



Article Diiron Aminocarbyne Complexes with NCE⁻ Ligands (E = O, S, Se)

Giulio Bresciani ^{1,2,*}, Stefano Zacchini ^{2,3}, Guido Pampaloni ^{1,2}, Marco Bortoluzzi ^{2,4}, and Fabio Marchetti ^{1,2}

- ¹ Department of Chemistry and Industrial Chemistry, University of Pisa, Via G. Moruzzi 13, I-56124 Pisa, Italy
- ² Interuniversity Consortium for Chemical Reactivity and Catalysis, CIRCC, Via Celso Ulpiani 27, I-70126 Bari, Italy
- ³ Department of Industrial Chemistry "Toso Montanari", University of Bologna, Viale Risorgimento 4, I-40136 Bologna, Italy
- ⁴ Department of Molecular Science and Nanosystems, University of Venezia "Ca' Foscari", Via Torino 155, I-30170 Mestre, Italy
- * Correspondence: giulio.bresciani@dcci.unipi.it

Abstract: Diiron μ -aminocarbyne complexes [Fe₂Cp₂(NCMe)(CO)(μ -CO){ μ -CN(Me)(R)}]CF₃SO₃ (R = Xyl, [1a^{NCMe}]CF₃SO₃; R = Me, [1b^{NCMe}]CF₃SO₃; R = Cy, [1c^{NCMe}]CF₃SO₃; R = CH₂Ph, [1d^{NCMe}]CF₃SO₃), freshly prepared from tricarbonyl precursors [1a-d]CF₃SO₃, reacted with NaOCN (in acetone) and NBu₄SCN (in dichloromethane) to give [Fe₂Cp₂(kN-NCO)(CO)(μ -CO){ μ -CN(Me)(R)}] (R = Xyl, 2a; Me, 2b; Cy, 2c) and [Fe₂Cp₂(kN-NCS)(CO)(μ -CO){ μ -CN(Me)(CH₂Ph)}], 3 in 67–81% yields via substitution of the acetonitrile ligand. The reaction of [1a^{NCMe}]CF₃SO₃ with KSeCN in THF at reflux temperature led to the cyanide complexes [Fe₂Cp₂(CN)(CO)(μ -CO){ μ -CNMe(R)}], 6a–c (45–67%). When the reaction of [1a^{NCMe}]CF₃SO₃ with KSeCN was performed in acetone at room temperature, subsequent careful chromatography allowed the separation of moderate amounts of [Fe₂Cp₂(kSe-SeCN)(CO)(μ -CO){ μ -CN(Me)(Xyl)}], 4a, and [Fe₂Cp₂(kN-NCSe)(CO)(μ -CO){ μ -CN(Me)(Xyl)}], 5a. All products were fully characterized by elemental analysis, IR, and multinuclear NMR spectroscopy; moreover, the molecular structure of *trans*-6b was ascertained by single crystal X-ray diffraction. DFT calculations were carried out to shed light on the coordination mode and stability of the {NCSe-} fragment.

Keywords: organometallic chemistry; diiron complexes; aminocarbyne ligand; selenocyanate; substitution reactions

1. Introduction

Pseudohalides of the general formula $\{NCE\}^-$ (E = O, S, Se) are ubiquitous and largely employed in coordination chemistry, and they may behave as ligands through either the nitrogen or chalcogen atom according to the nature of the metal center [1–11]. Metal binding of $\{NCO\}^-$ through the nitrogen atom (isocyanate) is predominant over the *O*coordination (cyanate), and activation of the isocyanate ligand by nucleophilic addition to the unsaturated carbon atom may be subsequently viable [12,13]. On the other hand, coordination linkage isomerism is often observed with complexes comprising an $\{NCE\}^$ ligand (E = S, Se) [14–17], and coordination switching from nitrogen (isothiocyanate) to sulfur (thiocyanate) may be achieved upon heating or UV irradiation [18].

A variety of selenocyanate/isoselenocyanate complexes have been synthesized [19–22], and thermogravimetric analyses on cobalt(II)-selenocyanate complexes evidenced stability up to 200 $^{\circ}$ C [23].

Notwithstanding, the metal coordination of the {NCSe}⁻ group might be unstable, resulting in decomposition into the cyanide ion and selenium atom; this reaction was previously reported with copper(II) acetate [24,25] and was exploited for catalytic purposes in cyanation reactions. Selenocyanate to cyanide degradation was also observed in human



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Copyright: © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). blood, and the ferric ion is typically involved in this process [26]. However, to the best of our knowledge, the clean isolation of a metal–cyanide derivative following activation of the coordinated {NCSe}⁻ moiety is a rare feature. Note that the reverse reaction, i.e., the cyanation of elemental selenium, is typically conducted to prepare selenocyanate compounds [27,28].

The $\{Fe_2Cp_2(CO)_x\}$ skeleton (x = 2–3) is a versatile scaffold for building organometallic architectures that take advantage of metal–metal cooperativity [29–32], and it can be used to explore novel reactivity patterns exploiting an earth-abundant metal element [33–35].

In this framework, in the last 20 years, our research has focused on the chemistry of diiron complexes with a bridging aminocarbyne ligand [36–38], and we previously reported the synthesis of isocyanate and isothiocyanate derivatives from a labile acetonitrile precursor (Scheme 1) [39]. The reaction with tetrabutylammonium thiocyanate at room temperature selectively afforded the *N*-coordinated product with the Cp ligands mutually *trans* oriented with respect to the Fe–Fe axis, and then thermal conversion into the *cis* isomer was observed without affecting the isothiocyanate moiety. On the other hand, the installation of the isocyanate ligand proceeded at high temperature only. We also demonstrated in one case that methylation of the isothiocyanate ligand was straightforward, affording a thiomethyl-nitrile species.



Scheme 1. Synthesis of diiron μ -aminocarbyne complexes with isocyanate and isothiocyanate as the terminal ligands and *S*-methylation reaction. Xyl = 2,6-C₆H₃Me₂.

Here, we extended the chemistry of diiron aminocarbyne complexes with pseudohalide ligands, including reactions with a selenocyanate source. Structural and thermodynamic aspects were elucidated by means of DFT calculations.

