

Perspective

Construction of Synthetic Models for Nitrogenase-Relevant NifB Biogenesis Intermediates and Iron-Carbide-Sulfide Clusters

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Abstract: The family of nitrogenase enzymes catalyzes the reduction of atmospheric dinitrogen (N_2) to ammonia under remarkably benign conditions of temperature, pressure, and pH. Therefore, the development of synthetic complexes or materials that can similarly perform this reaction is of critical interest. The primary obstacle for obtaining realistic synthetic models of the active site iron-sulfur-carbide cluster (e.g., FeMoco) is the incorporation of a truly inorganic carbide. This review summarizes the present state of knowledge regarding biological and chemical (synthetic) incorporation of carbide into iron-sulfur clusters. This includes the Nif cluster of proteins and associated biochemistry involved in the endogenous biogenesis of FeMoco. We focus on the chemical (synthetic) incorporation portion of our own efforts to incorporate and modify C_1 units in iron/sulfur clusters. We also highlight recent contributions from other research groups in the area toward C_1 and/or inorganic carbide insertion.

Keywords: nitrogenase; biomimetic; FeMoco; NifB-co; carbide; biogenesis

1. Introduction

1.1. Structural Determination of the Resting Nitrogenase Active Site

The nitrogenases are a class of enzymes that catalyze ammonia synthesis from dinitrogen and are the only biological systems known to catalyze the $N_2 \rightarrow 2 \text{ NH}_3$ conversion. In contrast to the H₂-consuming Haber-Bosch process, dinitrogen fixation by nitrogenase (N2ase) is achieved via coupled proton/electron transfer directed toward the substrate [1]. Three variants of nitrogenase are known: Mo-dependent, V-dependent, and Fe-type. The Mo-dependent nitrogenase, however, exhibits the highest rate of catalytic activity and has been the focus of the majority of biocatalytic dinitrogen-reduction research [2]. The crystal structure of the molybdenum-containing nitrogenase was first reported in 1992 by Rees et al. to 2.9 Å resolution (Figure 1). This structure revealed a homodimeric enzyme in which each monomer consists of two components: (1) nitrogenase (MoFe protein in Mo-dependent nitrogenase) in which the catalytic active site is contained, and (2) nitrogenase reductase (Fe protein), which is an electron-transfer protein [3,4]. The active site contained in the MoFe protein was initially described as a two-component active site comprised of two clusters: a 4Fe:3S cluster and a 1Mo:3Fe:3S cluster bridged by three non-protein ligands (designated as two sulfides and one unresolved site). This iron-molybdenum cofactor (FeMoco) is tethered to the protein by two ligands coordinated to the two distal metal sites: a cysteine residue (Cys^{α 275}) located at the distal iron site and a histidine residue (His^{α 442}) ligated to molybdenum. The molybdenum site is also coordinated by two oxygens from a bidentate homocitrate (HC) ligand. However, the six proximal Fe sites in this proposed structure appear to be supported by a non-planar, trigonal coordination sphere uncharacteristic for biological iron in a resting state enzyme.



A decade following the 1992 report, a higher resolution structure (1.16 Å resolution) of FeMoco by Einsle, Rees et al., revealed the presence of a central, interstitial hexacoordinate light (2*p*) atom residing in the tetrahedral hole of the six proximal Fe sites [5,6]. Einsle and Rees initially postulated the site to be a nitride based on electron density integration studies. However, they suggested that carbon and oxygen could not be unambiguously discounted.



Figure 1. Progression of structure refinement for the iron-molybdenum cofactor (FeMoco) from structural X-ray diffraction (XRD) data and specialized Electron Paramagnetic Resonance (EPR) and X-ray emission (XES) data [3–8].

In 2011, two independent research efforts concurrently ascertained the identity of the central atom to be a carbide rather than a nitride (Figure 1). Rees, Einsle et al. confirmed the carbide experimentally by utilizing two separate experiments: (1) improved structural resolution (1.0 Å) for electron density integration analysis and (2) comparative pulsed EPR-ESEEM (Electron Paramagnetic Resonance, Electron Spin Echo Envelope Modulation) on isotopically labeled (¹³C or ¹⁵N) MoFe protein (resting state S = 3/2) in which significant hyperfine coupling was observed in the ¹³C-labeled (I = 1/2) protein [7]. These monumental findings were printed onto a single page in the mid-November 2011 issue of Science. In the same issue, a publication of equal importance by DeBeer et al. utilized a combination of valence-to-core X-ray emission spectroscopy (V2C-XES) and computationally calculated spectra for cofactors bearing various interstitial atoms (C⁴⁻, N³⁻, and O²⁻) to determine the identity of the interstitial atom [8]. The capability of V2C-XES to distinguish between light atoms within a multi-metallic cluster was demonstrated in a preceding study by DeBeer, Holland et al. comparing synthetic and computational model compounds bearing various interstitial atoms to differentiate between Fe₆ hexacoordinate C, N, and O sites [9]. The results of these studies identified the interstitial atom in FeMoco as a purely inorganic carbide. Thus, the concurrent findings by the Einsle and DeBeer research groups unambiguously established the presence of an interstitial carbide in the N₂ase cofactor.

1.2. Biological Synthesis of the FeMoco Cluster

The biogenesis of nitrogenase is accomplished by a set of enzymes coded in the "nitrogen-fixing" (Nif) genes. NifS and NifU provide the initial conversion of cysteinyl thiolates and iron incorporation to produce the two inorganic Fe₄S₄ iron-sulfur clusters to be transferred onto NifB for carbide insertion. The presence of a CXXXCXXC signature motif consistent with the presence of *S*-adenosylmethionine (SAM) led Ribbe and Hu to postulate the utilization of a SAM-dependent mechanism by NifB [10,11]. Using isotope-labeling experiments in conjunction with CW (Continuous-Wave) and pulse EPR, the origin of the carbide was determined to be a methyl group from SAM during cofactor assembly by NifB [11–13]. Following initial transfer from SAM onto a sulfide in the (Fe₄S₄)₂ K-cluster (Figure 2), the CH₃ group undergoes H-atom abstraction by an additional equivalent of SAM, which is followed by continued, iterative deprotonation/dehydrogenation to form the L-cluster. The utility of SAM as

a methyl transfer agent toward sp^2 and sp^3 C sites has been previously reported, such as upon the tryptophanyl bicyclic ring [14] or in thienamycin biosynthesis [15]. Subsequent maturation by the NifEN assembly protein leads to substitution of a distal Fe site with Mo(HC) before final delivery into NifDK to achieve the final M-cluster [16–18].



