocyanate species, and provide clues to the amplitudes of motion of C in MCN and N in MNC. The observed $^{12}$C/$^{13}$C isotopic frequency ratios provide a reliable predictor of cyano and isocyanate structures, and allow distinction of radical and closed shell systems. Broadly, cyano systems display ratios near 1.0210 or greater, while isocyanate systems have smaller ratios. CCSD and CCSD(T) computations provide generally reliable estimates of isotopic frequency ratios, consistent with this behavior. The ultimate comparison for C atom movement in a vibration is to compare the $^{12}$C/$^{13}$C frequency ratios between the vibrations in $^{12}$C−$^{13}$O (1.02252, Table 1 observed ratio for CO in solid argon) with the ratios for $^{12}$C/$^{13}$C in the antisymmetric stretching modes for $^{16}$O$^{12}$C$^{16}$O and $^{16}$O$^{13}$C$^{16}$O: Figure S5 shows the featured antisymmetric vibration in OCO spectra which are matrix site split into two bands so we have two values for the carbon dioxide $\nu_3$ vibration, where the frequencies are given in Figure S5. The 12/13 ratios are 2339.2/2273.6 = 1.02885 and 2345.1/2279.5 = 1.02878: These numbers should be the same, but there are very small differences in the anharmonicity in the two isotopomers we are comparing. We round to four decimal places and get 1.0288 which is just an inking larger than the 1.0225 CO value given above. So the C atom moves more in the antisymmetric mode in OCO and in M−C−N by comparison than it does in C−N just as it moves more in OCO than in CO. Thus, the special character of the really M−N−C antisymmetric N stretching mode in the isocyanide sustains more N movement rather than in the MN−C vibration of the cyanide.

3. Shifts in isocyanate stretching frequencies can be rationalized: The observed N−C stretching frequencies for MCN's are similar for the three metals (range 2162−2120 cm$^{-1}$) as are MNC isocyanate stretching frequencies (range 2075−2033). The isocyanate stretching frequency for CdNC (2069.8 cm$^{-1}$) is just 4.8 cm$^{-1}$ less than that of ZnNC (2074.6 cm$^{-1}$), which may be attributed to coupling with the heavier Cd metal and an increase in the motion's reduced mass. The observed HgN−C frequency (2032.7 cm$^{-1}$) is shifted by a greater amount, 37.1 cm$^{-1}$, toward the red. This may be due to the greater relativistic effect for Hg (as modeled by the corrected...
pseudopotential and the basis aug-cc-pVTZ-pp (33) leading to stronger Hg–N interaction and a still greater reduced mass for this asymmetric N motion between Hg and C. The pattern is echoed in the assigned isocyano HMNC frequencies, 2097.8 for M=Zn, a small red shift to 2088.0 for Cd, and a more substantial shift to 2066.2 cm\(^{-1}\) for Hg.

4. **CCSD(T) calculations are useful but must be evaluated with care**: CCSD(T) calculations with an extensive basis (specifically aug-cc-pVTZ for Zn with a small pseudopotential...
(pp) for Cd and Hg) are useful in assigning features of vibrational spectra for H–M cyanides and isocyanides. Relativistic effects, which would be significant for Hg and perhaps also for Cd, are represented in calculations only by the pseudopotential in the basis aug-cc-pVTZ-pp. Such calculations fall short of best results for certain kinds of vibrations. M–H frequencies are overestimated in both CCSD and in CCSD(T). Cyano stretches for MCN radicals are sometimes overestimated, (see discussion below) while isocynano counterparts are accurately represented. HMCN and HMNC vibrations are well-described, more accurately described.

5. **A correlation diagram permits easy evaluation of vibrational assignments:** The correlation between computed and assigned frequencies is shown in Figure 11. The trendline suggests a slight overestimate by MOLPRO (< 2 %) for tetraatomics HM (C, N). Gaussian estimates for MCN are outliers, specifically points 1 (ZnCN), 2 (CdCN) and 3 (HgCN).

6. **Computed frequencies for C–N and M–H stretches,** and for hydrides (Figure 11) show uniform small decreases in value for the sequence H (Zn–Cd–Hg) cyanides. For other systems the pattern is not so simple.

7. **We would like to close with a comparison of the C–N and M–H stretching frequencies** for the molecules investigated here. Figure 12 depicts those observations.

5.5. **The ca. 100 cm⁻¹ Difference between MCN and MNC Radical Modes (see Table 10)**

We observed a nearly constant difference between cyano and isocynano stretching frequencies in a previous study. Summarized in Table 9 are cyano and isocynano frequencies computed by CCSD(T)/aug-cc-pVTZ (with pseudo potential for Cd and Hg). The highest frequencies suffer from the erroneously high values.

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**Figure 11.** Correlation diagram for calculated (vertical axis) and assigned (horizontal axis) stretching frequencies for cyano, isocyan, and M–H modes. Calculations are conducted with the model CCSD(T)/aug-cc-pVTZ with a small pseudopotential (pp) for Cd and Hg. Gaussian calculations are coded orange for triatomics and blue for tetra-atomics. Molpro calculations are coded yellow for triatomics and gray for tetra-atomics. Trendline is for Molpro calculated frequencies vs assigned frequencies for tetra-atomics. Data for this diagram is included in Supplementary Information (Table S3).
for the simple cyanides. These IR-active modes must be of symmetry $\sigma_u$.