2. Results and Discussion

The diiron aminocarbyne complexes $[1a-d]CF_3SO_3$ were prepared from commercially available $[Fe_2Cp_2(CO)_4]$ following the appropriate literature procedures (see Experimental). To evaluate the influence of the aminocarbyne ligand on the substitution reactions herein discussed, we considered a series of R substituents bearing different steric and electronic properties (Scheme 2). In fact, previous findings indicated that R may significantly affect the chemistry of this class of complexes [37]. Compounds $[1a-d]CF_3SO_3$ were converted into the respective acetonitrile adducts [40] using the trimethylamine-*N*-oxide (TMNO) strategy, which is often reliable with cationic complexes based on the {M₂Cp₂(CO)₃} core



with M = Fe or Ru [41–44]. The resulting derivatives $[1a^{NCMe}-1d^{NCMe}]CF_3SO_3$ were used in all cases as freshly prepared reactants.

Scheme 2. Installation of chalcogen cyanate ligands on diiron μ -aminocarbyne complexes and thermal selenocyanate to cyanide conversion (Xyl = 2,6–C₆H₃Me₂; Cy = C₆H₁₁, cyclohexyl; Ph = C₆H₅). Yields in parentheses.

The reactions of [1a^{NCMe}-1c^{NCMe}]CF₃SO₃ with sodium cyanate were performed in acetone at room temperature and afforded the neutral complexes 2a-c in 67-81% yields directly as cis isomers, in alignment with the major stability of this configuration that is usually exhibited by complexes based on the $[Fe_2(CO)_2Cp_2]$ frame [36]. Thus, the IR and NMR spectra of the previously reported complex 2b (see Scheme 1) matched the literature data [39]. The IR spectra of 2a and 2c (in CH_2Cl_2 , 2300–1500 cm⁻¹ spectral region) were similar to that of 2b, consisting of three bands related to the coordinated cyanate ligand and the terminal and bridging carbonyl ligands (e.g., at 2242, 1986, and 1818 cm^{-1} , respectively, in the case of 2a). The major stability of N-coordination (compared to O-coordination) of the $\{NCO\}$ moiety was as expected for a low-valent iron center [45-47]; otherwise, the preference for O-coordination of potential N- and O-donors is commonly observed with high-valent metal complexes [48-50]. Note that O-coordination towards the Fe¹ center in related systems has been rarely observed and only as part of a coordination by means of multidentate hydrocarbyl ligands [51,52]. In **2c**, the band due to the μ -CN moiety fell at 1540 cm⁻¹, in alignment with some double bond character (vide infra). In **2b**, the corresponding band was at 1578 cm^{-1} .

The NMR spectra of **2b** displayed one set of resonances, while the NMR spectra of **2a** and **2c** contained two sets of resonances ascribable to the α and β isomers, differing in the orientation of the *N*-substituents with respect to the cyanate ligand. This kind of isomerism was previously reported for diiron and diruthenium complexes of the type $[M_2Cp_2(L)(CO)(\mu-CO)\{\mu-CN(Me)(R)\}]^{0/+}$ with $R \neq Me$ [36,37]. The α isomer (R pointing

to NCO) was slightly prevalent with respect to the β isomer (R pointing to terminal CO) in the case of **2c** (α/β ratio = 1.3), whereas the bulkier xylyl group made the α form much more favorable in **2a** (α/β ratio = 4.5). Notably, that rotation of the amine group around the carbyne-nitrogen axis was inhibited due to the substantial C=N double bond character [36,37]. Ongoing from a to b, the *N*-Me resonance shifted to high fields by ca. 0.3 ppm in the ¹H NMR spectra. The salient ¹³C NMR feature was provided by the carbyne carbon, resonating within the 329–338 ppm range [36,53].

Motivated by our interest in the chemistry of carbamato species [54–56] and based on the documented reactivity of isocyanates [12,57,58], we tested the reactivity of **2a**,**c** with a range of alcohols and amines, but no reaction occurred even under high-temperature conditions. The substantial inertness of the isocyanate ligand in **2a** was observed even towards strong electrophiles (i.e., methyl triflate and trimethylsilyl triflate).

The reaction of the benzyl-aminocarbyne complex $[1d^{NCMe}]CF_3SO_3$ with tetrabutylammonium thiocyanate was conducted in dichloromethane at room temperature and led to **3**, which was finally isolated in 80% yield. The IR spectrum of **3** (in CH₂Cl₂) displayed carbonyl absorptions at 1970 and 1810 cm⁻¹, which were indicative of the *trans* configuration of the Cp ligands. Moreover, the {NCS} group was *N*-coordinated to the iron center, on account of an infrared absorption at 2114 cm⁻¹. This value was very close to that previously detected in other {Fe-NCS} species [2,15,39]. The NMR spectra of **3** pointed out the occurrence of α/β isomerism, with the α isomer prevailing. The same behavior was observed when related methyl- and benzyl-aminocarbyne complexes were used (Scheme 1) [39].