Figure 2. Proposed biosynthetic mechanism for carbide formation, insertion of the '9th' sulfide, and heterometal substitution prior to maturation of the MoFe protein as proposed by Ribbe and Hu [11–13,16–18]. Image adapted with permission from Joseph, C., Cobb, C. R., Rose, M. J. Single-Step Insertion of Sulfides and Thiolate into Iron Carbide-Carbonyl Clusters: Unlocking the Synthetic Door to FeMoco Analogues. *Angew. Chemie Int. Ed.* **2020**. *Accepted, not yet published*. Copyright *Wiley-VCH GmbH*. Reproduced with permission.

Initial studies with nitrogenase utilized dithionite as the reductant, convoluting the origin of the "9th" sulfide into the cluster. As such, Ribbe et al. surveyed a suite of biologically relevant sulfur-containing sources—sulfides (S^{2–}), sulfites (SO₃^{2–}), sulfates (SO₄^{2–})—and substituted Eu^{II}-EGTA as a reductant in their biogenesis studies to establish the role of an external sulfur source for M-cluster maturation [18]. It was demonstrated that only SO₃^{2–} led to the formation of the L cluster. Additionally, when radiolabeled ³⁵SO₃^{2–} was utilized, the radiolabel could be detected downstream in the extracted L-cluster. In biosynthetic experiments performed in the absence of SAM, no significant retention of the radiolabel was detected, suggesting that sulfide transfer likely occurs after incubation with SAM. Finally, incubation conditions in which SAM is present while SO₃^{2–} is absent leads to the formation of an undefined L*-cluster (tentatively assigned as an Fe₈S₈C cluster).

1.3. Multimetallic Co-Operativity toward Dinitrogen Activation

Initial insight into the nitrogenase mechanism was primarily founded upon extensive kinetic studies conducted throughout the 1970s to 1980s. Lowe and Thorneley famously proposed a rudimentary kinetic scheme whereby the resting-state N₂ase cofactor (designated E_0) undergoes four iterative protonation/reduction events (E_1 – E_4), at which point, the stage is set for substrate binding. Concurrent with the liberation of one equiv H₂, N₂ binds and proceeds to be reduced to two NH₃ as the cofactor continues to undergo subsequent protonation and reduction. Liberation of the second equivalent NH₃ returns the cofactor to its resting state, E_0 [1]. The precise geometric and electronic structural details of the E_1 – E_4 intermediates are still mostly unknown. However, recent spectroscopic and computational information about the E_1 state has begun to emerge, suggesting that the E_1 step involves reduction of an Fe site in conjunction with protonation of a belt-sulfide [19,20].

The location and mode of substrate-binding was a topic of considerable debate throughout the last two decades. The differences in N_2 -reducing activity between the M-cluster, V-cluster, and L-cluster

nominated the heterometal (i.e., Mo in FeMoco) as a likely site for substrate binding. This conclusion gained additional support with the publication in 2003 of Schrock's synthetic, Mo-centered complex as the first report of a synthetic and functional N₂-reducing catalyst with an N₂ase-relevant metal center [21]. In 2013, however, Peters published a synthetic Fe complex bearing a 2p donor atom (B) in its ligation sphere and capable of catalytic N_2 conversion [22]. Experimental and computational evidence with FeMoco and FeVco began to implicate iron as the likely N2-binding site. Inhibition of FeMoco with carbon monoxide (CO) and crystallization of the resulting MoFe protein revealed a structure in which the S2B belt sulfide—which bridges Fe2 and Fe6—is substituted with a bridging CO [23]. Additionally, treatment of FeMoco with KSeCN demonstrated similar lability of the belt sulfides: 1 equiv KSeCN under argon atmosphere resulted in substitution of S2B with Se [24]. Subsequent catalytic turnover with N₂ produced structures in which the Se migrated to the 5A or 3A positions. Crystallization and X-ray diffraction (XRD) of vanadium-dependent N₂ase during turnover presented a FeVco structure in which the S2B position was occupied by a small bridging ligand (presumably N or NH_x, even though O or OH_x cannot be discounted) [25]. Finally, the computational work surrounding the Mo site suggests that one of the oxygen donors dissociates from Mo during the E_2 - E_4 steps, leaving a possible open coordination site. However, energy calculations suggest that activation of N_2 at this site results in a significantly higher increase in energy and that N_2 binding at Fe_2 and Fe_6 is much more thermodynamically favorable (though still endothermic) [26].

Furthermore, in early 2020, a ground-breaking crystallographic structure by Ribbe and Hu under physiological turnover conditions depicted a substrate-bound nitrogenase intermediate, which revealed details regarding the putative substrate binding sites as well as binding modes during turnover [27]. While the exact identities of the substrates remain unconfirmed (N₂ vs. N₂H₂ vs. N₂H₄), the structural data demonstrates displacement of the belt sulfides to allow for substrate binding at the medial Fe sites. The two FeMoco clusters in the structure exhibit distinct substrate-binding modes, such that the substrate presents in a 'side-on' binding mode in the chain A cluster or an 'end-on' mode in the chain C cluster. In either case, the dinitrogen substrate adopts a bridged binding mode between two Fe sites, highlighting the significant role of multi-iron co-operativity toward the mechanism of dinitrogen activation and reduction by nitrogenase.

1.4. Synthetic Model Compounds in FeMoco Biomimetics

Throughout the past two decades of nitrogenase research, studies utilizing synthetic model compounds have contributed greatly to the early mechanistic and electronic discussions surrounding nitrogenase in lieu of highly sophisticated spectroscopic and computational techniques, which have only recently become available for studying the complicated active site. The Schrock catalyst, a Mo complex featuring a sterically protected substrate-binding site (see Figure 3) initially invigorated the debate to determine the binding site of FeMoco [21]. Nishibayashi has reported pincer-based monomeric and dimeric Mo complexes, which catalytically reduce N₂ [28,29]. A pincer motif was later used by the same group to develop a catalytically active Fe-centered complex as well [30]. The Peters research group has published a series of Fe complexes featuring a tripodal ligand set with various axial donor atoms. The Si-ligated complex was first shown to bind N₂, followed by reports of B-ligated and C-ligated complexes that could catalytically perform the reduction [22,31,32]. Spectroscopic and mechanistic studies with these compounds have demonstrated the axial donor atom modulates covalency with the Fe center throughout catalysis to support the various oxidation states and substrate coordination modes. Results from these studies were used to hypothesize a stabilizing role for the central carbide in FeMoco, performing similar modulations throughout turnover. The proposed hybrid distal-alternating mechanism determined for Peters' compounds was similarly extended to FeMoco as a possible mechanistic pathway [33]. Moreover, the tripodal donor set was later adapted to include a structurally relevant S donor as a bridging thiolate [34]. Synthetic work pursued by the Holland group has demonstrated activation of dinitrogen by multi-metallic cooperation between two or four iron sites, presenting biomimetic complexes for the di-iron activation of N₂ displayed in the Ribbe and

Hu crystal structure [27]. Additional synthetic work with an iron-carbonyl-thiolate complex utilizes a tridentate ligand set and is capable of N₂ activation [35].



