Coordination Chemistry Reviews 441 (2021) 213927

Contents lists available at ScienceDirect

Coordination Chemistry Reviews

journal homepage: www.elsevier.com/locate/ccr



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ARTICLE INFO

Article history: Received 4 December 2020 Received in revised form 9 March 2021 Accepted 14 March 2021

This contribution is dedicated to our mentor and friend Maurizio Peruzzini on the occasion of his retirement.

ABSTRACT

This review comprehensively covers the coordination chemistry of elemental phosphorus during the years 2011–2020. The main part is dedicated to the activation of white phosphorus mediated by early and late transition metals and its subsequent reactivity once in the coordination sphere of a metal. The burgeoning interest towards early TMs has opened the way to new coordination topologies and types of reactivity. Meanwhile, the coordination chemistry of P4 towards late transition metal elements has also attracted significant attention. Numerous studies have targeted transformations of P₄-derived phosphorus ligands and the subsequent release of said ligand fragments. The final part of this review focuses on the black allotrope of phosphorus, which offers a polydentate phosphorus surface able to interact with transition metals forming new functionalized 2D materials.

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https://doi.org/10.1016/j.ccr.2021.213927

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1. Introduction

Phosphorus (element 15 in the periodic table) is essential to life and human society [1,2]. It is one of the six "biogenic elements", which are needed in large quantities to create life [3]. Inorganic and organic phosphorus-containing molecules are ubiquitous in living organisms and nature. With that said, the industrial utilization of phosphorus for synthetic materials has prospered for over a century. A multitude of phosphorus compounds are produced by industry and have important commercial applications in agriculture, chemistry, and pharmaceuticals [4].

The high fundamental and industrial significance of elemental phosphorus has inspired extensive studies of its reactivity and its structural features. A range of different allotropes of phosphorus are known, among which, the three most common are white, red, and black phosphorus (Scheme 1). White phosphorus is the most reactive allotrope and, undoubtedly, the most industrially relevant form of elemental phosphorus [5]. Its structure consists of discrete tetrahedral P₄ molecules, with single P-P bonds. It is produced

commercially by heating phosphate rock in an electric furnace in the presence of silica and suitable reducing agents, mainly coke [6,7].

The production of organophosphorus compounds generally relies on the white allotrope as a starting material. This is disadvantageous as the process is energy intensive, requires highly corrosive/hazardous reagents, proceeds via harmful intermediates and generates large amounts of chemical waste. Consequently, it has been a goal in contemporary research to find methods for P₄ transformations that replace this unsustainable and environmentally harmful process. Such efforts towards a comprehensive, efficient, and less-polluting transformation of P₄ have been matter of interest for over four decades, being extensively studied by many research groups. On an academic level, alternatives such as the direct reaction of P_4 with main group elements [8–14] have been envisaged, but these processes usually suffer from low selectivity and low efficiency, thus their transfer to an industrial level looks doubtful. Other P-sources [15,16], such as phosphoric acid [17], traditionally utilized to produce phosphate fertilizers, could be

Abbreviations: Acac, acetylacetonate; BAr^F₄, tetrakis[3,5-bis(trifluoromethyl)phenyl]borate; BDI, NN'-bis(2,6-diisopropylphenyl)-3,5-dimethyl-β-diketiminate; ^{dipp}BIAN, 1,2-bis(2,6-diisopropylphenylimino)acenaphthene; MesBIAN, bis(mesitylimino)acenaphthenediimine; tBu, tert-butyl; tBuXPhos, PtBu₂(2',4',6'-triisopropylbiphenyl); Bu, butyl; BP, black phosphorus; BPQD, black phosphorus quantum dots; bpy, 2,2'-bipyridine; Bz, benzyl; CHD, cyclohexadiene; CHT, cycloheptatriene; CNAr^{Dipp2}, 2,6-Bis(2,6diisopropylphenyl)phenyl isocyanide; CNAr^{Mes2}, 2,6-Bis(2,4,6-trimethyl)phenyl isocyanide; CNAr^{Trip2}, 2,4,6-tris(2,6-diisopropylphenyl)phenyl isocyanide; Cot, cyclooctatetraene; Cod, cyclooctadiene; Cp, cyclopentadienyl; Cp*, pentamethylcyclopentadienyl; Cp'', 1,3-di-tert-butylcyclopentadienyl or 1,3-di-(trimethylsilyl)cyclopentadienyl; Cp^{'''} or Cp^R, 1,2,4-tris-tert-butylcyclopentadienyl; Cp^{BIG}, pentakis(4-n-butylphenyl)cyclopentadienyl; Cp^{Bn5}, Pentabenzylcyclopentadienyl; Cp^{4iPr}, C₅HiPr₄; Cy, cyclohexyl; D-A, Diels-Alder; DB18C6, dibenzo-18-crown-6 ether; DCM, dichloromethane; DFT, density functional theory; DIB, 1,3-diisopropylbenzene; DMAP, 4-*N*,*N*-dimethylaminopyridine; DMB, 2,3-dimethyl-1,3-butadiene; DME, dimethoxyethane; DMF, dimethylformamide; Dmp, 2,6-(2,6-di-isopropyl)phenyl; Dipp, 2,6diisopropylphenyl; DippForm, N,N'-bis(2,6-diisopropylphenyl)formamidinate; DIP₂pyr, 2,5-bis{N-(2,6-diisopropylphenyl)iminomethyl}pyrrolyl; dppe, 1,2-bis(diphenylpho sphino)ethane; dppm, 1,1-bis(diphenylphosphino)methane; dppp, 1,3-bis(diphenylphosphanyl)propane; dtbbpy, 4,4'-di-tert-butyl-2,2'-bipyridine; ENDOR, Electron-Nuclear Double Resonance Spectroscopy; EPR, electron paramagnetic resonance; ESI, electrospray ionization; Et, ethyl; ETM, early transition metal; EXSY, exchange spectroscopy; FAI, [FAI{OC₆F₁₀(C₆F₅)}]⁻; Fc, ferrocenyl; HMPA, hexamethylphosphoramide; HOMO, highest occupied molecular orbital; HSAB, Hard and Soft (Lewis) Acid and Base; IDipp, 1,3-bis(2,6-diisopropylphenyl)imidazolin-2-ylidene; IMes, [1,3-bis(2,4,6-trimethylphenyl)imidazolin-2-ylidene; IPh, 1,3-diphenylimidazolin-2-ylidene; JohnPhos, PtBu₂(o-biphenyl); LA, Lewis acid; LMCT, ligand to metal charge transfer; LTM, late transition metal; LUMO, lowest unoccupied molecular orbital; MALDI, matrixassisted laser desorption/ionization; MAS, magic-angle spinning; Me, methyl; Mes, Mesityl, 1,3,5-trimethylbenzene; MO, molecular orbital; MRI, magnetic resonance imaging; MS, mass spectroscopy; nacnac, HC(C(Me)NC₆H₃iPr₂)₂; NBO, natural bond orbital; NHC, *N*-heterocyclic carbine; ^{Me}NHC, 1,3,4,5-tetramethylimidazolin-2-vlidene; NMP, N-methyl-2-pyrrolidone; NMR, Nuclear Magnetic Resonance; NN^{fc}, [(η⁵-C₅H₄SiMetBu₂)₂Fe; Np, neopentyl; NPA, natural population analysis; n-Pr, normal propyl; iPr₂Im, 1,3-di-isopropylimidazolin-2-ylidene; iPr, isopropyl; Ph, phenyl; PHDI, bis(2,6-diisopropylphenyl)phenanthrene-9,10-diimine; PhPP₂Cy, PhP(CH₂CH₂PCy₂)₂; Ppy, 2-(2pyridyl)phenyl; rt, room temperature; SQUID, superconducting quantum interference device; TEF, $[Al{OC(CF_3)_3}_a]$; TEM, transmission electron microscopy; Tf, trifluoromethanesulfonate; THF, tetrahydrofuran; Tipp, 2,4,6-triisopropylphenyl; TM, transition metal; TMDPDD, 3,4,8,9-tetramethyl-1,6-diphosphabicyclo(4.4.0)deca-3,8 -diene; TMS, trimethylsilyl; Tol, toluene; tppme, 1,1,1-tris(diphenylphosphinomethyl)ethane1; TPPMS, sodium salt of *m*-monosulfonated triphenylphosphine; Tren^{TIPS}, N (CH₂CH₂NSii/Pr₃)₃; Triphos, 1,1,1-tris(diphenylphosphinomethyl)ethane; Ts^{Tol}, HC(SiMe₂NAr)₃; UV, ultraviolet light; Vis, visible light; vtms, Me₃SiCH=CH₂; WCA, weakly coordinating anions; XAS, X-ray absorption spectroscopy; XPS, X-ray photoelectron spectroscopy; XRD, X-ray diffraction analysis.



Scheme 1. Structures of the most common allotropic forms of elemental phosphorus. Focus of this work are white and black phosphorus.

considered, but this field is still in its infancy for the synthesis of organophosphorus species. Therefore, in the current research climate, the degradation of P₄ into P₁-P₄ fragments still constitutes the most important pathway towards the synthesis of organophosphorus compounds [2,11,18]. Also, its aggregation into P_n (P_n \geq 5) frameworks could provide further insights into the understanding of the structure and bonding in metal phosphides [19–21]. In this scenario, an eco-benign functionalization of P₄ emerges as a challenging task to afford the desired P-derivatives selectively. Accordingly, the development of such protocols, and comprehension of the mechanisms involved in the reactions [22] is an important goal in both academia and industry.

To achieve such a goal, one of the strategies has been to study the coordination of white phosphorus towards transition metals (TMs). By incorporating the P_4 molecule into the coordination sphere of a TM, an activated [M]-(P)_x species can be obtained, hence facilitating the transformation of the phosphorus motifs formed. This could lead to new reaction pathways to circumvent the use of hazardous/corrosive phosphorus halides in the synthesis of organophosphorus compounds. Alternatively, such activation could be a straightforward route to access polyphosphorus cages/ frameworks [19,23], obtaining building blocks for coordination polymers and supramolecular chemistry (*e.g.* the family of pentaphosphaferrocenes) [24], or synthesizing P-containing inorganic derivatives/phosphide materials [19,20].

When white phosphorus is heated or irradiated with UV light, the polymeric red allotrope is formed (Scheme 1) [12]. The precise structure of red phosphorus is unknown, although it is commonly thought to be an amorphous arrangement of different building units. By heating the amorphous red allotrope, other forms of elemental phosphorus can be obtained. These correspond to crystalline forms of red phosphorus, of which, for example, the structures of fibrous phosphorus and violet phosphorus are known. There are five proposed structural forms of red phosphorus, with commercially available amorphous red phosphorus being type I, fibrous phosphorus being type IV, and violet phosphorus being type V [12,25]. The molecular structures of types II and III have not been determined so far. Fibrous phosphorus, or type IV, and violet phosphorus (also known as Hittorf's phosphorus, type V) are assumed to be intermediates in the process to obtain black phosphorus from amorphous red phosphorus [26,27]. Both fibrous phosphorus and violet phosphorus share similar structural features, being composed of pentagonal P₈ and P₉ cages connected

by P_2 dumbbell-like linkers [28]. The main difference between them is the crosswise linkage of the strands through the P_9 units in violet phosphorus, or their parallel arrangement under formation of double tubes in fibrous phosphorus [12,28].

The third fundamental allotrope, black phosphorus (BP, Scheme 1) [29], is the most thermodynamically stable form of the element. In fact, from white to black, the phosphorus allotropes become progressively more stable [30]. BP can be obtained either by heating red phosphorus at low pressures in the presence of a suitable catalyst (e.g. SnI₄) [31,32] or, alternatively, by heating white phosphorus at elevated pressures (12000 bar). Like the red allotrope, BP is polymeric, although by contrast, it has an ordered layered structure, resembling that of graphite. Weak van der Waals interactions hold together the layers. As with graphite, a monolayer of BP can be obtained by mechanical or liquid phase exfoliation [33]. This exfoliated monolayer of BP is known as phosphorene, by analogy with graphene, and is a 2D material that exhibits atypical physical, chemical, and biomedical properties [34–37]. Phosphorene [38,39] has ignited a renaissance for black phosphorus in recent times [40], giving rise to enormous scientific interest.

2D black phosphorus is now a protagonist next to silicene and germanene in the post-graphene era. The possibility to access phosphorus-containing materials by reaction of exfoliated black phosphorus has consequently gained widespread attention. However, since phosphorene is highly reactive and unstable towards light, oxygen and water (exposure to air results in facile formation of phosphates) [41], scarce information regarding its interaction with transition metals was available, until recently. Attention has been directed towards the design of efficient strategies to perform protective modification of exfoliated black phosphorus, aimed at increasing its stability. Such modifications could enable the development of black phosphorus or phosphorene-based optoelectronic devices, among other interesting applications in materials science.

Throughout the history of chemical research, a wide array of information on the chemistry of elemental phosphorus towards transition metals has appeared in the literature. In previous decades, extensive studies have been summarized in several reviews [6,7,42,43], and the field is still very much in the academic zeitgeist today. Most of those reviews explicitly focused on the coordination chemistry of white phosphorus towards early [6] or late [7,44,45] transition metals (TMs). As well, photochemical [46,47] or electrochemical [48–52] pathways for the selective transformation of P_4 have also been developed recently. In addition, strategies for the functionalization of phosphorene have also been reviewed lately [38].

Nevertheless, these published reviews are either outdated, most of them having appeared before 2011, or they focus on other aspects of P₄ chemistry neglecting its coordination chemistry, which has continued to thrive. We felt that the rapid developments in the field necessitated an update covering the period 2011–2020. Prior contributions will not be covered in this work, although some are included when relevant for a particular discussion. Most of these exciting studies make use of a stoichiometric amount of metal, affording novel and oftentimes an unpredictable variety of phosphorus-containing structural motifs. For the moment, there are very few contributions which have achieved the long soughtafter catalytic functionalization of P₄ [53,54] yielding either aryl phosphines or aryl phosphates. It is also worth mentioning the emerging interest towards the coordination of white phosphorus by early transition metals, lanthanides and actinides, reactivity that previously was in the remit of late TMs. Another intriguing development is the study of P_n ligands, directly derived from P₄ and a suitable transition metal complex. It has been shown that they can be used as building blocks in the construction of either supramolecular assemblies or 1D and 2D coordination polymers

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giving access to fascinating hybrid organic/organometallic frame-works [55].

To help the reader navigate this voyage through phosphorus coordination chemistry, the review has been divided into sections. Section 2 has been dedicated to the coordination chemistry of white phosphorus towards early transition metals (groups 3–7), including examples with lanthanides and actinides, while Section 3 provides an overview of the contributions with late transition metals (groups 8–11). Section 4 showcases the emerging field of surface coordination of exfoliated black phosphorus, which transcends the classical coordination chemistry of phosphorus compounds.

2. Coordination chemistry of white phosphorus by early transition metals

2.1. Group 3 metals

2.1.1. Scandium

In the last decade, activation of white phosphorus with rare earth elements and actinides has seen important breakthroughs, although the field remains less developed with respect to late transition metals. Group 3 elements, whose coordination chemistry is dominated by M(III) metal complexes (d⁰), are amongst the hardest Lewis acidic metal cations. According to the HSAB theory, their matching with soft nucleophiles, such as phosphorus-containing ligands (phosphines or phosphides) is rather poor making the synthesis of rare-earth complexes bearing P-based ligands challenging [56]. However, their coordination chemistry with O-, N-, and Cbased ligands is vast and well developed. Nonetheless, in 2012 Diaconescu *et al.* have disclosed the first example of the activation of white phosphorus [57] using dinuclear scandium(III) arene complexes, $[(NN^{fc})Sc]_2(\mu_2,\eta^4:\eta^4-C_{10}H_8)$ (1), and $[(NN^{fc})Sc]_2(\mu_2,\eta^6:\eta^6-C_{14}H_{10})$ (2), $(NN^{fc} = 1,10$ -fc $(NSitBuMe_2)_2$, and fc = ferrocenylene). As shown in Scheme 2, 1 and 2 are synthesized from the same iodide precursor 3 by reduction with KC₈ in presence of naphthalene or anthracene respectively. They feature a binuclear inverted-sandwich motif where the middle-deck is occupied by the reduced arene moiety. It was also possible for the authors to convert 1 into 2 by heating at 50 °C in the presence of anthracene.

Due to being excellent two-electron reductants, **1** and **2** reacted with P₄ giving mixtures of two different polyphosphorus clusters **4** and **5**, having either a $[P_8]^{4-}$ or a $[P_7]^{3-}$ Zintl-type anion at the core and $[(NN^{fc})Sc]^+$ units at the edges (Scheme 3) [58].

The differing solubilities of **4** and **5** allowed for facile separation and their structures were unambiguously elucidated by X-ray crystallography, see Fig. 1.

Compound **4** represents the first example of Zintl-type anion, $[P_7]^{3-}$ obtained directly from the activation of P₄, without using an alkali metal or its equivalent as a reducing agent [12]. Both the starting complexes **1/2** and the products **4/5** displayed short Fe-Sc bond (2.80 Å), evidencing a weak Fe-Sc donor–acceptor interaction which is deemed to be essential for the stability of the complexes. ³¹P NMR analysis of **4** shows the presence of three different phosphorus atoms in 3:1:3 ratio, corresponding to three bridging atoms at the edges (P_{edge}), one phosphorus atom at the apex (P_{apex})



Scheme 2. Synthesis of the triple-decker complexes 1 and 2 from the common precursor 3.



Scheme 3. Reaction of 1 and 2 with P₄.



Fig. 1. X-ray crystal structure of a) 4, b) 5. Hydrogen atoms have been omitted for clarity; adapted from Ref. [58].

and a cyclo-P3 at the bottom (Pbottom) forming an AA'A"MM'M"X' spin system. This indicates that the bullvalene-type bond fluxionality, typically seen in alkaline earth polyphosphides such as Li₃P₇, is not present in **4** at room-temperature [59]. The bond distances between the phosphorus atoms are similar (P_{apex} - P_{edge} = 2 .201 Å, Pedge-Pbottom = 2.197 Å, Pbottom-Pbottom = 2.201 Å) and very close to the P-P bond distance found in P₄ (2.20 Å) [60]. In addition to this, compound 4 features relatively short Sc-P_{edge} distances (2.730 and 2.747 Å), which gives the complex a prevalently covalent character. It should be noted that the formation of either P₇ or P₈ clusters by reductive coupling of white phosphorus is quite rare in early transition metal chemistry [61]. Interestingly, when using the yttrium-arene complex $[(NN^{fc})Y(THF)]_2(\mu_2,\eta^4:\eta^4 C_{10}H_8$), only the Y_3P_7 species **6** is obtained. This compound shows a particularly shortened Pedge-Pbottom of 2.176 Å and longer Y-P distances (2.91 and 2.95 Å) with respect to Sc₃P₇, which can be addressed by the increased electropositivity of yttrium and therefore the complex has more ionic character. This P₄ activation protocol was extended also to analogous lanthanum and lutetium complexes such as $[(NN^{fc})La(THF)]_2(\mu_2,\eta^4:\eta^4-C_{10}H_8)$, and $[(NN^{fc})$ Lu(THF)]₂(μ_2 , η^4 : η^4 -C₁₀H₈), which reacted selectively with P₄ at room temperature affording the corresponding Zintl-type compounds 7 and 8 [62]. Fig. 2 shows the molecular structures of compounds 6, 7 and 8.

The related Zintl-type complexes **4**, **6**, **7** and **8** feature an inverse correlation between the ionic radius of the metal and the P_{edge} - P_{bottom} distance, and an increasing ionic character as the electronegativity of the metal decreases, a trend that culminates with La. The ³¹P NMR spectrum of La₃P₇(**7**) shows a single broad peak at room-temperature indicating the presence of the bullvalene-type tautomerization of the polyphosphide unit, which is not present in **4**, **6** and **8**.

By reacting **4** or **6** with Me₃Sil [58], the metal-free derivative $(Me_3Si)_3P_7$ was obtained and the resulting metal iodide byproduct was used to prepare the starting metal-arene complexes thus opening a new way towards the metal-mediated synthesis of similar phosphorus compounds directly from P_4 (Scheme 4).

2.1.2. Yttrium

As in the case for scandium, activation of P₄ by yttrium complexes has only been achieved recently. Beside **6** described in Section 2.1.1 an analogous complex characterized with a $[P_7]^{3-}$ Zintl anion core was obtained in a two steps process from elemental phosphorus by Zhou and co-workers [63]. The reaction of P₄ with two equivalents of the metal alkyl complex LYR(THF) [**9**, L = *N*,*N*⁻ di(2,6-diisopropylphenyl)-1,4-diazabutadiene, R = 2-*N*,*N*⁻ dimetylbenzyl] followed by addition of DMAP (4-*N*,*N*⁻ dimethylaminopyridine) proceeds via the migration of the benzyl moieties from the metal centers to two adjacent phosphorus atoms of P₄, affording the rare-earth complex (LY·DMAP)₂ [μ_2 , η^3 : η^3 -1,2- R_2 -*cyclo*-P₄] (**10**) (Scheme 5) where the *cyclo*-P₄ core bridges the two yttrium centers and assumes a rare twisted conformation (Fig. 3).

Interestingly, treatment with HMPA (hexamethylphosphoramide) promotes further migration of the benzylic residues affording $(LY \cdot HMPA)_{3}P_{7}$ [11, L = N,N'-di(2,6-diisopropylphenyl)-1, 4-diazabutadiene] with concurrent loss of the tribenzylphosphine 12 (Scheme 6, Fig. 4a). Such a Zintl-type complex is structurally comparable to the polyphosphides described above, except that in this case the Y atoms are bound to four P atoms with distances ranging from 3.002 to 3.317 Å. After prolonged heating of **11** in toluene, a P-P migration of the benzyl residue occurs, yielding the cluster $(LY \cdot DMAP)_2[\mu_2, \eta^2: \eta^3 - cyclo - P_3PR_2]$ (13, R = 0dimethylaminobenzyl), where the disubstituted phosphorus atom sits exo with respect to the remaining unsubstituted cyclo-P₃ moiety, see Fig. 4b. It is noteworthy to add that addition of benzylpotassium to 13 completes the extrusion of the exo-P atom through elimination of tribenzylphosphine. This leads to the formation of the ion-paired complex $(LY \cdot THF)_2 [(\mu_2, \eta^3: \eta^3 - cyclo - P_3)]$ K (14), which possesses an unsubstituted cyclo-P₃ moiety and, after



Fig. 2. Molecular structures of a) 6, b) 7 and c) 8. Hydrogen atoms have been omitted for clarity; adapted from Refs. [58] (6) and [62] (7 and 8).



Scheme 4. Synthesis of organophosphorus derivative mediated by ${\bf 4}$ and ${\bf 6}$ in the presence of Me₃Sil.



Scheme 5. Reaction of P₄ with 9.

addition of 18-crown-6, its ion-sequestered derivative [K(18crown-6)][LY(THF)₂(μ_2 , η^3 : η^3 -*cyclo*-P₃)] (**15**) is formed, Fig. 4c. Compound **14** could also be formed by direct addition of KR (R = N,N'-dimethyl-2-benzyl) to the *cyclo*-P₄ complex **10**, indicating that the nucleophilicity of KR might promote a non-thermal benzylic migration followed by elimination of tribenzylphosphine. As such, this process represents a remarkable 2-step [3 + 1]-fragmen tation process of P_{4_1} in which one phosphorus atom is also derivatized into an organophosphine.

2.1.3. Lanthanides and actinides

2.1.3.1. Lutetium. The formation of the cyclic $[P_3]^{3-}$ anion commonly occurs via white phosphorus activation in the presence of early or late transition metal complexes. As such, P₄ follows a [3 + 1]-fragmentation pathway which usually enables the isolation of either the P₃ or the P₁ moiety. One-pot, simultaneous trapping of both fragments is much rarer and is limited to few examples, mainly with late transition metals [64-66]. That was until Zhang and co-workers achieved this by employing early transition metal (ETM) complexes such as yttrium and lutetium metallacyclopentadienes 16, 17 and 18 (Scheme 7) [67]. These complexes react readily with 0.5 equivalents of P₄ in tetrahydrofuran at room temperature displaying dual dinucleophile/diene character. Similar to the known reactivity of 1,4-dilithium-1,3-butadienes with P₄ [68], the P₁ fragment is captured through the formation of phospholyl lithium 19, while the remaining P₃ moiety is involved in a Diels-Alder reaction with the diene ligand. This results in the exclusive formation of the exo-products 20, 21a and 21b (Scheme 7).

As shown in Fig. 5, the final products bear an unusual substituted bicyclo[4,1,0]triphosphaheptanide framework where the metal is *endo* with respect to the η^3 -P₃ ligand and retains the M–C bonds with the terminal carbon atoms of the C₄ fragment (**20**: Y-C1 = 2.409 Å, Y-C2 = 2.402, Y-P1 = 2.711 Å; **21a**: Lu-C1 = 2.367 Å, Lu-C2 = 2.353 Å, Lu-P1 = 2.664 Å; **21b**: Lu-C1 = 2.392, Lu-C4 = 2.375 Å, Lu-P1 = 2.639 Å).

Interestingly, the Lu-P1 bond in compounds **21a** and **21b** is markedly shorter than M–P bonds commonly found in known



Fig. 3. Solid state structure of 10. Hydrogen atoms have been omitted for clarity; adapted from Ref. [63].

lutetium phosphides and polyphosphides, while the Lu-P2 and Lu-P3 bond distances (**21a**: Lu-P2 = 2.976 Å, Lu-P3 = 2.992 Å; **21b**: Lu-P2 = 3.000 Å, Lu-P3 = 2.986 Å) are appreciably longer (Lu₃P₇, **8** displays average Lu-P bond = 2.893 Å). Additionally, the structural and spectroscopic features found for the *cyclo*-P₃ moiety in **20**, **21a** and **21b** are different compared to the known *cyclo*-P₃ complexes derived from the activation of white phosphorus with late-transition metals (LTM) such as Co [69], Rh, Ir [70], Ni [71], Pd and Pt [72]. In contrast with the mentioned ETM, in LTM com-



Scheme 6. Selective conversion of the cyclo-P₄ unit in 10 by alkyl migration.

plexes, the metal center resides opposite to the organic ligand occupying one vertex of the tetrahedron formed with the cyclo-P₃ fragment. In addition to this, in ETM complexes the M-P and P-P distances are averagely longer and the three P atoms are not equidistant. These features reflect in the ³¹P NMR spectra of the complexes, which usually show two different sets of signals for the cyclo-P₃ fragment, while in LTM complexes only one signal was observed, usually having ${}^{1}J_{(P-P)}$ coupling constant one order of magnitude smaller. DFT calculations have confirmed the existence of a weak interaction between the newly formed π system and the 5d orbitals of the lanthanide metal center. Reminiscent of the reactivity shown by the cyclo-P₃ unit in LTM towards M (CO)₅ where M = Cr, Mo, W, Re [73,74], Zhang et al. investigated the behavior of **20**, **21a** and **21b** towards {W(CO)₅(THF)}. The coordination of the tungsten carbonyl occurred exclusively at P1 with concurrent displacement of the Li(THF)₃ countercation, which indicates the prevailing nucleophilic character of P1 over P2 and P3. Nevertheless, coordination to P2 was also observed when employing two equivalents of $\{W(CO)_5(THF)\}$. Furthermore, the oxidation of 20 with *p*-benzoquinone was studied. In this process, the P_4 molecule is released along with the formation of the aromatic phospholyl lithium.

2.1.3.2. Samarium. Low-valent samarium species have been extensively employed in the reduction of transition-metal complexes bearing phosphides or polyphosphides moieties. The first direct activation of P₄ with a samarium(II) species, not involving other transition metals, was achieved in 2009 by Roesky and coworkers. Decamethylsamarocene, Cp*₂Sm(II), acting as a single electron donor, was shown to reductively couple two P₄ units affording the realgar-type ionic compound $[{Cp*_2Sm}_4P_8]$, the first isolated polyphosphide of the rare-earth elements [75]. To further explore how the steric demand of the reducing agent affects the activation of P₄, As₄ and As₄S₄, Roesky and coworkers employed the bulky samarium amidinate complex [(DippForm)₂Sm(THF)₂] $(22, \text{ DippForm} = \{(2,6-iPr_2C_6H_3)NCN(2,6-iPr_2C_6H_3)\}^{-}\}$. As shown in Scheme 8, the reaction of 22 with white phosphorus afforded red crystals of the dinuclear complex [{(DippForm)₂Sm}₂(μ_2 , η^4 : $\eta^{4}-P_{4}$] (23) in high yields (94%) [76].

The added steric hindrance of the amidinate ligands prevents the assembly of the $[P_8]^{4-}$ cage without affecting the efficiency of the reduction. Complex **23** features an inverse inorganic sandwich structure with a naked *cyclo*-P₄²⁻ moiety as the middle deck, representing the first example of an f-element to display such a structure, see Fig. 6. The *cyclo*-P₄²⁻ ligand is almost perfectly rectangular-shaped being characterized by regular P-P bonds (ranging from 2.1443 Å to 2.1618 Å) and internal P-P-P angles close to 90°. The Sm-P bonds range from 3.041 Å to 3.157 Å, which is comparable to known samarium phosphide complexes [75].

It should be mentioned that the $cyclo-P_4^{2-}$ moiety, being isolobal to the cyclobutadiene dianion $[C_4H_4]^{2-}$, consists of a 6π -aromatic system. Due to its high stability, the formation of $[cyclo-P_4]^{2-1}$ ligands occurs readily in the activation of P₄ by alkaline metals such as Cs [77], by ETM such as Zr [78], Nb [79], Ta [80] and LTM, either in monometallic complexes or by bimetallic cooperation [81,82]. In an attempt to further explore the ability of divalent samarium complexes to promote P-P bond formation, in 2013 Roesky and co-workers reacted at high temperatures the samarocenes $[Cp_2Sm(THF)_2]$ (24a) and $[\{C_5Me_4(n-propyl)_2\}Sm]$ **24b** with the well-known pentaphosphaferrocene $[Cp^*Fe(\eta^5-P_5)]$ (25a) [83]. This resulted in the formation of a 2 electron-2 center P-P bond between two $[Cp^*Fe(\eta^5-P_5)]$ units through a double single-electron reduction process mediated by Sm(II), which afforded the dimeric core $[{Cp^*Fe(\eta^4-P_5)}_2]^{2-}$. The latter is stabilized by two $[(C_5Me_4R_2Sm]^+$ (R = Me, *n*-Pr) cations forming the products 26a and 26b (Scheme 9) which constitutes the first exam-



Fig. 4. Molecular structures of a) 11, b) 13, c) 15. Hydrogen atoms have been omitted for clarity; adapted from Ref. [63].