The reactivity of the diiron aminocarbyne complexes with a selenium compound was investigated for the first time by allowing [1a^{NCMe}-1c^{NCMe}]CF₃SO₃ to react with potassium selenocyanate in acetone. When these reactions were conducted at room temperature, complicated mixtures of products were afforded, including modest amounts of 6a-c. Careful alumina chromatography on the mixture arising from [1a^{NCMe}]CF₃SO₃ allowed the separation of three components, which were spectroscopically analyzed and thus identified as complexes 4a (15% yield), 5a (13%), and 6a (48%). Complexes 4a and 5a comprised Seand N-coordinated selenocyanate ligands, respectively, while **6a** was a cyanide adduct. The infrared absorption for the pseudohalide ligand was detected at 2113 (4a) and 2109 cm^{-1} (5a); in general, the stretching vibration of a metal-coordinated {SeCN} group occurred at higher frequencies when it was Se-coordinated rather than N-coordinated [21,59–62]. The ⁷⁷Se NMR spectra of 4a and 5a clearly pointed out the different coordinations of the {SeCN} moiety. Thus, two signals were recognized in the 77 Se spectrum of 4a, at -232.5 (major) and -246.9 ppm (minor) [63,64], while the ⁷⁷Se spectrum of 5a displayed a unique signal at -340.2 ppm. This picture was consistent with the literature data reported for other complexes and the general trend whereby ⁷⁷Se NMR shielding increases from {M-SeCN} ongoing to {M-NCSe} [65]. The ¹H and ¹³C NMR spectra of 4a displayed two sets of resonances, attributed to *cis* and *trans* isomers, while the ¹H and ¹³C NMR spectra of 5a closely resembled those related to the homologous complex featuring a *cis* arrangement of the Cp rings and an N-coordinated NCS ligand [39]. In the ¹³C NMR spectrum of 4a, the resonance for the selenocyanate ligand occurred above 128.7 ppm, whereas the iso-selenocyanate resonated at 108.4 ppm in the 13 C NMR spectrum of **5a** [63,66,67]. The signal for the aminocarbyne carbon was upfield shifted in 5a (340.4 ppm) compared to 4a (345.4 ppm in the *trans* isomer), suggesting a different degree of back-donation from the diiron backbone to the carbyne in **4a** and **5a** [36]. Formation of *cis/trans* mixtures is believed to be consequent to rotation around the Fe–Fe bond (Adams–Cotton mechanism) [68,69], which is operative during the nitrile substitution process. In the majority of the cases, trans isomers based on the $Fe_2Cp_2(CO)_x$ (x = 2, 3) scaffold are kinetic and less thermodynamically favored products, which might be observed due to a combination of electronic and steric effects [36,37,39,70]. In the case of 4a and 5a, stability studies revealed that the *trans* to *cis* route was not viable in boiling THF solution, whereas formation of cyanide complexes was observed (vide infra). Evidence for the formation of (iso)selenocyanate complexes (IR

spectroscopy) was supplied by the room temperature reactions of [**1b**,c^{NCMe}]**CF**₃**SO**₃ with KSeCN, but attempts to isolate and characterize the products failed.

With the aim of elucidating structural aspects, DFT calculations were carried out on the products obtained from the reaction of [1a^{NCMe}]⁺ with selenocyanate, taking into account the spectroscopic outcomes. Views of the most stable cis and trans isomers of 4a and of the most stable *cis* isomer of **5a** are provided in Figure 1; all structures exhibited an α arrangement of the substituents on the aminocarbyne moiety with respect to the seleniumcontaining ligand. Selected computed bond lengths are summarized in the caption of Figure 1. The Se-coordination of the selenocyanate anion was meaningfully bent, with computed Fe2-Se-C angles between 107° and 114°. On the other hand, the alternative *N*-coordination was almost linear, with the computed Fe2–*N*–C angle in *cis*-5a around 180°. The C–N distance in selenocyanate was scarcely affected by the coordination mode, while on the other hand, the C–Se bond was elongated by more than 0.03 Å when the bonding to iron occurred with the selenium atom. A comparison of the computed bond lengths between *cis*-4a and *cis*-5a revealed that the Fe2–C bond lengths were negligibly affected by the selenocyanate coordination mode. In accordance with the experimental outcomes, the $v_{\rm CN}$ stretching of selenocyanate was predicted at slightly higher wavenumbers for the 4a isomers (unscaled values 2322 cm⁻¹ for *cis*-4a and 2316 cm⁻¹ for *trans*-4a) with respect to 5a (unscaled value 2314 cm⁻¹). From a thermodynamic point of view, *cis*-4a was less stable than *cis*-5a by about 7.7 kcal mol⁻¹, and the Gibbs energy difference between *trans*-4a and *cis*-4a was about 4.7 kcal mol⁻¹ in favor of the *cis* isomer. The relative Gibbs energy values, therefore, indicated that the reaction of [1a^{NCMe}]⁺ with selenocyanate afforded a mixture of kinetic products and that conversion of one product into another could take place as promoted by alumina during chromatography. The preference for N- rather than Se-coordination was computationally established for the mono iron systems [3].

Complex **6a** displayed an infrared band at 2090 cm⁻¹, accounting for iron *N*-coordinated cyanide receiving a significant back-donation [71–73]; the carbonyl ligands manifested themselves as two IR bands at 1959 and 1808 cm⁻¹, suggesting the *trans* configuration of the Cp ligands [74]. In fact, the *cis* isomer of **6a** was previously synthesized from the room temperature reaction of [**1a-NCMe**] with NBu₄CN [75], and its IR spectrum in the same conditions (CH₂Cl₂ solution) consisted of three absorptions at 2091 (C \equiv N), 1982 (CO), and 1804 cm⁻¹ (CO). The ¹H NMR spectrum of **6a** revealed the presence of a minor amount of the *cis* isomer (<15%).

It appeared that the isolation of **6a** from the reaction of $[2a^{NCMe}]CF_3SO_3$ with KSeCN at room temperature was the result of the preliminary coordination of selenocyanate, followed by a rearrangement giving **6a** and releasing one atom of selenium. To confirm this hypothesis, we performed the same reaction in tetrahydrofuran at reflux temperature; in this condition, **6a** was the only isolated product accompanied by the formation of a black solid (presumably elemental selenium). The elimination of selenium probably followed activation of the *Se*-C bond, therefore compounds of type **4a** were most likely involved in the reaction. The computed Gibbs free energy variation for the reaction $trans-4a \rightarrow trans-6a + 1/8 \text{ Se}_8$ was negative by 8.0 kcal mol⁻¹, in alignment with a thermodynamically favorable process. Reactants and products are depicted in Figure 2.



