Bis-[3]Ferroacenophanes with Central $\geq$ E–E’ $<$ Bonds (E, E’=P, SiH): Preparation, Properties, and Thermal Activation


A series of bis-[3]ferroacenophanes of the general type Fe(C$_5$H$_5$)$_2$E–E(=E’=E”(C$_5$H$_5$)$_2$)Fe (E = P, SiH and E’ = PrBu, NneoPentyl, NSi(CH$_3$)$_2$) with an isolobal molecular framework have been prepared and characterized by heteronuclear NMR spectroscopy and X-ray crystallography. The thermal dissociation behavior with respect to homolytic fission of the central bond generating phosphorus centered radicals was investigated using EPR spectroscopy and quantum chemical calculations.

1. Introduction

Recently, phosphorus modifications such as black phosphorus and especially its exfoliated version, phosphorene, attracted substantial interest because of their unique electronic and thermoelectric material properties.[1] While in most crystalline phosphorus modifications the relative orientation of the phosphorus lone-pairs is constrained by the three-dimensional lattice, molecular organopolyphosphorus frameworks can give rise to rather complex mixtures of diastereomers. Nevertheless, organopolyphosphorus frameworks may be regarded as molecular models for sections of phosphorus modifications such as the $(>\text{P})_2\text{P}($P$<$)$_2$ motif found in the rings and trenches of black phosphorus, or as connector units between the orthogonal or parallel tubes in Hittorf’s and Ruck’s variants of violet phosphorus (Chart 1).[2] However, the mobility at the central bond of the $(>\text{P})_2\text{P}($P$<$)$_2$ fragments is limited in the elemental allotropes by the rigidity of the solid matrix, while molecular compounds containing this particular unit may open a way to investigate the properties of this unique structural unit as well. In this respect, unsupported $(>\text{P})_2\text{P}($P$<$)$_2$ fragments, where the central P–P bond is not reinforced by additional bonding between the two halves of the fragment, would be attractive models to study the reactivity of this particular bond.

In previous work, we have shown that formally linking two terminal phosphorus atoms by a ferrocene fragment can provide a suitable template for the assembly of such cyclic structures.[3] As a consequence of their conformational constraints and rigidity, ferrocenophanes with phosphorus-rich fragments in the ansa-bridge can often be isolated as single diastereomers.[3a–c,4]

Beyond its beneficial structural features, the ferrocene unit usually provides reversible redox activity, where oxidation at iron may entail electron transfer with the phosphorus atoms.[5] Such oxidized systems could be related to cationic polyphosphorus compounds as described by Weigand and Burford et al.[6]
2. Results and Discussion


We anticipated that the bis-[3]ferrocenophane 4 (Scheme 1) would be readily accessible by reductive coupling of two molecules of P-halo-[3]ferrocenophane 1. However, reaction of 1 with metals such as magnesium or lithium under various conditions did not furnish the envisaged product, but yielded minor amounts of ferrocenophane 2 instead, which had previously been obtained from the same starting material and LiAlH₄. Trying to avoid a radical mechanism, which possibly occurred in the reduction of 1 with metals, we attempted to create the central PP bond by employing a heteropolar approach. To this end, we reacted lithium phosphanide 3 with one equiv. of 1 (Scheme 1). As intended, this reaction produced a yellow-orange solution of the hexaphospha-bis-[3]ferrocenophane 4 (Scheme 1), which was purified by recrystallization from toluene. Single-crystal X-ray diffraction analysis revealed that compound 4 crystallizes in a monoclinic space group (P 2₁/n). The crystal contains isolated molecules (Figure 1) showing the envisaged bis-ferrocenophane structure with orthogonal orientation of the ferrocene units (torsion angle between the Cp(cent)–Fe–Cp(cent) axes of both Fc units about
94°) featuring tilt angles of 3.48(3)° and 4.0(1)° respectively. On the whole, the structural features of the ansa-rings in each half of 4 match those of the trans-isomer of the corresponding mono-[3]ferrocenophanes we had reported earlier. It has to be noted that the distances between the central phosphorus atoms in the ansa-bridges and the closest iron centers (P2–Fe1 3.7313(7) Å, P5–Fe2 3.7542(7) Å) are somewhat shorter than the sum of the van-der-Waals radii (4.34 Å)[25], but longer than in ferrocene (Fe–P; X–Cl: 3.5523(3) Å, Br: 3.544(3) Å)[26].