Figure 3. Selected functional synthetic compounds for N₂ activation and/or reduction from the Schrock, Nishibayashi, Peters, and Holland research groups [21,22,28–35].

Functional synthetic model compounds capable of catalytically reducing or activating dinitrogen in various modes were particularly impactful toward early mechanistic discussions surrounding nitrogenase. In recent years, structural models (Figure 4) have also been extensively utilized as synthetic references in studies probing for geometric and electronic structure information about FeMoco and FeVco. The carbonyl-supported hexa-iron cluster $[Fe_6(\mu_6-C)(CO)_{16}]^2$ —which contains an interstitial, 6-coordinate carbide and has been featured prominently throughout our own research program—was utilized along with its nitride congener $[Fe_6(\mu_6-N)(CO)_{16}]^-$ as a means of calibrating X-ray emission spectra to distinguish between 2p atoms in multi-metallic frameworks, thereby, setting the stage for identification of the interstitial carbide in FeMoco [9].

In many respects, FeMoco is structurally similar to other synthetic and biological iron sulfur clusters. The average Fe–Fe distance of 2.63 ± 0.04 Å places individual iron sites within the range to induce a bonding interaction between metal centers. Similarly, the average Fe–Mo distance of 2.684 ± 0.008 Å is typical for reported synthetic heterometal-iron-sulfur clusters [36]. Fe–S contacts exhibit an average distance of 2.25 ± 0.03 Å. Finally, the Fe–C bond is not a known motif in iron-sulfur clusters outside of the nitrogenases. Moreover, synthetic iron-sulfur clusters found in the Cambridge Structural Database (CSD) do feature Fe-coordinated N and O sites, but examples with carbon have remained elusive. Nonetheless, the average Fe–C distance of 2.00 ± 0.02 Å observed in FeMoco fall within the range of distances displayed in compounds with N (Fe–N range: 1.871-1.977 Å) or O (Fe–O range: 2.052-2.190 Å) incorporated donor atoms. Synthetic iron sulfur clusters developed by the Holm group and featuring heterometallic Mo³⁺ and V³⁺ sites assisted in the assignment of spin states at individual metal sites in the resting-state M-clusters and V-clusters [37–39]. Notable work by the Agapie group has demonstrated

electronic and catalytic modulation of iron cluster sites performed by substitution of various interstitial light atoms (O, F) [40,41]. Synthetic structural modelling work from the Tatsumi group has presented a highly-relevant dicuboidal, vanadium-containing FeS cluster and an astonishingly remarkable cluster featuring belt-thiolates and an interstitial oxide [42,43]. Remarkably, Rauchfuss has recently published the first known multi-iron cluster to feature an interstitial carbide and one or multiple inorganic sulfides [44,45].



Figure 4. Selected synthetic structural models for the nitrogenase active site, including works published by the Holm, Agapie, Tatsumi, and Rauchfuss [37,38,40–49].

Synthetic development of mono-metallic and bimetallic iron complexes with C-donors is of potential importance for understanding the role of the Fe-C bond in controlling the electronic interactions between the metal sites in FeMoco. These complexes can be isolated with low-coordinate, high-spin ferrous, and ferric iron centers such as the complexes reported by Power, Fürstner, and Re [50–52]. Additionally, Yogendra et al. recently reported a set of dimeric, yldiide-bridged iron complexes accessed from the known monomers. Magnetic susceptibility and spectroscopic studies of these dimers show intense antiferromagnetic coupling between iron sites and pronounced negative charge localization on the C ligand [53]. The Holland group reported a series of β -diketiminate complexes with various bridging C-donors. Among these, an alkylidene-bridged and sulfide-bridged dimer was shown to be competent toward N₂ to NH₄⁺ conversion with dissociation of the sulfide ligand.

The presence of the carbon donor ligand ostensibly facilitates improvement of the N₂ reduction activity of the complex along with sulfide lability [54].

An alkyl-ligated Fe_4S_4 cluster published by Suess et al. demonstrates the dramatic electronic shift enforced upon the cubane cluster by the coordination of the electron-donating R group [48]. Notably, a variety of new tripodal chelating ligands were employed to enforce 3:1 site differentiation and stabilize a range of alkylated cluster oxidation states (Figure 5) [48,49]. The success and versatility of their approach represent new progress in the rational control of multi-metallic cluster chemistry, for which the ligand-control toolbox has historically been limited compared to monometallic and bimetallic inorganic synthetic work. As such, these findings open new doors toward modeling and understanding Fe_4S_4 -alkyl intermediates of enzyme active site clusters, especially the radical SAM family [55].



Figure 5. Fe₄S₄-alkyl clusters stabilized by chelating tripodal ligand scaffolds recently reported by Suess et al.

1.5. A Two-Fold Strategy toward Achieving FeMoco Construction

Undoubtedly, the field of structural modelling in the context of nitrogenase has been invaluable in advancing the structural, electronic, and mechanistic understanding of the nitrogenase active site. However, much synthetic work remains unaccomplished. As of this writing, the synthetic 'Holy Grail' for FeMoco modeling of achieving a paramagnetic iron cluster with sulfides/thiolates, and interstitial carbide continues to be pursued. Additionally, synthetic modelling work applied to the FeMoco biogenesis—such as the intermediate steps involved in carbide formation—are scarce if not absent completely. Through the work summarized in this perspective, we sought to expand upon the compendium of structural models in progression toward incorporation of important structural and electronic attributes of FeMoco (Figure 6) into synthetic clusters.



Figure 6. Target structural and electronic attributes of interest in FeMoco for biomimetic modelling.