Figure 1. DFT-optimized structures of *cis*-4a, *trans*-4a, and *cis*-5a (C-PCM/PBEh-3c, acetone as continuous medium). Color map: Fe, green; Se, dark yellow; O, red; N, blue; C, grey; hydrogen atoms omitted for clarity. Selected computed bond lengths (Å): *cis*-4a, Fe1–C(CO) 1.766, Fe1–C(μ -CO) 1.994, Fe1–C(carbyne) 1.856, Fe1–C(Cp, average) 2.113, Fe2–Se 2.440, Fe2–C(μ -CO) 1.839, Fe2–C(carbyne) 1.855, Fe2–C(Cp, average) 2.117, Fe1—Fe2 2.490, *Se*–C 1.813, C–N(SeCN) 1.159, C–N(carbyne) 1.298; *trans*-4a, Fe1–C(CO) 1.760, Fe1–C(μ -CO) 2.077, Fe1–C(carbyne) 1.842, Fe1–C(Cp, average) 2.121, Fe2–Se 2.449, Fe2–C(μ -CO) 1.821, Fe2–C(carbyne) 1.864, Fe2–C(Cp, average) 2.134, Fe1–Fe2 2.535, *Se*–C 1.809, C–N(SeCN) 1.160, C–N(carbyne) 1.299; *cis*-5a, Fe1–C(CO) 1.763, Fe1–C(μ -CO) 2.002, Fe1–C(carbyne) 1.847, Fe1–C(Cp, average) 2.108, Fe2–N 1.916, Fe2–C(μ -CO) 1.840, Fe2–C(carbyne) 1.856, Fe2–C(Cp, average) 2.122, Fe1–Fe2 2.484, *Se*–C 1.776, C–N(SeCN) 1.162, C–N(carbyne) 1.295. Selected computed angles (°): *cis*-4a, Fe1–C(carbyne)–Fe2 84.3, Fe2–Se–C 107.5; *trans*-4a, Fe1–C(carbyne)–Fe2 86.3, Fe2–Se–C 113.5; *cis*-5a, Fe1–C(carbyne)–Fe2 84.3, Fe2–N–C 18.0. Cartesian coordinates are provided in the Supplementary Materials.



Figure 2. DFT-optimized structures of *trans*—4a, *trans*—6a, and Se₈ (C-PCM/PBEh–3c, acetone as continuous medium). Color map: Fe, green; Se, dark yellow; O, red; N, blue; C, grey; hydrogen atoms omitted for clarity. Selected computed bond lengths (Å): *trans*—6a, Fe1–C(CO) 1.760, Fe1–C(µ–CO) 2.089, Fe1–C(carbyne) 1.839, Fe1–C(Cp, average) 2.119, Fe2–C(CN) 1.893, Fe2–C(µ–CO) 1.802, Fe2–C(carbyne) 1.858, Fe2–C(Cp, average) 2.123, Fe1–Fe2 2.523, C–N(CN) 1.162, C–N(carbyne) 1.300; Se₈, Se–Se(average) 2.319. Cartesian coordinates are provided in the Supplementary Materials.

Similarly, the thermal reactions of $[2b,c^{\text{NCMe}}]CF_3SO_3$ with KSeCN provided a direct route to the cyanide complexes 6b-c (53–67% yields). These results confirmed the low thermal stability of the selenocyanate complexes with respect to the corresponding cyanide derivatives. The IR spectra of 6b-c were quite similar to that of 6a, thus suggesting the predominance of *trans* species. The NMR spectra of 6b-c displayed two sets of resonances. These were attributed to *trans* and *cis* isomers in the case of 6b, with the former largely prevalent (83%), and to α and β isomers in the case of 6c (isomer ratio 1.3). The diagnostic ¹³C resonance for the cyanide ligand occurred within the range 139.7–141.4 ppm. Former computational studies highlighted the higher stability of the *trans* isomer of 6b with respect to the corresponding *cis* isomer [76].

The X-ray structure of *trans*-6b was determined by single crystal X-ray diffraction (Figure 3). This represented a very rare case of the $[Fe_2Cp_2(L)(CO)(\mu-CO)\{\mu-CN(Me)(R)\}]^n$ (L = mono-anionic ligand, n = 0; L = neutral ligand, n = 1+) complex possessing a *trans* geometry. The overall structure and bonding parameters were very similar to those reported for $[Fe_2Cp_2(NCS)(CO)(\mu-CO)\{\mu-CN(Me)_2\}]$ [39]. The Fe(2)–C(4) interaction [1.895(2) Å] was longer than Fe(1)–C(1) [1.758(2) Å] since the terminal cyanide ligand is a poorer π -acceptor than terminal CO. This caused a considerable asymmetry of the bridging μ -CO ligand, with Fe(1)–C(2) [1.999(2) Å] being significantly longer than Fe(2)–C(2) [1.863(2) Å]. This asymmetry was less marked in the bridging μ -CNMe₂ ligand [Fe(1)–C(3) = 1.889(2) Å; Fe(2)–C(3) = 1.850(2) Å]. A similar trend was previously observed in *cis*-[Fe₂(C₅H₄Me)₂(CN)(CO)(μ -CO)(μ -CO)(μ -CN(Me)₂]] [77].



Figure 3. View of the molecular structure (ORTEP drawing) of *trans*-6b. Displacement ellipsoids are at the 30% probability level. Selected bond lengths (Å) and angles (°): Fe(1)–Fe(2) 2.5166(4), Fe(1)–C(1) 1.758(2), Fe(1)–C(2) 1.999(2), Fe(2)–C(2) 1.863(2), Fe(1)–C(3) 1.889(2), Fe(2)–C(3) 1.850(2), Fe(2)–C(4) 1.895(2), C(1)–O(1) 1.144(3), C(2)–O(2) 1.173(3), C(3)–N(1) 1.299(3), C(4)–N(2) 1.149(3), N(1)–C(5) 1.473(3), N(1)–C(6) 1.480(3), Fe(1)–C(1)–O(1) 178.4(2), Fe(1)–C(2)–Fe(2) 81.25(9), Fe(1)–C(3)–Fe(2) 84.61(9), Fe(2)–C(4)–N(2) 177.3(2), C(3)–N(1)–C(5) 123.7(2), C(3)–N(1)–C(6) 122.91(19), C(5)–N(1)–C(6) 113.33(18).