As anticipated, the structural features of the P5-skeleton of 4 bear close similarity to the P,P-P,P-fragments linking the tubular subunits in Hittorf’s (violet) phosphorus.[27–30] Both species exhibit a central P–P bond that adopts a gauche conformation (dihedral angles 83.9° in Hittorf’s P, 92.8(1)° in 4) and is shorter than the adjacent bonds (2.178 vs. 2.199 to 2.206 Å in Hittorf’s P[28], 2.229(1) Å vs. 2.262(1) to 2.274(1) Å in 4). The same bond length pattern was also predicted computationally for the ω897XD/6-31+G* optimized structure of 4 (see Supporting Information) and elusive parent P3H5[31] and observed in other known molecules with P,P–P,P structural motifs.[32]

Solution 31P NMR spectra of 4 display line broadening effects and higher order patterns. The resonance of the central phosphorus atoms appears in the ambient temperature spectrum as a symmetric multiplet centered at −44.1 ppm (in toluene-d8), while the terminal phosphorus atoms give rise to a broad, structureless signal which extends from 0 to 30 ppm and exhibits maxima around 8 and 25 ppm. Assuming that the observed line broadening arises from a dynamic process, we performed a variable temperature NMR study on a CDCl3 solution of 4, as the solubility in this solvent is higher than in toluene.[15] As expected, the broad resonances transform eventually into two well separated multiplets at 18.8 and −4.0 ppm upon cooling to −45 °C, and the observed spectral pattern could be successfully simulated as an AA′MM′XX′ spin system (Figure 2). NMR experiments above room temperature revealed coalescence of the broad signals due to dynamic averaging of the AA′ and MM′ signals.

The 1H NMR spectrum of 4 displays at −45 °C signals attributable to two distinguishable tBu-groups and two Cp-rings, which undergo pairwise coalescence upon warming and have at room temperature merged into narrow dynamically averaged resonances. Dynamic lineshape simulations allowed us to relate the observed spectral changes to a single dynamic process. Determination of the activation parameters from an Eyring plot based on the fitted rate constants yielded values of ΔH$^{\ddagger}$ = 54.9(11) kJ mol$^{-1}$ and ΔS$^{\ddagger}$ = 17.7(4) J K$^{-1}$ mol$^{-1}$ (see the Supporting Information for details). We attribute the observed dynamics to a librational motion of the two Fe(CpP(tBu))P fragments relative to each other, which involves mutual interconversion of the two terminal phosphanyl fragments attached to each central phosphorus atom. Similar processes have been observed for tBu-P,P-PBu, and acyclic analogues of 4 reported by Kovacs[33] and Fritz[16] or in a symmetrically substituted bis-diazaphosphole, respectively.[17] It should be noted that ΔH$^{\ddagger}$ of the librational process comes close to the calculated ω897XD/6-31G* inversion barrier at the central phosphorus of 71.1 kJ/mol. This value is significantly smaller than the usual inversion barriers for the phosphorus atoms in phosphanes (146 kJ/mol for PH3),[19] or the terminal phosphorus atoms in 4 (133.5 kJ/mol), respectively. A lowering of the inversion barrier with respect to these reference values is, however, quite common for branched oligophosphanes.[19]

A comparison to the aforementioned related hexaphosphanes reveals that the 31P NMR chemical shifts for Weigand’s compound (−41.8 ppm (2 P), 13.2 ppm (4 P))[30] match those observed for 4 in the high temperature limit, while Wiberg’s silylated system resonates at higher field.[20] The acyclic congeners reported by Kovacs[33] and Fritz[16] give rise to 31P NMR spectra which display similar signals as 4 at low temperature and show also a similar temperature dependence.