To this end, a two-fold strategy was employed in our nitrogenase biomimetics research program (Scheme 1). First, we took inspiration from the biomechanistic pathway proposed by Ribbe and Hu for FeMoco biogenesis. A carbonyl-supported iron sulfide cluster K₂[Fe₃S(CO)₉] was synthesized and alkylated with various alkyl halides (R = Me, ⁱPr, Bz). The resulting μ_3 -thiolato cluster was subsequently treated with a radical initiator TEMPO ((2,2,6,6-tetramethylpiperidin-1-yl)oxyl) to emulate the H atom abstraction mechanism propagated by SAM upon the K cluster. We additionally pursued a converse strategy utilizing a family of carbonyl-supported iron clusters featuring interstitial carbides—the first of which was reported by Braye et al. in 1962 [56]. The hexa-iron cluster (NEt₄)₂[Fe₆(μ_6 -C)(CO)₁₆] reported by Churchill et al. [46,47] serves as a synthetic starting point toward highly FeMoco-relevant structural models, demonstrating flexibility between multiple redox states and versatility for heterometal substitution and sulfide incorporation [44,57,58]. Here, a strategy was explored for ligand CO→S substitution by using electrophilic sulfur upon the soft anionic clusters.



Scheme 1. A two-fold synthetic approach toward achieving structural analogues of the L-cluster, precursor to FeMoco. First, inspiration was taken from the biosynthetic conversion of the K-cluster $(2 \times \text{Fe}_4\text{S}_4)$ to the L-cluster (Fe₈S₉C) by NifB. As such, a methyl group was installed upon an Fe₃S cluster using MeI. TEMPO was then utilized as a radical initiator to emulate similar radical abstraction of hydrogen observed during L-cluster synthesis. Alternatively, an anionic Fe₆C cluster featuring an interstitial carbide was found to be a useful synthetic starting point for sulfur incorporation via electrophilic sulfur reagents.

2. Synthetic Manipulations of CO-Supported FeS and FeC Clusters

2.1. Biogenesis-Inspired Approach to FeSC Cluster Construction

As discussed above, the biogenesis of the M-cluster in nitrogenase has garnered much interest since the identification of the interstitial carbide in 2011 [7,8]. The unique motif of a 6-coordinate C^{4-} bound to the six iron centers present in FeMoco/FeVco are the only reported examples in biological systems [59]. It is highly likely, however, that the all-iron analogue of the enzyme also contains

a carbide. Few examples exist in chemical systems as well with the notable exception of the synthetic carbide clusters (*vide infra*).

It was shown that carbide formation at the K-cluster requires two equivalents of SAM including one for the methyl group donation (which eventually becomes the carbide) and a second one for the radical abstraction of a hydrogen atom [11]. The initial binding site of the methyl group was also traced by liquid and gas chromatographic methods HPLC and GCMS, revealing that the methyl group first binds to a sulfide in the native enzyme [12]. This is counterintuitive, as one would expect an all-iron ligated carbide to initially bind to an iron center. However, biological Fe oxidation states are likely not nucleophilic enough to form Fe–C bonds. Considering this, it is then understandable that nitrogenase utilizes a radical mechanism for carbide formation. Furthermore, the proposed mechanism indicates that transfer of the methyl group occurs prior to radical H atom abstraction and subsequent deprotonation/dehydrogenation.

These results provided the stimulus for our initial biomimetic approach for investigating the reactivity of methyl-substituted iron-sulfur clusters with a radical initiator. The synthetic cluster $K_2[(\mu_3-S)Fe_3(CO)_9]$ (Scheme 2) was chosen due to its versatility in substituting R groups using common electrophiles that can be installed on the S^{2–} 'bridgehead,' making it a suitable model system for investigating –SCH₃ reaction mechanisms associated with iron-containing clusters. The methylation of $K_2[SFe_3(CO)_9]$ was achieved by a reaction with CH₃I in THF at 0 °C, affording the monoanionic K[(CH₃S)Fe₃(CO)₉] (1). The resulting bond metrics were similar to those previously observed in NEt₄[(CH₃S)Fe₃(CO)₉] [60]. It would be expected that radical hydrogen abstraction reagents would readily abstract an H-atom from the anion. However, because the low valent Fe clusters typically demonstrate a proclivity toward forming bridging hydrides, we elected to utilize a relatively mild H-atom abstraction reagent. Thus, (2,2,6,6-tetramethylpiperidin-1-yl)oxyl (TEMPO) was selected due to its mild O–H bond dissociation energy [61].



Scheme 2. Synthetic scheme depicting the formation of 1 and its reaction with the radical initiator, TEMPO, to form 2.

The reaction of the methylated Fe₃S cluster and crystallization of the charged portion revealed an aggregation product in which the methyl group had been displaced, yielding the formula [K(benzo-15-crown-5)₂]₂[(SFe₂(CO)₁₂)₂Fe(CO)₂] (**2**). This reactivity indicated the occurrence of two key transformations: (*i*) S–CH₃ bond breakage to generate the inorganic sulfide, and (*ii*) the combination of two Fe₃ units (and extrusion of one Fe center) to form an S₂Fe₅ framework. Additionally, two-electron oxidation of the Fe centers was observed (one electron from each progenitor cluster of **1**). The resultant formation of **2** is unaffected by dramatic changes in reaction conditions including solvent identity (THF, MeCN, FPh) or temperature (–78 °C or ambient temperature), showcasing the low kinetic barrier present in reactions with these types of iron clusters such that the same thermodynamically stable product is recurringly generated, highlighting the uniqueness of the biogenesis in FeMoco assembly.

This process does bear some resemblance to the biosynthesis of the FeMoco active site, wherein the addition of SAM causes oxidation of the SAM-motif-bound $[Fe_4S_4]^{1+}$ in NifB to an oxidized EPR silent intermediate [62]. In this process, S^{2-} and C^{4-} are ultimately generated as charged ligands, implying oxidation of the iron centers en route to the intermediate L cluster. Unfortunately, the incorporation of any carbon (or carbide) motif to the synthetic cluster **1** was not observed in the generation of **2**, even though it was demonstrated that a radical electron can aid in the fusion of two iron-sulfur units and aid in the gradual oxidation of the Fe centers.

2.2. Sulfide Incorporation into Iron-Carbide Clusters: Initial Attempts

Our initial foray into ligand substitution upon (NEt₄)₂[Fe₆(μ_6 -C)(CO)₁₆] began with the utilization of a bidentate, anthracene-scaffolded bis-phenylthiolate ligand set (Scheme 3). The choice of the anthracene scaffold was motivated by the observation that the 5.1 Å distance between the 1 and 8 positions on the anthracene backbone closely correlated with the 5.0 Å S···S distance between distal sulfide sites in FeMoco. However, treatment of the [Fe₆C]^{2–} cluster with disodium 1,8-bis(phenylthiolato) anthracene does not induce any color change in the reaction mixture (Scheme 4). In 2015, Figueroa et al. reported on isocyanide analogues of iron carbonyls, noting that isocyanides—similar to carbonyls—act as excellent π -acceptors, but additionally exhibit stronger σ -donating ability relative to carbon monoxide. This increases the reactivity of their host metal toward substrates including N₂ [63]. Following this line of research, [Fe₆C]^{2–} was treated with 2,6-dimethylphenyl isocyanide, resulting in a reaction solution dominated by a dark green color—indicative of decomposition to Fe₃(CO)₁₂ and other intractable products.