3. Experimental

3.1. Materials and Methods

Reactants and solvents were purchased from Alfa Aesar, Merck, Strem, or TCI Chemicals and were of the highest purity available. Diiron complexes [1a–d]CF₃SO₃ were prepared according to the literature [78,79]. Reactions were conducted under N₂ atmosphere using standard Schlenk techniques. Products were stored in air once isolated. Dichloromethane and tetrahydrofuran were dried using the solvent purification system mBraun MB SPS5, while acetonitrile was distilled from CaH₂. IR spectra of solutions were recorded using a CaF₂ liquid transmission cell (2300–1500 cm⁻¹) on a Perkin Elmer Spectrum 100 FTIR spectrometer. IR spectra were processed with Spectragryph software [80]. ¹H, ¹³C, and ⁷⁷Se NMR spectra were recorded at 298 K on a Jeol JNM-ECZ500R instrument equipped with a Royal HFX Broadband probe. Chemical shifts (expressed in parts per million) were referenced to the residual solvent peak in ¹H and ¹³C NMR spectra [81] and to an external standard (Me₂Se) in ⁷⁷Se NMR spectra. NMR spectra were assigned with the assistance of ¹H–¹³C (*gs*-HSQC and *gs*-HMBC) correlation experiments [82]. NMR signals due to secondary isomeric forms (where it is possible to detect them) are italicized. Elemental analyses were performed using a Vario MICRO cube instrument (Elementar).

Cyanate complexes [Fe₂Cp₂(k*N*-NCO)(CO)(μ -CO){ μ -CN(Me)(R)}] (R = Xyl, 2a; Me, 2b; Cy, 2c).

3.2. General Procedure

A solution of $[1a-c]CF_3SO_3$ (*ca* 0.4 mmol) in MeCN (20 mL) was treated with Me₃NO·2H₂O (1.1 eq.) and the resulting mixture was stirred for 1 h, during which progressive color darkening was observed. The conversion of the starting material into the acetonitrile adduct $[1^{NCMe}]^+$ was checked by IR spectroscopy, as is routine for this type of reaction [75]. Volatiles were removed under vacuum to give a brown residue, which

was dissolved in deaerated acetone (30 mL), and NaOCN (3.0 eq.) was added to this solution. The resulting mixture was stirred for 18 h at room temperature and then the solvent was evaporated under reduced pressure. The resulting solid was dissolved in the minimum volume of CH_2Cl_2 and this solution was charged on an alumina column. Impurities were separated using neat CH_2Cl_2 and neat THF as eluents, and then a brown fraction corresponding to **2a**–**c** was collected with MeCN. The solvent was removed under reduced pressure, and the residue was suspended in hexane for 24 h. After filtration, the separated solid was dried under vacuum.

[Fe₂Cp₂(kN-NCO)(CO)(µ-CO){µ-CN(Me)(Xyl)}], 2a (Figure 4).



Figure 4. Structure of [Fe₂Cp₂(kN-NCO)(CO)(µ-CO){µ-CN(Me)(Xyl)}], 2a.

From [1a]CF₃SO₃ (268 mg, 0.432 mmol), Me₃NO·2H₂O (53 mg, 0.47 mmol), and NaOCN (187 mg, 1.30 mmol). Dark brown solid, yield 157 mg (75%). Anal. calcd. for C₂₃H₂₂N₂O₃Fe₂: C, 56.83; H, 4.56; N, 5.76. Found: C, 56.66; H, 4.62; N, 5.68. IR (CH₂Cl₂): $\tilde{\nu}$ /cm⁻¹ = 2242 w-br (NCO), 1986 vs. (CO), 1818 s (µ-CO). ¹H NMR (CDCl₃): δ /ppm = 7.40–7.28 (m, 3 H, C₆H₃); 5.09, 4.97, 4.45, 4.29 (s, 10 H, Cp); 4.78, 4.47 (s, 3 H, NMe); 0.66, 2.10, 2.06, 1.91 s, 6 H, C₆H₃Me₂). ¹³C{¹H} NMR (CDCl₃): δ /ppm = 338.2 (µ-CN); 264.6 (µ-CO); 211.5 (CO); 148.4 (C-*ipso*); 133.0, 132.8, 130.4, 129.2, 129.1 (C₆H₃); 129.8 (NCO); 88.2, 88.1 (Cp); 53.6 (NMe); 19.2, 18.1 (C₆H₃Me₂). Isomer ratio (α/β) = 4.5.

[Fe₂Cp₂(kN-NCO)(CO)(μ-CO){μ-CN(Me)(Me)}], 2b (Figure 5) [39].



Figure 5. Structure of [Fe₂Cp₂(k*N*-NCO)(CO)(μ-CO){μ-CN(Me)(Me)}], **2b** [39].

From **[1b]CF₃SO₃** (109 mg, 0.205 mmol), Me₃NO·2H₂O (27 mg, 0.25 mmol), and NaOCN (40 mg, 0.61 mmol). Brown solid, yield 66 mg (81%). Anal. calcd. for C₁₆H₁₆N₂O₃Fe₂: C, 48.53; H, 4.07; N, 7.07. Found: C, 48.72; H, 4.15; N, 7.00. IR (CH₂Cl₂): $\tilde{\nu}$ /cm⁻¹ = 2238 vs. (NCO), 1981 vs. (CO), 1804 s (µ-CO), 1578 w (µ-CN). ¹H NMR (CDCl₃): δ /ppm = 4.74, 4.62 (s, 10 H, Cp); 4.6 *, 4.21 (s, 6 H, NMe₂); * Hidden by Cp resonance.

[Fe₂Cp₂(k*N*-NCO)(CO)(μ-CO){μ-CN(Me)(Cy)}], 2c (Figure 6).



Figure 6. Structure of [Fe₂Cp₂(kN-NCO)(CO)(μ-CO){μ-CN(Me)(Cy)}], 2c.