2.2. Synthesis of P5(SiH)-bis-[3]ferrocenophane

A variation of the structural motif in hexaphospha-bis-[3]ferrocenophane 4 can be achieved by formal substitution of the central phosphorus atoms by isolobal units such as a silyldyne (SiH) fragment. While the attempt to synthesize a pentaphospha-bis-[3]ferrocenophane 6 (Scheme 2) by coupling of

![Scheme 2. Synthesis of bis-[3]ferrocenophanes 6 and 7.](image)

![Figure 2. Measured (black traces) and simulated (red traces, see SI for details) 31P{1H} NMR spectra of 4 recorded at −45 °C (bottom) and 0 °C (top).](image)
lithium phosphanide 3 with the previously published hydridochlorosilane 5\(^{[10]}\) failed, salt metathesis of 5 with the recently reported potassium phosphanide 3\(^{[1]}\) readily afforded 6 as yellow solid (Scheme 2).

A single-crystal X-ray diffraction study confirmed the molecular structure of 6, which is isostructural to that of 4 (Figure 3). The sum of PSiP angles around Si1 (319.7(3)°) is larger than the sum of PPP angles around P2 in 4. Furthermore, the central Si1-P23 bond (2.302(3) Å) is no longer the shortest Si–P bond of the scaffold, exceeding contacts Si1-P11 (2.264(3) Å) and Si1-P12 (2.269(3) Å). Compared to other known P-Si-P bridged [3]ferrocenophanes (P-Si 2.2380(5)–2.255(1) Å),\(^{[20]}\) these bonds are slightly elongated.

The \(^{31}\)P NMR spectra of 6 reveal a similar temperature dependence as in case of 4. The central phosphorus atom of the triphosphane part of the \(\text{P}_2\text{(SiH)}\text{-P}_3\) backbone appears in the room temperature spectrum as a triplet of triplets at \(-106.6 \text{ ppm with splittings of 163 Hz and 16 Hz due to coupling with the adjacent and remote ferrocene bonded phosphorus atoms, which themselves give rise to broad resonances between }-59 \text{ to } -34 \text{ ppm (}\nu_{11} \approx 1300 \text{ Hz, Si-bound P-atoms}\)\(^{[19]}\) and 2 to 20 ppm (\(\nu_{12} \approx 1600 \text{ Hz, P-bound terminal phosphorus atoms})\). The broad signals sharpen when the temperature is raised, and below room temperature eventually decoalesce into four multiplets, which display at \(-40 \text{ °C the signal pattern of a nearly first-order AFHMX spin system (Figure 4). The marked differences in the geminal (\(J_P\)) couplings between the three silicon-bound phosphorus atoms reflects presumably the dependence of the coupling interaction on the relative orientation of the phosphorus lone-pairs and has previously also been observed for the cis-trans-isomers of 5.\(^{[19]}\) The sensitivity of \(J_{PP}\) couplings towards conformational changes is well known both in general\(^{[21]}\) and in the chemistry of polyphosphorus compounds,\(^{[22]}\) and the observation of large magnitudes for “through space” couplings has been attributed to the overlap of the phosphorus lone-pairs in space.\(^{[22]}\) Similar temperature dependent changes are observable in the \(^{31}\)P NMR spectra of 6. The \(^{31}\)P NMR spectra displays at room temperature two signals attributable to \(tBu\)-groups and a complex signal pattern for the cyclopentadienyl groups and a complex signal pattern for the cyclopentadienyl protons, which merge into a single resonance (\(tBu\)) or the pattern of four signals expected for a single Cp-unit at higher temperature, respectively. The dynamically induced changes are, as in the previous cases, attributable to a librational motion of the two \(\text{P}_2\text{Si(H)}\text{-P}_3\) fragments. We did not determine the activation barrier for this process, but the calculated energy

**Figure 4.** \(^{31}\)P-\(^{1}\)H NMR spectrum of 6 at \(-40 \text{ °C with expansions of the individual multiplets. Signals labelled with an asterisk are due to impurities / decomposition products (Fe(CpPH(Bu))\(_3\)). Red traces represent the result of a spectral simulation as an AFHMX spin system; simulation parameters \(U\) given as absolute values: }\(\delta_1 = 14.0 \text{ (P-PBu)}, \delta_2 = 4.3 \text{ (P-PBu)}, \delta_3 = -40.3 \text{ (Si-PtBu)}, \delta_4 = -57.3 \text{ (Si-PtBu)}, \delta_5 = -109.1 \text{ (>PSiH)} <), \gamma_{11} = 139, \gamma_{12} = 165, \gamma_{13} = 135, \gamma_{14} = 113, \gamma_{15} = 32.\)**