Scheme 7. Activation of P_4 with **16**, **17** and **18** by a formal [3 + 1] fragmentation process.

ple of a mixed 3d/4f polyphosphide complexes bearing a rare P_{10} bridging moiety (Scheme 9).

As shown by the molecular structure in Fig. 7, after reduction the two *cyclo*-P₅ moieties assume an envelope type conformation whereby the two off-plane P atoms are involved in a newly formed single P-P bond (P–P bond length is 2.209 Å in **26a** and 2.199 Å in **26b**). The P-P distances in the *cyclo*-P₅ ring are in the range 2.137– 2.186 Å in **26a** and 2.114–2.189 Å in **26b**, suggesting partial double bond character. Considering that each iron atom is bound to the *cyclo*-P₅ in an η^4 fashion, the latter can be formally considered as a tetraphosphabutadienediide ligand formally donating six electrons.

As the Sm atom in **26a** and **26b** is in the III oxidation state, as confirmed by characteristic NIR absorption, the polyphosphide unit is one of the very few examples of a $[P_{10}]^{4-}$ anion (see Section 3.1.1). It should be mentioned that a neutral P_{10} ligand was first found in the following complexes $[(Cp''M)_4P_{10}]$ (M = Rh,



Fig. 5. Solid state structure of a) 20, b) 21a. Hydrogen atoms are omitted for clarity; adapted from Ref. [67].



Scheme 8. Activation of P₄ with 22.



Fig. 6. Solid state structure of 23. Hydrogen atoms and 2,6-diisopropylphenyl groups are omitted for clarity; adapted from Ref. [76].

 $Cp'' = 1,3-tBu_2C_5H_3$, M = Co, $Cp'' = 1,3-(SiMe_3)_2C_5H_3$), synthesized by Scherer in 1992 [84].

2.1.3.3. Ytterbium. The activation of P₄ with rare earth metals has been investigated by A. Trifonov *et al.* by using the reduced-arene Yb(II) complex, $[Cp^{Bn5}Yb(DME)_2(\mu_2,\eta^4:\eta^4-C_{10}H_8)]$ (**27**, $Cp^{Bn5} = pen$ tabenzylcyclopentadienyl), which can be conveniently synthetized by salt metathesis using $[YbI(DME)_2]_2(\mu_2,\eta^4:\eta^4-C_{10}H_8)$ and $Cp^{Bn5}K$ [85]. Compound **27** shares similarities with other reduced-arene complexes of rare earth metals. It features an inverted sandwich motif whereby the twisted dianionic naphthalene ligand occupies the middle deck and bridges two metal centers, each one coordinated in an η^4 mode. It should be noted that the two ytterbium atoms appear in a divalent oxidation state, as confirmed by ¹H and ¹³C{¹H} NMR analysis allowing for two potential reductive sites in addition to $(C_{10}H_8)^{2-}$. The reaction of **27** with P₄ affords the trinuclear Yb(II) complex $[Cp^{Bn5}Yb(DME)]_3P_7$ (**28**, Scheme 10) [86].

The molecular structure of **28** was elucidated by XRD analysis (Fig. 8). Similar to other M_3P_7 complexes (M = Sc, Y, La, Lu) described by Diaconescu [58,62] and Zhang [67], see Sections 2.1.1, 2.1.2 and 2.1.3, **28** bears a central Zintl-type $[P_7]^{3-}$ cage ligand which coordinates three Yb(II) atoms at the edges of the middle rim in an $\mu,\kappa^2:\kappa^2:\kappa^2$ mode.

The average Yb-P bond length (2.93 Å) is comparable to analogous yttrium and lutetium $[P_7]^{3-}$ polyphosphides (respectively 2.93 Å [58] and 2.89 Å) [62]. ¹H NMR spectroscopy showed the expected set of signals related to the organic ligands, thus confirming the diamagnetism of **28** and the divalent state of Yb ions. The ³¹P{¹H} NMR spectrum collected either at room temperature and at -60 °C, displayed only a broad singlet at δ -174.2 ppm. This is possibly due to the presence of a fast bullvalene-type tautomerization typically associated to more ionic $[P_7]^{3-}$ polyphosphides.

2.1.3.4. Thorium. Following the same strategy used for the reductive activation of white phosphorus with low-valent rare earth metals, in 2016 Mills and co-workers achieved the unprecedented Th(III)-mediated activation and complexation of P₄ [87]. The reaction of complex [Th(Cp'')₃] **(29**, Cp'' = C₅H₃(SiMe₃)₂–1,3) with P₄ affords the binuclear derivative [{Th(Cp'')₃}₂ ($\mu,\kappa^1:\kappa^1-P_4$)] **(30**) featuring the reduced (*cyclo*-P₄)^{2–} ligand in an unique coordination mode (Scheme 11). However, the reaction proved to be rather slow (72 h) with low yields (19%).

However, it is noteworthy to add that the analogous U complex $[U(Cp'')_3]$ did not show any reactivity with elemental phosphorus even in forcing conditions (reflux, 16 h). This was attributed to the greater electropositivity of Th(III) compared to U(III). The ³¹P {¹H} NMR analysis of **30** showed two low field signals at δ = 227.59 ppm (t, ${}^1\!J_{\rm (P-P)}\approx$ 400 Hz) and δ = 328.86 ppm (t, ${}^1\!J_{\rm (P-}$ $_{P)} \approx 400$ Hz) and two unresolved broad peaks at δ = -246.55 (br m, $\Delta v \approx 2000$ Hz) and 10.35 ppm (br m, $\Delta v \approx 2000$ Hz). Meanwhile the low field resonances correlate well with the ionic compound $[Cs_2P_4]^{2-}$ [77], the broad signals could not be confidently assigned. The peculiar hapticity of the cyclo-P₄ moiety, attributed to the steric demand of the bulky Cp^{''} coordinated to the metal, predominantly involves the sp²-hybridized lone pairs on phosphorus. This is confirmed by XRD analysis of **30** (Fig. 9), which shows P-P distances (average 2.051(9) Å) almost identical to a P=P double bond (2.04 Å) [88], whereas coordination in an η^4 fashion usually displays longer P-P bonds (2.16–2.14 Å) with greater σ -character (see for instance Section 2.1.3.2).

It is also worth mentioning that the calculated $\delta(P,P)$ delocalization index, which can be regarded as a measure of the aromaticity of the system, was found to be 1.33, close to the formal value of



Scheme 9. Complexes 24a and 24b react with 25a yielding the polyphosphides 26a and 26b.



Fig. 7. Solid state structure of 26a. Hydrogen atoms are omitted for clarity; adapted from Ref. [83].

1.50 expected for a 6π -aromatic systems. Thus, the $[cyclo-P_4]^{2-}$ moiety in **30** can be seen as a largely unperturbed, planar aromatic 6π -electron moiety. Finally, the elongation of the Th-Cp_{centroid} distances going from **29** to **30** (respectively 2.519 and 2.543 Å) is in line with the formal Th(IV) assignment of **30** [89]. The Th-P distances (2.919 Å) are similar to those observed in known thorium phosphides also derived from the reaction with P₄ [90].

2.1.3.5. Uranium. Although there are only a few examples of the activation of P₄ with uranium containing complexes that have been reported in the last decade, the variety of coordination patterns in these scant examples is quite remarkable. In 2011, Green and coworkers investigated the reactivity of P₄ with the U(III) species $[U(\eta^5-C_5Me_5)\{\eta^8-C_8H_6(SiiPr_3)_2-1,4\}_2(THF)]$ (**31**, Scheme 12) [91]. This compound, upon U(III)-promoted two-fold reduction of P₄, yields the dimeric complex **32** $[U(\eta^5-C_5Me_5)\{\eta^8-C_8H_6(SiiPr_3)_2-1,4\}]_2(\mu_2,\eta^2:\eta^2-P_4)$ as black crystals, in which the bridging $[P_4]^{2-}$ unit is tilted with respect to the U-U axis as clearly shown in Fig. 10. Nonetheless, the P-P distances (2.148–2.152 Å) are almost identical and are in line with other dianionic $[P_4]^{2-}$ ligands (*vide supra*). A ³¹P{¹H} NMR study revealed a signal at δ + 718 ppm, suggesting **32** is paramagnetic. Similarly, the ³¹P{¹H} resonance of the $[P_4]^{2-}$ complex $[U\{N(tBu)Ar\}_3]_2(\mu 2, \eta^4: \eta^4-P_4)$ was found to be +794 ppm [92].

DFT calculations were in close accordance with the observed hapticity of $[P_4]^{2-}$ in **32**, despite the absence of bulky substituents on the adopted molecular model $[U(\eta^5-C_5H_5)(\eta^8-C_8H_8)]_2(P_4)$. As suggested by the MO analysis, the singular $\mu_2, \eta^2: \eta^2$ binding mode of P_4 results from a combination of both steric (wedge-shaped nature of the UCotCp fragment) and electronic requirements (the tilt-



Fig. 8. Molecular structure of 28. Hydrogen atoms and benzyl groups on the cyclopentadienyl ligands are omitted for clarity; adapted from Ref. [86].

ing of P₄ respect to U-U vector, allows both σ and π orbitals of P₄ being involved in U-P bonding interactions).

U(III) complexes have also been involved in the formation of higher phosphides by catenation of P₄. By reacting the triamide complex [U(Tren^{TIPS})] [**33**, Tren^{TIPS} = N(CH₂CH₂NSiiPr₃)₃] with P₄, Liddle and co-workers [93] isolated the corresponding *cyclo*-P₅ dinuclear inverted-sandwich complex [{U(Tren^{TIPS})}₂(μ_2 , η^5 : η^5 -*cyclo*-P₅)] (**34**, Scheme 13 and Fig. 11 for X-ray structure).

Compound 34 emerges as one of the few actinide complexes bearing a cyclo-P₅ fragment together with the two samarium complexes [Cp*Fe(cyclo-P₅)Sm(DIP₂pyr)]₂ and [Cp*Fe(cyclo-P₅)Sm (DIP₂pyr)-(THF)₂] (DIP₂pyr = 2,5-bis{N-(2,6-diisopropylphenyl)imi nomethyl}pyrrolyl) [94], even though in the latter complexes it assumes an envelope-like conformation. It should be mentioned that the cyclo-P₅ phosphide, isolobal with the cyclopentadienyl ring, is a formal mono-anionic ligand usually found in association with electron rich d-metals [7], with whom it shares σ or π bonding interactions. Nonetheless, UV-Vis and SOUID magnetometry measurements agreed with the presence of U(IV) in 34. Accordingly, both uranium atoms display a $5f^2$ configuration, therefore suggesting a twofold reduction of the cyclo-P₅ moiety, making the ligand dianionic. Further investigations of the electronic structure of the complex through DFT calculations have revealed that the cyclo-P₅ ligand in **34** could be approximated to a dianion bridging the two metal centers via two polarized δ -bond, which are formed by the donation of uranium δ -symmetry 5f orbitals to δ symmetry frontier molecular orbitals of the cyclo-P₅.

Liddle and co-workers have also investigated the behavior of reduced-arene uranium complexes towards elemental phosphorus, a strategy that has proved successful for group 3 and 4f-block metal complexes [95]. When treating the diuranium(V)-arenetetraanion complex [{U(Ts^{Tol})}₂(μ_2 , η^6 : η^6 -C₆H₅CH₃)] (**35**, Ts^{Tol} = HC(SiMe₂NAr)₃; Ar = 4-MeC₆H₄) with 1.1 equivalents of P₄, the corresponding Zintl-type complex [{U(Ts^{Tol})}₃(μ,κ^2 : κ^2 : κ^2 -P₇)] (**36**) was obtained, see Scheme 14.



Scheme 10. Activation of P₄ by complex 27 yielding the trinuclear phosphide 28.



Scheme 11. Reaction of P₄ with complex 29.



Fig. 9. Solid state structure of 30. Hydrogen atoms are omitted for clarity; adapted from Ref. [87].

It should be mentioned that the precursor **35**, is best described as two U(III) centers bridged by a dianionic toluene moiety [96]. The magnetic moment of **36**, measured by Evans method in C_6D_6 was found to be 4.67 μ_B at 25 °C, which is a value of 2.42 μ_B per uranium atom. This is in good agreement with other U(IV) complexes [97], thus confirming a multi-electron redox process occurring between the precursor $\mathbf{35}$ and P_4 . In Fig. 12 it is shown that the molecular structure of **36** features a central $[P_7]^{3-}$ trianion bridging three [U(Ts^{Tol})]⁺ fragments, where each U center is coordinated to two P atoms on the upper rim of the Zintl-type scaffold. This represents the first example of fragmentation and catenation of P₄ promoted by uranium. U-P bond lengths in 36 are in the range of 2.948-3.031 Å, 0.1-0.2 Å longer than the sum of the covalent radii of uranium and phosphorus (2.81 Å) [58], possibly due to the steric hindrance of the $[U(Ts^{Tol})]^+$ unit and the bridging coordination mode.

A common parameter used to quantify the ionic character of heptaphosphanortricyclane cages is the Q value [98], which for ionic systems is usually in the range 1.3–1.4. XRD data allowed to determine the Q value for **36** (1.39), indicating the prevailing ionic character of the uranium cluster. As observed for other Zintl-type complexes of ETM (see Scheme 4, Section 2.1.1), the $|P_7|^{3-}$ core can be transferred to a large array of electrophiles, such

as metal salts (LiCl), sp³-C electrophiles (MeI), silyl chlorides (Me₃-SiCl) and sp²-C electrophiles (ArI), yielding the corresponding P_7R_3 substrate (R = Li, Me, Me₃Si, Ar) and the uranium halide complex.

2.2. Group 4 metals

2.2.1. Titanium

There are few examples of the titanium mediated activation of white phosphorus [6]. Nonetheless, it should be mentioned that TiCl₄ has been employed as a Lewis acid for the direct arylation of P₄ using ArCl, given a mixture of PClAr₂ and PCl₂Ar but not the desired PAr₃ [99]. In 2010, Cummins and co-workers [100] developed a protocol to synthesize aryl and alkyl tertiary phosphines, PR₃, from P₄ exploiting the known ability of P-P bonds to scavenge C-centered radical species. As showed by previous investigations, carbon radicals can be efficiently produced using the tris-amide Ti(III) complex $[Ti(N[tBu]Ar)_3]$ (**37**, Ar = 3,5-C₆H₃Me₂, Scheme 15), which is a potent one-electron reducing agent and halideabstractor towards both C-sp² and C-sp³ halides [101,102]. However, compound **37** does not react with P₄ directly, lowering the possibility of unwanted side reactions to occur. Reacting excess amounts of PhBr and $[Ti(N[tBu]Ar)_3]$ in presence of P₄ gives the corresponding PR₃ in good yields (72% for X = Cl and R = Ph). Interestingly, the intermediate P₂Ph₄ is selectively obtained (80% yield) by using sub-stoichiometric amounts of halide and Ti(III) complex, allowing the synthesis of mixed phosphines, PR₂R'. It should be added that, by increasing the steric demand on the organic halide, Cummins and co-workers successfully expanded the reaction scope to polyphosphorus substrates. Addition of MesBr resulted in the formation of P₃Mes₃ (61% yield), whereas when using DmpI the butterfly-type bicyclophosphine cis,trans-DmpP₄Dmp was obtained in 78% isolated yield (Mes: 2,4,6-Me₃-C₆H₂, Dmp: 2,6- $Mes_2-C_6H_3$).

2.2.2. Zirconium

In 1988, Scherer and co-workers [103] found that zirconocene, $[Cp''_2Zr(CO)_2]$ (**38**, $Cp'' = \eta^5-C_5H_3tBu_2-1,3$) is able to insert into one P-P edge of the P₄ tetrahedron forming the side-on coordinated adduct $[Cp''_2Zr(\kappa^2-P_4)]$ (**39**, Scheme 16). It should be mentioned that the analogous compound $[Cp^*_2Zr(\kappa^2-P_4)]$ was only detected in solution, failing any attempt of isolation indicating the need



Scheme 12. Activation of P₄ with complex 31.



Fig. 10. Solid state structure of 32. Hydrogen atoms are omitted for clarity; adapted from Ref. [92].



Scheme 13. Activation of P₄ by [U(Tren^{TIPS})] 33.



Fig. 11. Solid state structure of **34**. Hydrogen atoms are omitted for clarity; adapted from Ref. [93].



Fig. 12. Molecular structure of 36. Hydrogen atoms are omitted for clarity; adapted from Ref. [95].

for bulkier ligands. The central tetraphosphabicyclo[1.1.0]butane motif in **39** displays P-P distances ranging from 2.178 to 2.239 Å, with the bond between the two backbone P atoms being slightly shorter (2.178 Å). As shown in Scheme 16, treating **39** with tBuC \equiv P [104] afforded complex [Cp''_2Zr(κ^2 -P₃CtBu)] (**40**) as the major product among other compounds, such as [Cp''_2Zr{ κ^2 -P₆(CtBu)_2]] (**41**) displaying a ZrP₆C₂ cuneane-type framework, closely resembling the Hittorf's phosphorus allotrope [105], see Fig. 13, and the Zr-free substrate **42**.

Even though the mechanism of formation of **40** is not fully elucidated, it formally arises from the displacement of a P₂ unit in **39** with a molecule of $tBuC\equiv P$. In turn, **41** can be viewed as the product of double addition of $tBuC\equiv P$ to **39**. Similarly, the polyphosphorus compound **42** can be considered as a $tBuC\equiv P$ tetramer which is extended by a P₂ unit. DFT calculations showed that steric repulsion between Cp'' and $tBuC\equiv P$ plays a dominant role in directing the reaction towards the formation of the preferred regioisomer **40**, where the bulky *tBuC* fragment sits on the bridgehead position and points away from the Zr center.

DFT calculations [106] carried out on **39** show that the HOMOs are largely located on the P_4 fragment. Additionally, the P atoms directly attached to Zr are more negatively charged (-0.20) with respect to the bridgehead P atoms (+0.11) and will preferentially coordinate to Lewis acids. Steric repulsion between the LA and the hindered zirconocene moiety would in principle direct the coordination towards the bridgehead P atoms, as observed above. When [W(CO)₅(THF)] and **39** are mixed in equimolar amounts,



Scheme 14. Activation of P₄ by diuranium complex 35.



Scheme 15. [Ti(N[tBu]Ar)₃] acts as a halide abstractor through a single electron reduction process. This allows the formation of C-radical species in-situ which react with P₄ to afford a range of phosphines or phosphides.

the $[W(CO)_5]$ fragment coordinates exclusively to the Zr-attached P atoms yielding compound $[Cp''_2Zr(\mu_2,\kappa^2:\kappa^1-P_4)\{W(CO)_5\}]$ (**43**, see Scheme 17). Compound **43** was subsequently calculated to be 6.11 kJ/mol more stable than its bridgehead-P coordinated counterpart. Interestingly, the second equivalent of $[W(CO)_5(THF)]$ still coordinates to the more electronegative P atom giving a mixture of the mono **43** and twofold coordinated adduct $[Cp''_2Zr(\mu_3-\kappa^2:\kappa^1:\kappa^1-P_4)\{W(CO)_5\}_2]$ (**44**), in 1:4 ratio respectively (see Scheme 17 and Fig. 14 for solid-state structures of **43** and **44**). The three-fold coordinated complex $[Cp''_2Zr(\mu_4-\kappa^2:\kappa^1:\kappa^1:\kappa^1-P_4)\{W(CO)_5\}_3]$ (**45**) was formed as the major product by using a large excess of LA (8 eq). It should be mentioned that these conditions did not allow for the formation of the fourfold coordinated adduct but instead the compound, $[(CO)_4W(\mu_5-\eta^4:\kappa^1:\kappa^1:\kappa^1\cdotP_4)\{W(CO)_5\}_4]$ (**46**), arising from the detachment of the zirconocene moiety, was detected.

In contrast, the more sterically demanding $[CpMn(CO)_2(THF)]$ is expected to coordinate predominantly on the bridgehead P, which was calculated to be 9.08 kJ/mol more stable than the species coordinated to the metal-bonded P atom. This was confirmed experimentally, as one equivalent of $[CpMn(CO)_2(THF)]$ coordinates exclusively on the bridgehead P atoms giving complexes **47** (see Fig. 15 for X-ray structure) and **48**. Addition of two equivalents of $[CpMn(CO)_2(THF)]$ gives the two mono coordinated complexes (P-backbone **47** and P-apical **49**) and the twofold coordinated complex **48** in 10:0.5:1 ratio as indicated by ${}^{31}P{}^{1}H{}$ NMR (Scheme 18).

In contrast with [W(CO)₅], the addition of one equivalent of [Fe₂(CO)₉] gives both regioisomers of [Cp''_2Zr(μ 2, κ ²: κ ¹-P₄)(Fe (CO)₄] in almost an equimolar ratio (1:1.3) as observed by ³¹P NMR analysis (compound **50** and **51**, Scheme 19). Only **51** could be isolated after flash column chromatography and its solid-state structure is shown in Fig. 16.

After having evaluated the coordination behavior of **39** towards highly electrophilic transition metals fragments, Lewis acidic main group compounds, such as AlMe₃ and AlEt₃, were investigated (Scheme 20). In the case of AlMe₃, the reaction exclusively afforded the complex $[Cp''_2Zr(\mu_3-\kappa^2:\kappa^1:\kappa^1-P_4){AlMe_3}_2]$ (**52**), which bears two AlMe₃ molecules both bound to a P-coordinated Zr, regardless of the stoichiometry (1:1, 1:2, 1:5). The reaction employing the bulkier AlEt₃ gave the monocoordinated derivative $[Cp''_2Zr(\mu 2,\kappa^2:\kappa^1-P_4)]$ (AlEt₃] (**53**), where the LA is again attached to Zr bound P. The Al-P bond (2.577 Å) is shorter than in **52** (Al-P: 2.596 Å) (see Fig.17), which are both longer compared to a covalent Al-P bond (2.37 Å) indicating high dative bond character [88].

A tripodal aryloxide-coordinated zirconium hydride complex has been employed by Kawaguchi *et al.* to activate P₄ among other small molecules [107]. The reaction of complex [(O₃C)Zr(THF)₃] [**54**, (O₃C)³⁻ = tris(2-oxy-3,5-di-*tert*-butylphenyl)methane] with NaBHEt₃ leads to the formation of the hydride dimer [{Na (DME)}₂{(O₃C)Zr}₂(μ ,H)₃] (**55**), see Scheme 21. Reacting **55** with P₄ in toluene leads to the release of one equivalent of PH₃. The remaining *cyclo*-P₃ fragment replaces the bridging hydrides affording the complex [(Na(THF)₅]-[{Na(THF)₂}₂((O₃C)Zr}₂(μ , η ³: η ³-*cyclo*-P₃)] (**56**), after crystallization from a THF solution (Scheme 21). Complex 56 is a contact ion-pair, where the countercation [Na(THF)₅]⁺ balances the anionic complex [{Na(THF)₂}₂{(O₃-C)Zr}₂(μ , η ³: η ³-*cyclo*-P₃)]⁻. Complex **56** features an inverse sandwich structure, where the two Zr centers are bridged by a trihapto *cyclo*-P₃ unit, to which a formal –3 charge is assigned.

An XRD analysis of **56** (Fig. 18) shows that each Zr center is hepta-coordinated with a distorted mono-capped octahedral geometry, with the bridgehead C atom of the $[O_3C]^{3-}$ ligand occupying the capped position. The *cyclo*-P₃ moiety displays P-P single bonds of 2.226 Å and coordinates the two staggered $[(O_3C)Zr]$ units in an η^3 mode. These features are well representative of known late transition metals complexes $M_2(\mu_2,\eta^3:\eta^3-cyclo-P_3)$ with M = Ni, Co, Pd, Pt [69–74].

2.3. Group 5 metals

2.3.1. Vanadium

Low-valent vanadium species can be considered as powerful reducing agents due to their ability to undergo multiple electron processes, and therefore is an excellent candidate for the activation of white phosphorus. However, the reactivity of known V(II) complexes is often impaired by the saturation of the coordination sphere of the metal. A strategy to increase stability without imped-



Scheme 16. Synthesis of 39 and its subsequent reactivity with the phosphalkyne.



Fig. 13. Molecular structure of **41**. Hydrogen atoms are omitted for clarity; adapted from Ref. [104].

ing reactivity is to mask its coordination sites with an arene [108,109]. In 2010, Mindiola and co-workers reduced the previously synthesized $[(nacnac)VCl(Ntolyl_2)]$ (57) [110] with KC₈ or 0.5% Na/Hg in benzene, successfully affording the arene masked V(II) complex [(nacnac)V(Ntolyl₂)] (58), see Scheme 22. XRD analysis of **58** showed unambiguously the elimination of the chloride and the presence of an η^3 -coordination of Ntolyl₂ towards V, which is responsible of the stabilization of the highly reactive V (II) center. It should be noted that nacnac ligands generally are not effective in stabilizing electron-rich metals due to their tendency to undergo two-electron cleavage. Compound 58 is able to activate P₄ in a three electron reduction giving the diamagnetic vanadium(V) complex [(nacnac)V(η^3 -cyclo-P₃)(Ntolyl₂)] (59), the first example of a vanadium complex bearing a [cyclo-P₃]³⁻ ligand [111], whose solid state structure is shown in Fig. 19. The twosteps process (reduction of 57 and treatment with P₄) can be con-



Scheme 17. Compound 39 is reacted with 1, 2 and 8 eq of [W(CO)₅(THF)] affording prevalently P-edge coordinated products.



Fig. 14. Molecular structure of a) 43 and b) 44. Hydrogen atoms are omitted for clarity; adapted from Ref. [106].



Fig. 15. Molecular structure of **47**. Hydrogen atoms are omitted for clarity; adapted from Ref. [106].

veniently performed in a one pot reaction as shown in Scheme 22. When performed with a reduced quantity of P₄ (0.25 eq of P₄), the reaction affords the dinuclear complex [(nacnac)V(Ntolyl₂)]₂(μ 2, η^2 : η^3 -*cyclo*-P₃) (**60**) in 33% yield. The latter could be obtained in quantitative yields by reacting the mono-nuclear *cyclo*-P₃ complex **59** with one equivalent of **58**. Similarly, reduction of the starting aryloxy-coordinated complex [(nacnac)VCl(OAr)] (**61**, \neg OAr = 0 {2,6-*i*Pr₂C₆H₃}) with 0.5% Na/Hg in presence of P₄ gave complex [(-nacnac)V(η^3 -*cyclo*-P₃)(OAr)] (**62**). This could also be synthesised by heating the dinitrogen complex, **63** to 90 °C for ten minutes in presence of 2 equivalents of P₄ (Scheme 22) [112].

The molecular structure of 59 (Fig. 19a) features V-P bond lengths between 2.443 and 2.452 Å and an average intermolecular P-P distance of 2.15 Å, which is in the range of known [cyclo-P₃]³⁻ complexes. The cyclo-P₃ moiety in complex 62 has similar bond lengths. The vanadium center in 59 assumes a pseudo tetrahedral geometry having three N-atom donors and the cyclo-P₃ occupying the fourth position. The solid-state structure of **60** (Fig. 19b) displays a triple-decker geometry where the two V atoms occupy the two vertex of the trigonal bipiramidal V_2P_3 core. Consistent with the $\mu_2, \eta^2: \eta^3$ binding mode, one V atom is η^3 coordinated to the cyclo-P₃ ring, having V-P distances in the range 2.472–2.50 2 Å, while the other V atom displays a remarkably elongated V-P bond (2.928 Å), in accordance with an η^2 binding. The $\mu_2, \eta^2: \eta^3$ coordination mode is uncommon among *cyclo*-P₃ complexes; tBu₂)₂], prepared by Scherer in 1991 [90] and a few d⁸ metal complexes, such as [(triphos)Ni(cyclo-P₃)Pt(PPh₃)₂]⁺ [113], and [Au{M $(tppme)(cyclo-P_3)_{2}^{\dagger}$ (M = Co, Rh, Ir) [114]. Interestingly, in the case of **60**, the $\mu_2,\eta^2:\eta^3$ binding mode seems to average out to a $\mu_2, \eta^3: \eta^3$ -cyclo-P₃ coordination mode in solution, as the ¹H NMR spectrum did not show any inequivalence between the vanadium



Scheme 18. [CpMn(CO)₂(THF)] reacting with 39 preferentially at the P-backbone.



Scheme 19. Reaction of 39 with [Fe₂(CO)₉] affords respectively the P-edge and P-backbone complexes 50 and 51.



Fig. 16. Molecular structure of **51**. Hydrogen atoms are omitted for clarity; adapted from Ref. [106].

bound ligands. Compound **60** is paramagnetic as confirmed by Evans method (S = ½ at room temperature with μ_{eff} = 1.80 μ_B) and EPR gave an anisotropic *g* value \approx 2. Unexpectedly, the solution state ³¹P NMR spectrum of **59** and **62** showed significantly downfield signals corresponding to the P₃ moiety, with very broad resonances at δ = 85.0 and 125.0 ppm respectively, whereas the *cyclo*-P₃ motif usually displays signals within the range of –170 ppm and –220 ppm (see Section 2.1.3). This behavior was rationalized through magic-angle spinning (MAS) analysis and a detailed DFT study. It was noted that a particularly effective d(V)-p(*cyclo*-P₃) overlap in an applied external magnetic field generates a deviation to the metal-to-ligand density currents which are ultimately responsible of the deshielding phenomena.

2.3.2. Niobium

Although the selective and atom-efficient activation/functionalisation of white phosphorus remains elusive, recently great efforts have also been invested in the post-activation manipulations of polyphosphides-containing ETM complexes. For example, Cummins *et al.* reduced the niobium complex[$Cl_2Nb(ODipp)_3$] (**64**) in the presence of white phosphorus [115] to yield $[Na(THF)_3][(\eta^3 - P_3)Nb(ODipp)_3]$ (**65**) (Dipp = 2,6-diisopropylphenyl). They then demonstrated that **65** acts as a P₃ reservoir whereby the synthon $[cyclo-P_3]^-$ can be transferred to an organic fragment as well as to a different metallic complexes [116]. This can be achieved by reacting the nucleophilic complex **65** with SnClPh₃ (or other chlorides such as SiMe₃Cl, SiPh₃Cl and Ph₃CCl) affording the novel neutral adduct $[(ODipp)_3Nb(\mu_2,\eta^2:\eta^1-Ph_3SnP_3)]$ (**66**) via salt metathesis. In turn, the reaction of **66** with pyridine N-oxide liberates the reactive triphosphirene intermediate Ph₃SnP₃ **67**, which can be trapped with 1,3-cyclohexadiene (CHD) to form the corresponding Diels-Alder (D-A) adduct [Ph₃SnP₃(C₆H₈)] (**68**, Scheme 23, see Fig. 20**a** for solid-state structure of **68**).