Scheme 3. S•••S distances between the 1 and 8 positions on an anthracene scaffold (left) and the distal sulfides in FeMoco (right).

Another attempted effort to incorporate sulfur into carbidocarbonyl iron cluster was in the direct reaction of the *nido*-Fe₅ neutral cluster [Fe₅(μ_5 -C)(CO)_{15}] with elemental sulfur (S₈). However, performing the reaction at room temperature or 60 °C results in the formation of a small amount of an insoluble and ferromagnetic black powder (presumably metallic iron) and an unconverted [Fe₅C]⁰ band. Increasing the temperature to 90 °C significantly decomposes [Fe₅C]⁰ into the same black powder, and reaction at 120 °C results in total decomposition. Alternatively, a literature report found that elements from the pnictogen series were successfully reacted with [Fe₅C]⁰ in the presence of H₂SO₄ at 100 °C for 30 min to generate a tri-iron methylidene cluster with the formula Fe₃(μ_3 -CH)E(CO)₉ (E = As, Sb, Bi) [64]. Elemental arsenic and bismuth were found to work for this reaction. Interestingly, the Sb cluster could only be achieved with SbCl₅. Efforts to replicate these conditions with elemental sulfur produced a dark red-orange solution, which afforded only crystals of the non-carbide-containing cluster Fe₃S₂(CO)₉. Similarly, using selenium powder under the same conditions gave crystals of Fe₃Se₂(CO)₉.

Finally, the utilization of triphenyl antimony sulfide (Ph₃Sb=S) as a thermoneutral sulfur atom donor was considered due to its behavior as a weak sulfur-donating ability and the absence of S–S bonds such as those seen in elemental sulfur (Scheme 5) [65]. Preparation of the reagent for the in situ and transfer into an ice-cold toluene suspension of $[Fe_6C]^{2-}$ resulted in a highly convoluted mixture of compounds, thereby obstructing the growth of crystals suitable for X-ray diffraction.



Scheme 4. Various reaction conditions attempted to pursue controlled ligand substitution onto iron carbide clusters (**top**) and schematic representation of reaction conditions reported in Reference [64] (**bottom**).



Scheme 5. In situ synthesis of triphenyl antimony sulfide and reaction conditions for sulfur donation into $[Fe_6C]^{2-}$ cluster.

2.3. Electrophilic Sulfur Sources: Isolation of Sulfide-Containing Carbidocabonyl Iron Clusters

Because of the undesirable results obtained from using conventional Lewis basic ligands (sulfides, thiolates) with the anionic and neutral iron clusters, we sought an alternative strategy for achieving sulfide incorporation. At the time, the only report of successful ligation by a sulfur-based donation set upon carbidocarbonyl iron clusters was a 1988 publication by Shriver et al. describing the synthesis and isolation of SO₂-ligated Fe₆ and Fe₅ clusters [66]. However, the use of these SO₂ clusters as isolable intermediates in pursuit of a sulfide-ligated iron-carbide clusters was a suspected 'dead-end' in the absence of a strategy for successful reduction and oxide removal from SO₂. We do note, however, that Rauchfuss et al. has reported achievement of an S-ligated cluster via an SO₂ intermediate [44]. Nonetheless, the observation that, upon ligation, SO₂ behaves electrophilically (therefore, oxidizing the anionic cluster) and the additional observation that the gallery of reported heteroleptic carbidocarbonyl iron clusters in the CSD feature initial oxidation prior to ligand incorporation (NO, H⁺ \rightarrow H⁻, Lewis acidic Cu-, and Au-bearing reagents) [36] prompted us to seek out similarly electrophilic sulfur-based ligand sets.

As such, a synthetic strategy centered around S_2Cl_2 was deemed appropriate for sulfide incorporation [67]. While SCl₂ is more structurally analogous to SO₂ (Figure 7), the reported instability of the reagent and its spontaneous conversion to S_2Cl_2 marked it as less preferable. Treatment of the hexa-nuclear cluster (NEt₄)₂[Fe₆(μ_6 -C)(CO)₁₆] with S₂Cl₂ afforded a mixture of products (Scheme 6). We initially characterized the asymmetric, sulfide-bridged 'dimer of clusters' $(NEt_4)_2[(CO)_{15}(\mu_6-C)Fe_6](\mu_4-S)[Fe_5(\mu_5-C)(CO)_{13}]]$ (3). The dimer was structurally identified by XRD as featuring a four-coordinate bridging sulfide tethered between a six-iron and a five-iron cluster including each cluster with an interstitial carbide. Crystal structure data (Figure 8) reveals Fe-S distances [2.203(3), 2.189(3), 2.188(3), 2.169(2) Å] that are noticeably short for a CO supported {Fe₂(μ_4 -S)Fe₂} motif. Exploration of this motif in the CSD [36] produces an average of 2.25 ± 0.01 Å. The average Fe–S bond distance of 2.19 ± 0.01 Å in 3 is, thus, remarkably short. The longest bond length in 3 at 2.203(3) Å is shorter than the shortest recorded CSD Fe–S bond (2.215 Å) in an $\{Fe_2(\mu_4-S)Fe_2\}$ motif. In comparison to the bond metrics found in FeMoco [68], the Fe–S bond falls short of the average Fe–S bond found in the active site (2.25 \pm 0.03 Å). The average Fe–Fe bond distance of 2.65 \pm 0.06 Å in 3 is unremarkable. Thus, the presence of the sulfide does not induce any notable elongation or compression of Fe–Fe bond lengths in Fe₅–Fe₈. The average Fe–C bond distance in the six-iron unit of cluster **3** is 1.89 ± 0.01 Å when compared to 1.881 ± 0.005 Å in the published Fe₆ cluster [57]. In the 5-iron unit of cluster 3, the equatorial iron sites average an Fe–C bond distance of 1.86 ± 0.01 Å, while the axial iron resides at an elongated bond distance of 1.95(1) Å. This is strikingly similar to the published Fe₅ neutral cluster $[(Fe_{eq}-C_{avg} = 1.88 \pm 0.02 \text{ Å}; Fe_{ax}-C = 1.949(7) \text{ Å}]$. The Fe–C bonds found in 3, however, are shorter than the FeMoco average of 2.00 ± 0.02 Å.