2.3. Synthesis of \(\text{P}_2\text{(SiH)}_2\text{-bis-[3]Ferrocenophane}\)

The synthesis of \(\text{Si–Si}\) coupled tetraphospha-bis-[3]ferrocenophane 7 is accomplished by lithium mediated coupling of literature known hydridochlorosilane 5\(^{[10]}\) at elevated temperature (170 °C). Bis-[3]ferrocenophane 7 is obtained as yellow solid in moderate yield (Scheme 2).

The \(^{31}\)P NMR spectrum of 7 in toluene solution at room temperature displays two resonances of an AB spin system at \(-44.9 \text{ ppm and } -50.1 \text{ ppm (}\gamma_{PP} = 123 \text{ Hz), which exhibit dynamic broadening when the temperature is raised and are close to coalescence at 75 °C (cf. Si for details). The }\(^{1}\)H NMR spectrum displays at room temperature two signals attributable to \(tBu\)-groups and a complex signal pattern for the cyclopentadienyl protons, which merge into a single resonance (\(tBu\)) or the pattern of four signals expected for a single Cp-unit at higher temperature, respectively. The dynamically induced changes are, as in the previous cases, attributable to a librational motion of the two \(\text{P}_2\text{Si(H)}\text{-P}_3\) fragments. We did not determine the activation barrier for this process, but the calculated energy
barrier for P inversion of 79.9 kJ/mol (ωB97XD/6-31G*) matches the value for the analogous process in 6.

A single crystal XRD study revealed that 7 is likewise isostructural to 6 and 4. The sum of P–Si–(P,Si) angles around each silicon atom (Si1 328.9(1)°, Si2 327.5(1)°) is even larger than in 6. The Si1-Si2 distance is 2.3828(9) Å, which is even longer than in a sterically challenged disilane (2.354(3) Å).[24] All other bond lengths and angles in 7 are comparable to 4 and 6 and will not be discussed in detail (Figure 5).

Figure 5. Molecular structure (left) of 7 and conformation of its P,Si(H)-Si(H) P₂ motif (right). Ellipsoids are drawn at 30% probability level.


Nitrogen-containing bis-ferrocenophanes are accessible starting from 1,1′-diaminoferrocenes. Base-promoted condensation of 13 a with PCl₅ followed by reductive coupling of the spectroscopically detectable 2-chloro-1,3,2-diazaphospha-[3]ferrocenophane 14 a[25] with magnesium yields tetraaza-bis[3]ferrocenophane 8. Finally, diaza-bis-[3]ferrocenophane 9, which can be regarded as a link between 4 and 8, is obtained via salt metathesis between 14 b[25] and 3 (Scheme 3). Both 8 and 9 were isolated as yellow solids in reasonable to good yields.

The ³¹P NMR spectrum of 8 displays the single line of an A₂ spin system. The ³¹P NMR spectrum of 9 displays two diastereomers featuring A₂B₈ (major isomer) and ABMX (minor isomer) type spin systems (Figure 6). The ¹H NMR spectrum appeared at first glance rather complicated, but the signals of the major component were readily identified and assigned from two-dimensional ¹H COSY and NOESY NMR spectra. Tracking characteristic NOESY correlation signals arising from mutual chemical exchange between both species then allowed us to assign the ¹H NMR signals of the minor component, and to establish that both species undergo reversible dynamic interconversion on a second time scale. The number of ¹H and ³¹P NMR signals and the coupling pattern in the ³¹P NMR spectrum exposes both components as isomers that are distinguished by different effective symmetry (C₃ or C₂ for the major and C₁ for the minor isomer). These findings cannot be explained by assuming simply the presence of different rotamers as in 4, 6, and 7, but are compatible with an assignment to two diastereomeric bis-ferrocenophanes featuring cis-(major isomer) and trans-alignment (minor isomer) of the tBu-groups in the PPP-ferrocenophane ring, respectively. To explain the observed patterns of ²Jₚₚ coupling constants, we hypothesize that the major isomer adopts a conformation that is characterized by a close proximity between the lone-pairs on all three terminal phosphorus atoms which could provide substantial through-space contributions[23b–d] to the coupling between the diaminophosphorus and both tBu-substituted phosphorus atoms (Figure 5a). The trans-isomer can formally be generated by inversion of one tBu-unit, which would quench any through-space contributions to ²Jₚₚ couplings involving this phosphorus atom (Figure 5b). Similar conformational influences on ²Jₚₚ coupling constants are well established in the literature.[21,23b–d] DFT calculations predict that the cis-isomer is more stable, but the energy difference between the stereoisomers is small (7.5 kJ/mol). The ωB97XD/6-31G* inversion barrier for isomerization of
the cis- into the trans-isomer is 66.9 kJ/mol, which is compatible with a reversible interconversion between both stereoisomers at ambient temperature. The inversion barriers of the phosphorus atoms in 9 are low within the P₃ unit (66.9, 69.5 and 85.4 kJ/mol), unlike the inversion of the phosphorus atom in the NPN unit (146.0 kJ/mol).