As usually observed in similar cycloadditions involving P=P moieties, the endo product is exclusively formed due to decisive secondary orbital interactions taking place. The authors were able to observe via ³¹P NMR spectroscopy the formation of the C₅H₅NOcoordinated complex at -10 °C prior to the release of Ph₃SnP₃. A detailed DFT study carried out by Morello and co-workers further elucidated the mechanism of this transformation [117]. The authors demonstrated that the reaction proceeds through a redox-neutral Bayer-Villiger oxygen insertion into the Nb-P bond, is the rate-determining step of the reaction which $(+23.7 \text{ kcal mol}^{-1})$. Subsequently, the formation of the strong Nb=O bond leads to the concomitant formation of the proposed $(\eta^1-P_3)Ph_3Sn$ intermediate (221267.7 kcal mol⁻¹), which then reacts in situ with CHD. The energy barrier associated with the D-A reaction is calculated to be as low as +4.8 kcal mol⁻¹. The isolable compound **68** can be considered a $[cvclo-P_3]^-$ carrier featuring a "protected" P=P bond as well as a labile Sn-P bond and can be used to deliver the cyclo-P₃ unit to a suitable acceptor. The reaction of **68** with the Wilkinson's catalyst [ClRh(PPh₃)₃], formed the corresponding $[(\eta^3 - cyclo - P_3)Rh(PPh_3)_3]$ (69) (Scheme 23), featuring a symmetric cyclo-P₃ moiety with a resonance at δ = -191 ppm and an octahedral geometry for the Rh(III) center (Fig. 20b), similar to the previously reported $[(\eta^3 - cyclo - P_3)Rh(triphos)]$ $(triphos = CH_3C(CH_2PPh_2)_3)$ [118].

More recently, Scheer *et al.* demonstrated that the anionic complex **65** can undergo ring expansion if treated with a suitable electrophilic P-atom donor, to access [*cyclo*-P₄]^{3–} complexes [119]. The reaction of **65** with the phosphinidene complex [Cp*P{W(CO)₅}₂]



Scheme 20. Coordination behavior of 39 towards AlMe₃ and AlEt₃.



Scheme 21. Reduction of 54 with NaBHEt₃ affords dinuclear tri-hydride complex 55 which in turn reacts with P₄ forming the *cyclo*-P₃ complex 56 with concomitant loss of PH₃.



Fig. 17. Molecular structure of a) 52 and b) 53. Hydrogen atoms are omitted for clarity; adapted from Ref. [106].



Fig. 18. Molecular structure of **56**. Hydrogen atoms and $[Na(THF)_5]^*$ are omitted for clarity; adapted from Ref. [107].

 $(Cp^* = C_5Me_5)$ at -78 °C in presence of 18-crown-6 afforded [Na (18-crown-6)(THF)₂][{W(CO)₅}₂{ μ_3 , η^3 : κ^1 : κ^1 -P₄Cp*}Nb(ODipp)₃] (**70**) in excellent yield (Scheme 24).

As elucidated by the XRD analysis of **70** (see Fig. 21), the [*cyclo*- P_4]³⁻ moiety adopts a butterfly-like conformation where the newly inserted P atom is out of the plane of the other P atoms and one of the W(CO)₅ fragments has migrated to the former P₃ unit. Bond lengths within the *cyclo*-P₄ range from 2.2105(15) Å to 2.2302 (14) Å, slightly longer compared to the precursor [*cyclo*-P₃]³⁻ in **65** (2.194–2.205 Å).

The ³¹P NMR spectrum of **70** reveals four different sets of signals at δ = 105.7, 73.6, 38.1 and 4.3 ppm, all downfield shifted compared to the precursor **65** that features a singlet at δ = -204.8 ppm. Additionally, the signals at δ = 38.1 and 73.6 ppm show satellites for P-W coupling with ¹J_{P-W} = 237 Hz and 138 Hz respectively, due to the different hybridization of the two P atoms (sp³ and sp²). Compound **65** not only allows access to higher congeners but also to P₂-containing complexes. Velian and Cummins showed that [IMo(N[*t*Bu]Ar)₃] could abstract a P⁻ atom from the [*cyclo*-P₃]⁻ unit in **65** by salt metathesis [120,121]. The reaction affords [PMo (N[*t*Bu]Ar₃)] (**71**) in 79% yield, along with the binuclear complex **72** in 71% yield, see Scheme 25. The authors suggested that **72** could arise from the dimerization of the fragment [Nb(P₂)(ODipp)₃] (**73**) as the result of a 2(3–1) process which involves the starting [*cyclo*-P₃]⁻ moiety in **65**.

XRD analysis of **72** (Fig. 22) revealed that it bears a square *cyclo*-P₄ ligand with bond lengths/angles virtually identical to the known zirconium dimer [{(L)Zr}₂($\mu_2, \eta^4: \eta^4-P_4$)] (L = PhP(CH₂SiMe₂-NSiMe₂CH₂)₂-PPh) [**78**]. Nonetheless the [*cyclo*-P₄]^{4–} ligand in **72** occupies a disordered arrangement between the two Nb centers. The main crystalline component (75% occupancies) features an almost perfectly square planar P₄ fragment (torsion angle 0.15



Scheme 22. Synthesis of the cyclo-P₃ containing complexes 59 and 62.



Fig. 19. Molecular structures of cycloP₃ complexes 59 and 60. Hydrogen atoms are omitted for clarity; adapted from Ref. [112].

(9)°) which is coordinated to the metal atoms in an μ_2 , η^3 : η^3 mode. Notably, the internal *cyclo*-P₄ is tilted with respect to the Nb-Nb axis as observed for [U{(η^5 -C₅Me₄H)(μ^8 -C₈H₆SiiPr₃)₂-1,4)(μ , η^2 : η^2 -P₄)] (see Section 2.1.3.5). The P-P bond lengths (ranging from 2.230 to 2.250 Å) are considerably longer compared to dianionic [*cyclo*-P₄]²⁻ ligands found in the ionic salt CsP₄·2NH₃, (2.146 Å, [77]), or binuclear ETM complexes. This characteristic, in conjunction with the diamagnetic nature of **72**, indicates that the P₄ deck can be formulated as a [*cyclo*-P₄]⁴⁻ ligand. However, the structure representative at 20% occupancies shows a more rectangular shaped P₄ moiety coordinated in an μ_2 , η^2 : η^2 fashion whose structure recalls the phosphide ligand in the U(III) complex **32** mentioned above (see Section 2.1.3.5). DFT calculations indicates that the two coordination modes are separated by a small energy barrier (1.5 kcal mol⁻¹) suggesting fluxionality of the ligand. This correlates well with the ³¹P{¹H} NMR spectrum of **72**, in which no different P atoms are detected (singlet at δ = +124 ppm). In order to intercept the reactive species **73** prior to dimerization, the reaction was repeated in the presence of a large excess of 1,3-CHD affording the corresponding D-A adduct **74**. The relative instability of **74**, due to the presence of both *retro*-Diels-Alder equilibrium and concomitant dimerization of Nb-P₂, has prevented its isolation. It is worth mentioning that the double D-A adduct **75** could be observed (45% spectroscopic yield) when using pyridine-N oxide to cleave niobium from **74**. Pursuing a complementary (2 + 1) strategy, Cummins and co-workers also

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Scheme 23. Reductive activation of P₄ with Nb complex 64 and transformation in 69.



Fig. 20. Molecular structure of a) 68 and b) 69. Hydrogen atoms are omitted for clarity; adapted from Ref. [116].

demonstrated that Nb-[*cyclo*-P₃]⁻ complexes can be obtained by addition of the terminal phosphide complex [Na(OEt₂)]₂{PNb[N (Np)Ar]₃}₂ (**76**) to a suitable source of P₂, of which white phosphorus represents the most convenient and readily available choice [122]. When performed in a non-coordinating solvent (Et₂O), the reaction of **76** with one equivalent of P₄ led to the formation of the complex [Na(OEt₂)][(η^3 -P₃)Nb[N(Np)Ar]₃ (**77**) in 56% yield (³¹P NMR singlet at δ = -223 ppm). This process was further optimized by adding 4 equivalents of 12-crown-4 to the crude mixture, from which the monomeric cation-sequestered complex [Na(12-crown-4)₂][(η^3 -P₃)Nb[N(Np)Ar]₃] (**78**) could be obtained in an



improved 83% yield (³¹P NMR singlet at δ = -183 ppm in d_5 -pyridine) (Scheme 26).



Fig. 21. Molecular structure of the monoanion in complex 70. Hydrogen atoms are omitted for clarity; adapted from Ref. [119].



Scheme 25. Synthesis of the tetraphosphorus complex 72 and of the phosphide complex 71 via a 2(3-1) process and Diels-Alder cycloaddition product 75.



Fig. 22. Molecular structure of 72. Hydrogen atoms are omitted for clarity; adapted from Ref. [120].

The solid-state dimeric structure of **77** reported in Fig. 23 is retained in solution as suggested by the ³¹P NMR analysis of the complex. At room temperature **77** still possesses a pseudo three-fold symmetry due to fast scrambling of the Na-P interactions (singlet at $\delta = -223$ ppm) which freezes out at cryogenic temperature (2:1 doublet-triplet pair at $\delta = -178$ ppm and $\delta = -203$ ppm respectively, showed at -80 °C).

A different product was formed when performing the reaction in THF, which furnished the formal (4 + 1) addition complex [Na (THF)₆]{(N[Np]Ar)(η^4 -P₅)Nb(N[Np]Ar)₂} (**79**) in 71% yield with minimal observed formation of the *cyclo*-P₃ adduct **77** (Scheme 27).

The cyclopentaphosphane ligand $[RP_5]^{4-}$ in **79** possesses an envelope-like conformation with a P atom carrying the migrated anilide ligand, and the other four P atoms coordinated to the metal in an η^4 fashion (Fig. 24).

In addition to the cyclo-P₅ fragment, the remaining two anilide ligands interact with niobium in an η^3 fashion (Nb-C 2.5 Å) saturating the coordination sphere of the metal. Such interactions, although weaker than a canonical Nb-C single bond (usually below 2.3 Å), impede free rotation of the anilide fragments giving the complex an overall C_1 symmetry which translates into five different P atom environments and three different anilide ligands as observed by ³¹P NMR and ¹H NMR spectroscopy, respectively. The main factor determining the observed product selectivity is the aggregation state of the phosphide source in solution. When using coordinating solvents such as THF, the starting dimeric phosphide complex **76** splits into two low-coordinate monomeric units, which add to the P₄ molecule to form **79**. Conversely, in non-polar solvents, 76 is believed to retain its aggregation state and reacts with P_4 via a twofold (2 + 1) addition to give the dimeric complex 77. The ability of the terminal phosphide 76 to act as a single P⁻ atom donor, has also been exploited by Krummenacher and Cummins towards the generation of the phosphaethynolate salt [NaOCP] [123]. To accomplish this, 76 was first reacted with tris(pentafluorophenyl)borane, affording the capped complex [Na (OEt)₂][B(C₆F₅)₃PNb(N[Np]Ar)₃] (80) in 87% yield, see Scheme 28 and Fig. 25.

Upon treatment with a stoichiometric amount of CO₂, the red solution containing **80** turned bright yellow. ³¹P NMR analysis evidenced the consumption of **80** (δ = 514 ppm) in favor of a new product with a diagnostic resonance at δ = -393 ppm that could be attributed to the known [Na(dme)]₂[OCP]₂ (**81**). Along with **81**, the authors identified the boron-capped oxo-niobium complex [B (C₆F₅)₃][ONb(N[Np]Ar)₃] (**82**) deriving from the formal multiple bond metathesis between the two reagents (Scheme 28). As previously observed, the formation of a strong Nb=O bond is proposed to be the thermodynamic driving force for the CO₂ activation. This is proposed to occur via the nucleophilic attack of the phosphide on the carbonyl, which forms the PCO₂⁻ adduct or the cyclic Nb-P-C (O)-O intermediate [124]. The latter, previously isolated in a process involving the phosphide **76** and acylchlorides [125], undergoes [2 + 2] fragmentation to give the final products **81** and **82**.



Scheme 26. Reaction of P₄ with the dimer 76 in Et₂O affords the cyclo-P₃ compound 77. Yields are improved by addition of 12-crown-4 prior isolation (compound 78).

The groups of Arnold and Maron reacted P₄ with the carbonylated precursors [(BDI)(NtBu)M(CO)₂] [M = Nb (**83**), M = Ta (**84**), BDI = 2,6-diisopropylphenyl- β -diketiminate], yielding niobium and tantalum inverted sandwich complexes [(BDI)(NtBu)M]₂(μ_2 , η^3 : η^3 -P₄) [M = Nb (**85**), M = Ta (**86**)]. Such compounds were also obtained by hydrogenolysis of the bis-dimethyl precursor [(BDI) (NtBu)Nb(Me)₂] in presence of P₄ [126] (Scheme 29).

The reaction proceeded much faster in the case of Nb, this is explained by the higher energy barrier for CO dissociation in the case of Ta. Interestingly, both species **85** and **86** are air-stable and were characterized in solution by ¹H and ¹³C NMR spectroscopy, resulting highly symmetric diamagnetic species. The ³¹P NMR spectra in solution did not show any signals for both com-



Fig. 23. Molecular structure of **77**. Hydrogen atoms are omitted for clarity; adapted from Ref. [122].

plexes, thus ³¹P MAS NMR spectroscopy was required. The niobium complex **85** displayed two sets of signals (δ_{iso} = +251(2) ppm and δ_{iso} = +84(2) ppm) in agreement with the $\mu_2, \eta^3: \eta^3$ coordination mode adopted by the cyclo-P₄ ligand which was also observed in the solid-state structure (Fig. 26a). The ³¹P MAS NMR spectrum of 86 was difficult to interpret because of line-broadening and only a signal at δ_{iso} = +208(8) ppm was observed. The molecular structure of **86** was elucidated by XRD, see Fig. 26**b**. The cyclo- P_4 core in both complexes is close to a perfect planar square, the average P-P bond lengths in 85 and 86 are 2.233 Å and 2.248 Å respectively, very similar to those reported for $[\{(L)Zr\}_2(\mu,\eta^4:\eta^4-P_4)]$ (L = PhP $(CH_2SiMe_2NSiMe_2CH_2)_2$ -PPh) (2.240 Å) and $[(ODipp)_3Nb]_2(\mu_2,\eta^3)$: η^3 -P₄) (2.232 Å, vide supra), formally described as containing a tetraanionic $[cyclo-P_4]^{4-}$ core. Complexes **85** and **86** differ from these examples as the $[cyclo-P_4]^{4-}$ is sandwiched between the two metals in an asymmetrical manner, as evidenced by the large difference between the M–P distances: one short (Nb-P = 2.511 Å, Ta-P = 2.500 Å), and two average bond lengths (Nb-P = 2.700 Å and 2.702 Å, Ta-P = 2.691 Å and 2.674 Å). Another route to synthesize 85 involved the hydrogenation of the dimethyl precursor [(BDI)Nb (NtBu)Me₂] 83 in the presence of P₄. As before this protocol furnished the desired triple decker 85 as the major species but proved to be less selective. Among the by-products observed, the trinuclear niobium complex {[(BDI)NbNtBu]₃(µ-P₁₂)} (87), Scheme 30, was isolated bearing a rare P₁₂ cluster with an unprecedented architecture of a cyclo-P₅ ring and a P₇ cage featuring a nortricyclane structure as shown by XRD analysis (Fig. 26c).

The P-P bond distances vary between 2.146 and 2.290 Å, comparable with other phosphide cages and in agreement with P-P single bonds. To date, there are only two examples of P₁₂ clusters, though possessing different topology, NHC-capped P₁₂ [127] and $\{[(Cp')Co]_3(\mu-P_{12})\}$ [128]. To assess the basicity of the [*cyclo*-P₄]^{4–} core in **85**, it was reacted with [HOEt₂][B(C₆F₅)₄], resulting in the



Scheme 27. Synthesis of complex 79.

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Fig. 24. Solid state structure of the monoanion in the salt complex **79.** $[Na(THF)_6]^+$ moiety and hydrogen atoms are omitted for clarity; adapted from Ref. [122].



Scheme 28. Capping of 76 with B(C₆F₅)₃ and subsequent reaction with CO₂.

protonation of the less sterically hindered β -diketiminate backbone instead of the phosphide moiety, see Scheme 31, yielding {[(BDI")(NtBu)Nb]₂(μ 2, η ³: η ³-P₄)}(BAr^F)₂ (**88**) in 82% yield.

Oxidation with one or two equivalents of Ag[B(Ar^F)₄] afforded the complexes {[(BDI)(NtBu)Nb]₂(μ_2 , η^3 : η^3 -P₄)}{B(Ar^F)₄} (**89**), and {[(BDI)(NtBu)Nb]₂(μ_2 , η^3 : η^3 -P₄)}{B(Ar^F)₄} (**90**), respectively, the latter displaying a sharp singlet at δ = +516 ppm in the ³¹P NMR spectrum. It would seem the oxidation state of the internal *cyclo*-P₄ greatly affected the structure of the complex, particularly the interatomic P-P bond distances. The mono-oxidized form, **89**, displayed shorter P-P bonds (average, 2.209(4) Å) with respect to **85**. The P-P bond lengths are shortened further in the two-fold oxidized complex **90** (averagely 2.182(5) Å), which is consistent with known *cyclo*-P₄²⁻ complexes (Scheme 32).



Fig. 25. Solid state structure of the anion of 80. Hydrogen atoms are omitted for clarity; adapted from Ref. [123].



Scheme 29. Activation of P₄ with Nb and Ta complexes 83 and 84.

2.3.3. Tantalum

The formation of inorganic fullerene-like structures, arising from the supramolecular assembly of CuX units (X = Cl, Br, I) together with [Cp*Fe(η^5 -P₅)] (Cp* = C₅Me₅) was established by Scheer and co-workers [129]. This approach was extended to Ta-complexes [Cp''Ta(CO)₂(η^4 -P₄)] (Cp'' = 1,3-C₅H₃tBu₂) containing the *cyclo*-P₄ ligand [130]. Recent studies carried out on the [Cp^R-Ta(CO)₂(η^4 -P₄)]/CuX system [Cp^R = 1,4-C₅H₃tBu₂ (**91**), or Cp^R = Cp''' = 1,2,4-C₅H₂tBu₃ (**92**), X = Cl, Br) have shown that distinct non-classical fullerene structures are accessible depending on both the halide used and on the specific Cp^R ligand [55]. These supramolecular assemblies are formed upon layering a CuX solution in CH₂Cl₂/CH₃CN on a CH₂Cl₂ solution of [Cp^RTa(CO)₂(η^4 -P₄)] (see Scheme 33).

Structures ranging from **93a** to **93d** were obtained from the precursor **91** or **92** upon the addition of CuX = CuCl, CuBr. In this assembly, six *cyclo*-P₄ units are bound to eight CuX units in a 1,2,3,4-coordination mode. Each Cu atom is tetrahedrally coordinated by three P atoms and one terminal halide. The inorganic core consists of 32 atoms and features a Cu₈ cube with *cyclo*-P₄ rings above its faces. When switching to iodide, two distinct products were isolated upon layering a CH₃CN solution of CuI onto a solution of **91** in toluene, $[{Cp''Ta(CO)_2(\eta^4-P_4)}_{10}{Cu_1_4I_10(\mu,I)_2(\mu_3,I)_2}]$ **(94)** and $[{Cp''Ta(CO)_2(\eta^4-P_4)}_{5}{Cu_1_2I_3(\mu,I)_8(\mu_4-I)(CH_3CN)_5}]$ **(95)** (see Scheme 33). The molecular structure of **94**, shown in Fig. 27**a**, consists of an unprecedented nano-capsule 'peanut'shaped motif, made up of ten *cyclo*-P₄ units bound to eight CuI units and two Cu₃I₃ units. Compound **95** also possesses a hollow structure, consisting of five *cyclo*-P₄ units bound to three neutral



Fig. 26. Solid-state structures of a) 85 b) 86 and c) 87. Hydrogen atoms are omitted for clarity; adapted from Ref. [126].

 $\{Cu_2I_2CH_3CN\}\ units, one cationic \{Cu_2I(CH_3CN)_2\}^+$ and one anionic $\{Cu_4I_5\}^-$ unit (Fig. 27**b**).

When an additional *tert*-butyl group is added in the Cp ring (i.e. using **92** in place of **91**), the reaction with Cul yielded a new 2D coordination polymer $[Cp'''Ta(CO)_2(\eta^4-P_4)\{Cu_4(\mu_3,I)_4\}]_n$ (**96**), in which Ta-phosphide moieties are linked to $\{Cu_4(\mu_3,I)_4\}$ heterocubane units in a 1,2,3,4-coordination mode.

2.4. Group 6 metals

2.4.1. Chromium

In 2005 the Power group reported the first stable compound featuring a quintuple Cr-Cr bond, L_2Cr_2 where $L = C_6H_3$ -2,6(C_6H_3 -

2,6-*i*Pr₂)₂ [131]. Following this study, other examples of Cr₂ units with the same bond order have appeared in literature, such as **97** (see Scheme 34) reported by Kempe [132], where the bimetallic core, stabilized by a chelating N-ligand, displays Cr-Cr distances of 1.750 Å. In a collaboration between Scheer's and Kempe's groups [133] it was shown that Cr₂ units can be suitable platforms for the activation of P₄ as well as other pnictogens and pnictides (i.e. As₄ and AsP₃). Reacting **97** with P₄ in THF at room temperature yielded the *cyclo*-P₄²⁻ derivative (**98**) in excellent yield (97%, see Scheme 34).

The X-ray structure of **98** (Fig. 28**a**) revealed a nearly squareplanar phosphorus arrangement with P-P distances ranging from 2.168 to 2.183 Å, consistent with P-P bonds of a *cyclo*- P_4^2 -ligand.



Scheme 30. Activation of P₄ with Nb and Ta complexes 83 and 84.



Scheme 31. Protonation of **85** occurs on the ligand backbone leaving the [*cyclo*-P₄]^{4–} core unaltered.



Scheme 32. Oxidation of complex 85 with $Ag[B(C_6F_5)_4]$.

Opposite P1 and P3 atoms are bonded in a η^1 fashion to different Cr atoms, while P2 and P4 bridge the two Cr centers. It is worth noting that the Cr-Cr distance in **98** (1.866 Å) is indicative of quadruple bond. The elongation of the Cr-Cr distance signifies an oxidation of the Cr₂ core occurs upon reaction of **97** with P₄, which in turn is reduced to give the planar, aromatic *cyclo*-P₄²⁻ fragment in **98**. The reaction was repeated using As₄ and AsP₃, yielding the unprecedented *cyclo*-As₄²⁻ (68%) and *cyclo*-AsP₃²⁻ ligands (69%).

Aiming to investigate the reactivity of P_4^{-1} lone pairs, complex **98** was reacted with an excess of [W(CO)₅(THF)], which gave **99** exclusively (19%, see Scheme 34). X-ray diffraction analysis of **99** revealed the coordination of the W(CO)₅ fragment to a terminal phosphorus atom, see Fig. 28**b**. However, the ³¹P NMR spectrum showed a broad pseudo triplet at δ = 250 ppm (uncoordinated P atoms) and a very broad signal between δ = 210 and δ = 230 ppm for P-W atoms, suggesting **99** possesses dynamic behavior in solution due to a switching of the {W(CO)₅} moiety between bridging and terminal positions on the *cyclo*- P_4^{-1} ring. This discrepancy between solid state and solution behavior was rationalized *via* DFT calculations, which indeed evidenced a relatively small energy difference existing between the two coordination modes of the {W(CO)₅} fragment.

2.4.2. Molybdenum

Over the past decade only a few examples of the activation of P_4 involving the use of molybdenum complexes have been reported. In 2013, Braunschweig and co-workers, reported on the reaction of Mo metallophane **100** with P_4 in toluene at 80 °C, resulting in the formation of **101** as an air and moisture-sensitive red solid in 30% yield (Scheme 35) [134].

This represents a unique mode of activation for P_4 mediated by an ansa-metallocene complex. The ${}^{31}P{}^{1}H$ NMR spectrum of **101** showed only one resonance at δ = -370.9 ppm, highly deshielded in comparison to other three-membered rings containing P-P bonds [135]. The solid-state structure (see Fig. 29) confirms the formation of a three-membered ring consisting of a P₂ fragment and the central molybdenum center, with concomitant cleavage of two Mo-Si bonds and formation of two Si-P bonds. This enables an appreciable release of strain as the angle between the Si-C_{inso} axes and the C₅H₄ ligand decreases from 48.7° and 48.9° in 100 to 19.4° and 19.1° in **101**, respectively. Likewise the tilt angle between the two C₅H₄ ligands increases, from 20.48(7)° to 43.08° for 100 and 101, respectively. The P-P bond length (2.172 Å) in 101 is shorter compared to a P-P single bond (2.21 Å) [60] but longer than a P=P double bond found in organic diphosphenes such as $(tBu)_{3}C_{6}H_{2}$ -P=P-C₆H₂ $(tBu)_{3}$ (2.031 Å) [136]. Wiberg bond indices and computational analysis (NBO) revealed that all the bonds involved in the three-membered cyclic core of 101 are single despite the existence of some degree of delocalization over the ring and π - π interaction between the two P atoms.

The pursuit of air-stable platforms for the safe storage, transport and controlled release of white phosphorus is a relevent challenge in contemporary research. One valuable route is the encapsulation of P₄ in supramolecular cages [137] or porous materials. In this regard, Scheer et al. successfully intercalated P4 molecules in the porous structure of activated carbon [138] and the resulting P_4/C could be used as a source of P_4 either in solution or in the gas phase. In 2019, Figueroa and co-workers found that low-valent Mo complexes stabilized with bulkier isonitrile ligands could be used as a storage/release system for white phosphorus [139]. The zero-valent Mo complexes of formula [Mo(η^2 -C,H: κ^1 - $N-(HN(i-Pr)_2)(CO)_2(CNAr^{Dipp2})_2$ (102, $Ar^{Dipp2} = 2,6-(2,6-(i-Pr_2))_2$ C_6H_3) were reacted with P₄ selectively affording the mononuclear derivative $[(\eta^4-P_4)Mo(CO)_2(CNAr^{Dipp2})_2]$ (103) in near quantitative yield, as shown in Scheme 36. It is worth noting that 103 is kinetically stable for days upon air exposure and warming up to 90 °C.



Scheme 33. Reactivity of 91 and 92 with Cu(l) halides. Each cyclo-P₄ unit is additionally bonded to an hexo Cp^RTa(CO)₂ fragment, omitted for clarity.



Fig. 27. Core motifs of structures a) **94** and b) **95** (b). Each *cyclo*-P₄ unit in the frame is additionally bonded to a [Cp^RTa(CO)₂] fragment protruding outside of the shell (omitted for clarity); adapted from Ref. [55].



Scheme 34. Synthesis of complexes 98 and 99. The number above the Cr-Cr bond refers to the bond order.

The solid-state structure of **103** (see Fig. 30), shows a rectangular planar *cyclo*-P₄ ligand with P-P bond lengths between 2.148(2) and 2.180(2) Å. The observation of a single NMR resonance at δ_{31P} = 113 ppm confirmed the highly symmetrical structure remains in solution. DFT calculations best formulated the *cyclo*-P₄ moiety as a six π -electron dianion ligand, indicating that a formal two-electron reduction of P₄ occurs upon reaction with **102**.

The presence of high-lying σ P-P HOMOs and low-lying P-Mo anti-bonding LUMOs suggested that release of P₄ could be achieved through light irradiation of **103** as a result of a ligand to metal

charge transfer process (LMCT). However, the energy needed for this process (irradiation at $\lambda = 284$ nm), also triggers the photo degradation/polymerization of P₄ [140,141]. With the aim to reduce the energy gap involved in the LMCT transition and releasing P₄ without subsequent polymerization, the divalent complex [MoI₂(CO)₂(CNAr^{Dipp2})₂] (**104**, precursor of **102**) was reacted with P₄ (Scheme 37). The reaction afforded the desired P₄-adduct [(η⁴-cyclo-P₄)MoI₂(CO)(CNAr^{Dipp2})₂] (**105**) in excellent yield (90%), with loss of one CO and retention of both iodide ligands, with one occupying the *trans* position respect to the centroid *cyclo*-



Fig. 28. Molecular structures of a) 98 and b) 99. Hydrogen atoms are omitted for clarity; adapted from Ref. [133].

P₄, see Fig. 31. The ³¹P NMR spectrum is desymmetrized giving an AA'BB' spin pattern centered at δ = 75 ppm. DFT calculations on a model substrate of **105** showed that it has a smaller HOMO-LUMO gap and therefore a LMCT transition accessible at longer wavelengths. As such, irradiation of **105** in benzene together with 2% DME by a low power violet led source (18 W, λ = 385–395 nm) gave dissociation of P₄ and its replacement by a DME molecule in the coordination sphere of Mo. ³¹P NMR spectroscopy showed the reaction was completed after only 20 min as 95% of free P₄ returned to solution. It should be added that the newly formed complex [MoI₂(CO)(k_1 -O-DME)(CNAr^{Dipp2})₂] (**106**), coordinates the dissolved white phosphorus only when the mixture is heated at 80 °C, giving **105**, demonstrating the complete reversibility of the process (Scheme 37).

The mononuclear complexes **103** and **105** were further investigated with the aim to elucidate the aromaticity of the *cyclo*-P₄ moiety, its bonding situation towards the metal center, and how the valence of the latter influences the structural features of the complexes [142]. Thus, **105** was treated with KC₈ in THF, followed by addition of dibenzo[18-crown-6] (DB18C6), which yielded red crystal of [K₂(dibenzo-18-crown-6)][(η^4 -P₄)Mo(CO)(CNAr^{Dipp2})₂] (**107**, Scheme 38).

XRD analysis (Fig. 32) revealed a three-legged piano stool geometry in which both iodide ligands are lost while one K⁺ is partially sequestered and is involved in a secondary interaction with a P atom of the *cyclo*-P₄ ligand. Conversely, the second K⁺ is coordinated to the C \equiv N and the aryl moieties of Ar^{Dipp2} and cannot be removed from the coordination sphere of the metal even using dibenzo[18-crown-6] in large excess. The ³¹P NMR spectrum of **107** showed a singlet at δ = 112.4 ppm which is relatively close to the signals exhibited by the precursor **105** (δ = 90 ppm and δ = 60 ppm), suggesting that the dianionic nature of the [*cyclo*-



Scheme 35. Synthesis of complex $101\ \mbox{by}$ activation of $\mbox{P}_4\ \mbox{mediated}$ by the Mo complex 100.



Fig. 29. Solid-state structure of 101. Hydrogen atoms are omitted for clarity; adapted from Ref. [134].