Figure 7. Schematic indication of the dipole in electropositive sulfur reagents.



Scheme 6. Synthetic scheme depicting the electrophilic sulfurization of the Fe₆C dianion cluster by either S₂Cl₂ or elemental sulfur to afford a μ_4 -S cluster (**3** or **4**) and a {CS}^{4–} cluster (**5** or **6**). The non-carbide cluster [Fe₃S₂(CO)₉] (not depicted) was also characterized as a side product in both reactions. Image adapted with permission from Joseph, C., Cobb, C. R., Rose, M. J. Single-Step Insertion of Sulfides and Thiolate into Iron Carbide-Carbonyl Clusters: Unlocking the Synthetic Door to FeMoco Analogues. *Angew. Chemie Int. Ed.* **2020**, *Accepted, not yet published.* Copyright *Wiley-VCH GmbH*. Reproduced with permission.



Figure 8. Thermal ellipsoid plots (50% probability) of $(NEt_4)_2\{[(CO)_{15}(\mu_6-C)Fe_6](\mu_4-S)[Fe_5(\mu_5-C)(CO)_{13}]\}$ (3). Orange = Fe, Gray = C, Yellow = S, Maroon = O. Two NEt₄⁺ cations have been removed for clarity. Image adapted with permission from Joseph, C., Cobb, C. R., Rose, M. J. Single-Step Insertion of Sulfides and Thiolate into Iron Carbide-Carbonyl Clusters: Unlocking the Synthetic Door to FeMoco Analogues. *Angew. Chemie Int. Ed.* **2020**, *Accepted, not yet published.* Copyright *Wiley-VCH GmbH*. Reproduced with permission.

However, the generation of side products and the presence of chloride in the synthesis of the sulfide-bridged, asymmetric dimer led us to seek more straightforward reaction conditions. Since this cluster can be thematically formulated as an Fe₆ cluster, an Fe₅ cluster, a sulfur bridge, and an overall 2^- charge, we attempted to stoichiometrically construct the compound by introducing elemental sulfur into a solution of $(Et_4N)_2[Fe_6(\mu_6-C)(CO)_{16}]$ and $[Fe_5(\mu_5-C)(CO)_{15}]$. To our surprise, crystal structural data for this charged species in this reaction revealed a symmetric dimer of Fe₅ clusters bridged by the same 4-coordinate sulfide motif (Figure 9) with the formula $(NEt_4)_2[Fe_5(\mu_5-C)(CO)_{13}]_2(\mu_4-S)]$ (4). The Fe–C bond distances of this cluster are similar to the Fe₅ component of **3**, exhibiting an average Fe_{eq}–C bond distance of 1.87 ± 0.01 Å and an average Fe_{ax}–C bond distance of 1.961 ± 0.005 Å. The average Fe–S distance of 2.175 ± 0.005 Å, however, is notably shorter than the already short distance found in **3** and is far shorter than the Fe–S distances observed in FeMoco.



Figure 9. Thermal ellipsoid plots (50% probability) of $(NEt_4)_2\{[Fe_5(\mu_5-C)(CO)_{13}]_2(\mu_4-S)\}$ (4). Orange = Fe, Gray = C, Yellow = S, Maroon = O. Two NEt_4^+ cations have been removed for clarity. Image adapted with permission from Joseph, C., Cobb, C. R., Rose, M. J. Single-Step Insertion of Sulfides and Thiolate into Iron Carbide-Carbonyl Clusters: Unlocking the Synthetic Door to FeMoco Analogues. *Angew. Chemie Int. Ed.* **2020**, *Accepted, not yet published.* Copyright *Wiley-VCH GmbH*. Reproduced with permission.

Our efforts to fully characterize the product profiles for the reactions described above prompted us to separate and purify the remaining neutral compounds. The major component of the pentane-soluble portion is a deep, bright red compound. Isolation and crystallization of the bright-red compound generated the well-known cluster Fe₃S₂(CO)₉ [69,70]. More remarkably, crystals from the red-orange Et₂O solution revealed a remarkable CO-supported iron-sulfur cluster featuring a "carbide-like" site, multiple sulfur atoms, and a 'dangler' iron (Figure 10), resulting in the formula [{Fe₄(κ_2 S- κ_4 C)(CO)₁₀}(μ_3 -S)(μ_3 -S)Fe(CO)₃] (5).



Figure 10. Thermal ellipsoid plots (50% probability) of $[{Fe_4(\kappa_2 S - \kappa_4 C)(CO)_{10}}(\mu_3 - S)(\mu_3 - S_2)Fe(CO)_3]$ (5). Orange = Fe, Gray = C, Yellow = S, Maroon = O. Image adapted with permission from Joseph, C., Cobb, C. R., Rose, M. J. Single-Step Insertion of Sulfides and Thiolate into Iron Carbide-Carbonyl Clusters: Unlocking the Synthetic Door to FeMoco Analogues. *Angew. Chemie Int. Ed.* **2020**, *Accepted, not yet published.* Copyright *Wiley-VCH GmbH*. Reproduced with permission.

A thorough search through the CSD for {Fe₄(μ_4 -C)} motifs exclusively returns clusters in which the irons sites adopt a 'butterfly' geometry about the carbide, marking the planar geometry displayed in **5** as unique. While the average Fe–Fe bond distance (2.66 ± 0.09 Å) is typical of iron-carbonyl-carbide clusters, the average Fe–C distance of 1.97 ± 0.04 Å is remarkably longer than average for this type of cluster and, instead, is very close to the average Fe–C distance found in FeMoco (2.00 ± 0.02 Å). This elongation is—in part—an artifact of the displaced position of the carbide from the Fe₄ plane, such that the carbide actually resides 0.59 Å above the least-squares plane derived from the positions of the four Fe atoms. Lastly, the Fe–S distances found in **5** are notable as the average distance of 2.27 ± 0.03 Å, which is considerably elongated when compared to **3** and **4**, placing it closer to that of FeMoco (2.25 ± 0.03 Å). Of particular note, if 'belt sulfides' are excluded from the FeMoco average, the similarity is even more pronounced as the average Fe–S of 2.27 ± 0.03 Å in FeMoco is exactly on par with that of **5**. Two of the S sites (S3, S4) are bonded (2.048(2) Å), resulting in the presence of a persulfide S–S bond.