Single crystal XRD studies of 8 and 9 show a twisted arrangement with respect to the P–P bond for 8, while 9 adopts a nearly anti-periplanar orientation (Figures 7 and 8). The P–P distance of the central bond is 2.3019(11) Å for 8 and 2.281(3) Å for 9, both exceeding the respective bond length in Hittorf’s phosphorus[7–8] but well within the range observed for bulkily substituted tetraamino-diphosphanes (2.24–2.34 Å)[26]. The pyramidal coordination geometry around the phosphorus atoms of 8 (sum of bond angles 313(1)°) is slightly flattened compared to 4 (sum of bond angles 304–308°) and becomes even more planarized in 9 (sum of angles 322(1)°/323(1)°). The sums of bond angles at the nitrogen atoms indicate a quasi-planar geometry for these atoms in both 8 (357(1)°/359(1)°) and 9 (352(2)°/347(2)°).

The observed conformation of crystalline 9 suggests that all phosphorus lone-pairs in the P₃ motif are oriented gauche to each other. The presence of short P–P distances (2.204(3)/2.212(3) Å) and gauche-orientation of the lone-pairs on adjacent phosphorus atoms in the PPP-ferrocenophane ring compared to an increased P–P distance and an anti-periplanar orientation of the lone-pairs in the central bond are suitable to explain the large deviation between ¹JPP coupling constants in the ansa-ring (360–464 Hz) and the central bond (241 Hz), respectively. If 4 can serve as model compound for the connecting units in Hittorf’s phosphorus, then 9 shows a similar connection mode as in fibrous red phosphorus (Ruck’s Phosphorus).[2a,b,4]

2.5. Thermal Homolytic Bond Activation of bis-[3] Ferrocenophanes

Thermally induced homolytic cleavage of the central bond in sterically encumbered diphosphanes (X₂P–PX₂) is an attractive and traceless way to generate phosphorus centred radicals. This methodology works particularly well for the generation of amino-phosphanyl radicals (X₆P₄; X=RN) which entails a propensity for bond fragmentation in sterically strained tetraaminodiphosphanes.[17b,2a,26] In this context, we explored the potential of the above mentioned bis-[3]ferrocenophanes to undergo thermally induced homolytic cleavage of the central P–P, P–Si or Si–Si bond, respectively.

The experimental studies were carried out by heating solutions of bis-ferrocenophanes 4, 6–9 in suitable inert solvents (toluene, mesitylene) in the cavity of an EPR spectrometer in order to detect any radicals resulting from a homolytic bond fission process. The temperature was generally raised in 10 °C steps up to a maximum of 165 °C. In case of 4, visible EPR signals started to emerge at approximately 90 °C and grew continuously in intensity until the maximum temperature of 165 °C was reached. The temperature dependent changes were reversed when the temperature was lowered again, and the signals disappeared again below 90 °C. The observed spectral pattern (Figure 9) can be decomposed into two signals with relative intensities of approx. 1:3 that do not vary perceptibly over the temperature range studied.