 $P_4]^{2-}$ ligand was not affected by the reduction. This is also the case for the average P-P bond length within the $[cyclo-P_4]^{2-}$, which does not appreciably change after the reaction (2.157(5) Å in **105** vs 2.160(12) Å in **107**).

Analogous results were obtained when treating **103** with KC₈. ³¹P NMR spectroscopy evidenced the formation of two compounds, one corresponding to $K_2[(\eta^4-P_4)Mo(CO)(CNAr^{Dipp2})_2]$ (**108**), analogous to **107** without dibenzo[18-crown-6], with a resonance at δ_{31P} = 105 ppm, the other resulting from the loss of one CNAr^{Dipp2} ligand, $K_2[(\eta^4-P_4)Mo(CO)_2(CNAr^{Dipp2})]$ (**109**), resonating at δ_{31P} = 121 ppm, Scheme 39. Notably, **108** could be fully converted to **109** when treated with CO.

The similar ³¹P NMR chemical shifts found in **103**, **108** and **109**, suggested the *cyclo*-P₄ ligand remained intact upon reduction with KC₈. As with **107**, XRD analysis of **109** (see Fig. 33) showed a three-legged stool geometry with almost unchanged P-P bond lengths (2.160 Å vs 2.163 Å for **109** and **103** respectively). Conversely, a marked increase of the Mo-(η^4 -P₄) distance in **108** and **109** (2.112 Å and 2.102 Å respectively) compared to **103** (2.075 Å), is representative of an increase of electron density at the metal center as confirmed also by computational analysis.

Complexes **108** and **109** possess a Mo-C bond distance for both the isocyanide and CO ligands that are 0.1 Å and 0.2 Å shorter than those for **103** and **105** respectively. This agrees with v_{CO} and v_{CN} IR



Scheme 36. Coordination of P₄ to the zero-valent Mo complex **103**.



Fig. 30. X-ray structure of $[(\eta^4-P_4) [Mo(CO)_2(CNAr^{Dipp2})_2]$ (103). Hydrogen atoms are omitted for clarity; adapted from Ref. [139].

bands that are lower in energy in **108** and **109** than those in **103** and **105** and support the increased π -backbonding interaction in the salts. Thus, the $(cyclo-P_4)^{2-}$ ligand can be viewed as a Hückel-type aromatic system, bound to a zero-valent Mo fragment in **108** and **109**.

Molybdenum-phosphide complexes, due to their unique coordinating properties, have also been extensively investigated in relation to the formation of 2D or 3D frameworks. In their seminal work [143], Scheer and co-workers have been committed in building up coordination polymers involving non-covalent bonding interactions between three components: 1) metal-phosphide complexes; 2) Lewis acidic metal salts coordinated to weakly coordinating anions (WCA) and 3) organic linkers. With this in mind, it was observed that the reaction of $[Cp_2Mo_2(CO)_4(\eta^2-P_2)]$ (110) with Ag[Al{OC(CF_3)_3}] forms the dimeric cluster 111 (Scheme 40), which has a dynamic coordination behavior in dichloromethane solution. It has been proposed that 111 is in equilibrium with its two monomeric forms, where a P₂ ligand can be coordinated to Ag⁺ as η^1 -end on (112) or η^2 -side on (113). The two coordination modes are separated by a modest energetic barrier (9 kJ mol⁻¹ from 113 to 112 as calculated by DFT methods) [144].

Such $\eta^2 \rightarrow \eta^1$ equilibrium opens up a free site on the Ag centers and the possibility of connecting Ag[Cp₂Mo₂(CO)₄(η^2 -P₂)]⁺₂ units with an appropriate linker, giving rise to new organometallicorganic hybrid materials. The reaction of **111** with the ditopic organic linker *trans*-1,2-di(pyridine-4-yl)ethene **114** in CH₂Cl₂ gives the two coordination isomers **115** and **116** shown in Scheme **41** [144].

Compound **115** is a one-dimensional chain-like polymer where the $\{Ag_2[Cp_2Mo_2(CO)_4(\eta^2-P_2)]_4\}^{2+}$ units are connected to each other through the organic linker, **114**. In particular, the sixmembered Ag_2P_4 motif is retained and assumes an almost planar conformation (folding angles 2.23(7)° and 1.99(7)°) while in **111** the Ag_2P_4 units assume a chair conformation with folding angles of 20.69(2)°. The linkers are perpendicular to the plane of the Ag_2P_4 units and *trans* with respect to the Ag-Ag axis and the Ag (I) centers exhibit a distorted tetrahedral geometry (see Fig. 34 for X-ray structure of **115**). Conversely, **116** consists of a discrete molecular assembly where two Ag_2P_4 units are connected by two linker molecules which are in a *cis* arrangement (see Fig. 35 for X-ray structure of **116**) relative to the Ag-Ag axis. Compound **116**



Scheme 37. Reversible process of coordination and release of P₄.



Fig. 31. X-ray structure of $[(\eta^4-cyclo-P_4)Mol_2(CO)(CNAr^{Dipp2})_2]$ (105). Hydrogen atoms are omitted for clarity; adapted from Ref. [139].

is the first example of a metallaparacyclophane assembled using P_n ligand complexes, cationic metal salts and organic linkers [145].

It was possible to form 2D networks by replacing Ag[Al{OC $(CF_3)_3\}_4$] with $[Cu(CH_3CN)_4][BF_4]$. Following the same strategy as above, the reaction of $[Cu(CH_3CN)_4][BF_4]$ with the diphosphide complex ligand **110** was carried out and, independent of the stoichiometry, provided the dimeric complex $[Cu_2\{Cp_2Mo_2(CO)_4(\eta^2 - P_2)\}_4][BF_4]_2$ (**117**), see Scheme 42. Compound **117** was able to coordinate the rigid *trans*-bipyridine linker **114** which replaced the η^2 -side on phosphide complex ligands affording the poorly soluble coordination polymer **118**. The X-ray structure of **118** is shown in Fig. 36, and consists of a 2D network of $Cu_2[Mo]_4P_4$ nodes with $[Mo] = [CpMo(CO)_2]$ and the Cu_2P_4 six-membered ring nearly planar with a folding angle of 3.61°. The Cu(I) centers feature a distorted tetrahedral coordination geometry with N-Cu–N angles of 110.32(16)°, which creates rhomboid shaped cavities in the nanometer range with maximum diameter of 2.20 nm.

The coordination properties of dimer **117** (see Scheme 43) were further investigated by using a series of flexible linkers based on 4,4'-bipyridine (119, 120) and 3,3'-bipyridine (121, 122) [146]. Regardless of the length and position of the spacer between the pyridyl fragments, only 1D coordination polymers were selectively formed in moderate to high yields (65 to 94%), see compounds 123, 124, 125, 126, in Scheme 43. Two different morphologies were observed which are characterized by a different number of Cu(I) centers at the nodes of each ring of the chain: 2 Cu(I) centers for 123 and 125, 4 Cu(I) centers for 124 and 126 (Fig. 37). Compounds 123 and 125 can be described as infinite dinuclear metallacyclic units connected to each other by Mo₂P₂ ligands, while in 124 and 126 there is a tetrametallic repeating unit, where the Mo₂P₂ ligands are part of the metallacycle. Consequently, the new onedimensional organometallic-organic polymers possess a virtual porosity having cavities with circular (123-125) and rectangular meshes (126) of maximum diameter 1.47 nm.



Fig. 32. X-ray structure of $[K_2(dibenzo[18-crown-6][(\eta^4-P_4)Mo(CO)(CNAr^{Dipp2})_2]$ (**107**). Hydrogen atoms are omitted for clarity; adapted from Ref. [142].

Higher order polyphosphorus molybdenum complexes have also been considered for the formation of novel coordination polymers. The hexaphosphabenzene complex $[(Cp^*Mo)_2(\mu, \eta^6; \eta^6 - P_6)]$ (**127**), ($Cp^* = C_5Me_5$) first isolated in 1985 by Scherer *et al.* [147], bears great potential in terms of the possible variety of coordination patterns, although the research on this field has been limited by the scarce efficiency of known protocols to obtain it. Synthesis of the triple decker complex $[(Cp^*Mo)_2(\mu_2,\eta^6:\eta^6-P_6)]$ was improved to a remarkable 64% yield by Scheer et al. [148] after prolonged heating of $[Cp^*Mo(CO)_{2/3}]_2$ in the presence of P₄ in refluxing diisopropylbenzene. Interestingly, the reaction also yielded trace amount of a novel binuclear complex $[(Cp^*Mo)_2(\mu_2, P_{10})]$ (128), displaying a folded $[P_{10}]^{2-}$ ligand with four short P-P bonds (2.140(2) Å) of double bond character. It is one of the few known P₁₀ ligand derived from P₄, see Section 2.1.3.2. As shown by the solid state structure in Fig. 38, the Mo centers are coordinated to the P=P moieties in an η^2 : η^2 mode and to the edge-P atoms in an κ^1 mode.

The coordination behavior of **127** was investigated with Ag[TEF] and Cu[TEF] (TEF = [Al{OC(CF₃)₃}₄]⁻. Addition of Ag[TEF] or Cu[TEF] to a DCM solution of **127** resulted in the oxidation of the starting material to afford the paramagnetic derivative **129** (μ_{eff} = 1.67 μ_B), which was detected through ESI mass analysis (see Scheme 44). EPR analysis of **129** showed a broad signal with no hyperfine coupling at 77 K at g_{iso} = 2.024 (solid state) or g_{iso} = 2.024 (DCM solution). Suitable crystals for XRD analysis were obtained by exchanging the anion (TEF) for another WCA, [FAl{OC₆F₁₀(C₆F₅)}₃]⁻ ([FAI]) affording compound **130**. According to the solid-state structure of **130**, the oxidation of the Mo center impacts the bonding situation of the *cyclo*-P₆ deck, thus it can be now regarded as



Scheme 38. Reduction of complex 79 with KC₈ affords complex $[K_2(dibenzo[18-crown-6][(\eta^4-P_4)Mo(CO)(CNAr^{Dipp2})_2]$ (107).



Scheme 39. Reduction of 103 with KC₈ occurs at the metal center leaving the cyclo-P₄ fragment intact.



Fig. 33. X-ray structure of the dianionic portion of $[K_2(dibenzo18-crown-6)][(\eta^4-P_4)Mo(CO)_2(CNAr^{Dipp2})]$ **109**. Hydrogen atoms are omitted for clarity; adapted from Ref. [142].

two P_3 fragments joined by elongated P-P bonds (2.247 and 2.294 Å, bisallylic distortion).

The addition of Cu[TEF] and Ag[TEF] in a toluene/DCM solution of **127**, allowed for the formation of the coordination derivatives **131** and **132** respectively. The ³¹P{¹H} NMR spectrum showed a broad singlet (ω_{ν_2} = 126 Hz) at δ = -290.4 ppm for **131** and a broad singlet (ω_{ν_2} = 155 Hz) at δ = -294.0 ppm for **132**, suggesting **127** possesses a highly dynamic coordination behavior towards Cu⁺ and Ag⁺. Single crystal X-ray analysis (Fig. 39) shows that **131** and **132** are isostructural and the central metal cation Cu(I) and Ag(I) respectively, is coordinated to two *cyclo*-P₆ moieties in an η^2 side-on mode, which results in a distorted square planar geometry.

When the reaction was performed in toluene only, the addition of Cu[TEF] to **127** provided a new coordination adduct, **133** with a distorted tetrahedral coordination geometry around the Cu(I) center and shorter Cu-P bonds with respect to **131** (Scheme 44 and Fig. 40), while the coordinating P-P bonds are longer.

Although 131 and 132 form a layered structure in the crystal lattice with alternating anionic [TEF] and cationic $\{M[(Cp^*Mo)_2(\mu, \mu)]$ $\eta^6:\eta^6-P_6]_2$ (M = Ag, Cu) sheets, they do not form 2D networks. By reacting 127 with Tl[TEF], see Scheme 44 below, a 2D coordination polymer 134 was obtained and elemental analysis was consistent with a Tl⁺/**127** ratio of 1:2. XRD analysis of **134**, showed that Tl^+ coordinates three cyclo-P₆ units in an η^2 side-on fashion displaying a bipyramidal trigonal geometry. This allows the coordination to propagate through the cationic sheets of the crystal lattice and to form a supramolecular network which can be regarded as a graphene-like structure. The ³¹P{¹H} NMR spectrum of **134** showed no signal at room temperature, a broad peak ($\omega_{1/2}$ = 1303 Hz) at δ = -305.5 ppm was observed when cooling the solution down to 193 K. Additional studies were carried out by Scheer and coworkers using Cu(I) salts of more coordinative anions, such as halides [149]. Treatment of 127 with CuBr afforded black crystals of the polymer [{(CpMo)₂(μ , $\eta^{3:3:1:1:1:1}$ -P₃)(μ , $\eta^{3:2:1:1:1:1}$ -P₃)}Cu₄(μ , Br_{4}_{n} (**135**). This consists of a 2D network formed by four P atoms of the cyclo-P₆ deck, each one coordinating two copper atoms in an η^1 : η^1 mode. The Br⁻ anions bridge the copper atoms completing the tetrahedral environment around Cu(I). Notably, the cyclo-P₆ moiety undergo not only a marked bisallylic distortion, with distances between the two P₃ fragments of 2.652(3) Å and 2.444(3) Å, but also a considerable distortion from planarity, with one P atom of the two P₃ fragments out of plane (envelope conformation). This leads the Cp ligands to be inclined by 9.26° with respect to the parallel conformation in 127. Despite this, polymer 135 still forms layers whereby 127 units are bound together by CuBr at the level of the cyclo-P₆ middle-deck (see Fig. 41).

Similar results were obtained by employing Cul. Layering a CH₃CN solution of Cul on a CH₂Cl₂ solution of **127** caused the formation of black crystals of a 2D coordination polymer of formula [{(CpMo)₂(μ , $\eta^{3:3:1:1}$ -P₃)(μ , $\eta^{3:2:1:1}$ -P₃)]_{0.84}{(CpMo)₂ (μ , $\eta^{6:6:1:1:1:1}$ -P₆)]_{0.16}Cu₂(μ ,I)₂]_n (**136**)]. As with **135**, in **136** the *cyclo*-P₆ deck coordinates with Cu(I) in a 1,2,4,5 mode with only one copper atom bound to each ligating phosphorus atoms. Moreover, the *cyclo*-P₆ moiety shows some degree of disorder as



Scheme 40. Synthesis of 111 and the proposed coordination equilibrium.



Scheme 41. Synthesis of **115** and **116** by assembly of **111** with bidentate ligand **114**, X = [Al{OC(CF₃)₃]₄]

it displays two different coordination patterns: $[\{(CpMo)_2 (\mu, \eta^{3:3:1:1}-P_3)(\mu, \eta^{3:2:1:1}-P_3)\}Cu_2(\mu, I)_2]_n$ **136a** and $[\{(CpMo)_2 (\mu, \eta^{6:6:1:1:1:1}-P_6)Cu_2(\mu, I)_2]_n$ **136b** with occupancy factors of 0.84 and 0.16 respectively. **136a** displays both bisallylic distorsion of the P₆ deck and the envelope conformation as with **135**. However, the minor component, **136b** shows a planar *cyclo*-P₆ deck with shortening of the P-P bonds upon coordination to Cu without a notable bisallylic distorsion (bond lengths in the range of 2.000–2.132 Å). Owing to their inherent insolubility, **135** and **136** were only characterized by ³¹P MAS spectroscopy. Six different signals were observed spread over a window of 900 ppm, consistent with a distorted P₆ moiety.

Ruiz and co-workers reported the activation of P₄ by the triply bonded anion $[Mo_2Cp_2(\mu-PCy_2)(\mu-CO)_2]^-$ (**137**) under mild conditions [150], yielding the anoinc diphosphorus complex $[Mo_2Cp_2(\mu2,PCy_2)(CO)_2(\mu2,\kappa^2:\kappa^2-P_2)]^-$ (**138**) (see Scheme 45).

The anionic **138** features a tetrahedral Mo_2P_2 core, where the P_2 ligand symmetrically bridges the two Mo centers. The PCy₂ ligand

lies in the plane defined by a Mo₂P face, with the remaining Cp and CO ligands filling the coordination sphere of the metal. The computed P-P distance is 2.09 Å, much shorter than the value of 2.21 Å found in P₄, suggesting the presence of some additional π (P=P) character. The intrinsic nucleophilic nature of 138 translates into high reactivity towards electrophiles such as CH₃I and CISnPh₃ see Scheme 45. Methylation takes place at a basal P atom, yielding the neutral diphosphenyl complex $[Mo_2Cp_2(\mu-PCy_2)(CO)_2(\mu 2,\kappa^2:$ κ^2 -P₂-Me)] (139), whose X-ray structure is shown in Fig. 42a. The P-P-C bond angle measures at 122.8(1)° and is consistent with Me⁺ addition to the P lone pair, as also revealed by theoretical calculation, with no effect on the P-P bond. The reaction with SnPh₃Cl takes place in an analogous fashion, yielding [Mo₂Cp₂(µ-PCy₂) $(CO)_{2}{\mu^{2},\kappa^{2}\cdot\kappa^{2}-P_{2}(SnPh_{3})}$ (140) (see Fig. 42b). However, compared to 138, the P-P distance in 140 is more elongated, approaching the typical value of a P-P single bond and a P-P-Sn bond angle of 80.3(1)°. As clarified by DFT calculations, interaction with SnPh₃⁺



Fig. 34. Section of the cationic coordination polymer of 115 in the solid state. Hydrogen atoms, Cp and CO ligands on Mo, are omitted for clarity; adapted from Ref. [144].



Fig. 35. Molecular structure of the tetracation of 116 in the solid state. Hydrogen atoms, Cp and CO ligands on Mo are omitted for clarity; adapted from Ref. [144].

involves filled orbitals on the anion having partial π (P=P) character, whose orientation gives the best overlap when P-P-Sn = 90°, thus explaining both the unexpectedly acute P-P-Sn bond angle and the observed P-P bond elongation. This rare coordination mode described by Ruiz *et al.*, was thoroughly explored by reacting **138** with various XER₃ electrophiles (X = halogen, E = C, Si, Ge, Sn, Pb, R = H, alkyl or aryl) [151]. In addition to this, the nuchleophilic character of the diphosphorus unit in **138**, was further investigated through the reaction with chlorophosphines ClPR₂ (R = Cy, Ph, *t*Bu) and transition-metal halide complexes such as [FeClCp(CO)₂], [MoCpl(CO)₃], [ZrCl₂Cp₂], [MnBr(CO)₅] and [ReCl(CO)₅] [152], as well as with 16-electron metal carbonyl fragments [ML_n(THF)] (ML_n = Fe(CO)₄, MnCp'(CO)₂, Mo(CO)₅, W(CO)₅) [153]. Similar studies were carried out to investigate the reactivity of **139** towards 14- and 16-electron metal carbonyl fragments [154,155], with the aim to form novel heterometallic clusters. In particular, the reaction of **139** with [Fe₂(CO)₉] or [W(CO)₄(THF)₂] in a 1:1 ratio yielded the stable trinuclear clusters [Mo₂MCp₂(μ_3 -P)(μ -PCy₂)(μ_3 -PMe)(CO)_X] [M = Fe, x = 5 (**141**); M = W, x = 6, **141a**], see Scheme 46 and Fig. 43**a**. Repeating the reaction with an excess of [Fe₂(CO)₉], the main product was the tetranuclear phosphide- and phosphinidene-bridged derivative [(Fe₂Mo₂Cp₂)(μ_4 -P)(μ -PCy₂)(μ_3 -PMe)(CO)₉] (**142**) along with small amounts of **141**, see Scheme 46. In both cases, there is cleavage of the P-P bridge with a Fe(CO)₃ moiety sitting on the MoP₂ face of the Mo₂P₂ tetrahedral core. Additionally, in **142** a Fe(CO)₄ unit is coor-



Scheme 42. Synthesis of 2D coordination polymer 118.



Fig. 36. Structure of the dicationic repeating unit of 118 in the solid state. Hydrogen atoms, Cp and CO ligands on Mo are omitted for clarity; adapted from ref. [144].

dinated to the naked P atom. Similarly, the reaction of 139 with an equimolar amount $[MnCp'(CO_2)(THF)](Cp' = C_5H_4Me)$ or $[W(CO)_5(-$ THF)] gave $[(ML_n)Mo_2Cp_2(\mu_3-P)(\mu-PCy_2)(\mu_3-PMe)(CO)_x]$ (ML_n =-MnCp', x = 4, **143**; ML_n = W, x = 7, **143a**). These compounds were found to be unstable and decompose at room temperature or under gentle heating with loss of the methylphosphinidene (PMe) unit, yielding the corresponding trinuclear complexes [(ML_n)Mo₂Cp₂(- μ_3 -P)(μ -PCy₂)(CO)_x] (ML_n = MnCp', x = 4, **144**; ML_n = W, x = 7, 144a). In turn, complexation with an excess of the 16-electron metal fragments $[MnCp'(CO)_2(THF)]$ (Cp' = C₅H₄Me) led to the formation of tetranuclear diphosphenyl-bridged [Mn₂Mo₂Cp₂Cp'₂(µ- PCy_2 (μ - κ^2 : κ^1 : κ^1 - P_2Me)(CO)₆] (**145**) along with a small amount of the trinuclear derivative 143 and the "PMe"-extruded derivative 144. Compound 145 also proved to be unstable, forming the trinuclear complex 144 overtime. Similarly, when 139 was treated with 3.5 eq of [W(CO)₅(THF)], it afforded both the tetranuclear derivatives $[Mo_2W_2Cp_2(\mu-PCy_2)(\mu,\kappa^2:\kappa^1:\kappa^1:\kappa^1-P_2Me)(CO)_{12}]$ (146) and

 $[Mo_2W_2Cp_2(\mu_3-P)(\mu-PCy_2)(\mu_3-PMe)(CO)_{10}]$ (147) in 3:2 ratio, along with small amounts of 144a (Scheme 46). Complexes 145 and 146 feature a trigonal-planar phosphide ligand, bridging the two Mo centers in addition to the exocyclic ML_n moiety. Notably, the short Mo-Mo distance (2.749(1) Å in 145 and 146 is consistent with a double Mo=Mo bond, as also confirmed by DFT calculations. Fig. 43 depicts the X-ray structures of 141, 141a and 144a.

Lastly, the reaction of **139** with metal carbonyl dimers $[M_2(CO)_X]$ where M = Mn, Re, Co; x = 10, 8, was investigated and led invariably to tetranuclear clusters, via the cleavage of the diphosphenyl moiety and incorporation of a partially decarbony-lated dimetal unit. In particular, the reaction of **139** with $[Co_2(-CO)_8]$ occurs at room temperature giving the 64-electron tetranuclear cluster $[Co_2Mo_2Cp_2(\mu-PCy_2)(\mu_4-PMe)(\mu-CO)(CO)_6]$ **147**, including the formation of two new Mo-Co bonds. As shown in Fig. 44, the molecular structure of **147** displays a square planar Co_2Mo_2 core symmetrically capped by a naked phosphide and a

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Scheme 43. Synthesis of 1D organometallic-organic polymers 123–126 by coordination of the flexible linkers 119, 120, 121 and 122 to the dimer 117. [Mo] = [CpMo(CO)₂].



Fig. 37. Structure of the dicationic repeating unit of a) 123, b) 124, c) 125, d) 126 in the solid state. Hydrogen atoms, Cp and CO ligands on Mo are omitted for clarity; adapted from Ref. [146].

methylphosphinidene ligand giving an overall octahedral M_4P_2 structure, with PCy_2 and CO ligands bridging two Mo centers and two Co centers, respectively.

The dimolybdenum agostic methyl-bridged complex [Mo₂Cp₂(- μ , κ^1 : η^2 -CH₃)(μ ,PtBu₂)(μ ,CO)] reacts with P₄ affording the blue triphosphorus complex [Mo₂Cp₂(μ , η^3 : η^3 -P₃)(μ ,PtBu₂)] via decarbonilation and methylphosphinidene elimination [156]. XRD analysis of [Mo₂Cp₂(μ , η^3 : η^3 -P₃)(μ ,PtBu₂)]revealed that the P₃ chain is symmetrically coordinated (P-P = 2.15 Å) to the two Mo centers, which in turn are triply bound to each other (Mo-Mo = 2.622 Å). Noteworthy, the internal P atom appears to be extremely shielded

as it resonates at the lowest ^{31}P chemical shift recorded to date (δ = -625.5 ppm).

Roesky and co-workers [157] investigated the reaction between the neutral complex [{CpMo(CO)₂}₂(μ_2 , η^2 -P₂)] (110) [158], and lanthanide complexes of the type [Cp*₂Ln(THF)₂] (Ln = Sm (24a), Yb (148), yielding different polyphosphide species (see Scheme 47). In particular, the reaction of 110 with 148 gave product 149, featuring a central {P=P}²⁻ ligand, end-on coordinated to two {CpMo (CO)₂}₂ fragments. The reaction of 110 with samarocene complex, 24a led to compound 150 as the major product (see Fig. 45 for Xray structure) in addition to small amounts of 151 and 152, which



Fig. 38. X-ray structures of 128. Hydrogen atoms are omitted for clarity; adapted from Ref. [148].

display a *cyclo*- P_4 and a *cyclo*- P_5 core units, respectively (see Scheme 47).

2.4.3. Tungsten

Ruiz and co-workers [159] applied the same functionalization strategy as seen with **137** (see Section 2.4.2) to tungsten by treating the analogous ditungsten complex Na[W₂Cp₂(μ ,PCy₂)(μ ,CO)₂] (**153**) with P₄ at 60 °C in THF. The reaction afforded the novel diphosphorus-bridged ion pair Na[W₂Cp₂(μ , κ^2 : κ^2 -P₂)(μ ,PCy₂) (CO)₂]⁻ (**154**), although it could not be isolated. To confirm the identify of **154**, it was treated with Mel affording the more stable methyldiphosphenyl complex [W₂Cp₂(μ , κ^2 : κ^2 -P₂)(μ ,PCy₂)(CO)₂]⁻ (**155**, Scheme 48).

The IR spectrum of **153** revealed four C-O stretching bands deriving from an equilibrium between the contact (1824 and 1695 cm⁻¹) and solvent separated (ν = 1824 and 1757 cm⁻¹) ion-pairs at room temperature. The ³¹P NMR analysis showed only resonances for the PCy₂ ligand and no signals were present for the κ^2 : κ^2 coordinated P₂ moiety. In turn, the IR spectrum of **155** showed two C-O absorbances (ν = 1860 and 1837 cm⁻¹) which are consistent with the distorted transoid conformation of the

 $M_2(CO)_2$ oscillator. The bridging methyldiphosphenyl ligand displayed two multiplets at δ = -166.0 ppm for (PMe) and δ = -390.1 ppm for (P) with large mutual coupling constants (${}^1J_{PP}$ = 467 Hz), which is common for κ^1 diphosphene complexes [160]. This is consistent with the short P-P bond (2.085 Å) found in the corresponding Mo complex **139**.

In an attempt to further study the reactivity of low-valent ETMphosphide complexes and to explore the possibility of the complete functionalization of the phosphide ligand, Cummins and coworkers [161] carried out the reduction of the precursor *mer*-Cl₃W(ODipp)₃ (**156**) with Na/Hg amalgam in the presence of P₄ in toluene, affording the neutral aryloxide complex (η^3 -P₃)W (ODipp)₃ (**157**) in 10% yield (Scheme 49). Compound **157** is valence-isoelectronic with [(η^3 -P₃)Nb(ODipp)₃]⁻ **65** (see Section 2.3.2.). The purpose was to assess the influence of the ionic charge of the complex on the transferability of the *cyclo*-P₃ ligand to main-group acceptors.

The ³¹P NMR spectrum of **157** displayed a shielded signal at δ = -156 ppm with ¹⁸³W satellites (¹J_{183W-31P} = 9 Hz). XRD analysis of **157** (see X-ray structure in Fig. 46a), showed shorter W-P and P-P distances (2.439–2.447 Å and 2.174–2.180 Å respectively) compared to **65** attributed to the smaller covalent radius of W with respect to Nb [58]. **157** proved to be remarkably less nucleophilic compared to the anionic **65**. The reaction of **157** with either PCl₃ or AsCl₃ in THF resulted in the displacement of the aryloxide by the chloride with no evidence of *cyclo*-P₃ transfer to the electrophile, affording (η^3 -P₃)W(Cl)(ODipp)₂(THF) (**158**), (see Fig. 46b for X-ray structure). This differs significantly to the reaction of **65** with AsCl₃, which forms AsP₃, suggesting that salt elimination aids in the formation of EP₃ species (E = As or P).

2.5. Group 7 metals

2.5.1. Manganese

The Scheer group reported the preparation of Mn complexes bearing intact P_4 tetrahedra as ligands, in the form of both mononuclear and dinuclear complexes [162]. Remarkably, these species represent the first examples of Mn complexes bearing E_x ligands (E = pnictogen). Following some unsuccessful attempts aimed at the activation of P_4 using cymantrene-derived fragments



Scheme 44. Reaction of 127 with Ag(TEF), Cu(TEF) and Tl(FAI) is highly dependent on the reaction media.



Fig. 39. Solid state structures of a) the cation of 131, and b) the cation of 132. Hydrogen atoms are omitted for clarity; adapted from Ref. [148].



Fig. 40. Molecular structure of the cation of **133**. Hydrogen atoms are omitted for clarity; adapted from Ref. [148].

such as $[Cp^{R}Mn(CO)_3]$ (R = H, Me), $[Cp^{BIG}Mn(CO)_2(THF)]$ (**159**, $Cp^{BIG} = C_5(C_6H_4^nBu)_5)$ was prepared, which possesses superior Lewis acidity and therefore is expected to react with P₄ upon displacement of the weak THF ligand. The reaction of **159** with 5 equivalents of P₄ gave the mononuclear complex $[Cp^{BIG}Mn(CO)_2(\kappa^1-P_4)]$ (**160**), which features an intact P₄ molecule end-on coordinated to Mn (see Scheme 50).

The molecular structure of complex **160** is shown in Fig. 47**a**. The bonds between the apical (P1) and the basal (P2, P3, P4) atoms (2.145–2.165 Å) are about 7 pm shorter than the basal P-P bonds (2.211–2.232 Å), as previously observed in related κ^1 -P₄ complexes of Group 6 and 8 [163–169]. The ³¹P{¹H} NMR spectrum of **160** in toluene *d*₈, features a quartet at δ = -305.7 ppm and a doublet at δ = -490.0 ppm with a ¹J_{PP} = 224 Hz, corresponding to the apical P1 atom and the three basal atoms, respectively.