We propose that the roughly 100 cm$^{-1}$ difference between the cyano and isocyano isomer frequencies arises from the distinct differences in charge distributions between cyano and isocyano fragments. The uniform distribution in CN (which accounts for the weak IR absorption of its stretch) can be associated with stronger bonding (reflected in its higher stretching frequency) than the isocyano fragment. The persistence of the 100 cm$^{-1}$ difference suggests that the cyano and isocyano vibrations are not strongly coupled to the other modes of these molecules.

In this work we observe a similar gap near 100 cm$^{-1}$ between HMCN and HMNC, as shown in Table 9. However the CCSD(T) cyano stretches for MCN radicals produce a considerably larger differences (Table 10). This complicated the assignment of those stretches. Fortunately the isotopic ratios allowed us to arrive at an unambiguous assignment, as described in the experimental sections above.

An early warning that the MCN system might be exceptional was the ludicrous MP2/aug-cc-pVTZ prediction of a CN stretch in ZnCN over 3000 cm$^{-1}$. In the same basis for Zn, MP2 was well behaved for the closed shell hydrides. CCSD and related models (QCISD, BD) gave results consistent with the 100 cm$^{-1}$ rule of thumb. We have no definite explanation for the MCN anomaly. Within Gaussian 09 and 16, a change to Ahlrichs basis sets (def2-TZVPP) did not alter the anomalies. Oddly, the implementation of CCSD(T) in Molpro produced cyano – isocyano stretching frequency difference (Table 11) near 100 cm$^{-1}$ as expected.
Che m P hys C h e

The Zn–H stretches still elude accurate estimation. CCSD(T)/aug-cc-pVTZ frequencies for Gaussian and Molpro are shown in Table 12.

We observed anomalies in CCSD(T) calculations of cyano stretching frequencies. As a direct result of this major discrepancy CASPT2 (7,6) calculations were conducted. They provided agreement within 20 cm⁻¹ and improved the accuracy of the CCSD(T) calculations by fixing the electronic structure to the appropriate configurations.

6. Conclusions

Laser ablated Group 12 metals react with hydrogen cyanide to first form the MCN and MNC molecules and then their monohydrides HMCN and HMNC. The ease of hydrogen removal from HCN leads to a high yield of the cyanyl radical, CN, and provides a direct route to the formation of the simple metal cyanides and isocyanides. The hydrides grow markedly on annealing the solid argon matrix to 10, 15 or 20 K, owing to metal cyanides and isocyanides. The hydrides were, of course, not radical reactions are strongly exothermic according to this work. Hongmin Li acknowledges Postdoctoral support from China Scholarship Council (File No. 20180260104) and the German Academic Exchange Service (File No. 91695363) for a CSC-DAAD Postdoctoral Fellowship in Berlin. We gratefully acknowledge the Freie Universität Berlin for the allocation of computing resources. We thank the ERC Project HighPotOx for continuous support. Open access funding enabled and organized by Projekt DEAL.

Conflict of Interest

The authors declare no conflict of interest.

Keywords: cyanides · electronic structure · Group 12 · isocyanides · zinc

References

[1] D. V. Lanzisera, L. Andrews, J. Am. Chem. Soc. 1997, 119, 6392–6398.
[2] D. V. Lanzisera, L. Andrews, J. Phys. Chem. A 1997, 101, 9660–9665.
[3] X. Chen, Q. Li, L. Andrews, Y. Gong, J. Phys. Chem. A 2018, 122, 7099–7106.
[4] D. V. Lanzisera, L. Andrews, J. Phys. Chem. A 1997, 101, 9666–9672.
[5] X. Chen, Q. Li, L. Andrews, Y. Gong, J. Phys. Chem. A 2017, 121, 8835–8842.
[6] T. M. Greene, W. Brown, L. Andrews, A. J. Downs, G. V. Chertihin, N. Runeberg, P. Pyykkö, J. Phys. Chem. 1995, 99, 7925–7934.
[7] X. Wang, L. Andrews, J. Phys. Chem. A 2004, 108, 11006–11013.
[8] X. Wang, L. Andrews, Phys. Chem. Chem. Phys. 2005, 7, 750–759.
[9] L. Andrews, X-F. Wang, Y. Gong, T. Schloder, S. Riedel, M. J. Franger, Angew. Chem. Int. Ed. 2012, 51, 8235–8238; Angew. Chem. 2012, 124, 8339–8363.
[10] L. Andrews, H.-G. Cho, Organometallics 2006, 25, 4040–4053.
[11] a) T. Vent-Schmidt, L. Andrews, S. Riedel, J. Phys. Chem. A 2015, 119, 2253–2261; b) T. Vent-Schmidt, J. Metzger, L. Andrews, S. Riedel, J. Fluorine Chem. 2015, 174, 2–7.

Table 12. Collected M–H stretching frequencies calculated by MOLPRO and Gaussian CCSD(T)/aug-cc-pVTZ (Zn) and aug-cc-pVTZ-pp (Cd, Hg), with assigned observed frequencies.

| M     | H–M | Isocyanide | H–M | Difference | Software |
|-------|-----|------------|-----|------------|----------|
| HZnCN | 2052| 2008 (2011)| 2074| 22         | Molpro   |
|       | 1996|            | 1972| 19 (17)    | Gaussian |
| HCdCN | 1908| HCdNC      | 1925| 17         | Molpro   |
|       | 1907|            | 1949| 42         | Gaussian |
|       | 1865|            | 1884| –19        | As assigned |
| HHgCN | 2149| HHgNC      | 2211| 62         | Molpro   |
|       | 2176|            | 2244| 68         | Gaussian |
|       | 1996|            | 1972|            | As assigned |

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