Spectral simulations allowed to describe each signal as a doublet of triplets resulting from hyperfine coupling with one

![Figure 7. Molecular structure (left) of 8 and conformation of its N₃P-P₃ motif (right). Thermal ellipsoids for the heavy atoms are drawn at 30% probability level.](image)

![Figure 8. Molecular structure (left) of 9 and conformation of its N₃P-P₃ motif (right). Thermal ellipsoids for the heavy atoms are drawn at 30% probability level.](image)

![Figure 9. Measured EPR spectrum (violet trace) of a solution of 4 at 150 °C in mesitylene and result of a spectral simulation (green trace).](image)
and two phosphorus nuclei, respectively (more intense signal: \( g = 2.0012, A = 40.1 \) \( G \) (1 \( P \)), 65.4 \( G \) (2 \( P \)); less intense signal: \( g = 2.0011, A = 70.6 \) \( G \) (1 \( P \)), 79.5 \( G \) (2 \( P \))). The hyperfine splitting of both signals is in accord with the presence of triphospha-[3]ferrocenophanyl radicals 11 (Scheme 4), the de-

tection of which indicates that the expected bond homolysis process has taken place. The observation of more than one signal can be accounted for by the presence of different stereoisomers. Anticipating that the \( \text{CP}_2\text{C} \)-bridge (where \( C \) denote the \( \text{P} \)-substituted \( \text{C} \)-carbon atoms) adopts a similar non-planar conformation as in 4, one can envisage two diastereomers featuring cis- or trans-orientation of the \( \text{P} \)-Bu-substituents relative to the “ring” structure defined by the two \( \text{Cp} \)-centroids and the \( P_3 \) bridge, respectively. DFT calculations support the identity of radical 11. The calculated (\( \omega\text{B97XD/6-31} + \text{G}^* \)) Gibbs free energy difference between the two isomers is only 0.8 \( kJ/mol \), and the barrier for planar inversion at one of the phosphorus atoms is only 65.7 \( kJ/mol \). The calculated spin density (Figure S1) verifies that the radical center is localized at the central phosphorus atoms of both isomers. The calculated hyperfine coupling constants (for more information see Table S2 in SI) are in agreement with the experimentally observed values and give further support for the proposed constitutional assignments of the isomeric species. Similar conformational isomerization processes as observed here have been reported in related cases.[11,14]

The observed temperature dependence of the EPR spectra suggests that the radicals 11 recombine at lower temperature and exist in a dynamic equilibrium with the diphospha. Similar reversible equilibria have precedence for amino-diphosphanes.[17b,20,28a–f] Prolonged heating of toluene solutions of 4 under inert conditions affords detectable amounts of secondary phosphophane 2, suggesting that radicals 11 may be irreversibly quenched by hydrogen abstraction, e.g. from the solvent. Triphospha 2 is also formed in the reaction of 4 with tributyltin hydride. Even if the tin-containing reaction product could not be identified, formation of the phosphophane can be accounted for by a radical promoted reductive hydrogenation of the central \( \text{P} \)\( \text{P} \) bond in 4.

HT EPR measurements on toluene solutions of 9 revealed that the signals attributable to radical 11 already appeared at ca. 60 °C. Interestingly, signals attributable to the complementary NPN radical 12 could not be detected. The EPR studies suggest that homolytic cleavage of the central \( \text{P} \)\( \text{P} \) bond in 9 occurs even more readily than in case of 4 but the postulated aminophosphanyl radical is immediately quenched (Scheme 5).