When compound **159** was reacted with<5 equivalents of P₄, the dinuclear complex $[{Cp^{BIG}Mn(CO)_2}_2(\mu,\kappa^1:\kappa^1-P_4)]$ (161) was formed alongside 160. Complex 161 can be obtained in pure form either by reacting 159 with 0.5 equivalents of P₄ or reacting 160 with 1 equivalent of 159 (see Scheme 50). A remarkable property of 160 and 161 is their stability both in the solid state and in solution at room temperature, a feature not observed before for neutral P_4 complexes. Complex **161** (see Fig. 47b) features an intact P_4 tetrahedron shared by two {Cp^{BIG}Mn(CO)₂} fragments in an endon coordination. Notably, the Mn-P distance in complexes 160 and **161** is identical within the experimental error, indicating that second coordination is as strong as the first one. In agreement with the X-ray structural analysis, ³¹P{¹H} NMR measurements carried out at 193 K showed an A₂X₂ spin system, with two triplets at δ = -478.7 ppm and -250.2 (¹J_{PP} = 159 Hz), assigned to free and Mn-coordinated P atoms respectively. Upon heating to room temperature, these signals broaden as a result of the tumbling of the P₄ unit. Thus, the P₄ tetrahedron can act as a ligand binding metal fragments either in an end-on or a side-on fashion. In both cases



Fig. 41. Section of the 2D polymeric network 135. [CpMo] units and hydrogen atoms are omitted for clarity; adapted from Ref. [149].

the P₄ molecule is kept 'intact' and its reactivity is influenced by the coordinated metal fragments.

Scheer and co-workers succeeded in the synthesis of a rare heterocubane cage [{CpMn}₄(μ_3 ,P)₄] (**162**), isolated via the thermolysis of $[CpMn(\eta^6-CHT)]$ (CHT = cycloheptatriene) and P₄ at 205 °C in 1,3-diisopropylbenzene (DIB), see Scheme 51 [170]. Using a stoichiometric amount of P₄, reddish-brown needle-like crystals of 162 could be obtained with a yield of 73%. X-ray diffraction analysis revealed a highly symmetric geometry (see Fig. 48a), in which the Mn atoms sit at the vertices of a tetrahedron, whose faces are capped by the P atoms, resulting in a heterocubane Mn₄P₄ structure. DFT calculations revealed Mn-Mn and Mn-P bond order of 0.33 and 0.92 respectively. Considering the existence of three Mn-Mn bonds of order 1/3 for each Mn atom, the number of its valence electrons can be calculated classically to be 18. The high symmetry of 162 is also evidenced by NMR spectroscopic analysis. Both the ¹H and ¹³C{¹H} NMR spectra of **162** show only one singlet at δ = 4.58 ppm and δ = 82.7 ppm respectively, which are typical for the cyclopentadiene ring. The ³¹P{¹H} NMR spectrum of **162** shows an extremely low-field shifted singlet at δ = 1079.8 ppm. Bridging phosphide ligands are expected to be available for further coordination. The availability of four naked P atoms in 162 should allow for its use as a building block in coordination chemistry. Indeed, the reaction of 162 with Cu(I) salts, afforded 1D coordination polymers $[{(CpMn)_4(\mu_3,P)_4}(CuX)]n (X = Cl; Br)$ with a zigzag-like chain structure. As a proof of concept, complex 162 was reacted with $[CpMn(CO)_2(THF)]$, yielding $[{(CpMn)_4(\mu_3,P)_4}(CpMn(CO)_2]_n]$ species, with n = 1, 2, 3, 4 depending on the amount of [CpMn(CO)₂(-THF)] used. The molecular structure of $[{CpMn}_4(\mu_3, P)_4{CpMn}]$ $(CO)_{2}_{n}$ (163, n = 1) is shown in Fig. 48b. As revealed by X-ray



Scheme 45. Activation of P₄ by 137 affords 138 which reacts with electrophilic species, giving complexes 139 and 140.



Fig. 42. X-ray structures of a) 139 and b) 140. The different bond angle P2-P3-X (X = C, Sn) is clearly visible. Hydrogen atoms are omitted for clarity; adapted from Ref. [150].



Scheme 46. Reaction of 139 with mononuclear carbonyl fragments ML_n resulting in the formation of tetra- and trinuclear complexes.


Fig. 43. X-ray structure of a) 141 and b) 141a, featuring a cleaved diphosphorus bond. X-ray structure of complex c) 144a reveals a [PW(CO)₅] ligand bridging the two Mo centers, evidencing the absence of the extruded "PMe" fragment. Hydrogen atoms are omitted for clarity; adapted from Ref. [154].



Fig. 44. Molecular structure of **147.** Hydrogen atoms and Cy groups (except the C¹ atoms) are omitted for clarity, adapted from Ref. [155].

studies, an increase in the number of attached {CpMn(CO)₂} fragments lead to a volume expansion of the Mn_4P_4 unit, characterized by a progressive elongation of Mn-Mn and Mn-P bonds (from av. Mn–P 2.235 Å and av. Mn–Mn 2.713 Å in **162** to av. Mn–P 2.258 Å and av. Mn–Mn 2.757 Å when n = 4).

Scheer and co-workers [171] devised the synthesis of unprecedented Mn triple decker complexes featuring central P_n ligands, using the precursor [Cp^{BIG}Mn(CHT)] (**164**, Cp^{BIG} = C₅(4-*n*BuC₆H₄)₅, CHT = cycloheptatriene) and reacting it with P₄. When **164** is heated in toluene with excess P₄, the CHT ligand is eliminated and complexes [{Cp^{BIG}Mn}₂(μ 2, η ⁵: η ⁵-P₅)] (**165**) and [{Cp^{BIG}Mn}₂(- μ 2, η ²: η ²-P₂)₂] (**166**) are formed (see Scheme 52).

Being chemically alike, **165** and **166** could not be separated by TLC and even co-crystallize at the same crystallographic positions. This occurrence causes a triple decker complex with a disordered middle deck. Upon careful analysis of the atom occupancies in the central deck, a 7:3 mixture of **165** and **166** was ascertained. The structure of **165**, which features a *cyclo*-P₅ ligand, is shown in Fig. 49.

The P–P bond lengths in **165** have an average value of 2.20 Å, in good agreement with other known triple-decker complexes [172] containing a *cyclo*-P₅ middle-deck (typically 2.15 Å – 2.28 Å). Nota-

bly, the Mn-Mn distance is 2.795 Å, which is in the usual range for Mn-Mn single bonds. DFT calculations of **165** indicated a Mn-Mn Wiberg bond index of 0.6, confirming a bond between the two metal centers. Compound **166** features two coplanar P₂ units, each one bridging in a side-on fashion to the two Mn atoms. The ³¹P{¹H} NMR spectrum of **165** revealed only one broad singlet at δ = 396.9 ppm. This feature was explained by DFT calculations, which showed **165** is paramagnetic with a doublet ground state. EPR measurements were then carried out at 77 K, showing a well resolved anisotropic multiplet. This arises from the coupling of the unpaired electron with two Mn centers having spin of 5/2 each. The g_{iso} value was identified to be 1.923 (simulation), slightly smaller than the value of 1.957 calculated via DFT.

2.5.2. Rhenium

Within the last decade a great deal of research related to the activation of P₄ has focused on employing reactive low-valent and coordinatively unsaturated TM complexes with the aim to promote multi-electron reductive couplings between the two species. In 2019, Arnold and co-workers [173], successfully synthesized the neutral complex $\text{Re}(\eta^5-\text{Cp})(\text{BDI})$ (BDI = N,N'-bis(2,6-diisopropy) phenyl)-3,5-dimethyl- β -diketiminate) (**168**), a rare example of an isolable unsaturated Re(II)-d⁵ complex, obtained in good yields (76%) from the oxidation of the anionic Re(I) precursor Na[Re(η^{5} -Cp)(BDI)] (167) with AgOTf (Scheme 53). Due to the paramagnetic and reduced nature of 168, confirmed by the measure of its magnetic moment (μ_B = 1.67) with the Evans method, 168 was expected to be an excellent candidate for the reductive activation of P_4 . Upon addition of P_4 to **168** in toluene at room temperature the novel Re(V) complex $(\eta^3$ -cyclo-P₃)Re $(\eta^5$ -Cp)(BDI) (169) was formed in good yield (64%) [174]. ³¹P NMR analysis of 169 displayed two highly shielded and well resolved resonances at $\delta = -$ 206.9 ppm and a δ = -297.3 ppm (a triplet and a doublet respectively with a ${}^{1}J_{PP}$ = 285 Hz), consistent with a formal trianionic [*cy* $clo-P_3$]³⁻ ligand. The molecular structure of **169** (see Fig. 50) revealed P-P bond length in the range from 2.127 to 2.144 Å, standard for cyclo-P₃ moieties. The redox properties of 169 were investigated through cyclovoltammetry, which indicated an oxidation peak at -0.25 V (E_{1/2} vs Fc/Fc+, THF, 0.3 M [nBu₄N][PF₆]), associ-



Scheme 47. Synthesis of compounds 149, 150, 151 and 152.



Fig. 45. X-ray structure of 150. Hydrogen atoms are omitted for clarity; adapted from Ref. [157].

ated to the redox couple Re(V)/Re(VI). For this reason, **169** was treated with AgOTf, which afforded the Re(VI) complex [$(\eta^3$ -cyclo-P₃)Re(η^5 -Cp)(BDI)][OTf] (**170**, Scheme 53).

Interestingly, there are no remarkable differences in the Re-P and P-P bond distances between **169** and **170**. The oxidation seemed to affect only the Re-N distance of one N atom of the BDI ligand (Re-N1 gets 0.12 Å smaller going from **169** to **170**). The electronic structure of **169** was investigated by EPR and ENDOR (Electron-Nuclear Double Resonance Spectroscopy), which, in agreement with DFT calculations, indicated that the spin density of this complex is partially localized on the Re center but resides mainly on the BDI ligand, with very little spin density on the *cyclo*-P₃ fragment. This explains the observed change of Re-N distance between **169** and **170**.

3. Coordination chemistry of white phosphorus by late transition metals

The coordination chemistry of white phosphorus by late transition metals has attracted considerable attention for almost five decades now, ever since the first examples of rhodium, iridium, or nickel complexes featuring activated or intact P_4 or P_4 -derived *cyclo*- P_3 ligands were reported by the groups of Ginsberg [175,176] and Sacconi [69,177] in the early seventies. The range and topology of the reported P_4 -derived phosphorus moieties coordinated to late TM has, since then, grown significantly. This chemistry was comprehensively reviewed in the early 2010 by the group of Peruzzini [7], and some updates have also been included in other



Scheme 48. Reaction of the dinuclear tungsten complex 153 with P₄ and subsequent alkylation with MeI yielding the complex 155.



Scheme 49. Reaction of complex 156 with P₄ afforded the neutral complex 157. Unfortunately, any attempt of further reaction with PCl₃ or AsCl₃ gave complex 158 bearing an intact *cyclo*-P₃ moiety.



Fig. 46. X-ray structure of a) 157, and b) 158. Hydrogen atoms are omitted for clarity; adapted from Ref. [161].



Scheme 50. Synthesis of neutral tetrahedro-P₄ complexes 160 and 161.

reviews [42,43]. Further contemporary (2011–2020) insights in this field are presented herein, with subsections according to the metal groups. In the period under review, significant advances have been made on understanding the activation of the P_4 tetrahedron by late TM, and a variety of phosphorus frameworks have been reported. The reactivity of P_4 towards Earth-abundant metals, such as iron, cobalt, and nickel, has been particularly important, and striking examples have also been reported, for example, with ruthenium, copper, and gold. For the description of the transformations of white phosphorus by late transition-metal complexes, in general, the contributions have been organized according to the degree of degradation/aggregation of P_4 in each case.

3.1. Group 8 metals

Within the late transition metals, group 8 P₄-derived complexes, mainly of iron and ruthenium, account for most of the contributions in the period under review. Regarding iron, remarkable



Fig. 47. Molecular structure of compounds a) 160b) and b) 161. 4-n-butylphenyl groups on the Cp^{BIG} ligands have been omitted for clarity; adapted from Ref. [162].



Scheme 51. Synthesis of heterocubane Mn₄P₄ (162).

examples include, but not limited to, the synthesis of butterfly- P_4 complexes which have been used as metalloligands to develop a rich coordination chemistry of the tetraphosphorus moiety. Furthermore, different groups have achieved the activation of white phosphorus at iron to form *cyclo*- P_4 or *catena*- P_4 motifs, as well as a variety of aggregates. Whereas the chemistry of ruthenium- P_4 complexes is largely dominated by the coordination of the intact tetrahedron and its hydrolytic functionalization, which contrasts with the stability of the free molecule in water. Osmium complexes of white phosphorus are the least represented within this group. The only example is a study of the behavior in solution, on an NMR timescale, of a compound featuring the intact P_4 tetrahedron in its coordination sphere.

3.1.1. Iron

During the period 2011–2020, an early example of the ironmediated transformation of P_4 was the catalytic synthesis of phosphates described by Armstrong and Kilian [54]. In this system, P_4 is

oxidized to PI₃ and then reacted with phenols under aerobic conditions. While a wide variety of transition-metal compounds were initially evaluated for the reaction, the use of iron(III)diketonates (171, see Scheme 54, top) as catalysts were found to be more efficient, and it was determined that the addition of iodine was required, to generate substoichiometric amounts of PI₃. Remarkably, no acid waste is produced, air is used as a green oxidant, and full conversion to the phosphates with high selectivity was achieved without using chlorine. However, a minimum of 25 mol% catalyst loading was required. It was determined that this high amount of iron catalyst is necessary, since it is involved in at least two key transformations: it favors the formation of the phosphate from O=PI(OPh)₂ over the possible hydrolysis, and increases the rate of reoxidation of HI to form iodine. From the mechanistic investigations performed in the case of phenol using Fe(acac)₃ (**171a**, acac = 2.4-pentanedionate) as a catalyst (Scheme 54, bottom), it was concluded that initial oxidation of P_4 by iodine to form Pl₃ is crucial to maintain the catalytic reaction. The nucleophilic substitution of O=PI(OPh)₂ with phenol to yield the final O=P (OPh)₃ was found to be the rate-determining step. This system represents a proof of concept of the production of phosphates from P₄ without using PCl₃ as intermediate.

In the period under revision, to the best of our knowledge, only a few examples of iron complexes featuring the intact P₄ tetrahedron as ligands were reported [178,179]. Peruzzini and coworkers have long studied the chemistry of white phosphorus towards transition metal precursors [7,42]. In 2012, this group investigated the fluxional processes in solution of [L_nM(η^1 -P₄)] complexes, featuring intact P₄ ligands [178]. Generally, the ³¹P {¹H} NMR spectra of such compounds showcase an MQ₃ spin system, in which a doublet and a quartet are observed for the "free" and the metalated P-atoms on the coordinated P₄-tetrahedron, respectively. However, the broadened ³¹P signals for the tetraphos-



Fig. 48. Molecular structure of a) 162 and b) 163, highlighting the Mn₄P₄ core. Hydrogen atoms are omitted for clarity; adapted from Ref. [170].



Scheme 52. Synthesis of compounds 165 and 166 from thermolytic activation of P₄ with 164.

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Fig. 49. Molecular structure of 165. Hydrogen atoms are omitted for clarity; adapted from Ref. [171].

phorus ligand were indicative of fluxional processes, for which a tumbling motion of the P₄-ligand in solution, corresponding to the dynamic $\eta^1 \rightarrow \eta^2 \rightarrow \eta^1$ change in its coordination mode, was proposed. To investigate the nature of such dynamic processes, a series of rhenium [180], iron, ruthenium (see Section 3.1.2) and osmium (see Section 3.1.3) complexes featuring the intact P₄-cage, as well as three bimetallic RuPt species bearing a bridging butterfly-P₄ scaffold (vide infra) were studied in solution, by variable-temperature (VT) NMR and ³¹P{¹H} exchange spectroscopy (EXSY). The iron complex $[Cp^*Fe(dppe)(\eta^1-P_4)]^+$ (**172a**, dppe = 1,2-bis(diphenylphosphino)ethane (Ph₂PCH₂CH₂PPh₂), Scheme 55) was obtained after reaction of [Cp*Fe(dppe)Cl] (173) with a chloride scavenger, followed by addition of P₄. Such species is analogous to the previously reported monometallic complex $[CpFe(dppe)(\eta^1-P_4)]^+$ (**172b**, Scheme 55), by the same group [169]. For compound **172a** in solution, the main dynamic process observed at room temperature was the dissociation of the tetraphosphorus ligand. In the ³¹P{¹H} EXSY experiment at higher temperatures (T = 45 $^{\circ}$ C), weak exchange peaks attributed either to thermal activation of the tumbling motion of the coordinated P₄ molecule or multiple P₄ dissociation/coordination processes, were observed. Due to the decomposition of the complex at temperatures higher than 45 °C, it was not possible to determine the activation parameters of the dynamic process in solution [178].

Recently, the group of Krossing reported complexes featuring intact P₄-cages coordinated to iron centers [179]. Unlike the previ-



Fig. 50. X-ray structure of 169. Hydrogen atoms are omitted for clarity; adapted from Ref. [174].

ous examples [169,178], in which electron-rich iron centers coordinate the tetrahedron, the work by Krossing [179] describes the use of the electron-poor fragments $[CpFe(CO)_2]^+$ and [CpFe(CO)] (PPh_3) ⁺ and the effect exerted by them on the bounded P₄-cage. Treatment of the iron precursors [CpFe(CO)(L)Br] (174, L = CO, PPh₃) with a strong halide scavenger in the presence of white phosphorus (2 equiv.), afforded the η^1 -P₄ containing cationic complexes $[CpFe(CO)(L)(\eta^{1}-P_{4})]^{+}$ (L = CO (175), PPh₃ (176), Scheme 55). An excess of P₄ (2 equiv.) was necessary to prevent the formation of the cationic dimer, $[{CpFe(CO)_2}_2(\mu-Br)]^+$. Their molecular structures (see Fig. 51) demonstrated an important electronic influence, evidenced by the Fe-P bond distance in both complexes: the replacement of the CO ligand with PPh₃ shortened the bond length from 2.209(2) Å in 175 to 2.178(2) Å in 176. Furthermore, the P-P bond lengths within the cage in both compounds are remarkably shorter (2.144 Å in 175 and 2.133 Å in 176, average values) than in the free white phosphorus molecule (2.21 Å), which is reflected also in a strong blueshift in the Raman spectra of the complexes. The phenomenon was attributed to the stronger back donation from the iron center to the P₄ ligand in **176** with respect to **175**. On the NMR timescale, a slow exchange of the phosphorus atoms for the P₄ tetrahedron in both complexes was observed, and a correlation between the electron density at the metal and the activation energy of the exchange process was established. Theoretical studies, along with the experimental results indicated that the coordination of the P₄-cage to electron-deficient iron centers, as in **175** and **176**, gives rise to an umpolung in the Fe-P_{cage} bond



Scheme 53. Activation of P₄ with neutral Re complex 168 and subsequent oxidation with AgOTf.



Scheme 54. General reaction for the direct synthesis of phosphates from P₄ mediated by iron(III)-diketonates (top), and proposed catalytic cycle (bottom) [54].



Scheme 55. Monometallic iron complexes featuring an intact P₄ ligand by the groups of Peruzzini (top) [169,178] and Krossing (bottom) [179].

and, thus, such species can be described as phosphonium complexes [179].

The activation of the white phosphorus tetrahedron can lead either to the formation of other P₄ moieties or to its fragmentation/reaggregation. Extensive studies by the group of Scheer have led to a series of P_n ligands, with the most popular being the *cyclo*-P₅ ring that replaces a Cp ligand in $[Cp^RFe(\eta^5-P_5)]$ (**25**). These have been largely used as building blocks in coordination and supramolecular chemistry, in particular after P_n aggregation [129,181–187]. Several groups have agreed that, among the common P₄-activation scaffolds, the first step in the degradation at transition metals is the formation of the tetraphosphabicyclo[1.1. 0]butane unit, usually known as a butterfly-P₄ ligand. This framework is obtained after cleavage of one of the P-P bonds in the P₄ tetrahedron.

The reactivity of white phosphorus with the family of iron complexes $[Cp^{R}Fe(CO)_{2}]_{2}$ (**177**) has been a matter of interest for over two decades by different research groups [188–194]. Several P₄ activation, fragmentation and reaggregation products, obtained after photolysis, thermolysis and, more recently under ambient conditions, have been showcased. After seminal work by Scherer and co-workers [7,188,189], effort has been devoted to the study of analogous systems which could enable the activation of white



Fig. 51. X-ray structures of complexes **175** (left) and **176** (right), featuring intact P₄ tetrahedron ligands; adapted from Ref. [179].

phosphorus under milder conditions and in more selective pathways. For example, Scheer and co-workers reported the more bulky analogue, $[Cp^{BIG}Fe(CO)_2]_2$ ($Cp^{BIG} = pentakis(4-n-1)$ butylphenyl)cyclopentadienyl, 177a) which activates small molecules such as P_4 , As_4 , P_4S_3/P_4Se_3 and CS_2 at room temperature [190]. For the E_4 tetrahedra (E = P, As), the reaction proceeds selectively at room temperature, yielding the product of homolytic cleavage (through 17 valence electron (VE) metal-centered radicals) in almost quantitative yield (Scheme 56). The molecular structure of the iron-butterfly-P₄ complex [{Cp^{BIG}Fe(CO)₂}₂(μ , κ ^{1:1}-P₄)] (178a) was determined (Fig. 52). The P-P bond lengths between the "wing tip" P-atoms and the bridgehead ones are within the range of single bonds (2.209 Å and 2.234 Å), and are similar to those in the P₄ tetrahedron, while the bond between the bridgehead atoms is shortened (2.172 Å). Regardless of the stoichiometry or reaction conditions used, it was not possible to further activate the P₄ unit. Heating 178a in solution led to the formation of $[Cp^{BIG}Fe(\eta^5-P_5)]$ (**25b**) and $[(Cp^{BIG}Fe)_2(\mu,\eta^4:\eta^4-P_4)]$ (179a). $[Cp^{R}Fe(\eta^{5}-P_{5})]$ (25) and $[(Cp^{BIG}Fe)_{2}(\mu,\eta^{4}:\eta^{4}-P_{4})]$ (179) had been previously obtained after thermolysis of 177 in the presence of P₄ at elevated temperatures (T > 190 °C) (Cp^R = Cp^{BIG} [191]; Cp^{'''} = 1,2,4 tris(tert-butyl)cyclopentadienyl (177b) [188]). Addition of P₄ to **178a** in solution, before heating, leads to the formation of 25b and 179a in the same ratio and yields as before [191]. This indicates that $[{Cp^{R}Fe(CO)_{2}}_{2}(\mu,\kappa^{1}:\kappa^{1}-P_{4})]$ (178) is the initial product of P₄ activation by Cp^RFe fragments. Furthermore, if **178b** is reacted with yellow arsenic (As₄), an analogous process takes place, yielding the mixed P_nAs_m ligand analogues [Cp^{'''}Fe(η^5 - P_nAs_{5-n}] (25d) and [{Cp'''Fe}₂($\mu,\eta^4:\eta^4-P_nAs_{4-n}$)] (179c), which were the largest mixed E_n motifs synthesized at the time [192].

The procedure described for **178a** [190], can be extended to the synthesis of other L_nM-butterfly-E₄ [L_nM = Cp*Cr(CO)₃, Cp^{'''}Fe (CO)₂; E = P, As)] complexes featuring different Cp^R ligands [193], including the previously reported [{Cp^{'''}Fe(CO)₂}₂(μ , κ^1 : κ^1 -P₄)] (Cp^{'''} = 1,2,4-tri-*tert*-butylcyclopentadienyl, **178b**, Scheme 56) [188]. Irradiation of **178a** with UV light (see Scheme 57**a**) promotes the release of two CO ligands, with selective formation of [{Cp^{BIG}Fe}{Cp^{BIG}Fe(CO)₂}(μ , η^4 : η^1 -P₄)] (**180**) [194]. Similar photolytic reactions were carried out on the butterfly-As₄ analogues of **178a** and **178b**, causing rearrangement or reaggregation to form a variety of structural As_n motifs (*n* = 4–6) in the case of the Cp^{BIG} derivatives [194], and an As₈ *cuneane* tetrametallic structure for the Cp^{'''} analogue [193].

Further investigations on the reactivity of the iron butterfly compound **178b** include its protonation (Scheme 57**a**) using [(Et₂-O)_nH]X (X = BF₄⁻, n = 1; [Al(OC(CF₃)₃)₄]⁻, n = 2) [195]. According to variable temperature (VT) ³¹P NMR and computational studies, these reactions yielded the compounds [{Cp^{''}Fe(CO)₂}₂ ($\mu,\kappa^1:\kappa^1-P_4$)H]X (**181a,b**, X = BF₄⁻; [Al(OC(CF₃)₃)₄]⁻) in which the protonation has occurred at a wing tip P atom. The asymmetrical



Scheme 56. Activation of P_4 at $[Cp^RFe(CO)_2]_2$ (**177**, $Cp^R = Cp^{BIG}$; $Cp^{\prime\prime\prime}$).

substitution of one P atom in the butterfly-P₄ moiety was confirmed via X-ray crystallography, although a good refinement was not possible due to the inherent poor quality of the crystals. DFT calculations also ruled out the possibility of protonation at a bridgehead P atom.

Scheer and co-workers have developed a rich array of coordination chemistry and reactivity for **178b**, taking advantage of its potential as a bidentate ligand (see Scheme 57**b**). Computational studies (DFT methods at the BP86/def-SVP level of theory) [195] performed on the model complex [{CpFe(CO)₂}₂($\mu,\kappa^1:\kappa^1-P_4$)] (**178c**) revealed that the position of the filled molecular orbitals should permit a substantial interaction between the wing tip P atoms on the butterfly moiety and a Lewis acid. Furthermore, the LUMO (lowest unoccupied molecular orbital) is located precisely between these wing tip P atoms and, thus, could allow back-

bonding from the Lewis acid to the butterfly complex. Depending on the stoichiometry used, the reaction of **178b** with [Cu(MeCN)₄] BF₄ (**182**) led to the formation of two different copper complexes containing **178b** as a metalloligand (Scheme 57b): the monoadduct $[{Cp'''Fe(CO)_2}_2(\mu^3,\kappa^1:\kappa^1,\kappa^2-P_4){Cu(MeCN)}]BF_4$ (183, ratio 178b: Cu⁺ = 1:1) and the spiro compound $[{(Cp'''Fe(CO)_2)_2(\mu^3,\kappa^1;\kappa^1,\kappa^2-\mu^3)_2(\mu^3,\kappa^1;\kappa^2,\kappa^2-\mu^3)_2(\mu^3,\kappa^1;\kappa^2,\kappa^2-\mu^3)_2(\mu^3,\kappa^2;\kappa^2,\kappa^2)_2(\mu^3,\kappa^2;\kappa^2,\kappa^2)_2(\mu^3,\kappa^2;\kappa^2,\kappa^2)_2(\mu^3,\kappa^2;\kappa^2,\kappa^2)_2(\mu^3,\kappa^2;\kappa^2,\kappa^2)_2(\mu^3,\kappa^2;\kappa^2,\kappa^2)_2(\mu^3,\kappa^2;\kappa^2,\kappa^2)_2(\mu^3,\kappa^2;\kappa^2)_2(\mu^3,\kappa^2;\kappa^2)_2(\mu^3,\kappa^2;\kappa^2)_2(\mu^3,\kappa^2;\kappa^2)_2(\mu^3,\kappa^2)_2$ P_4 }₂Cu]BF₄ (**184**, ratio **178b**:Cu⁺ = 2:1). The isolation of these complexes confirms that the butterfly-P₄ compound **178b** can coordinate unsaturated metal centers through the wing tip P atoms, and that their spatial proximity enables it to act as a bidentate ligand [195]. Such compounds were fully characterized, including single-crystal X-ray diffraction (see Fig. 53). The effect of the steric bulk of the metalloligand 178b becomes evident when the Cu-P bond lengths in both complexes are compared. The Cu-P bond distances in these cases are 2.3224(7) Å and 2.2891(6) Å for **183** and in the range of 2.3630(8) Å to 2.4268(8) Å for **184**. Thus, in the monoadduct **183**, the ligand is more tightly bound to the copper center than in the spiro compound 184. This is due to the reduced steric bulk in the former which enables a closer interaction with the copper metal center. Likewise, the bond distances in 184 are slightly longer than in complexes bearing bidentate phosphine ligands, such as $[(dppe)_2Cu]ClO_4$ (2.259(2) Å-2.301(2) Å; dppe = bis(diphenylphosphino)ethane) [196]. Additionally, in both complexes the Fe-P bond lengths are shorter than in the parent compound 178b (2.2928(6) Å and 2.2845(6) Å in 183 or 2.3059(8) Å to 2.3380(8) Å in 184 versus 2.348(2) Å and 2.355(2) Å in 178b [188,190]). Small bite angles towards copper were also observed in both cases in the coordinated metalloligand chelate (75.2° in **183** and 72.6° (average) in **184**) [195].

Additional coordination studies of the intact butterfly-P₄ motif were similarly carried out [197]. To investigate the outcome of the reaction with a different transition metal, a less electron rich moiety was selected (Scheme 57b). After reaction of **178b** with [FeBr₂·dme] (**185**, dme = dimethoxyethane), coordination of the butterfly-P₄ ligand through the wing tip P atoms to the FeBr₂ unit occurs, analogously to the previously observed copper(I) complexes **183** and **184**. In the compound obtained, [{Cp^{''}Fe(CO)₂}₂($\mu_{3},\kappa^{1}:\kappa^{1},\kappa^{2}-P_{4}$ }[FeBr₂]] (**186**), according to the molecular structure determined by XRD (Fig. 54a), the P_{wing-tip}-Fe-P_{wing-tip} bond angle is more acute (70.3°) in comparison to the angles observed for the copper complexes **183** and **184** (*vide infra*). Thus complex **186** is an example of the potential of metalloligand **178b** to act as a small bite angle chelate for different Lewis acids. Surprisingly, the analogous reaction of **178b** with the labile species [Fe(MeCN)₆][PF₆]₂



Fig. 52. Molecular structure of 178b; adapted from Ref. [190].