Purification of the nonpolar fractions collected during the synthesis of **4** provides a related crystal structure of formula [{Fe₄(κ_2 S- κ_4 C)(CO)₁₀}(μ_3 -S)₂Fe(CO)₃] (**6**) (Figure 11). Average bond distances are quite similar to those in **5**, with the marked difference being the average Fe–S of 2.25 ± 0.03 Å in **6**, which is nearly identical with that of FeMoco. The key difference, however, displayed in **6** is the absence of a sulfur site (S4 in **5**), therefore, designating S3 as a second face-bridging sulfide (S²⁻) site (in addition to S2). The absence of the S4 site, however, does induce a break in the *C*_s chemical symmetry displayed in cluster **5**, such that the Fe5–Fe3 distance has shortened to 2.616(1) Å. While we found that synthesis of cluster **6** has, in fact, been previously reported from the reaction of Fe₃(CO)₁₂ with CS₂ at 80 °C [71], the conspicuous absence of this structure from the CSD coupled with the fact that the original report has seemingly remained in obscurity from the bioinorganic synthetic modelling community has hindered its utilization as a clear candidate for nitrogenase structural modeling.



Figure 11. Thermal ellipsoid plots (50% probability) of $[\{Fe_4(\kappa_2S-\kappa_4C)(CO)_{10}\}(\mu_3-S)_2Fe(CO)_3]$ (6). Orange = Fe, Gray = C, Yellow = S, Maroon = O. Image adapted with permission from Joseph, C., Cobb, C. R., Rose, M. J. Single-Step Insertion of Sulfides and Thiolate into Iron Carbide-Carbonyl Clusters: Unlocking the Synthetic Door to FeMoco Analogues. *Angew. Chemie Int. Ed.* **2020**, *Accepted, not yet published*. Copyright *Wiley-VCH GmbH*. Reproduced with permission.

In addition to the biologically relevant bond metrics observed in the two {CS}-containing clusters, we found that the dangler Fe site in cluster **5** also approaches similarly relevant electronic properties (i.e., approaching a ferrous oxidation state). X-ray photoelectron spectroscopy (XPS) of **5** could be fit as 3 distinct components at both the Fe 2p(3/2) region (711.6, 709.4, and 707.5 eV) and the Fe 2p(1/2) region (724.6, 722.2, and 720.5 eV) with integrated peak area ratios of 1:2:2, respectively (Figure 12). On the basis of symmetry, the Fe5 site could be assigned to the higher binding energy peak (highest valence state Fe site). This assignment was additionally supplemented by density-functional theory (DFT) calculations, indicating Fe5 to be the Fe site with the least electron density on the basis of Mulliken population analysis and back-bonding contribution into the coordinated COs. The assignment of XPS peaks at 711.6 eV and 724.6 eV additionally falls within the expected range for a ferrous and CO-supported iron site [72,73]. The conclusions drawn from the XPS data are also consistent with the higher CO stretching frequencies observed in the IR spectrum in which all ν_{CO} values range upward from 2002 cm⁻¹. These are significantly blue-shifted relative to **3** ($\nu_{CO} \ge 1973$ cm⁻¹) and **4** ($\nu_{CO} \ge 1991$ cm⁻¹) and is additionally consistent with the presence of a ferrous site, which is not typically observed in CO-supported FeC clusters.



735 730 725 720 715 710 705 700 Binding Energy (eV)

Figure 12. Observed high-resolution X-ray photoelectron spectrum (XPS) and component fitting of the iron 2*p* region of cluster **5**. Image adapted with permission from Joseph, C., Cobb, C. R., Rose, M. J. Single-Step Insertion of Sulfides and Thiolate into Iron Carbide-Carbonyl Clusters: Unlocking the Synthetic Door to FeMoco Analogues. *Angew. Chemie Int. Ed.* **2020**, *Accepted, not yet published*. Copyright *Wiley-VCH GmbH*. Reproduced with permission.

In many respects, the construction of clusters 5 and 6 elevates the family of carbonyl-supported, carbide-containing iron clusters into the realm of highly structurally relevant models for the nitrogenase active site. Comparison of bond metrics between the clusters with FeMoco (Table 1) conveys the approaching similarity in 5 and 6 toward that analogous FeMoco metrics. However, the presence of a true C–S bond [1.714(5) Å] does preclude the C from being an authentic C^{4–} carbide. Nonetheless, this unusual bonding motif (only three entries in the CSD for transition-metal-bound {C-S}—none of which contain iron) is reminiscent of the proposed biogenesis mode of carbide insertion into the M-cluster as postulated by Wiig, Hu, and Ribbe depicted in Figure 2 [11,12]. The presence of the $\{C-S\}^{4-}$ motif (i.e., tetra-deprotonated methylthiol) in 5 and 6, thus, serves as the first rudimentary structural model for intermediates in M-cluster biosynthesis. Such intermediates would likely be difficult to isolate for crystallographic characterization. On the other hand, comparative spectroscopic studies of synthetic cluster compounds have previously demonstrated utility in the determination of structural aspects of nitrogenase, such as the assignment of the central carbide (which itself utilized the Fe₆ cluster precursor described above as a reference compound) [8,9]. The assortment of iron clusters that have been developed by our lab now present a range of reference compounds available for spectroscopic characterization and comparison against the various stages of nitrogenase biosynthesis: Clusters 1 and 2 provide examples of comparable alkylated and non-alkylated FeS clusters as appropriate for early-stage NifB reactions, while clusters 5 and 6 bearing the {C-S} motif present model compounds for intermediates in late-stage NifB activity.

	3	4	5	6	FeMoco
Fe–Fe	2.65 ± 0.06	2.64 ± 0.05	2.66 ± 0.09	2.7 ± 0.1	$\textbf{2.63} \pm 0.04$
Fe–C	1.88 ± 0.03	1.88 ± 0.04	1.97 ± 0.04	1.96 ± 0.02	2.00 ± 0.02
Fe–S	$\textbf{2.19} \pm 0.01$	$\textbf{2.175} \pm 0.005$	2.27 ± 0.03	2.25 ± 0.03	2.25 ± 0.03

Table 1. Selected bond distance averages of the FeCS clusters described in this work compared with the corresponding averages found in the nitrogenase cofactor (FeMoco).