This hypothesis is corroborated by the observation that heating of a toluene solution of 9 to 100 °C under inert conditions results in controlled decomposition, which can be traced by \( ^{31}\text{P} \) NMR spectroscopy. With continuous heating, the signals of 9 fade away, while those of recombination product 4 and hydrogen abstraction product 2 together with two additional singlets at lower field start to grow in. One of the new signals at 110.4 ppm splits into a doublet with a \( J_{\text{PP}} \), coupling constant of 272 Hz when proton decoupling is switched off. Similar shifts and coupling constants were reported for secondary dianomophosphanes,[29] and we assign this signal therefore to the NPN analogue \( 2^+ \) of 2. The formation of \( 2^+ \) is in good agreement with the predicted exothermic outcome (\( \Delta \text{E} = -23.4 \) \( kJ/mol \) at the \( \omega\text{B97XD/6-31} + \text{G}^* \) level) of the reaction \( 2 \rightarrow 2^+ + 11 \), which implies that the hydrogen abstraction capability of NPN-based radical 12 is higher than that of PPP-based radical 11. The chemical shift of the second singlet at 165.0 ppm is slightly larger than those of bulkily substituted N-heterocyclic diphosphanes[10c–21] which resonate around 150 ppm. It is tempting to attribute the singlet to a symmetrical diphospha resulting from recombination of radical 12. However, all attempts to access \( 2^+ \) or the postulated symmetrical diphospha by reduction of chlorophosphate \( 14 \), using e.g. lithium aluminium hydride or metal mediated reductive coupling reactions, were as yet unsuccessful.

Similar HT EPR experiments on tetraamino-bis-ferrocenophane 8 and the silicon analogues 6 and 7 up to 120 resp. 250 °C do not result in formation of any detectable radicals. While the absence of the dissociation in case of 6 and 7 is in good agreement with their higher dissociation Gibbs free energy (175.7 \( kJ/mol \) for 6 and 225.1 \( kJ/mol \) for 7 at \( \omega\text{B97XD/6-31} + \text{G}^* \) level of theory and related to 25 °C) compared to 4 and 9 (93.3 \( kJ/mol \) and 100.8 \( kJ/mol \), respectively), the low calculated dissociation Gibbs free energy of 3.8 \( kJ/mol \) for 8 suggests that thermolytic cleavage should be energetically feasible. However, quenching and high reactivity of the N,P centered radical may preclude its practical observation just as in the case of 9.

2.5.1. Electrochemical Behavior of Bis-[3]ferrocenophanes

Intrigued by the energetic proximity of iron and phosphorus centered electronic states in the frontier orbital region of such compounds, we explored the electrochemical properties of the
bis-[3]-ferrocenophanes presented. Cyclic voltammetry measurements of 4 in THF with a platinum working electrode in the presence of TBAHFP as conducting salt revealed an irreversible oxidation process with its peak potential at 0.031(1) V (vs. Fc/Fc\(^-\)). Several further oxidations indicated by broad current responses can be observed at potentials around 1 V (vs. Fc/Fc\(^-\)). After several cycles, a second defined oxidation wave starts to emerge at ca. 0.2 V (Figure S4). We assign this wave to triphosphate 2, which shows an oxidation response of the ferrocene unit at 0.22(1) V (vs. Fc/Fc\(^-\)) in DCM (see SI for detailed information). This interpretation is consistent with the observation of detectable amounts of 2 in \(^{31}\text{P}\) NMR spectra recorded subsequent to the electrochemical experiments and implies that electrochemical oxidation of 4 entails formation of 2, probably via intermediate formation of radical 11 which should be prone to hydrogen abstraction from the solvent (see above). The dissociation of 4\(^{+}\) into 11 and the corresponding cation is somewhat endergonic (the Gibbs free energy is 36.0 kJ/mol at \(298\text{K}\) \(\text{atm}\)), which is significantly smaller than the 93.3 kJ/mol required for the dissociation of neutral 4 into two molecules of 11. For further elucidation, we performed dual electrode experiments on 4 using a rotating ring-disc electrode to determine the average lifetime of 4\(^{+}\). However, even at rotation rates up to 2000 rpm no reductive response could be detected at the ring, indicating a lifetime of the anticipated radical cation shorter than 0.02 seconds (Figure S6).