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(187) does not yield a monoadduct similar to 183, but the unprecedented homoleptic-like sandwich complex $[{(Cp'''Fe(CO)_2)_2(\mu_3, \eta^1)}]$ η^{1} : η^{4} -P₄)₂Fe|[PF₆]₂ (**188**) featuring 6π -electrons aromatic (planar) cyclo-P₄ (P₄R₂) ligands. These cyclo-P₄R₂ ligands are formed by isomerization, via P-P bond scission between the two bridgehead phosphorus atoms of the butterfly unit in 178b. Currently, such "carbon-free" all-phosphorus sandwich complexes are limited to a single further example: the decaphosphatitanocene dianion $[Ti(\eta^5$ -cyclo-P₅)₂]²⁻ reported by Ellis and co-workers [198]. The P-P bond distances in the *cyclo*- P_4R_2 ligands of **188** are in the range of 2.1406 Å to 2.1547 Å, which are between the values of typical P-P single (2.20–2.25 Å) and P–P double bonds (2.00–2.05 Å), as would correspond to such aromatic systems (see Fig. 54b for the molecular structure). Compound 188 can also be obtained upon reaction of **186** in the presence of **178b** and TIPF₆ (4 equivalents). However, **188** is not the sole product of the reaction. A mixture of compound **188** and the minor species $[{Cp''Fe(CO)_2}_2(\mu_3, n^1)]$ η :¹: η ⁴- P_4){Cp^{'''}Fe}]PF₆ (**189**), [{Cp^{'''}Fe(CO)₂}₂($\mu_3,\kappa^1:\kappa^1,\kappa^2-P_4$){Cp^{'''} Fe(CO)}]PF₆ (**190**) (Scheme 57**b**) and other side-products, which could not be characterized, was formed. 189 corresponds to an heteroleptic cyclo-P₄R₂ product, while **190** still bears the intact butterfly-P₄ scaffold. Compounds 189 and 190 were characterized

through single-crystal X ray diffraction (Fig. 54**c**-**d**, respectively). Complex **189** could be independently synthesized from **178b** in the presence of the dimer $[Cp^RFe(\mu,Br)]_2$ (**191a**, $Cp^R = Cp'''$) and TlPF₆ (3 equivalents). The authors highlighted that the stepwise transformation of the butterfly-P₄ motif is strongly dependent on the steric hindrance and electronic properties of the Fecompound used [197].

An isomer of the tetraphosphabutadiene compound **179b**, $[Cp'''Fe]_2(\mu,P_4)$ (**192**), featuring a severely distorted kite-like *cyclo*- P_4 unit, was synthesized from the iron-hydride complex $[Cp'''FeH_2]_2$ (**193**) and white phosphorus when heated to 100 °C for 10 min (Scheme 58) [199]. The authors highlighted that the path of activation of white phosphorus at a transition metal center with an odd number of electrons should proceed *via* scission of two P–P bonds of the P₄-cage, to yield a square-planar *cyclo*-P₄ motif, followed by a kite-like distortion, and final disproportionation of such a unit. It was proposed that complex **192** can be considered as a "snapshot" of the disproportionation process into P₁ and *cyclo*-P₃ units after activation of P₄ at Fe(I). Therefore, NMR characterization of **192** showed an A₂MX-spin system for the *cyclo*-P₄ motif consistent with the mentioned distortion, which persisted in the solid state, according to XRD. Two short (2.13 Å) and two long P–



Scheme 57. Reactivity of the iron-butterfly-P₄ complexes 178a,b. a) activation of the butterfly-P₄ scaffold, b) coordination chemistry of the butterfly-P₄ scaffold in 178b towards Lewis acids.



Fig. 53. X-ray structures of complexes a) 183 and b) 184, featuring the chelating metalloligand 178b; adapted from Ref. [195].

P distances (2.53 Å) were observed (Fig. 55 shows the molecular structure), with the longer ones in the range of values between single P–P bonds and a van-der-Waals contact. Heating a solution of **192** (75 °C, 7 days) afforded **179b** as major product, along with the phosphaallyl complex [{Cp^{$'''}Fe}_2(\mu,\eta^3:\eta^3-P_3)]$ (**194**) and **25d**, as minor product. Thus, the kite-like distorted complex **192** can be considered an intermediate in the formation of **179b** and as a source of P₁ fragments [199].</sup>

A butterfly-P₄ scaffold was also accessed via the activation of P₄ by Cp^R radicals, mediated by transition-metal species [200]. The one-step synthesis was described by the group of Scheer and gives rise to a variety of Cp₂^RP₄ organophosphorus compounds (**195**, Cp^R: $Cp^{BIG} = C_5(4-nBuC_6H_4)_5$, $Cp''' = 1,2,4-C_5H_2tBu_3$, Cp^* , $Cp^{4iPr} = C_5HiPr_4$). While attempting to synthesize the Cp^{BIG} analogue of a species previously assigned as $[Cp^{4iPr}Cu(\kappa^2-P_4)]$ (196) [201], the authors obtained yellow cube-shaped crystals. XRD analysis revealed that the crystals corresponded to the metal-free compound 195a, in which an edge of the P₄-cage has been cleaved, and two C-P bonds formed. The formation of {Cp^{BIG}} was confirmed through EPR measurements. Attempts to reproduce the reaction with other Cp^R compounds failed, this was attributed to the lack of stabilization of the alkyl substituted Cp^{R} radicals in comparison to those of Cp^{BIG} . The latter was posited to be a less reactive species, due to its high steric demand and mesomeric stabilization which prevents radical dimerization, allowing it to react with P₄. Since a crucial step in the formation of **195a** is the Cu(I)-mediated oxidation of $\{Cp^{BIG}\}^-$ to $\{Cp^{BIG}\}^-$, the use of a more flexible redox system was considered. Therefore, the reaction of $[Cp'''Fe(\mu,Br)]_2$ (**191a**) with P₄ (1 equiv.; Scheme 59a) afforded the carbon-substituted butterfly-P₄ compound

195b, along with the iron complex 179b and precipitated FeBr₂. However, after work up product 195b was only obtained in a low vield. The product distribution of the isomeric species, together with the presence of FeBr₂ provided information on the mechanistic pathway of the reaction and explains the low yield. It was proposed that in solution, 191a disproportionates into the complex fragments {Cp^{'''}Fe^I} and {Cp^{'''}Fe^{III}Br₂}. Two equivalents of the {Cp^{'''}Fe^I} fragment react with one P_4 molecule, thus forming **179b**, featuring a cisoid P_4^{2-} ligand. The two remaining 16 VE {Cp^{'''}Fe^{III}Br₂} fragments react with white phosphorus, presumably via initial coordination, enabling the transfer of radical {Cp^{'''}} species onto the P₄ molecule to afford product 195b, and reducing the Fe(III) center to Fe(II) (which precipitated as FeBr₂). Further modification of the system to generate the {Cp^{'''}Fe^{III}Br₂} fragments from NaCp^{'''} and FeBr₃ in situ and react them with P_4 (Scheme 59b), afforded compound **195b** in better yield without producing **179b**. The scope of this method could be extended to synthesize **195a**, as well as $Cp_{2}^{*}P_{4}$ (195c) and $Cp_{2}^{4iPr}P_{4}$ (195d), thus providing a general route for the transfer of carbon-centered radicals to P₄. Analysis of the molecular structures of compounds 195 revealed that all of them show an exoexo configuration. Careful insight into the NMR data of 195d also indicated that the chemical shifts observed for such compounds were nearly identical to the ones previously reported for the species assigned as $[Cp^{4iPr}Cu(\kappa^2-P_4)]$ (196) [201]. Furthermore, FD mass spectrometry analysis of the crude reaction mixture using the optimized method showed only the presence of the molecular ion peak of 195d (45%) and some phosphorus-free Cp^{4iPr}Fe complex fragments. It was concluded that the previous report [201] had mistakenly assigned compound 195d as the putative copper complex 196.

In 2017, the group of Lammertsma reported on the coordination chemistry of P₄ towards iron metalates, and their subsequent functionalization [202]. The reactivity of P₄ towards transition metal anions has been much less developed when compared with cations. To generate an anionic Fe- $(\eta^1 - P_4)$ adduct, a bulky nucleophilic metalate was reacted with P₄ and stabilized (trapped) by interaction with Lewis acids, in this case BAr_3 (Ar = C_6F_5 , Ph) (Scheme 60). The resulting Lewis acid-stabilized iron complexes $[Cp^*Fe(CO)_2(\eta^1-P_4)]^-$ (197) are the first examples of transition metal generated P₄-butterfly anions and show an unprecedented metalphosphido-borane interaction. Single-crystal X-ray diffraction of complex 197a (Fig. 56) confirmed the asymmetric substitution on the tetraphosphido moiety, caused by the coordination of the $[Cp^*Fe(CO)_2]$ ($Cp^* = C_5Me_5$) fragment on one side, and the borane on the other. The Fe-P coordination bond in **197a** is slightly shorter than for the previously observed analogous complex $[{Cp^{BIG}Fe(CO)_2}_2(\mu,\kappa^1:\kappa^1-P_4)]$ (**178a**) (2.3192 Å vs. 2.3397 Å in 178a) [190] whilst the P-B bond is longer (2.080 Å) than for organyl-substituted Li[Mes*P₄·B(C₆F₅)₃] (2.064 Å) butterfly-P₄ derivatives [203]. The anionic Lewis acid-stabilized butterfly-P₄ compounds 197 were reacted with [Me₃NH][BPh₄] to prove their utility as nucleophilic reagents, forming initially the intermediate wing-tip protonated species 198. Driven by the loss of the borane as the amine-borane adduct $Me_3N\cdot BAr_3$ (Ar = Ph, C_6F_5), the neutral bicyclo[1.1.0]tetraphosphabutane isomers exo,endo-199 and exo, exo-199 are formed. Calculations indicate that both isomers lie close in energy, and the evidence shows that the phosphanes decompose within 24 h, due to the lack of steric protection.

Figueroa and co-workers described the reactivity of the iron isocyanide complex $Fe(N_2)(CO)_2(CNAr^{Tripp2})_2$ (**200**, $Ar^{Tripp2} = 2,6-(2,4,6-(iPr)_3C_6H_2)_2C_6H_3$)) towards different substrates, including P₄ [204]. Treatment of compound **200** with 1 equivalent of white phosphorus led to the clean formation of a product, structurally characterized as the butterfly-P₄ complex $Fe(\kappa^2-P_4)(CO)_2(CNAr^{Trip-P^2})_2$ (**201**, Scheme 61). A similar oxidative addition was observed upon reaction, *e.g.*, with molecular hydrogen (H₂), yielding a dihydride complex.



Fig. 54. X-ray structures of a) 186, b) 188, c) 189 and d) 190; adapted from Ref. [197].



Scheme 58. Synthesis of complex $[Cp'''Fe]_2(\mu-P_4)$ (**192**), featuring a distorted kitelike *cyclo*-P₄ ligand, and its products after thermic reaction.

As already discussed, iron complexes with *cyclo*-P₄ ligands obtained after direct activation of white phosphorus, are quite rare, with known examples reported only by the groups of Scheer [197], Mézailles [205] and Wolf [206]. Compounds **188** and **189**, by Scheer and co-workers [197] were obtained from a butterfly-P₄ iron complex, in which the butterfly-P₄ moiety underwent isomerization (*vide supra*). Later, Mézailles and co-workers [205] reported the synthesis of the first stable end-deck *cyclo*-P₄ iron complex [($^{Ph}PP_{\Sigma}^{Cy}$)Fe(η^{4} -P₄)] (**202**) and investigated its reactivity towards electrophiles (Scheme 62). Complex **202** can be synthesized either



Fig. 55. X-ray structure of **192** featuring a kite-like *cyclo*-P₄ unit; adapted from Ref. [199].

in a one-pot (Scheme 62, path a) or stepwise reaction, reducing the Fe(II) precursor bearing the tridentate phosphine ligand *in situ* (**203**) to generate a coordinatively unsaturated Fe⁰ fragment, which is then reacted with P₄ (Scheme 62, path b). The formation of the *cyclo*-P₄ moiety was confirmed through X-ray diffraction (Fig. 57a), where an almost perfect square planar configuration of the P₄²⁻ ligand was observed (sum of P-P-P angles equal to 359.99°) with bond distances (2.135(1)–2.166(1) Å) shorter than P-P single bonds (2.21 Å), but longer than typical P=P bonds (*ca.* 2.0 Å). The Mössbauer spectrum is in accordance with an Fe(II) center in a singlet state, and this evidence suggests that the reduction of white phosphorus was promoted by an Fe⁰ species (two-electron transfer), thus generating the Fe(II) complex bearing a *cy-clo*-P₄²⁻ ligand (**202**) [142]. Additionally, nucleus independent



Scheme 59. Synthesis of $Cp_2^{\mathbb{R}}P_4$ via one-step Fe-mediated transfer of carbon-centered radicals: a) Synthesis of 195b mediated by $[Cp'''Fe(\mu-Br)]_2$ (191a) and b) Optimized synthesis using *in situ* generated { $Cp'''Fe^{[IIB}P_2$ } fragments.

chemical shift (NICS) DFT calculations confirm the aromaticity of the cyclo-P₄ moiety. The authors suggest that the coordination of the P₄²⁻ ligand to the Fe center results in its aromatization and constitutes the driving force for the reaction and the reason of the quantitative yield of the complex. The high stability of this species was confirmed through thermal treatment for days, making it a suitable platform for additional functionalization. Thus, the reactivity of 202 towards electrophiles was studied (Scheme 62, bottom). The reaction of **202** with $B(C_6F_5)_3$ in dichloromethane yields the P-borylated Lewis adduct **204**, in equilibrium with the end-deck cyclo-P₄ complex. Additionally, 202 reacts with CuCl through P-coordination on the cyclo-P₄ ligand, to form a tetrametallic Fe₂Cu₂ complex with two CuCl bridging moieties (205). The formation of both P-functionalization products was confirmed *via* X-ray diffraction (Fig. 57b-c). Treatment of **202** with [H(Et₂O)₂] $[BAr_4^F]$ (BAr_4^F = *tetrakis*(3,5-bis(trifluoromethyl)phenyl)borate) results in protonation at the metal center (**206**) as opposed to the cyclo- P_4 unit.

Wolf and co-workers reported another example of an iron complex bearing a tetraphosphacyclobutadiene (P_4^{2-}) ligand [206]. Upon reaction of $[Cp^RFe(\mu,Br)]_2$ (**191b**, $Cp^R = C_5(C_6H_4-4-Et)_5$) with potassium naphtalenide $(K(C_{10}H_8), 4 \text{ equiv.})$, followed by addition of P₄ (2 equiv.), $[\{Cp^RFe(\eta^4-cyclo-P_4)\}]^-$ (**207**, Scheme 63a) forms. This was isolated as a pure, crystalline solid (after addition of 2 equiv. 18-crown-6), albeit in a low yield (4%). Nonetheless, crystals suitable for X-ray diffraction were obtained, and the molecular structure of **207** determined (Fig. 58a). As in the example by Mézailles [205], the P-P bond distances (2.1558(6)–2.1722(6) Å) suggest that a delocalized planar P₄-square has been obtained. DFT calculations on a truncated model of **207**, $[(C_5Ph_5)Fe(\eta^4-cyclo-P_4)]^-$ (**207**') reproduced the structure of the former, and analysis of the frontier molecular orbitals suggests a d⁶ configuration for the iron center.

During these investigations, Wolf and co-workers [206] also achieved the functionalization of P_4 chains in the coordination sphere of 3d metalate anions for the first time. Reaction of $[Cp^{R}-Fe(\mu,Br)]_2$ (**191b**, $Cp^{R} = C_5(C_6H_4-4-Et)_5$) with Na/Hg (exc.) and P_4 , instead of yielding the *cyclo*-P₄ compound **207**, led to the new diiron complex $[Na_2(THF)_5{(Cp^{R}Fe)_2(\mu,\eta^4:\eta^4-P_4)}]$ (**208**) in moderate yield (67%) (Scheme 63b). Compound **208** is highly air sensi-



Scheme 60. Synthesis of a Lewis-acid stabilized Iron-butterfly- P_4 complex (**197**) and functionalization (protonation) of the P_4 -moiety.



Fig. 56. X-ray structure of the Lewis-acid stabilized complex $[Cp^*Fe(CO)_2\{\eta^{1}-P_4\cdot|B(C_6F_5)_3]\}]^-$ (**197a**); adapted from Ref. [202].



Scheme 61. Synthesis of the Fe-butterfly-P₄ complex $Fe(\kappa^2-P_4)(CO)_2(CNAr^{Tripp2})_2$ (201) by the group of Figueroa [204].



Scheme 62. Activation of P_4 at Fe, and reactivity of the *cyclo*- P_4 ligand towards electrophiles.

tive, decomposing slowly predominately to the protonated ferrate $[Na(THF)_{3}\{(Cp^{R}Fe)_{2}(\mu,\eta^{4}:\eta^{4}-P_{4})(H)\}]$ (**209**) when dissolved in THF. In the structure of **208** (Fig. 58**b**), two P₂ dumbbell units coordinate to the two Cp^R-iron fragments. The molecular structure reveals a weak interaction between these two units, and P-P bonds (P1-P2 = 2.0782(17) Å, P3-P4 = 2.0778(17) Å) shorter than other known compounds. The protonated ferrate 209 can be synthesized, in moderate yields, through the reaction of the diiron compound **208** with one equivalent of Et_3N ·HCl or $[H(Et_2O)_2][BAr_4^F]$, or even trace amounts of water. Crystallographic (Fig. 58c) and spectroscopic investigations show that the proton is highly mobile. Computational optimization indicates that in both of the minimum-energy isomers, the hydride ligand is bridging the iron center and a phosphorus atom in the P₄-chain. Additionally, treatment of 208 with an excess of Me₃SiCl releases the phosphorus scaffold from the iron center and generates a mixture of $P_7(SiMe_3)_3$, $HP(SiMe_3)_2$ and $P(SiMe_3)_3$ in a ratio of 10:1:1. This proves that upon reaction with suitable electrophiles, the functionalized polyphosphido fragments can be liberated from the metal center.

Wolf, Weigand and co-workers reported on the synthesis of heterobimetallic complexes with strongly reduced P_4 ligands

[207]. In their approach, a metalate was reacted with the previously reported [208] gallium tetraphosphido complex [(nacnac) $Ga(\kappa^2-P_4)$] (**210**, nacnac = CH[CMeN(2,6-*i*Pr₂C₆H₃)]₂), obtained by reaction of (nacnac)Ga¹ with P₄. This methodology has proven useful for the synthesis of iron and cobalt [207,209] (see Section 3.2.1) complexes containing P_n ligands. The subsequent reactions of **210** with bis(anthracene)metalates $[M(\eta^4-C_{14}H_{10})_2]^-$ [M = Fe (211), Co (212)] were studied (Scheme 64) [207]. The ferrate [K([18]crown-6)(thf)₂][Fe(η^4 -C₁₄H₁₀)₂] (**211**) [210,211], was reacted with **210** in a 1:1 ratio. Due to the labile anthracene ligands, 211 can be considered a source of "naked" Fe⁻ atoms. Dark crystals of a highly air and moisture sensitive compound were obtained although it could not be isolated in a pure form. However, its molecular structure could be determined by single-crystal X-ray diffraction analysis (Fig. 59), thus confirming the synthesis of $[K(18]crown-6){(\eta^4 C_{14}H_{10}$)Fe(μ , η^4 : κ^2 -P₄)Ga(nacnac)}] (**213**). The structure features a bridging P₄-chain in which the iron center is coordinated to the four phosphorus atoms, while the gallium center only binds to the terminal phosphorus atoms. The P-P bond distances are shorter than typical P-P single bonds, and the terminal P-P bonds (2.1166 (6) Å and 2.1131(7) Å) are shorter than the internal P-P bond (2.1801(7) Å). The long distance between the terminal atoms (3.5473(6) Å) confirms that the P₄ moiety is coordinated as an open P₄-chain.

Upon degradation of the P₄ molecule, other P_n ligands (n < 4)can be obtained. Such fragments can either coordinate to the metal as different structural motifs, or aggregate, giving rise to different P_n topologies. P_n frameworks with $n \ge 5$ are usually the result of the reaggregation of different P₄ degradation products. The fragmentation of P_4 at iron to yield two bridging P_2^{2-} ligands was reported by the group of M. Driess [212]. The β -diketiminatoiron(I) complex $[L^0$ Fe toluene] (**215a**, $L^0 = CH[CHN(2,6-iPr_2C_6H_3)]_2$, see Scheme 65, bottom) activates white phosphorus selectively, affording the iron(III) compound $[(L^0Fe)_2(\mu_2;\eta^2;\eta^2-P_2)_2]$ (216), which features two dumbbell P_2^{2-} ligands (Scheme 65a). The P=P bond distance (2.036(2) Å) observed in the molecular structure of the complex (Fig. 60a), confirms the double bond on the dumbbell ligands. One-electron reduction of the dinuclear complex 216 yields the mixed-valence anion $[(L^0Fe)_2(\mu_2:\eta^2:\eta^2-P_2)_2]^-$ (217) without affecting the structural features of **216**. Mössbauer spectroscopic studies, as well as DFT calculations, indicate that 217 is a rare example of a delocalized mixed-valent iron(II,III) pair.

Scheer and co-workers reported very different results on the activation of P_4 using similar albeit smaller β -diketiminato-iron(I) complexes (see Scheme 65b) under the same reaction conditions [213]. While Driess and co-workers found that complex **215a** cleaves the P_4 molecule to form two P_2 units (Scheme 65a) [212], the analogous compounds **215b-215d** react with P_4



Fig. 57. X-ray structures of a) 202 and its Lewis adducts b) 204 and c) 205; adapted from Ref. [205].

(Scheme 65**b**) to yield either tetranuclear $[(LFe)_4(\mu_4,\kappa^2:\kappa^2:\kappa^2:\kappa^2:\kappa^2)]$ P_8] (L = L¹ (**218a**), L² (**218b**)) complexes, featuring a realgar-type P_8 structural motif or the cyclo- P_4 dinuclear compound $[(L^3Fe)_2(\mu, \mu)]$ $\eta^4:\eta^4-P_4$] (**219**). According to DFT calculations, the dimerization of the hypothetical species $[(L^1Fe)_2(\mu, \eta^4: \eta^4-P_4)]$ should be endothermic, but analysis of the stability of the spin states of the complex and dimeric product, as well as the natural population analysis (NPA), indicate the presence of Fe(II) centers and a P_8^{4-} ligand scaffold. XRD of complexes 218 (Fig. 60b) revealed that in the P_8^{4-} ligand the P-P bonds lie within the typical single P-P bond distances, and that the Fe(II) centers reside on a distorted tetrahedral geometry. Hence, it was found that even small changes to the nature of the flanking groups on the β -diketiminato ligands have a profound effect on the type of P₄-activation product obtained. The ligands featuring dimethylphenyl substituents afforded the tetranuclear-P₈ products, **218**, whereas complexes **216** and **219** only form when the substituent on the ligands is the more sterically demanding diisopropylphenyl (dipp). Remarkably, the effect of the α -substituents on the backbone of the β -diketiminato ligand on the activation of P₄ was also demonstrated, since 216 features two separated P_2^{2-} units [212], whilst **219** bears a planar cyclo- P_4 motif (Fig. 60c). Such an outcome was attributed to the additional steric influence of the Me groups, which tend to push the dipp substituents closer to the metal center affecting the distances between the iron(II) centers [213].

In a collaboration between the groups of Peruzzini and Scheer, the reactivity of manganese and ruthenium complexes featuring coordinated intact P₄-tetrahedra towards low-valent βdiketiminato iron and cobalt compounds was described [81]. Among these were **215c** and its cobalt congener [L²Co toluene] (220a) (L² = CH[CHN(2,6-Me₂C₆H₃)]₂; Scheme 66). The selective cleavage of two P-P edges from the P4 moiety in the previously described compound $[{Cp^{BIG}Mn(CO)_2}_2(\mu, k^1:k^1-P_4)]$ [161, $Cp^{BIG} =$ $C_5(C_6H_4nBu)_5)$, see Section 2.5.1] by such β -diketiminato compounds, led to the formation of the π -delocalized P_4^{2-} complexes $[{Cp^{BIG}Mn(CO)_2}_2{ML^2}(\mu,\eta^1:\eta^1:\eta^4-P_4)]$ [M = Fe (221), Co (222)] (see Fig. 61 for the molecular structures of the complexes). This study becomes particularly interesting since the differences or similarities in reactivity between the free and coordinated P4 molecule can be examined directly. In its reactivity towards the β diketiminato complexes, the P₄ ligand in **161** behaved analogously to the free P₄ tetrahedron, forming cyclo-P₄ moieties. For 221, the effective magnetic moment (Evans method), along with EPR and ⁵⁷Fe Mössbauer measurements, suggested the presence of a lowspin iron(III) center with one unpaired electron. Cobalt complex **222** is diamagnetic and features an A_2X_2 spin system in the ³¹P



Scheme 63. Synthesis of a) $[{Cp^RFe(\eta^4-cyclo-P_4)}]^-$ (**207**, $Cp^R = C_5(C_6H_4-4-Et)_5)$ and b) a dinuclear ferrate bearing a bridging P₄ chain ligand, and its subsequent functionalization with electrophiles, by Wolf and co-workers [206].

 ${^{1}\text{H}}$ NMR spectrum. It was suggested that, unlike similar systems (*c.f.* complex **219**) [213], the formation of bimetallic inverted sandwich compounds with polyphosphorus middle-decks was prevented in this case by the [Cp^{BIG}Mn(CO)₂] fragments bonded to the *cyclo*-P₄ ligand.

Another example of metal-mediated fragmentationreaggregation of P₄ with anionic iron complexes, is the formation of iron polyphosphides described by the Wolf group [214]. By reacting the anionic iron complex [K(18-crown-6){Cp*Fe(η^4 -C₁₀H₈)}] (**223**) with P₄ (Scheme 67), two major products were obtained by fractional crystallization, in low to modest yield. These compounds corresponded to P₄ fragmentation and aggregation products, namely the mononuclear complex [K{([18]-crown-6)}₂(Cp*Fe(k²: k²-P₇)] (**224**) and the anionic cluster [K{([18]-crown-6)(thf)₂] [(Cp*Fe)₃(μ ,k²:k²-P₃)₂] (**225**). The molecular structures (Fig. 62) of these compounds show a P₇ norbornadiene-like framework in **224** and two *cyclo*-P₃ units in cluster **225**. Computational as well as spectroscopic studies correlate well with the findings from XRD.

Of the common P_n ligands ($n \ge 5$) formed by aggregation of different P-units, the *cyclo*-P₅ scaffold has attracted particular attention. Due to the isolobality between the C–H moiety and the phosphorus atom, the *cyclo*-P₅ motif can be viewed as an analogue of the cyclopentadienyl ligand. The coordination and supramolecular chemistry of the *cyclo*-P₅ ligand towards iron has been extensively investigated by Scheer and co-workers, particularly the family of pentaphosphaferrocene compounds [129,181–187]. However, in this review we will only summarize the reactivity of this class of compounds towards nucleophiles [215–218] and its redox chemistry [219]. For supramolecular assemblies featuring pentaphosphaferrocenes, see Refs. [129,181–187].



Fig. 58. X-ray structures of a) 207, b) 208 and c) 209; adapted from Ref. [206].



 $\begin{array}{l} \mbox{Scheme 64. Synthesis of the heterobimetallic anionic complexes } \{(\eta^4\mbox{-}C_{14}H_{10}) \\ M(\mu,\eta^4\mbox{:}\kappa^2\mbox{-}P_4)Ga(nacnac)\}^- \ [M = Fe\ (\textbf{213}),\ Co(\textbf{214})]. \end{array}$

Scheer and co-workers studied the reactivity of the pentaphosphaferrocene compound [Cp^RFe(η^5 -P₅)] (Cp^R = Cp*, **25a**) towards different main-group nucleophiles [215]. Although the reactivity was dependent on the specific nucleophile used, in general, an unprecedented functionalization of the *cyclo*-P₅ motif to generate η^4 -P₅ ferrate complexes was observed. Upon reacting **25a** with LiCH₂SiMe₃ or LiNMe₂ (Scheme 68a), formation of the mononuclear ferrate complexes **226** was reported, whereas the analogous reaction with NaNH₂ yielded the dinuclear species **227** (Scheme 68b). A mixture of **226c** and **227b** was obtained after



Fig. 59. X-ray structure of the anionic complex 213; adapted from Ref. [207].

reaction with LiPH₂, with the latter being a by-product. The ³¹P NMR spectra of complexes **226a** and **226b** shows an AMM'XX' spin system for the *cyclo*-P₅ moiety. The envelope conformation of the η^4 -P₅ ligand was confirmed through crystal structure analysis of **226a** (Fig. 63**a**). The four P-atoms in the η^4 -P₅ unit are nearly planar, with bond distances in the order of P-P double bonds (2.130 Å-2.162 Å). The molecular structures of the dinuclear complexes **227** were also confirmed through XRD (see Fig. 63**b** for the structure of **227b**).

In a collaboration between the groups of Scheer and Roesky [216], **25a** was reacted with low-valent silicon compounds of differing steric encumbrance. Upon treatment with (N-N)SiCl (3 equiv., N-N = PhC(NtBu)₂, see Scheme 68c), selective substitution of a P-atom and insertion of the silicon fragment into the all-phosphorus ring was evidenced through the formation of [Cp*Fe{ η^4 -P₄Si(N-N)}] (**228**) containing the first example of a silatetraphospha-cyclopentadienyl ligand (molecular structure)

shown in Fig. 63c). This was formed alongside the functionalized phosphasilene species **229** (Fig. 63d). It was proposed that **228** is obtained from an intermediate $[Fe](\eta^4-P_5)Si(N-N)(X)$ (230a, [Fe] = Cp^*Fe , X = Cl) similar to **226**, which could not be isolated. ³¹P{¹H} VT-NMR experiments indicate that the substituted cyclo-P₅ intermediate species **230a** is formed at -70 °C. To further investigate this species, 25a was treated with the bulkier (N-N)Si(N $(SiMe_3)_2$) (1 equiv., N-N = PhC(NtBu)_2) bearing the less labile N $(SiMe_3)_2$ group, affording the Si-substituted η^4 -P₅ compound 230b as structural proof of such a reaction intermediate (Scheme 68). Crystal structure analysis of compounds 228 and **230b** confirmed the transformation of the *cyclo*-P₅ ring in both cases (Fig. 63c,e). Additionally, the reaction of 25a with the Si(I) [(N-N)Si–Si(N-N)] compound (ratio 25a:Si(I) compound = 1:1) affords complex $[Cp^*Fe{\eta^4-P_5[Si(N-N)]_2}]$ (231), which is the product of double ring expansion of the *cvclo*-P₅ ring, thus generating a cvclo-Si₂P₅ ring. The solid state structure of **231** (Fig. 63f) confirms the formation of this seven-membered ring. Along with 231, complex 228 and the previously reported compound [(N- $N_{2}Si_{2}P_{2}$ (232) [217] were detected as by-products. The authors highlighted 231 as the largest structurally characterized coordinated silicon-polyphosphorus ring. This Si₂P₅ ring is analogous to the tropylium anion [216].