3. Extension of Electrophilic Sulfur Ligation toward Fine-Tuning of Cluster Thiolates

3.1. Electronic Effects of Fe-Alkyl Bonding in Iron-Sulfur Cubane Clusters

The alkylated clusters generated by the Suess group (vide supra) illustrate how binding of an alkyl fragment at a single iron site significantly alters the oxidation states of the irons within the cluster. Classically, a $[Fe_4S_4]^{2+}$ cluster with relatively weak-field and uniform ligands features an overall S = 0 spin state with four $Fe^{2.5+}$ sites in high-spin configurations featuring double exchange within two pairs of $[Fe_2S_2]^+$ units that antiferromagnetically couple with each other. This holds true for the chloride precursor of the $[Fe_4S_4]^{2+}$ -ethyl cluster and for many thiolate-ligated protein-bound $[Fe_4S_4]^{2+}$ clusters. Suess et al. show that the introduction of a strong-field, electron rich alkyl ligand at the apical iron site polarizes the double-exchange interaction and introduces ferric character at the apical alkyl-bound iron. Correspondingly, it introduces ferrous character at its spin-aligned partner iron site with no overall change in charge [48].

This effect was replicated in the reported $[Fe_4S_4]^{3+}$ -methyl structure where the ferric character was found to be localized at the alkylated iron site. Furthermore, solution studies of this structure suggest rapid interconversion of the oxidation states at the three chelated iron sites, resulting in a shared oxidation state of essentially 2.67+. Overall, alkylation has a profound and unique effect on oxidation states of the irons within both the 2+ and the 3+ Fe₄S₄ clusters [49]. These two synthetic structures represent the first direct bioinorganic insight into Fe-alkyl bonding and the electronic attributes of Fe₄S₄-alkyl intermediates in nature. Perhaps most essential for the field, the range of chelating ligand scaffolds employed, and the demonstrated impact of remote steric effects on apical substrate binding in this class of clusters promise a drastic increase in the degree of synthetic control over iron-sulfur clusters [74]. This approach has already allowed for the isolation of increasingly elusive intermediates as well as the study of the reactivity of radical formation from alkyl intermediates [48,49,75], and suggests one possible route toward controlling the environment of reactive intermediates and active sites in other iron-sulfur metalloclusters as well.

3.2. New Routes toward Carbide-Containing Iron-Thiolato Clusters

The structure and reactivity of these alkyl clusters present novel findings relevant to radical SAM, but not directly relevant to the NifB intermediate. However, the growing family of chelating ligands and the control they afford over the oxidation states and reactivities of these clusters suggest new directions in rational cluster design that may yield novel insights into nitrogenase biogenesis intermediates [48,49]. Potential for any ligand-directed, cluster-based nitrogenase modeling remains to be borne out, but the recent results from Suess et al. suggest the stabilization of more reactive metalloclusters may be accessible through the design of suitable chelating ligand systems.

Historically, controlled ligation of sulfides and thiolates onto iron carbonyl carbide clusters has proven difficult to control and would generally result in cluster decomposition. However, upon the discovery that the [Fe₆] dianion was amenable toward substitution by an electrophilic sulfur source, we elected to extend the utility of this synthetic strategy toward the incorporation of a thiolato ligand. As a proof of concept, *p*-toluenesulfenyl chloride was reacted with the $[Fe_6]^{2-}$ cluster, yielding pink crystals of the first-ever reported thiolato-iron-carbide cluster (Figure 13) [67]. This controlled ligation of an iron-carbide cluster presents an as-yet-unrealized golden opportunity to finally engage this family of interstitial carbide clusters to be functionally "fine-tuned" with the introduction of an electronically

diverse range of ligands. As such, a new branch has sprouted in our own research program, and a pursuit for elegantly designed, functionally active carbide-containing iron clusters, which have, for so long, evaded the synthetic modeling community, may now be forthcoming.



Figure 13. Reaction conditions (left) to obtain (NEt₄)[Fe₅(μ ₅-C)(SC₇H₇)(CO)₁₃] and thermal ellipsoid plots (50% probability) of the thiolate-bearing cluster anion (right). Image adapted with permission from Joseph, C., Cobb, C. R., Rose, M. J. Single-Step Insertion of Sulfides and Thiolate into Iron Carbide-Carbonyl Clusters: Unlocking the Synthetic Door to FeMoco Analogues. *Angew. Chemie Int. Ed.* **2020**, *Accepted, not yet published*. Copyright *Wiley-VCH GmbH*. Reproduced with permission.

4. Closing Remarks

The pursuit for model compounds of structural relevance to FeMoco continues to inspire many creative research avenues for synthetic chemists. Still, the "Holy Grail" for synthetic FeMoco modeling—a paramagnetic iron cluster with inorganic sulfides, an interstitial carbide, and heterometal Mo—remains unaccomplished. Success in this pursuit will allow for insight into the role of the carbide as a structural and electronic modulator in nitrogenase and understanding into how the electronic properties of NifB affect carbide incorporation. Thus far, synthetic models focusing on FeMoco have shown that the carbide likely facilitates structural flexibility of the cluster and even promotes substrate binding and reduction (vide supra). Other modelling avenues have given insight into how such a C-donor ligand may promote very strong antiferromagnetic coupling in the cluster and play a significant electronic role in the cofactor.

In our own venture to progress ever closer toward achieving this goal, we have utilized carbidocarbonyl iron clusters as a synthetic starting point for structural modeling of the nitrogenase cofactor. While $[Fe_6(\mu_6-C)(CO)_{16}]^{2-}$ had previously served a crucial role in the assignment of the interstitial carbide in FeMoco (and later FeVco), further utilization of carbonyl-clusters for nitrogenase modelling is sparse. For our own research, a two-fold strategy toward pursuing nitrogenase-relevant synthetic clusters was employed. First, inspiration was taken from the radical-initiated biosynthetic mechanism as proposed by Ribbe and Hu in which an initial SAM methylates the K-cluster. The methylated FeS cluster is subsequently dehydrogenated by an additional equivalent of SAM. The pursuit of this strategy for carbide insertion led us to explore TEMPO-mediated reactions upon alkylated Fe₃S clusters. Although carbide insertion ultimately proved unsuccessful, the methylated precursor clusters were successfully isolated and characterized. Along with the Fe₃S precursor [Fe₃(μ_3 -S)(CO)₉], these clusters do provide a structurally interesting synthetic comparison for early-stage NifB reactions. In an alternative synthetic strategy, the carbide-containing Fe_6 cluster served as a starting point for sulfurization reactions. The reaction with electropositive sulfur reagents proved successful in achieving an FeCS cluster, and we were additionally pleased to discover two {C-S}⁴⁻-containing clusters, building upon our series of biogenesis-relevant models with appropriate compounds for late-stage NifB reactivity modeling. Future pursuits to close the gap between the early-stage and late-stage NifB models would see us aspire toward partially protonated/hydrogenated {SCH₂} and {SCH} structures [76].

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