From [1c]CF₃SO₃ (256 mg, 0.427 mmol), Me₃NO·2H₂O (52 mg, 0.47 mmol), and NaOCN (184 mg, 1.28 mmol). Dark brown solid, yield 133 mg (67%). Anal. calcd. for C₂₁H₂₄N₂O₃Fe₂: C, 54.35; H, 5.21; N, 6.04. Found: C, 54.29; H, 5.30; N, 6.08. IR (CH₂Cl₂): $\tilde{\nu}$ /cm⁻¹ = 2255 m (NCO); 1984 vs. (CO), 1818 s (µ-CO), 1540 w (µ-CN). ¹H NMR (CDCl₃): δ /ppm = 5.59, 4.8 * (m, 1 H, CH^{Cy}); 4.99, 4.90, 4.87, 4.81 (s, 10 H, Cp); 4.44, 4.08 (s, 3 H, NMe); 2.71–2.46, 2.22–2.11, 1.82–1.49, 1.39–1.21 (m, 10 H, CH₂^{Cy}). * Partially hidden by Cp resonances. ¹³C{¹H} NMR (CDCl₃): δ /ppm = 329.3, 328.9 (µ-CN); 266.4, 266.1 (µ-CO); 211.6, 210.9 (CO); 130.3, 130.0 (NCO); 88.6, 88.1, 87.2, 86.9 (Cp); 78.1 (CH^{Cy}); 45.9, 45.6 (NMe); 32.5, 31.3, 30.9, 28.6, 26.0, 25.9, 25.7, 25.2, 25.1, 24.4 (CH₂^{Cy}). Isomer ratio (α/β) = 1.3.

Thiocyanate complex [Fe₂Cp₂(kN-NCS)(CO)(µ-CO){µ-CN(Me)(CH₂Ph)}], 3 (Figure 7).



Figure 7. Thiocyanate complex [Fe₂Cp₂(kN-NCS)(CO)(µ-CO){µ-CN(Me)(CH₂Ph)}], 3.

A mixture of $[1d^{NCMe}]CF_3SO_3$, freshly generated from $[1d]CF_3SO_3$ (90 mg, 0.148 mmol) according to the procedure above, and NBu₄SCN (220 mg, 0.733 mmol) in CH₂Cl₂ (12 mL) was stirred for 3 h at room temperature. The final solution was directly charged on an alumina column, and elution with neat dichloromethane afforded the fraction corresponding to the title product. Thus, volatiles were removed under vacuum to give an orange solid. Yield 58 mg (80%). Anal. calcd. for C₂₂H₂₀Fe₂N₂O₂S: C, 54.13; H, 4.13; N, 5.74; S, 6.57. Found: C, 54.32; H, 4.04; N, 5.62; S, 6.49. IR (CH₂Cl₂): $\tilde{\nu}/cm^{-1} = 2114$ s (NCS); 1970 vs. (CO), 1810 s (μ -CO), 1535 w (μ -CN). ¹H NMR (CDCl₃): δ /ppm = 7.59–7.39 (m, 5 H, CH₂Ph); 6.90, 6.17, 5.87, 5.50 (d, ²J_{HH} = 14 Hz, 2 H, CH₂Ph); 4.92, 4.83, 4.61, 4.56 (s, 10 H, Cp); 4.47, 4.06 (s, 3 H, NMe). ¹³C{¹H} NMR (CDCl₃): δ /ppm = 341.5, 340.6 (μ -CN); 264.1, 263.2 (μ -CO); 212.3, 211.6 (CO); 141.2, 140.6 (NCS); 134.2–127.1 (CH₂Ph); 89.5, 89.4, 88.2 (Cp); 71.0, 70.5 (CH₂Ph); 49.8, 49.0 (NMe). Isomer ratio (α/β) = 2.

Reactivity of [1a-NCMe]CF₃SO₃ with KSeCN: isolation of 4a, 5a and 6a.

The acetonitrile adduct $[1a^{NCMe}]CF_3SO_3$ was prepared from $[1a]CF_3SO_3$ (176 mg, 0.283 mmol) and Me₃NO·2H₂O (35 mg, 0.31 mmol), as described above. Then, $[1a^{NCMe}]CF_3SO_3$ was dissolved in deaerated acetone (30 mL), and KSeCN (122 mg, 0.85 mmol) was added. The resulting mixture was stirred for 16 h at room temperature [final IR spectrum (CH₂Cl₂): $\tilde{v}/cm^{-1} = 2112$ m (SeCN), 2068 w, 1970 vs-br (CO), 1812 s-br (µ-CO)]. Volatiles were evaporated under reduced pressure, hence a solution of the residue in the minimum volume of dichloromethane was charged on an alumina column. Neat CH₂Cl₂ allowed the separation of a light green fraction corresponding to 4a, while CH₂Cl₂/THF (9/1 v/v) mixture was used to collect a red fraction corresponding to 5a. After removing impurities with THF, elution with MeCN/MeOH (9/1 v/v) led to the separation of a dark green fraction corresponding to 6a. For each fraction, the solvent was removed under reduced pressure, and the residue was suspended in hexane for 24 h. After filtration, the solid product was dried under vacuum.

[Fe₂Cp₂(k*Se*-SeCN)(CO)(μ-CO){μ-CN(Me)(Xyl)}], 4a (Figure 8).



Figure 8. Structure of [Fe₂Cp₂(kSe-SeCN)(CO)(µ-CO){µ-CN(Me)(Xyl)}], 4a.

[Fe2Cp2(kN-NCSe)(CO)(μ-CO){μ-CN(Me)(Xyl)}], 5a (Figure 9).



Figure 9. Structure of [Fe₂Cp₂(kN-NCSe)(CO)(μ-CO){μ-CN(Me)(Xyl)}], 5a.

Light brown solid, yield 23 mg (15%). Anal. calcd. for $C_{23}H_{22}N_2O_2Fe_2Se: C, 50.31$; H, 4.04; N, 5.10. Found: C, 50.15; H, 4.12; N, 5.18. IR (CH₂Cl₂): $\tilde{\nu}/cm^{-1} = 2113$ m (SeCN), 1970 vs. (CO), 1812 s (μ -CO). ¹H NMR (CDCl₃): δ /ppm = 7.42–7.30 (m, 3 H, C₆H₃); 5.07, 4.76, 4.38, 4.08 (s, 10 H, Cp); 4.92, 4.53 (s, 3 H, NMe); 2.57, 2.48, 2.44, 2.39 (s, 6 H, C₆H₃Me₂). ¹³C{¹H} NMR (CDCl₃): δ /ppm = 345.4, 342.1 (μ -CN); 263.1, 262.5 (μ -CO); 212.6, 211.1 (CO); 149.0, 148.9 (C-*ipso*); 136.1–128.7 (C₆H₃ + SeCN); 89.7, 89.2, 88.1, 87.8 (Cp); 54.7, 53.8 (NMe); 19.0, 18.9, 18.8, 18.5 (C₆H₃Me₂). ⁷⁷Se{¹H} NMR (CDCl₃): δ /ppm = -232.5, -246.9. Isomer ratio (*cis/trans*) = 1.7.