Scheer and co-workers also studied the iodination of the pentaphosphametallocenes $[Cp^{R}M(\eta^{5}-P_{5})]$ $[Cp^{R} = Cp^{*}; M = Fe$ (**25a**), Ru (**233**)] as a possible method for the formation of P-I cages [**218**]. By reacting the corresponding pentaphosphametallocene with iodine (Scheme 69), complexes of the general formula $[Cp^{*}M(k^{3}-P_{6}I_{6})]X$ [M = Fe, X = I (**234a**); $M = Fe, X = (I)_{0.5}(I_{3})_{0.5}$ (**234b**); M = Ru, X = I (**235**)] were obtained. Each complex contained a MP₆ moiety reminiscent of the common nortricyclane anion, P₇³⁻. The P-P bond lengths in the three compounds lie in the range of single bonds (2.160 Å-2.281 Å). When **25a** was reacted with an excess of I₂ (9 equiv.), in addition to **234b**, PI₃ is



Scheme 65. Activation of P₄ at Fe(I)-β-diketiminato complexes.



Fig. 60. X-ray structures of selected β -diketiminato-iron complexes featuring a) P₂, b) P₈ and c) *cyclo*-P₄ ligands derived from P₄, by the groups of Driess and Scheer; adapted from Refs. [212,213].

obtained. Due to the low solubility of **234a/b** in most conventional organic solvents, the more soluble $[Cp^{BIG}Fe(\eta^5-P_5)]$ (**25b**) was screened as a starting material. However, when **25b** was treated with I₂, only the selective formation of PI₃ was observed. When **234a** was dissolved in MeOH a reaction occurred, affording $[Cp^*Fe\{k^3-P_6(OMe)_6\}]X$ (**236**, Scheme 69), according to the ESI-MS and ³¹P{¹H} NMR spectra, and MeI, as proposed by the authors. The formation of **236** is accelerated when the reaction is carried out in the presence of NaOMe, but the procedure is less selective. Attempts to react the metallo-nortricyclane complexes with a variety of nucleophiles or reducing agents were unsuccessful. The oxidation of the pentaphospharuthenocene compound **233** with iodine also resulted in an aggregation process, yielding the analogous metallo-nortricyclane complex **235** (Scheme 69).

A rich redox chemistry with pentaphosphaferrocene complexes (25) has also been developed [219]. A study [220] of the frontier orbitals of 25 revealed that for the pentaphosphaferrocene, strong contributions of the P atoms in the HOMOs, and metal as well as phosphorus contributions in the LUMOs are expected. This in in contrast with its carbonaceous analogue ferrocene, in which the redox chemistry is mainly metal-centered. Thus, for pentaphosphaferrocene redox reactions give rise to structural changes with participation of P atoms. Scheer and co-workers [219] were able to isolate and characterize the dicationic species $[(Cp^*Fe)_2(\mu,\eta^4)]$ P_{10}]²⁻ (**238**) and [Cp*Fe(η^4 -P₅)]²⁻ (**239**) after reaction of **25a** with a range of oxidizing or reducing reagents (Scheme 70). The structural features of complexes 237-239 were determined through XRD (see, for example, Fig. 64a for the structure of 237). The results of this work highlight the differences in reactivity between ferrocene and 25a. While ferrocene mainly shows iron-centered redox behavior, pentaphosphaferrocene exhibits an iron-centered behavior strongly coupled with the P-atoms. Thus, tilted η^4 -P₅ motifs are obtained, and these can dimerize to yield P₁₀ moieties, although essentially retaining the structure at the metal center upon oxidation or reduction. The anionic products 238 and 239 were reacted with P₄, yielding norbornadiene-like polyphosphorus compounds (P₇ units), including the previously reported **224** [214] and the P₁₄ species **240** (Scheme 70**b**). By using this procedure, compound 224 was obtained in almost quantitative yield. Similarly, with 240 the P₁₄ ligand can be described as two P₇ cages linked through the apical P-atom, as evidenced from its molecular structure (Fig. 64b). The ³¹P{¹H} NMR of **240** shows signals that indicate chemical non-equivalence of the P-atoms directly coordinated to the iron centers [219].

The group of Mézailles described the synthesis of iron phosphide nanoparticles, which were obtained from *in situ* generated iron nanoparticles (using Fe(CO)₅ (**241**) as Fe(O) source) in the presence of white phosphorus [221]. The preferential formation of crystallized FeP (**242**) hollow nanoparticles was reported, upon heating amorphous non-hollow FeP nanoparticles at 250 °C. By using the theoretical exact stoichiometry (Fe/P = 2), or even lower amounts of white phosphorus (Fe/P = 4), either the Fe₂P (**243**) phase could not be obtained or nonpure Fe₂P hollow double-shell nanoparticles were formed, respectively. The authors indicated that the formation of Fe₂P may be possible by using an amorphous Fe–P intermediate in P-poor systems.

3.1.2. Ruthenium

The coordination of intact P_4 -cage to ruthenium centers [178,222–228], as well as its subsequent functionalization via hydrolytic methods [222–227] has dominated the P_4 chemistry of this metal, during the period under review.

In 2011, Peruzzini and co-workers described the dynamic behavior of P₄ ligands coordinated to ruthenium [222]. By reacting either the Ru(II) precursor trans-[Ru(dppm)₂(H)(η^2 -H₂)]BF₄ (**244**, dppm = $Ph_2PCH_2PPh_2$) [222] or $[Ru(dppm)_2(H)_2]$ (245a) with HBF₄-Et₂O in CH₂Cl₂ in the presence of white phosphorus [178], the cationic complex *trans*-[Ru(dppm)₂(H)(η^1 -P₄)]⁺ (**246a**) was obtained (Scheme 71a). Single-crystal XRD confirmed the η^{1} coordination of the P_4 tetrahedron (Fig. 65a), which was located trans to the hydride ligand in the octahedral Ru(II) center. VT ³¹P {¹H} along with ³¹P{¹H} EXSY NMR experiments indicated that, in solution, the P₄ ligand in complex 246a rotates freely about the Ru-P_{cage} bond. This process renders the four phosphorus atoms in the P₄ cage equivalent at room temperature. Complex 246a reacted with [(C₂H₄)Pt(PPh₃)₂] (247) in CH₂Cl₂, affording the heterobimetallic compound [{Ru(dppm)₂(H)}($\mu,\kappa^1,\kappa^2-P_4$){Pt(PPh₃)₂}]BF₄ (248a) in quantitative yield. Complex 248a is the product of inser-



Scheme 66. Activation of P₄ at a Mn-coordinated P₄ ligand.



 $\label{eq:Fig. 61. Molecular structures of [{Cp^{BIG}Mn(CO)_2}_2[ML^2](\mu,\eta^1;\eta^1;\eta^4-P_4)] [a] M = Fe \ (\textbf{221}), \ b) \ M = Co \ (\textbf{222})]; \ adapted \ from \ Ref. \ [\textbf{81}].$

tion of the $[Pt(PPh_3)_2]$ fragment into one of the basal P-P bonds of the Ru-coordinated P₄ tetrahedron [222].

The study of the dynamic P_4 cage behavior in solution was then extended to other $Ru(\eta^1-P_4)$ complexes [178]. Compound **246b**, featuring the bidentate ligand dppe and the bimetallic RuPt complex **248b**, were synthesized by a procedure similar to the one previously described for the analogue with dppm. Moreover, treating
$$\begin{split} & [Cp^{R}Ru(PP)Cl] \ (249) \ with \ P_{4} \ and \ a \ reductant \ forms \ the \ cationic \ Cp^{R}Ru \ complexes \ [Cp^{R}Ru(PP)(\eta^{1}-P_{4})]^{+} \ (Cp^{R} = Cp, \ PP = (PPh_{3})_{2}, \ 250a; \ Cp^{R} = Cp^{*}, \ PP = (PPh_{3})_{2}, \ 250b; \ Cp^{R} = Cp^{*}, \ PP = dppe, \ 250c; \ see \ Scheme \ 71b). \ These \ complexes \ as well \ as \ the \ bimetallic \ RuPt \ complex \ [{CpRu(PPh_{3})_{2}}](\mu,\kappa^{1},\kappa^{2}-P_{4}){Pt(PPh_{3})_{2}}]^{+} \ (251) \ were \ examined \ in \ the \ solution \ phase. \ 246b \ and \ complexes \ 250 \ were \ characterized \ through \ single-crystal \ X-ray \ diffraction \ (see \ Fig. \ 65-) \ displays \ Single-crystal \ X-ray \ diffraction \ (see \ Fig. \ 65-) \ displays \ Single-crystal \ X-ray \ diffraction \ (see \ Fig. \ 65-) \ displays \ Single-crystal \ X-ray \ diffraction \ (see \ Fig. \ 65-) \ displays \ Single-crystal \ X-ray \ diffraction \ (see \ Fig. \ 65-) \ displays \$$



Scheme 67. Fragmentation-aggregation of P₄ mediated by Cp*Fe⁻.



Fig. 62. X-ray structures of the iron polyphosphide complexes a) 224 and b) 225; adapted from Ref. [214].

b-e). VT NMR and EXSY experiments revealed that the octahedral complexes 246 are considerably more fluxional in solution than the piano-stool compound 250a or its osmium analogue [CpOs $(PPh_3)_2(\eta^1-P_4)]^+$ (252). Such observations were associated with the existence of a more ordered structure for the piano-stools, according to its less negative entropic term. Interestingly, by simply substituting the Cp ligand in 250a for the more basic Cp* in 250b, the dynamic process in solution for the P₄-cage ligand seems to be suppressed, which was explained in terms of the electronic character of the cyclopentadienyl ligand. The different behavior of the Cp^{*} derivative was also evidenced by the lack of reactivity of 250b towards hydrolysis, even at high temperatures. For 250c, featuring a bidentate diphosphine ligand, the tumbling movement of the P₄-tetrahedron was also detected in solution, in spite of the presence of the chelating diphosphine. The activation parameters for the dynamic processes were calculated using line-shape analysis and 2D-EXSY spectra. The results showed that subtle changes at the metal center affected the activation parameters for the slow tumbling motion of the P₄-cage in the piano-stool complexes. Remarkably, for compounds 246b and 250a, the dynamic process was also identified in the solid state by solid state MAS ³¹P NMR and ³¹P MAS EXSY experiments. The results indicated that the tum-

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bling motion of the P_4 -cage is still observable in the solid state and confirm that dissociative mechanisms are not responsible for the scrambling of the four P_4 atoms in solution [178].

Additionally, Peruzzini, Yakhvarov and co-workers reported the first example of a water-soluble ruthenium complex featuring a P₄ cage, coordinated in a η^1 -mode [223]. The complex [CpRu $(TPPMS)_2(\eta^1-P_4)|X$ (250d, TPPMS = sodium salt of mmonosulfonated triphenylphosphine, $Ph_2P(m-C_6H_4SO_3Na)$, X = PF_6^- or BF_4^- ; Scheme 72) was obtained after reacting [CpRu (TPPMS)₂Cl] (249d) with white phosphorus (1 equiv.) in the presence of a chloride scavenger (TIPF₆ or AgBF₄). In neat water, rapid decomposition of 250d occurs and H₃PO₃ and H₃PO₄ were observed to form exclusively. The complex underwent mild hydrolysis with a known excess of water in DMF (25 °C) to yield [CpRu(TPPMS)₂(PH₃)]⁺ (**253**), free phosphorous acid, H₃PO₃, phosphoric acid H₃PO₄ and molecular hydrogen. At higher temperatures (35 °C), the reaction proceeded differently since, under these conditions, 250d afforded the complexes [CpRu(TPPMS)₂{- $PH(OH)_{2}]^{+}$ (254) and $[CpRu(TPPMS)_{2}\{P(OH)_{3}\}]^{+}$ (255), as identified via ³¹P{¹H} NMR. Kinetic measurements (³¹P{¹H} NMR) indicated that the hydrolysis of 250d involves an associative mechanism, in which nucleophilic attack of water on the η^1 -P₄ ligand occurs in the transition state.

Stoppioni and co-workers [224] studied the activation of coordinated P₄ by treating the cationic ruthenium complex [{CpRu $(PPh_3)_2\}_2(\mu,\eta^1:\eta^1-P_4)](OTf)_2$ (**256**, OTf = OSO₂CF₃) with iodine (Scheme 73). Complex 256 features an intact P₄-tetrahedron as a η^1 : η^1 -P₄ bridge. In the presence of adventitious water, addition of iodine (1 equivalent) to 256 at low temperatures gives rise to the monocation $[{CpRu(PPh_3)_2}_2(\mu_1:\mu_3,\eta^1:\eta^1-P_4H_2I)]^+$ (257). This features a 1,3-dihydride-2-iodidecyclotetraphosphan-1-ide ligand, (P₄H₂I)⁻. Facile dissociation of the iodide from the tetraphosphorus ligand in polar solvents, yielded the elusive 1,3-dihydride-2,4-bicy clotetraphosphane[1.1.0] (P₄H₂ butterfly), also stabilized by metal coordination in complex 258. The coordinated P₄H₂ ligand is still susceptible to hydrolysis and, after addition of an excess of water, $[{CpRu(PPh_3)_2}_2 {\mu_1: \mu_3, \eta^1: \eta^1}$ the triphosphine complex PH₂PHPH₂](OTf)₂ (**259**), featuring a fully hydrogenated P₃H₅ ligand, and phosphorous acid, are formed. Remarkably, at this point in time both the P₄H₂ and P₃H₅ ligands observed in the molecular structures of 258 and 259 were unknown in coordination compounds [225,226]. ³¹P{¹H} NMR spectra of complexes 257-259 correlate well with the characterization in the solid state by single-crystal XRD (Fig. 66), thus concluding that the solid state structure is maintained in solution. In contrast to previous investigations by the same group [44], the iodine-mediated hydrolytic degradation of coordinated P4 in complex 256 does not yield hydroxypolyphosphane species, suggesting that a different reaction mechanism operates in this case. The authors highlighted that stable coordination of the P₄ tetrahedron is important to achieve controlled transformations/functionalizations of white phosphorus [224].

Recently, Caporali, Peruzzini and co-workers reported on the synthesis and hydrolysis of a ruthenium-P4 compound bearing the basic PMe₃ ligand [227]. Treatment of [CpRu(PMe₃)₂Cl] (249e) with a chloride scavenger (TlPF₆ or AgOTf), followed by addition of P₄ in equimolar quantities selectively afforded the compound $[{CpRu(PMe_3)_2}_2(\mu,\eta^1:\eta^1-P_4)]^{2+}$ cationic (260)Scheme 74). Attempts to obtain the mononuclear complex [CpRu $(PMe_3)_2(\eta^1-P_4)$ ⁺ were unsuccessful. As with other intact P₄ complexes [222], 260 exhibits a fluxional behavior in solution. Single crystal X-ray diffraction confirmed the molecular structure of 260 and revealed a conformation in which the two cyclopentadienyl rings are eclipsed. Upon addition of water (500 equiv.), 260 hydrolyzes in solution at ambient temperature yielding a mixture of species, of which the major products correspond to [{CpRu



Scheme 68. Functionalization of the cyclo-P₅ ligand in $[Cp^*Fe(\eta^5-P_5)]$ (25a) by nucleophiles (N-N = $[PhC(NtBu)_2]$).

 $(PMe_3)_2\}_2(\mu,\eta^1:\eta^1-P_2H_4)]^{2+}$ (261, 75%), $[CpRu(PMe_3)_2P(OH)_3]^+$ (262, 18%) along with the minor species $[CpRu(PMe_3)_2(PH_3)]^+$ (263), $[CpRu(PMe_3)_2\{PH(OH)_2\}]^+$ (264) and the free phosphonic acids. Better yields of the hydrolytic products were obtained with the complex bearing the more basic phosphine ligand PMe_3, 260, in comparison to the similar reaction with the analogous 256. The reaction proceeds in a more selective manner by treating 260 with MeOH. In this case, only 262 and $[CpRu(PMe_3)_2\{P(OMe)_3\}]^+$ (265) (ratio = 2:1) were obtained, with the former resulting from the water present in the methanol.

In 2017, another example of a Ru complex showing an intact P₄ tetrahedron was reported [228]. In this case, the complex was stabilized by N-heterocyclic carbene (NHC) ligands. Treatment of a THF solution of the unsaturated hydride species $[Ru(IMe_4)_4H]^+$ (266) with one equivalent of P₄, afforded $[Ru(IMe_4)_4(P_4)H]BAr_4^F$ (267). The η^1 -coordination of the P₄ cage *trans* to the hydride ligand was confirmed in the molecular structure determined by XRD (see Fig. 67). The high field ¹H NMR signal of the hydride ligand featured signs of fluxional behavior, showcasing a doublet at room temperature which resolved into a doublet of quartets $(J_{HP} = 266.1 \text{ and})$ 11.0 Hz) at low temperatures. The low-temperature ³¹P{¹H} spectrum showed a pattern like that observed for other intact-P₄ complexes (quartet, δ –394.6; doublet, δ = –468.6). The chemical shifts for the hydride ligand in 267 as well as a series of other $[Ru(IMe_4)_4(-$ L)H]^{0/+} species (L = vacant, H₂, N₂, CO, MeCN, O₂, SO₂, H⁻, F⁻ and Cl⁻) were modelled theoretically, finding a good correlation between experimental and computed data when a scalar relativistic correction is applied, which improves significantly after inclusion of spin-orbit effects.

In a collaboration between the groups of Caporali and Grützmacher [229], the halogenation of P₄ by the unsaturated 16 electron species [Cp*RuX(PCy₃)] [X = Cl (**268a**); X = Br (**268b**)] was investigated. When **268** was reacted with half an equivalent of P₄ (Scheme 75, featuring the X = Cl derivatives), the bimetallic complexes [Cp*Ru(PCy₃)($\mu_2, \eta^2: \eta^4-P_4X_2$)RuCp*] (**269**) featuring a pla-

nar P_4X_2 ligand, were obtained. It was proposed that upon interacting **268** and P₄ form an intermediate species presumably containing an intact P₄ tetrahedron in a bridging coordination mode to two Ru-centers. Subsequently, metal-promoted activation of the coordinated P₄ ligand, followed by cleavage of two P-P bonds and oxidative insertion into the Ru-X bond, gives rise to the unprecedented P₄X₂ motif, which can be described as a 1,4-dihalo tetraphosphabutadiene (X = Cl, Br). Upon reduction with magnesium, the chloro-substituents of the P₄Cl₂ ligand in complex 269a are eliminated as chloride anions, giving rise to two weaklyinteracting P₂ (dumbbell) moieties, in the triple-decker compound 270. Likewise, selective alkylation with *n*BuLi led to the tetranuclear complex $[(Cp^*Ru)_4(\mu_3,\eta^2;\eta^2;k^4-P_4nBu_2)_2]$ (271), in which the P₄Cl₂ ligand has transformed into two coplanar P₄nBu₂ scaffolds. Furthermore, addition of a chloride scavenger to 269a yielded the cationic, unstable species $[Cp^*Ru(PCy_3)(\mu_2,\eta^2;\kappa^4-P_4Cl)$ RuCp*][GaCl₄] (272), which loses the remaining chlorosubstituent to give again compound 270. Crystal structure analyses confirmed the molecular structures of complexes 269a-272 (Fig. 68). DFT calculations suggest that the P₄ motif in these compounds is electronically highly flexible and behaves as a noninnocent ligand.

Fabrizi de Biani, Ienco and co-workers reported the synthesis of triple-decker *cyclo*-P₃ bimetallic Ru-Co, Ru-Ru and Os-Co complexes [230]. These were obtained either by reaction between suitable ruthenium/osmium precursors with the known [(triphos)Co (η^3-P_3)] [273, triphos = CH₃C(CH₂PR₂)] [231], or by direct reaction with white phosphorus. The redox properties of the triple-decker complexes, namely [(triphos)Ru(μ , η^3 : η^3-P_3)Co(triphos)]²⁺ (274), [(triphos)Ru(μ , η^3 : η^3-P_3)Ru(triphos)]⁺ (275) and [(triphos)Os(μ , η^3 : η^3-P_3)Co(triphos)]²⁺ (276), were also studied and compared with analogous cobalt and nickel complexes. Of these, only 275 (Fig. 69) was obtained from direct reaction with white phosphorus, and its molecular structure was determined. Electrochemical studies of complexes 274–276 revealed several monoelectronic redox



Fig. 63. X-ray structures of selected products of functionalization of the *cyclo*-P₅ ligand in $[Cp^*Fe(\eta^5-P_5)]$ (**25a**): a) $[Cp^*Fe(\eta^4-P_5CH_2SiMe_3)]^-$ (**226a**), b) $\{[Cp^*Fe(\eta^4-P_5)]_2PH\}^{2-}$ (**227b**), c) $[Cp^*Fe(\eta^4-P_4Si(N-N)]$ (**228**), d) $[(N-N)Si(Cl) = P-Si(N-N)(Cl)_2]$ (**229**), e) $[\{Cp^*Fe(\eta^4-P_5)\}\{(N-N)Si(N(SiMe_3)_2)\}]$ (**230b**), f) $[Cp^*Fe(\eta^4-P_5[Si(N-N)]_2]$ (**231**) $(N-N = PhC (NtBu)_2)$; adapted from Ref. [**215**,216].



Scheme 69. Iodination of pentaphosphametallocene (M = Fe, Ru) complexes and reactivity or the iron derivative towards MeOH.

processes. These systems were compared with their dinickel and dicobalt analogues and, in general, cyclic voltammetry studies revealed that the triple-decker complexes are stable when their electronic count varies between 30 and 34 valence electrons. Interestingly, with the heavier analogue **275**, an unprecedented species with 29 VE was formed through electrochemical reactions.

3.1.3. Osmium

In the period of 2011–2020, complexes resulting from the coordination/activation of white phosphorus by osmium are the least represented among group 8 metals. As mentioned in the previous section, the triple-decker bimetallic Os-Co compound **276** was reported, and its redox properties were studied *via* cyclic voltammetry [230].

Dynamic behavior in solution was observed for the osmium(II) complex $[CpOs(PPh_3)_2(\eta^{1}-P_4)]^+$ (252), featuring an intact P₄-cage ligand [178]. This dynamic process was similar to that of its ruthenium analogue, 250a. In both cases, they contrasted with that of the iron complex 172, for which only the dissociation of the P₄-ligand in solution was identified. This behavior was also evidenced



Scheme 70. a) Redox chemistry of $[Cp^*Fe(\eta^5\text{-}P_5)]$ (25a) and b) reactivity of the reduced products towards $P_4.$



Fig. 64. X-ray structures of a) 237 and b) the P₁₄-ligand complex 240; adapted from Ref. [219].

by the slight differences observed on the activation parameters of the dynamic process of **250a** in comparison to those of **252**. Based on such observations, it was suggested that the Fe-P_4 bond for the

related complex **172** was kinetically more labile than for the Ru-P₄ and Os-P₄ analogues, **250a** and **252**, respectively.

3.2. Group 9 metals

Undoubtedly, in the period 2011–2020 for group 9 metals, the activation of white phosphorus by cobalt complexes have been the most prevalent. Such complexes have led not only to activated tetraphosphorus moieties, but even products of fragmentation/reaggregation and functionalization. Remarkable contributions have been made in this matter by the groups of Radius, Figueroa, Scheer, Driess and Wolf, among others. Least represented are the species derived from rhodium and iridium.

3.2.0.1. Cobalt

It is common for cobalt(I) complexes bearing labile ligands as potential vacant sites to be used in the activation of white phosphorus [232,233]. One such example was in 2012, when the stepwise degradation of the P₄ tetrahedron at a cobalt(I) center was achieved by the group of Radius [232]. Upon reacting the cobalt compound $[Cp^*Co(iPr_2Im)(\eta^2-C_2H_4)]$ (277, $Cp^* = \eta^5-C_5Me_5$, iPr_2- Im = 1,3-di-isopropylimidazolin-2-ylidene) with white phosphorus, a [Cp*Co(*i*Pr₂Im)] fragment inserts into the P₄-cage promoting cleavage of a P-P bond (Scheme 76a). The product, $[Cp^*Co(iPr_2Im)(\kappa^2-P_4)]$ (278) features a butterfly-P₄ ligand (Fig. 70a shows its molecular structure). Complex 278 reacts with another equivalent of 277 cleaving a second P-P bond to give the dinuclear compound, $[{Cp*Co(iPr_2Im)}_2(\mu,\kappa^2:\kappa^2-P_4)]$ (279), bearing a bridging cyclo-P₄⁴⁻ ligand. Single-crystal XRD (Fig. 70b) revealed that the P₄ ring shows a puckered conformation. This intermediate complex 279 undergoes further transformations when heated in toluene solution at 80 °C. After dissociation of one NHC ligand and rearrangement/opening of the cyclo-P₄⁴⁻ unit, two products are obtained, one featuring a *catena*-P₄, and the other two bridging dumbbell ligands, P_2^{2-} , $[Cp^*Co(\mu, \eta^4: \kappa^2-P_4)Co(iPr_2Im)Cp^*]$ (280) and the previously reported [234] [{ $Cp^*Co(\mu, \eta^2: \eta^2-P_2)$ }] (281a), respectively. Fig. 70c shows the molecular structure of complex **280**. Furthermore, by controlling either the $[Co]:P_4$ stoichiometry or the reaction conditions, compounds 278-281a can be selectively obtained from 277, in high yields. A complex multistep mechanism was proposed by the same group after studying the transformation with DFT methods [235]. It was found that the formation of the dinuclear complex 279, as well as the loss of the different NHC co-ligands (or CO, given that the analogous system was also modelled), were key steps in the reaction. Such loss was considered necessary since the vacant sites facilitate the transformations of the P₄-cage in the coordination spheres of the different cobalt species. According to the calculations, the presence of the donor NHC ligand is better for the overall reaction than with the carbonyl analogue, since the release of the former has a larger entropic effect than for the latter. These calculations highlighted the importance of the nature of the co-ligands on the outcome of P₄-activation reactions.

Figueroa and co-workers described the synthesis of the notable two-legged piano stool Co(I) complex $[Cp^*Co(N_2)(CNAr^{dipp2})]$ (**282**), featuring a sterically encumbering isocyanide and a labile N₂ ligand [233]. Such dinitrogen ligands can be displaced by Lewis bases under ambient conditions therefore, **282** can be seen as a stabilized source for the unsaturated 16 VE fragment $[Cp^*Co(CNAr^{dip$ $p2})]$. Reacting **282** with white phosphorus (ratio Co:P₄ = 1:1, Scheme 76b) yielded the butterfly-P₄ complex $[Cp^*Co(\kappa^2-P_4)$ (CNAr^{dipp2})] (**283**). The elongated P-P bond distance (2.606 Å) between the two P-bounded atoms is indicative of the cleavage of one of the edges of the tetrahedron, giving rise to the butterfly-P₄ ligand (see Fig. 71a). If the Co:P₄ ratio is 1:0.5, or if **283** is treated with an additional equivalent of **282**, the dinuclear



Scheme 71. Synthesis of $Ru(\eta^{1}-P_{4})$ and bimetallic $Ru(\mu,\kappa^{1},\kappa^{2}-P_{4})Pt$ complexes by Peruzzini and co-workers [178,222].

complex [{Cp*Co(CNAr^{dipp2})}₂($\mu_2, \kappa^2: \kappa^2-P_4$)] (**284**) is formed. In the latter, the mild activation of another P-P edge has occurred (Fig. 71b for the molecular structure) [233]. This behavior is analogous to the one observed in the activation of P₄ leading to complexes **278** and **279** [232].

Scheer and co-workers also reported on the activation of white phosphorus, through their investigations on the synthesis of the cobalt-cyclo-P₄ end-deck species $[Cp'''Co(\eta^4-P_4)]$ (285), referred to as "missing homologue" of the series of first-row sandwich complexes $Cp^{R}M(cyclo-P_n)$ [236]. **285** is obtained after reaction of the 14 valence electron Cp"'Co fragment source (286) with white phosphorus (Scheme 77a). The complex dimerizes irreversibly in solution to form two isomeric P₈ species $[(Cp'''Co)_2(\mu,\kappa^4:\kappa^2:\kappa^1-P_8)]$ (287). It should be noted that 287b isomerizes to 287a, thus suggesting a greater stability for the latter. Crystallographic analysis of 285 and 287 (Fig. 72) revealed that the cyclo-P₄ ligand in 285 is almost perfectly square planar and that the P₈ moiety in 287 is comprised by two bound P₅ and P₃ cycles. P-rich polyphosphorus complexes can be obtained by thermolytically promoted rearrangement of compound 285 and isomers 287 in solution. DFT calculations indicate that the solvent plays an important role on the synthesis of 285, since it affects the degree of dissociation of the cobalt precursor to generate the 14 VE fragment. Moreover, these theoretical studies indicated that the key intermediate in the synthesis of the cyclo-P₄ complex is posited to be an intermediate η^3 tetrahedron P_4 species [236].

Recent investigations by the group of Scheer have also shown that complex **285** undergoes ring expansion [119] and ring contraction reactions (see Scheme 77b) [237]. By reacting complex **285** with the electrophilic pnictogenidene complexes [Cp*E{W (CO)₅}₂] the *cyclo*-P₄E ring (E = P, As) species resulting from the insertion of the phosphinidene or arsinidene atom into the *cyclo*-P₄ unit, are obtained [119]. Two isomers of [Cp^{'''}Co($\mu_3, \eta^4: \eta^1: \eta^1-AsP_4Cp^*$]{W(CO)₅}₂] (**289**) were isolated. Compounds **288** are the result of a ring-expansion reaction from a *cyclo*-P₄ to a

substituted cyclo-P₅ motif coordinated in an η^4 -mode to the cobalt center, with 288a being the main product of reaction (ratio **288a**:**288b** = 1:0.19). The envelope conformation of the $cvclo-P_5$ ring was confirmed crystallographically, for both isomers (Fig. 73 shows the structure of 288a), and is structurally reminiscent of the ferrate complexes 226 (see Scheme 68, above) [215]. A possible driving force for the reaction is the easing of the ring strain for the *cyclo*- P_4 moiety. Isomers **289** differ only on the position of the arsenic atom within the cyclo-P₄As ring (Fig. 73b). According to ¹H and ³¹P NMR spectra, the three isomers are present in solution in the ratio 289a:289b:289c = 0.48:0.42:0.10. DFT calculations revealed that the reaction proceeds similarly for the phosphinidine and the arsinidene reagents, via initial formation of an adduct between compound **285** and $[Cp^*E\{W(CO)_5\}_2]$, followed by a shift of a $\{W(CO)_5\}$ fragment and the insertion of the pnictinidene into an unpolarized P-P bond of the cyclo-P₄ ligand to yield the ringexpanded products [119].