Hybrid structure 9 displays similar electrochemical behavior, although more redox events are detectable. The first oxidation occurs at a potential of \(-0.26(1)\) V (vs. Fc/Fc\(^-\)), which is in the expected range for amino-substituted ferrocenophanes,\(^{[25]}\) as an irreversible process (Figure S7). At higher potential, two quasi reversible oxidation processes (0.02(1) and 0.13 (1) V vs. Fc/Fc\(^-\)) were detected using differential pulse techniques. Further broad current responses above 0.4 V (vs. Fc/Fc\(^-\)) indicate consecutive reactions which are not interpretable. Both current responses at potentials near the ferrocene standard are comparable to hexaphosphate 4. As in the case of 4, repetitive measurements entail a distinct decrease of the signals accompanied by new signals at \(-0.66(1)\) V vs. Fc/Fc\(^-\) (for comparison, \(1,1^*\)-diaminoferrocene at \(-0.60\) V\(^{[26]}\) resp. \(1,1^*\)-bis(dimethylamino)ferrocene at \(-0.63\) V\(^{[21]}\) vs. Fc/Fc\(^-\) in CH\(_3\)CN and around 0.1-0.2 V). The latter is superimposed by the previously mentioned broad signals, but could be attributed to triphosphate 2. In accordance, \(^{31}\text{P}\) NMR experiments on the electrolyte solution after the measurements revealed the presence of triphosphate 2 and evidence for fluoro-analogue 14b.\(^{[2]}\) It is noteworthy that measurements with chloride-containing conducting salts like tetrabutylammonium chloride were precluded by instantaneous reaction with compound 9 forming chlorophosphate 14b. Both experimental findings imply a cleavage of the central P–P bond upon oxidation or by reaction with the conducting salt, which might well be heterolytic with the positive charge remaining on the N\(_2\)P scaffold. Analogously to the case of 4, oxidation of 9 to 9\(^{+}\) can be considered to permit facile dissociation of the central P–P bond – the Gibbs free energy for the decomposition of 9\(^{+}\) to 11 and 12\(^{+}\) was calculated as 61.1 kJ/mol (the alternative fragmentation of 9\(^{+}\) into 11\(^{+}\) and 12 was calculated as more endergonic by 86.6 kJ/mol) – which suggests to relate the electrochemical behavior to the occurrence of chemical decay processes (see Figure S2 for spin density distribution in 4\(^{+}\) and 9\(^{+}\)).

Similar to 4 and 9, diphosphate 8 reveals an irreversible first oxidation process (\(-0.21(1)\) V vs. Fc/Fc\(^-\); Figure S8). A second process follows at 0.11(1) V (vs. Fc/Fc\(^-\)), which appears as (quasi)-irreversible oxidation at first glance. Again, broad current responses are detectable at higher potentials which cannot be assigned. Differential pulse measurements indicate that the reductive response is independent from the second oxidation wave, which may be attributed to the occurrence of unspecified follow-up products. In case of the silicon based bis-[3]-ferrocenophanes 6 and 7, unspecific fragmentation of the main-group framework occurred, which precluded further interpretation. As a general feature of these compounds, their propensity to undergo inversion and other dynamic processes entails a variable energetic sequence of the frontier orbitals, which precludes a simple correlation between the electrochemical behavior and the HOMO/LUMO orbitals of the ground state.

3. Conclusions

In summary, we developed a series of bis-[3]-ferrocenophanes in which the ansa-units consist of isolobal trivalent fragments. Despite the number of potential P-stereogenic centers, the ferrocenophane motif limits the number of diastereomers observed for this series of compounds. As a characteristic feature, most compounds show restricted rotation around the central bond at ambient temperature, which leads to a characteristic temperature dependence of the respective NMR spectra. Quite remarkably, DFT calculations predict that the inversion barriers for some bridgehead phosphorus atoms are only slightly higher than the activation energies for the rotational processes. For bis-ferrocenophane 4, thermally induced homolytic cleavage of the central P–P bond of the P–P–PP \(_2\) fragment is observed at elevated temperature. Similarly electrochemical oxidation leads to the hydrogenated derivatives of the same radical. The barrier for thermal P–P fission is significantly reduced by formal change to a P–P–PN\(_2\) as in 9 (and N\(_2\)P–PN \(_2\) as in 8). Formal replacement of P with isolobal SiH units on the other hand does not lead to analogous silyl radicals up to a temperature of 250 °C. Based on the structural analogy of 4 with P–P–PP \(_2\) fragments in phosphorus modifications outlined above, our results impose the question whether similar homolytic bond rupture may occur in such solid state materials at elevated temperatures as well, although the dissociation barriers are to be expected at much higher energies owing to the reinforcement of the columnar structures in the lattice.

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Conflict of Interest

The authors declare no conflict of interest.

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