[Fe₂Cp₂(CN)(CO)(μ-CO){μ-CN(Me)(Xyl)}], 6a (Figure 10).



Figure 10. Structure of [Fe₂Cp₂(CN)(CO)(µ-CO){µ-CN(Me)(Xyl)}], 6a.

Dark red solid, yield 20 mg (13%). Anal. calcd. for $C_{23}H_{22}N_2O_2Fe_2Se: C, 50.31; H, 4.04; N, 5.10.$ Found: C, 50.46; H, 4.12; N, 5.22. IR (CH₂Cl₂): $\tilde{\nu}/cm^{-1} = 2109$ m (NCSe), 1960 vs. (CO), 1804 s (μ -CO). ¹H NMR (CDCl₃): δ /ppm = 7.35–7.27 (m, 3 H, C₆H₃); 4.85, 4.27 (s, 10 H, Cp); 4.83 (s, 3 H, NMe); 2.54, 2.42 (s, 6 H, C₆H₃Me₂). ¹³C{¹H} NMR (CDCl₃): δ /ppm = 340.4 (μ -CN); 263.3 (μ -CO); 213.1 (CO); 149.2 (C-*ipso*); 134.2, 132.6, 129.6, 129.3, 128.6 (C₆H₃); 108.4 (NCSe); 89.3, 88.9 (Cp); 53.4 (NMe); 18.5, 18.4 (C₆H₃Me₂). ⁷⁷Se{¹H} NMR (CDCl₃): δ /ppm = -340.2.

Green solid, yield 64 mg (48%). Anal. calcd. for $C_{23}H_{22}N_2O_2Fe_2$: C, 58.76; H, 4.72; N, 5.96. Found: C, 58.46; H, 4.90; N, 5.79. IR (CH₂Cl₂): $\tilde{\nu}/cm^{-1} = 2090$ w (C \equiv N), 1959 vs. (CO), 1808 s (μ -CO). ¹H NMR (CDCl₃): $\delta/ppm = 7.3$ –7.2 (m, 3 H, C₆H₃); 4.85, 4.8 *, 4.35, 4.29 (s, 10 H, Cp); 4.52, 4.45 (s, 3 H, NMe); 2.64, 2.52, 2.40, 2.23 (s, 6 H, C₆H₃<u>Me₂</u>). * Partially hidden by Cp resonance of major isomer. ¹³C{¹H} NMR (CDCl₃): $\delta/ppm = 340.3$, 337.0 (μ -CN); 261.6, 260.8 (μ -CO); 213.0, 211.5 (CO); 148.9, 148.5 (C-*ipso*); 140.2, 139.7 (C \equiv N); 134.4, 134.3, 132.9, 132.7, 130.4, 129.5, 129.3, 128.7, 128.5, 128.3 (C₆H₃); 89.5, 88.8, 87.3, 86.5 (Cp); 53.6, 52.9 (NMe); 18.7, 18.6, 18.5, 17.9 (C₆H₃<u>Me₂</u>). Isomer ratio (*trans/cis*) = 6. When the reaction of [**1a**^{NCMe}]CF₃SO₃ (179 mg, 0.28 mmol) with KSeCN (122 mg, 0.85 mmol) was conducted in THF at reflux temperature, complete conversion of the starting material into **6a** was checked via IR spectroscopy after 4 h; **6a** (¹H NMR) was isolated in 45% yield after alumina chromatography.

Reactivity of other acetonitrile complexes with KSeCN: evidence for the formation of selenocyanate derivatives and isolation of $[Fe_2Cp_2(CN)(CO)(\mu-CO){\mu-CN(Me)(R)}]$ (R = Me, 6b; R= Cy, 6c).

3.3. General Procedure

The acetonitrile adduct was prepared from [1b,c]CF₃SO₃ as described above, then it was dissolved in deaerated acetone (30 mL), and KSeCN (3.0 eq.) was added to this solution. The resulting mixture was stirred for 16 h at room temperature. The volatiles were evaporated under reduced pressure, thus the residue was dissolved in dichloromethane and this solution was charged on an alumina column. Neat CH₂Cl₂ allowed the separation of impurities, and then CH₂Cl₂/THF (2/1 v/v) mixture was used to collect a brown fraction. Elution with MeCN/MeOH (9/1 v/v) led to the separation of a dark green fraction corresponding to **6b–c**. For each fraction, the solvent was removed under reduced pressure, and the residue was suspended in hexane for 24 h. After filtration, isolated products (i.e., brown solid and green solid) were dried under vacuum.

(1) From [**1b**]**CF**₃**SO**₃ (226 mg, 0.426 mmol), Me₃NO·2H₂O (52 mg, 0.47 mmol), and KSeCN (184 mg, 1.28 mmol). Brown solid (yield 52%), $\tilde{v}/cm^{-1} = 2110$ m (SeCN), 1966 vs. (CO), 1804 s (μ -CO). **6b**, yield 23%.

When the reaction of $[1b^{NCMe}]CF_3SO_3$ (102 mg, 0.19 mmol) with KSeCN (122 mg, 0.85 mmol) was conducted in THF (or acetone) at reflux temperature, complete conversion

of the starting material into **6b** was checked via IR spectroscopy after 4 h; **6b** was isolated in 53% yield after alumina chromatography.

[Fe₂Cp₂(CN)(CO)(μ-CO){μ-CNMe₂}], 6b (Figure 11).



Figure 11. Structure of $[Fe_2Cp_2(CN)(CO)(\mu-CO)\{\mu-CNMe_2\}]$, 6b.

Green solid. Anal. calcd. for $C_{16}H_{16}N_2O_2Fe_2$: C, 50.57; H, 4.24 N, 7.37. Found: C, 50.44; H, 4.14; N, 7.28. IR (CH₂Cl₂): $\tilde{\nu}/cm^{-1} = 2089$ w (C \equiv N), 1962 vs. (CO), 1805 s (μ -CO), 1568 w (μ -CN). ¹H NMR (CDCl₃): δ /ppm = 4.84, 4.80, 4.76, 4.69 (s, 10 H, Cp); 4.33, 4.24, 4.21, 4.10 (s, 6 H, NMe₂). ¹³C{¹H} NMR (CDCl₃): δ /ppm = 337.0 (μ -CN); 261.5 (μ -CO); 212.3, 210.8 (CO); 141.2 (C \equiv N); 89.9, 89.0, 87.0, 86.5 (Cp); 52.3, 51.9 (NMe₂). Isomer ratio (*trans/cis*) = 5. Crystals of *trans*-6b suitable for X-ray analysis were obtained by slow diffusion of hexane layered on an acetone solution of 6b at -30 °C.

(2) From [1c]CF₃SO₃ (256 mg, 0.427 mmol), Me₃NO·2H₂O (52 mg, 0.47 mmol), and KSeCN (184 mg, 1.28 mmol). Brown solid (yield 41%), $\tilde{\nu}/cm^{-1}$ = 2110 w (SeCN); 1964 vs. (CO), 1803 s (μ -CO). 6c, yield 36%.

When the reaction of $[1c^{NCMe}]CF_3SO_3$ (186 mg, 0.30 mmol) with KSeCN (122 mg, 0.85 mmol) was conducted in THF at reflux temperature, complete conversion of the starting material into **6c** was checked via IR spectroscopy after 4 h; **6c** was isolated in 67% yield after alumina chromatography.

[Fe₂Cp₂(CN)(CO)(μ-CO){μ-CN(Me)(Cy)}], 6c (Figure 12).



Figure 12. Structure of [Fe₂Cp₂(CN)(CO)(µ-CO){µ-CN(Me)(Cy)}], 6c.

Green solid. Anal. calcd. for $C_{21}H_{24}N_2O_2Fe_2$: C, 56.29; H, 5.40; N, 6.25. Found: C, 56.44; H, 5.31; N, 6.38. IR (CH₂Cl₂): $\tilde{\nu}/cm^{-1} = 2089$ w (C \equiv N); 1959 vs. (CO), 1803 s (μ -CO); 1528 w (μ -CN). ¹H NMR (CDCl₃): δ /ppm = 5.33, 5.13 (m, 1 H, CH^{Cy}); 4.80, 4.78, 4.65, 4.64 (s, 10 H, Cp); 4.17, 4.04 (s, 3 H, NMe); 2.27–2.17, 2.10–1.83, 1.64, 1.34–1.24 (m, 10 H, CH₂^{Cy}). ¹³C{¹H} NMR (CDCl₃): δ /ppm = 336.1, 335.8 (μ -CN); 262.2, 261.8 (μ -CO); 212.6, 212.3 (CO); 141.4, 140.6 (C \equiv N); 90.2, 90.0, 89.2, 89.0 (Cp); 75.9, 75.3 (CH^{Cy}); 44.5, 44.2 (NMe); 32.0, 31.6, 31.5, 31.0, 26.2, 25.8, 25.7, 25.5, 25.3 (CH₂^{Cy}). Isomeric ratio (α / β) = 1.3.

4. X-ray Crystallography

Crystal data and collection details for *trans*-6b are reported in Table 1. Data were recorded on a Bruker APEX II diffractometer equipped with a PHOTON2 detector using Mo–K α radiation. The structures were solved by direct methods and refined by full-matrix least-squares based on all data using F^2 [83]. Hydrogen atoms were fixed at calculated positions and refined using a riding model.

Table 1. Crystal data and measurement details for trans-6b.

	trans-6b
Formula	$C_{16}H_{16}Fe_2N_2O_2$
FW	380.01
Т, К	100(2)
λ , Å	0.71073
Crystal system	Monoclinic
Space group	P2 ₁ /c
<i>a</i> , Å	8.7226(2)
<i>b,</i> Å	12.4479(3)
<i>c,</i> Å	14.1827(4)
eta , $^{\circ}$	107.2940(10)
Cell Volume, Å ³	1470.31(6)
Z	4
D_c , g·cm ⁻³	1.717
μ , mm ⁻¹	1.980
F(000)	776
Crystal size, mm	0.16 imes 0.15 imes 0.13
θ limits, °	2.222-26.996
Reflections collected	23,142
Independent reflections	3205 [R _{in} = 0.0474]
Data/restraints/parameters	3205/0/201
Goodness on fit on F ²	1.074
$R_1 \ (I > 2\sigma(I))$	0.0283
wR_2 (all data)	0.0677
Largest diff. peak and hole, e ${\rm \AA}^{-3}$	0.562/-0.444

5. Computational Details

Geometry optimizations were performed using the PBEh-3c method, which is a reparametrized version of PBE0 [84] (with 42% HF exchange) that uses a split-valence double-zeta basis set (def2-mSVP) [85,86] and adds three corrections considering dispersion, basis set superposition, and other basis set incompleteness effects [87–89]. The C-PCM implicit solvation model was added to PBEh-3c calculations, considering acetone as a continuous medium [90,91]. IR simulations were carried out using harmonic approximation, from which zero-point vibrational energies and thermal corrections (T = 298.15 K) were obtained [92]. The software used was ORCA version 5.0.3 [93].

6. Conclusions

New diiron aminocarbyne complexes with a terminal chalcogen-containing pseudohalide ligand were synthesized, and their stereochemistry and thermodynamic stability were investigated by IR and NMR spectroscopy and DFT calculations. Combined with previous findings, this work highlights that *N*-coordination generally prevails over the alternative coordination mode, although different kinetic products may be formed, and that the reactivity of the NCE⁻ ligand increases along the sequence O (inertness) < S (electrophilic addition) < Se (chalcogen elimination). In particular, we provide clear evidence for the formation of diiron cyanide complexes from the fragmentation of the selenocyanate fragment.

Supplementary Materials: The following supporting information can be downloaded at: https: //www.mdpi.com/article/10.3390/molecules28073251/s1, NMR spectra of products and DFT data. CCDC reference number 2225698 (*trans*-6b) contains the supplementary crystallographic data for the X-ray study reported in this paper.

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