Ring contraction of the *cyclo*-P₄ ring occurs when compound **285** is treated with two equivalents of 1,3,4,5-tetramethylimidazo lin-2-ylidene (^{Me}NHC) to afford [(^{Me}NHC)₂P][Cp^{'''}Co(η^3 -*cyclo*-P₃)] (**290**, see Scheme 77 and Fig. 73c) under mild conditions [237]. The reaction proceeds *via* selective NHC-induced phosphorus cation extraction from the *cyclo*-P₄ ring in **285**, generating the ionic compound **290** containing an anionic [Co](*cyclo*-P₃)⁻ ([Co] = Cp^{'''}Co) complex and a bis(NHC)-coordinated P^I cation. It was not possible to either identify or isolate reaction intermediates to support a mechanistic proposal for the formation of **290** [237].

Further chemistry on the [Co]-*cyclo*-P₄ complex **285**, specifically its reduction or reaction with nucleophiles (Scheme 77b), was reported by the same group [238]. To avoid the previously reported [236] irreversible dimerization, the reactions were carried out at room temperature or lower. Reduction of **285** using K, KH or KC₈ (THF at r.t.) yields the known [Co](*cyclo*-P₃)[–] anionic complex, previously obtained from compound **290** [237], and a P₈-aggregate, [(Cp^{'''}Co)₂(μ , η^3 : η^3 -P₈)]^{2–} (**291**), which was structurally characterized (Fig. 73d). The P₈-ligand consists of a *cis*-bicyclo[3.3.0]octane



Fig. 65. X-ray structures of ruthenium(phosphine) cationic complexes featuring intact P₄-tetrahedron ligands: a) 246a, b) 246b, c) 250a, d) 250b and e) 250c; adapted from Refs. [178,222].

core which serves as bridge between two {Cp^{'''}Co} fragments. Alternatively, by treating 285 with nucleophiles such as organolithium reagents, substituted cyclo-P₄ complexes 292 can be selectively obtained at low temperatures. For **292b** ($R = CH_2SiMe_3$, Fig. 73e), both the equatorial and the axial-CH₂SiMe₃ isomers were identified. These compounds are metastable and upon warming to room temperature, the aggregation products $\mathbf{293}$, featuring a P_8R ligand (R = tBu, CH_2SiMe_3) similar to that in **291**, are obtained (see Fig. 73f for the structure of **293b**). For **292a** (R = tBu), it was also possible to identify the novel diorgano-substituted cyclo- P_5R_2 species **294**, which is formed along with the known anion, 290 [237]. Likewise, the reaction of 285 with NaOH affords $[Cp'''Co(\eta^3-P_4(O)H)]^-$ (295) (Fig. 73g), which is presumably formed *via* attack of the OH⁻ group to a phosphorus atom, yielding a substituted P-OH moiety that undergoes tautomeric rearrangement to generate P=O and P-H bonds [238].

The use of low-valent transition metalates, including cobaltates, to activate P₄ has been investigated by the group of Wolf [207,214]. By reacting the cobaltate complex [K(OEt₂)₂{($^{dipp}BIAN$)Co(η^{4} -1,5-cod)}] (**296a**, $^{dipp}BIAN = 1,2$ -bis(2,6-diisopropylphenylimino)ace

naphthene, 1,5-cod = 1,5-cyclooctadiene) with P₄, the dianion $[(^{dipp}BIAN)Co]_2(\mu,\eta^4:\eta^4-P_4)]^{2-}$ (**297**, Scheme 78) is obtained [239]. The molecular structure of the dinuclear compound 297 was determined (see Fig. 74) indicating that, according to the P-P bond lengths, the P₄ motif can be described as a superposition of two mesomeric structures comprising two P_2^{2-} dianions and a P_4^{4-} unit. Cyclic voltammetry (CV) studies revealed that two reversible one-electron oxidation processes occur and therefore, preparative oxidation using [Cp₂Fe]BAr^F₄ allowed for the synthesis of species which are oxidized by one and two electrons. This led to a complete series of dinuclear cobalt tetraphosphido complexes [(^{dipp}BIAN) $Co_{2}(\mu,\eta^{4}:\eta^{4}-P_{4})]^{n-}$ [n = 2 (297),n = 1 (298),n = 0 (299)]. Both **298a** and **299a** feature a cyclic P_4^{4-} motif, while the isomers **298b** and **299b** bear an open-chain P_4^{4-} ligand. However, the open-chain isomer 299b was obtained only as a minor product in the singlecrystal X-ray structure with approximately 9% occupancy. Additionally, the difference between 298a and 298b arises from the fact that these correspond to the ion-contact and the ion-separated structures of compound 298, which does not affect the characterization in solution. The structures of this series of complexes demonstrate that



Scheme 72. Synthesis of a water-soluble tetraphosphorus complex and its reactivity towards hydrolysis.



Scheme 73. Nucleophilic addition of iodine to the dinuclear complex [{CpRu (PPh₃)₂}₂(μ , η^{1} : η^{1} -P₄)](OTf)₂ (**256**) and subsequent hydrolysis of the coordinated P₄ ligand.

redox reactions on the complex do not affect the oxidation state of the P_4^{4-} ligand but effect changes in its structure (Fig. 74). It was highlighted that the presence of the redox-active ligand ^{dipp}BIAN was necessary to achieve such a degree of reduction of the P_4 molecule, due to its exceptional electron-releasing and -accepting properties.

Likewise, the synthesis of a *cyclo*-P₄ cobaltate anion [(PHDI)Co $(\eta^{4}-P_{4})$]⁻ (**300**, PHDI = bis(2,6-diisopropylphenyl)phenanthrene-9, 10-diimine) was also reported by Wolf and co-workers [240]. Complex **300** was obtained after the reaction of [K(18c-6)(thf)_{1.5}] [(PHDI)Co(η^{4} -1,5-cod)] (**301**, PHDI = bis(2,6-diisopropylphenyl)p henanthrene-9,10-diimine, see Scheme 79, bottom), analogous to **296a** [239], with white phosphorus. Interestingly, a contrasting behavior was observed between the reactivity of complex **300** and that of dinuclear **297**, for which only outer-sphere electron-transfer reactions were identified [239]. The *cyclo*-P₄ unit in complex **300** can either be further coordinated to other metallic fragments, as in complex **302** (Scheme 79**a**), or selectively functionalized with diorganochlorophosphanes (R₂PCl, R = Cy,



Fig. 66. X-ray structures of $[{CpRu(PPh_3)_2}_2(\mu_1;\mu_3,\eta^1;\eta^1-P_4H_2I)]^*$ (**257**) and its subsequent products of hydrolysis, $[{CpRu(PPh_3)_2}_2\{\mu_1;\mu_3,\eta^1;\eta^1-P_4H_2]]^*$ (**258**) and $[{CpRu(PPh_3)_2}_2\{\mu_1;\mu_3,\eta^1;\eta^1-PH_2PHPH_2]\}^*$ (**259**). Hydrogen atoms have been omitted for clarity purposes; adapted from Ref. [224].

*t*Bu, Ph, Mes, N(*i*Pr)₂), yielding the neutral ring-expansion η^4 -*cyclo*-P₅R₂ products (**303**) quantitatively (Scheme 79b). Single-crystal X-ray diffraction analysis of **302** revealed that the *cyclo*-P₄ unit is almost perfectly square (see Fig. 75a), with bond distances (2.147 (7) Å on average) in-between the values expected for P-P single and P=P double bonds, thus confirming the presence of a P₄²⁻ ligand. The envelope conformation of the *cyclo*-P₅R₂ products **303** (see Fig. 75**b-c** for the molecular structures of **303a** and **303c**) is structurally reminiscent of complexes **226** (Scheme 68, above) and **288** (Scheme 77, below) [119].

The reactivity of complexes **303** with cyanide salts differs depending on the nature of the substituent, R (Scheme 79b). For the bulky R = Mes, after reacting with cyanide (1 equivalent) a rearrangement of the *cyclo*-P₅R₂ moiety leads to the cyclotetraphosphido complex **304** (see Fig. 75d for the molecular structure). Addition of a second equivalent of cyanide does not promote further conversion of **304**. In contrast, for the other derivatives, treatment of **303** with 2 equivalents of the cyanide salt results in the remarkable fragmentation of the *cyclo*-P₅R₂ ligand into P₃ and P₂ units, forming the anionic cyclotriphosphido cobalt complex **305**



Scheme 74. Synthesis and reactivity towards H_2O and MeOH of the cationic complex [{CpRu(PMe₃)₂}₂(μ , η^1 : η^1 - P_4)]²⁺ (**260**) by the group of Caporali, Peruzzini and co-workers [227].



Fig. 67. X-ray structure of 267. The counterion has been removed for clarity. Adapted from Ref. [228].

and releasing the corresponding 1-cyanodiphosphan-1-ide anions (**306**). Fig. 75**e-f** shows the molecular structures of complex **305** and the cyanodiphosphan-1-ide **306b**, respectively. The different outcome in reactivity between the $cyclo-P_5(Mes)_2$ complex and its analogues was attributed to steric shielding that hinders its [3 + 2] fragmentation. It was proposed that similar cyclotetraphosphido complexes could be key intermediates in the formation of species **305** and **306** [240].

Rectangular P₄ and *cyclo*-P₃ ligands coordinated to low-valent β -diketiminato cobalt complexes were also reported by the groups of Driess [82] and Scheer [241]. Driess and co-workers [82] described the synthesis (Scheme 80a) of the neutral *cyclo*-P₄ complexes [(LCO)₂(μ_2 : η^4 , η^4 -P₄)] [L: L⁰ = CH[CHN(2,6-*i*Pr₂C₆H₃)]₂ (**307a**); L¹ = CH[CMeN(2,6-Et₂C₆H₃)]₂ (**307b**)], which were obtained by treatment of either [L⁰Co-toluene] (**220b**) or [(L¹CO)₂] (**308**) with 0.5/1 equivalents of white phosphorus at room temper-

ature, respectively. The molecular structures of complexes 307a and 307b revealed that the cobalt centers reside in an octahedral geometry, and that the distances between both metal centers exclude any possible Co-Co interactions. Furthermore, the cyclo-P₄ ligand in such complexes can be described as rectangular with two long P-P bonds (2.289 Å for 307a and 2.298 Å for 307b) and two short P-P bonds (2.130 Å for 307a and 2.124 Å for 307b). Fig. 76a shows the molecular structure of 307b. Brokensymmetry DFT calculations, along with bond order analysis, agreed well with the experimental characterization and supported the proposal that the ligand in such complexes corresponds to a tetraphosphacyclobutadiene motif. One-electron reduction of complexes **307** using KC₈ (1 equiv.), led to the anionic compounds $[(LCo)_2(\mu_2, \eta^4; \eta^4 - P_4)]^-$ (**309**), which were isolated in good yields. Single-crystal X-ray diffraction showed that complexes 309a and **309b** feature structural characteristics similar to **307**. except the P-P distances are approximately the same giving a square P_4 ring as opposed to the rectangular one found in the latter. Theoretical calculations and additional characterization evidenced that compounds **309** feature a delocalized six π -electrons cyclo-P₄²⁻ ligand, with mixed-valent cobalt(I,II) centers [82].

Similarly, the group of Scheer reported [241] the related monoanionic complex $[(L^3Co)_2(\mu_2,\eta^4;\eta^4-P_4)]^ [L^3 = CH[CMeN]$ $(2,6-iPr_2C_6H_3)]_2$ (**309c**)] and the P₃-containing compound $[(L^3Co)_2(\mu_2;\eta^3,\eta^3-P_3)]^-$ (**310**), which were obtained after the reaction of P₄ with the formal Co(0) precursor $[K_2(L^3Co)_2(\mu_2:\eta^1,\eta^1-N_2)]$ (311, Scheme 80b). Compound 310 is the result of P-atom abstraction from the P₄-middle deck in **309c**. Single-crystal analysis of **309c** revealed the characteristic features of a planar cyclo- P_4^{2-} motif (Fig. 76b). This evidence, together with the results from magnetic measurements (Evans method and SQUID) led to the conclusion that, as for complexes **309a** and **309b**, the cyclo-P₄ ligand bridges mixed-valent Co(I) and Co(II) centers. For compound 310 (see Fig. 76c for the X-ray structure), such characterization indicated that the cyclo- P_3^{3-} bridges two Co(II) sites. Interestingly, reacting the Co(I) precursors [LCo·toluene] (L = L^2 (**220a**), L^3 (**220c**) or L^4 (220d), see Scheme 80, bottom) with P_4 yields the corresponding neutral species under different conditions. At room temperature, the reaction afforded compound $[(L^3Co)_2(\mu_2;\eta^4,\eta^4-P_4)]$ (**307c**) selectively (see Fig. 76d for the molecular structure), and the approach can be reproduced for the analogues 307d and 307e. The cyclo-P₄ unit in such complexes was described as a neutral P_4^0 ligand, and would be analogous to the examples reported by Driess, **307a** and **307b** [82]. Upon refluxing in toluene, complex **307c** underwent a thermally induced elimination of a P-atom, forming $[(L^{3}Co)_{2}(\mu_{2},\eta^{3}:\eta^{3}-P_{3})]$ (**312**). One-electron reductions of the neutral compounds bearing ligand L³ yielded the corresponding monoanionic complexes **309c** and **310** [241]. Altering the size of the β-diketiminato ligands in 220 does not affect the P₄ activation products [82,241], this behavior contrasts with that previously observed for the iron analogues **215** [212,213].

As mentioned above, the groups of Peruzzini and Scheer studied the reactivity of intact P₄-manganese or ruthenium complexes towards low-valent iron or cobalt compounds [81]. Besides the reactivity towards the β-diketiminato iron (**215c**) and cobalt (**220a**) compounds discussed in Section 3.1.1, the manganese complex **161** and the ruthenium complex **250a** were also reacted individually with the 14 VE cobalt fragment source **286** (Scheme 81). Such reactions afforded [{Cp^{BIG}Mn(CO)₂}₂{CoCp'''}(μ , η^1 : η^1 : η^4 -P₄)] (**313**) or [{CpRu(PPh₃)₂}{CoCp'''}(μ , η^1 : η^4 -P₄)]OTf (**314**), which correspond to a [Cp'''Co(η^4 -P₄)] motif coordinating to 16 VE Mn- or Ru- complex fragments. Remarkably, compound **314**, with one substitution at the *cyclo*-P₄ moiety, or the doubly substituted **313**, do not undergo further degradation or reaggregation as previously observed with the metastable compound **285** (*cf.* Scheme **77**, below) [236–238]. Hence, the intact P₄ ligand in complexes **161**



Scheme 75. Ruthenium-mediated halogenation of P₄ and subsequent functionalization of the resulting P₄Cl₂ complex.



Fig. 68. X-ray structures of complexes a) 269a, b) 270, c) 271 and 272; adapted from Ref. [229].



Fig. 69. Complex $[(triphos)Ru(\mu,\eta^3:\eta^3-P_3)Ru(triphos)]^+$ (275).

and **250a**, or the free P₄ molecule behaved in an analogous manner towards **286**, but the formation of the cyclo-P₄ motif using the intact-P₄ compounds becomes more advantageous. A similar behavior was also observed for the β -diketiminato iron (215c) and cobalt (220a) complexes discussed above. DFT calculations revealed that the formation of **314** proceeds through a potential intermediate in which a butterfly-shaped P_4 motif is bridging both the ruthenium and the cobalt centers, behaving as an η^3 coordinated ligand towards the latter while still η^1 -coordinated to the former. This would be an example of a rare triphosphaallyl ligand. In such putative species, the P₄ ligand has one of the P-P bonds cleaved. These observations indicate that the P₄ tetrahedron is not released from the coordination sphere of the starting metal complexes during the reaction. The structure then rearranges to the final cyclo-P₄ moiety. The theoretical results are in good agreement with the characterization in the solid state via X-ray diffraction (Fig. 77) [81].

Recently, Power, Wolf and co-workers [242] reported on the isolation of the unusual Co_2Sn_2 heterobimetallic cluster [Ar'SnCo]₂ (**315**, Ar' = C_6H_3 -2,6(C_6H_3 -2,6-iPr₂)₂), and its reactivity towards white phosphorus (Scheme 82). Selective insertion of the P₄ tetrahedron into the Co_2Sn_2 cluster core of **315** results in opening of the white phosphorus molecule to obtain a *catena*-P₄ ligand, which has been functionalized after migration of a tin-bound terphenyl substituent to a P-atom. The product [Ar'₂Sn₂Co₂P₄] (**316**), corresponds to the first molecular cluster compound containing phosphorus, cobalt and tin. Characterization through XRD (Fig. 78) revealed bond distances in the range of single P-P bonds (2.2005(8) to 2.1621(8) Å) for the P₄-chain ligand.

As previously mentioned, back in 1991 Barr and Dahl reported the synthesis of the complex [{Cp*Co(μ , η^2 : η^2 -P₂)}₂] (**281a**) in low yield, *via* the photolytic reaction of [Cp*Co(μ -CO)₂] with P₄ [234]. In 2010, Scheer and co-workers described [243] a procedure for the quantitative synthesis of [{Cp'''Co(μ , η^2 : η^2 -P₂)}₂] (**281b**, Cp''' = 1,2,4-*t*Bu₃C₅H₂), using [(Cp'''Co)₂(μ , η^4 : η^4 -C₇H₈)] (**286**) as a starting material in the presence of white phosphorus, in toluene at room temperature. Several polyphosphorus frameworks (*e.g.* P₈, P₁₂, P₁₆, and P₂₄) were also accessed after treatment of **286** with white phosphorus and adjusting the stoichiometry and/or reaction conditions [243]. More recently, Roesky, Konchenko, Scheer, and co-workers synthesized [244] [(Cp'''Co)₂(μ , κ^3 : κ^3 : κ^2 -P₄)SmCp^P₂] (**317**, Cp^R = Cp* (η^5 -C₅Me₅) or η^5 -C₅Me₄iPr; Scheme 83), featuring



Scheme 76. Stepwise activation of P₄ at Co(I) by the groups of a) Radius [232] and b) Figueroa [233].



Fig. 70. X-ray structures of complexes **278–280**, intermediates in the stepwise activation of P_4 at cobalt; adapted from Ref. [232].

a *catena*-P₄ ligand, *via* an unusual intramolecular P–P coupling process starting from **281b**. Such polyphosphide complexes were obtained after reduction of the transition metal by the divalent samarocene $[Cp_2^RSm(THF)_n]$ $[Cp^R = Cp^* (\eta^5-C_5Me_5)$ (**24a**) or $\eta^5-C_5Me_4iPr$ (**24b**)]. DFT calculations and analysis of the bonding situation in **317** revealed that the coupling of the P₂²⁻ units in **281b** to form a new P-P bond is not the result of the direct reduction of the P-atoms, but a consequence of the rearrangement of the positive charges between the two metal atoms. Hence, the aggregation of the P₂²⁻ ligands led to the formation of an acyclic P₄⁴⁻ motif [244]. Fig. 79**a** shows the X-ray structure of **317a**.

Similar reactivity was reported in a subsequent contribution by Scheer [245], in which P-P bond formation is induced via redox reactions. By reacting **281b** with weakly coordinating anion salts of Ag(I) (Scheme 83), the formation of two new P-P bonds was observed resulting in the cyclo- P_4^{3-} cationic complex $[(Cp'''Co)_2(\mu, \mu)]$ $\eta^{4}:\eta^{4}-P_{4}$]X (**318**, X = BF₄, FAI [FAI{OC₆F₁₀(C₆F₅)}]]). If the reaction is carried out in the presence of Ag[TEF] (2 equiv., [TEF] = [Al{OC $(CF_3)_3$, another electron is abstracted from the P₄ moiety, giving rise to the planar cyclo- P_4^{2-} triple-decker compound [(Cp'''Co)₂(μ , η^4 : η^4 -P₄)][TEF]₂ (**319**). The *cyclo*-P₄²⁻ in compound **319** has identical P-P bond distances [(2.236(2) and 2.239(2) Å], which suggest the presence of single bonds. The oxidations leading to 318 and **319** are fully reversible. Treating the oxidized products with KC₈ reforms complex 281b, in accordance with the cyclic voltammetry results. The use of a small excess of KC₈ (>1 equiv.) led to the formation of a different product, namely the cyclobutadiene-like P₄ middle deck compound $[K(18-c-6)(dme)_2][(Cp'''Co)_2(\mu, \eta^4: \eta^4-P_4)]$



Fig. 71. X-ray structures of a) 283 and b) 284; adapted from Ref. [233].

(320). However, when 281b was reacted with two equivalents of KC₈, the anionic product [K(18-c-6)(dme)₂][K(18-c-6][(Cp^{'''}Co)₂(μ , $\eta^3:\eta^3-P_4$)] (321) was isolated. The reduced compound features an open-chain P₄ ligand, which was described as butadiene-like, and was reminiscent of the trinuclear dicobalt-samarium complex 317. Fig. 79b-c show the molecular structures of 320 and 321, respectively.

The chemistry of heterobimetallic complexes featuring a highly reduced P₄-chain (catena-P₄) ligand similar to the previously discussed iron complex 213 (Scheme 64, Section 3.1.1.) has been reported on by the collaboration between the Wolf and Weigand groups [207]. In contrast to the synthesis of the iron complex, monitoring of the reaction via ³¹P{¹H} NMR spectroscopy revealed selective formation of the cobalt analogue $[K(dme)_2\{(\eta^4-C_{14}H_{10})\}$ $Co(\mu, \eta^4: \eta^2-P_4)Ga(nacnac)$] (**214**) (see Scheme 64). Dark-violet crystals of **214**, suitable for XRD, were analyzed, revealing the presence of a $Co(\mu, P_4)Ga$ core similar to that of the iron analogue (P- $P_{terminal}$ = 2.1095(9) Å and 2.1057(9) Å, P-P_{internal} = 2.1825(9) Å). In this case, with respect to the iron complex 213, an interaction was observed between the $K(dme)_2^+$ moiety, the anthracene ligand and one of the phosphorus atoms in the *catena*-P₄ ligand. NMR as well as UV-Vis data on this complex agree with the molecular structure determined by XRD. Complex 214 presents structural similarities with the samarium-cobalt complex 317 [244]. Reaction of the cobaltate precursor $[K(dme)_2][Co(\eta^4-C_{14}H_{10})_2]$ (212) [211,246,247] with an excess of $[(nacnac)Ga(\kappa^2-P_4)]$ (210) did not result in the substitution of the second anthracene ligand, yielding again **214** as the sole product.

Related heterodinuclear cobalt-gallium compounds bearing a *catena*-P₄ ligand, namely [K(dme)₂{(^RBIAN)Co(μ , η^4 : κ^2 -P₄)Ga(nac-nac)}] [^RBIAN = ^{dipp}BIAN, **322a**; ^{Mes}BIAN (bis(mesitylimino)acenaph thenediimine), **322b**; the molecular structure of **322b** is shown in Fig. 80a], were later reported [209]. Complexes **322** (Scheme 84a)



Scheme 77. Synthesis and functionalization of the cyclo-P₄ complex $[Cp'''Co(\eta^4-P_4)]$ (**285**) by Scheer and co-workers.

were obtained by reaction between $[K(Et_2O){(^RBIAN)Co(\eta^4-1,5$ cod)}] [^RBIAN = ^{dipp}BIAN (296a), ^{Mes}BIAN (296b)] and [(nacnac)Ga (κ^2-P_4)] (210), through oxidative addition of a P-P bond at the cobalt center. 322b can be used in the construction of organofunctionalized polyphosphorus cyclo-P5R2 ligands upon reaction with alkylchlorophosphanes (R_nPCl_{3-n}, Scheme 84**b**). Treatment of **322b** with dialkylchlorophosphanes (R_2PCl , R = iPr, tBu, Cy) selectively affords a series of η^4 -coordinated cyclo-P₅R₂ complexes (323). Such complexes resemble species 303, reported by Wolf and co-workers (see Scheme 79, below) [240], and have the same envelope conformation as in the iron complexes **226** and **227**, by the group of Scheer (Scheme 68, above; Fig. 80b shows the molecular structure of **323a**) [215]. During the course of the reaction the [Ga(nacnac)] fragment is released. A contrasting behavior was observed upon reacting 322b with the alkyldichlorophosphane tBuPCl₂, in which a cyclo-P₅ ligand is formed (**324**), and the chloride shifts to the P-coordinated gallium fragment (Fig. 80c). Monitoring of the reaction on an NMR scale, ³¹P{¹H} VT NMR studies, and DFT calculations gave insights into the mechanism of functionalization of the catena-P₄ motif by chlorophosphanes. Upon P-P condensation with *i*Pr₂PCl at low temperature, the two reaction

intermediates were identified. Of these presumably isomeric species, the more abundant was characterized via XRD (Fig. 80d), and its molecular structure confirms that the neutral disubstituted P₅-chain complex **325** is formed. In solution (C_6D_6) at room temperature, such intermediate irreversibly affords **323**, after dissociation of [Ga(nacnac)] (Scheme 84) [209].

In 2019 [248] Figueroa and co-workers reported on the synthesis of the zero-valent tetraisocyanide complex $Co(CNAr^{Mes2})_4$ (**326**, $Ar^{Mes2} = 2,6-(2,4,6-Me_3C_6H_2)_2C_6H_3$), a sterically encumbered analogue of the binary cobalt carbonyl complex $Co(CO)_4$, which undergoes associative ligand substitution reactions. To assess on the possible one electron metalloradical behavior of **326**, its reactivity towards a variety of substrates was tested. Reaction towards elemental sulfur (S₈) or phosphorus (P₄), among other substrates, revealed that such cobalt(0) complex reacts in a multielectron manner to produce formal Co(III) centers. By treating **326** with P₄ (Scheme 85), the diamagnetic *cyclo*-P₃ complex (η^3 -P₃)Co ($CNAr^{Mes2}$)₃ (**327**) was isolated. XRD (Fig. 81) and ³¹P{¹H} NMR (singlet at $\delta = -278$ ppm) confirm the presence of the *cyclo*-P₃ ligand. It was highlighted that, unlike the analogous (η^3 -P₃)Co (CO)₃ (**328**), **327** shows remarkable thermal stability in solution



Fig. 72. X-ray structures of a) $[Cp'''Co(\eta^4-P_4)]$ (**285**) and b) the dimeric $[(Cp'''Co)_2(\mu, \kappa^4; \kappa^2; \kappa^1-P_8)]$ (**287a**); adapted from Ref. [236].

at room temperature, and is even stable for days at 60 °C, attributed to its higher steric protection afforded by the bulky isocyanide ligands [248].

3.2.1. Rhodium

In 2006, Peruzzini, Midollini, Yakhvarov and co-workers investigated the interaction between the cationic species $[M(dppm)_2]^+$ [M = Rh (329), Ir (330); dppm = 1,1-bis(diphenylphosphino)methane (Ph₂PCH₂PPh₂)] and white phosphorus [249]. In 2011, Schoeller revisited the activation of P₄ at rhodium with such a system from a theoretical point of view through DFT calculations in the gas phase [250]. In particular, the formation of the inserted product [Rh(dppm)(Ph₂PCH₂PPh₂PPPP)]OTf (**331**), in which a new P-P bond formed between a dppm ligand and the white phosphorus molecule, was evaluated. According to the calculations, due to a conformationally controlled philicity (the transition metal fragment can behave as either a nucleophile or an electrophile, after twisting the four-membered (PCP)Rh rings) the insertion reaction into P₄ would occur through a small energy barrier. Therefore, it was suggested that product **331** is obtained from an $[M]-\eta^1-P_4$ complex (332), which undergoes ligand dissociation and reaction through a series of intermediates (Scheme 86). Such a dissociation was identified as the rate-determining step. Furthermore, an equilibrium between **332** and the butterfly-P₄ complex $[M]-(\eta^1:\eta^1-P_4)$ -[M] (**333**), stabilized by π -stacking with the phenyl rings of the dppm ligands, was proposed.

Caporali, Peruzzini and co-workers [251] investigated the activation of P₄ by the dimeric Rh(I) complex [RhCl(CO)(dppm)]₂ (**334**). Reaction of the dinuclear rhodium(I) compound with white phosphorus yielded the compound [Rh₂(CO)₂(μ ,dppm)₂(μ , η^2 -P₂)] (**335**, Scheme 87), which features a bridging P₂ unit. According to

its molecular structure, the P₂ ligand is oriented perpendicularly to the intermetallic vector, forming a Rh₂P₂ tetrahedrane core. The solid state structure (Fig. 82) additionally revealed a Rh-Rh single bond (2.723 Å), and a remarkably short P-P bond (2.081 Å). It was suggested that the formation of the P₂ unit can be viewed as the oxidative addition of the P₄ molecule to the rhodium centers, which resulted in its symmetrical cleavage. VT NMR spectroscopic screening of the reaction revealed the formation of a species at -40 °C, which was proposed to be the hypothetical butterfly-P₄ intermediate complex **336**, according to its spectroscopic data. Computational and molecular orbital (MO) studies on the truncated model $[Rh_2(CO)_2\{\mu, P(CH_3)_2CH_2P(CH_3)_2\}_2(\mu, \eta^2-P_2)]$ (335') were in good agreement with the molecular structure determined for complex 335. The MO qualitative analysis indicated that the HOMO of the complex has metal-metal bond character, in accordance with XRD findings. Comparison between the P₂ ligand and the isolobal units S_2^{2-} or C_2H_2 , showed that the phosphorus motif behaved more like the disulfide anion, rather than like acetylene. Hence, the authors described the ligand as a P_2^{4-} ligand, which should be considered as an eight-electron donor, and the metal centers in the product are formally Rh(II) [251]. The complex is diamagnetic due to the formation of a Rh-Rh bond.

3.2.2. Iridium

During the period reviewed in this contribution, the chemistry of iridium complexes towards white phosphorus was scarcely explored.

However, in 2019 Wolf and co-workers [53], reported on the direct catalytic transformation of white phosphorus into arylphosphines and phosphonium salts, using the iridium complex, [Ir $(dtbbpy)(ppy)_2$]PF₆ ([**337**]PF₆; dtbbpy = 4,4'-di-tert-butyl-2,2'bipyridine, ppy = 2-(2-pyridyl)phenyl) as a catalyst (see Scheme 88). The catalytic preparation of organophosphorus compounds was performed directly from P₄ and aryl iodides (C-P bond formation) using blue-LED visible light irradiation. Synthetically valuable triarylphosphines and tetraarylphosphonium salts were obtained in yields up to around 80%. Control experiments carried out in absence of iridium catalyst [337]PF₆, visible light irradiation (455 nm) or reductant (NEt₃), confirmed the role each one of them plays in the reaction. The type of product formed was strongly dependent on the steric nature of the aryl iodide used (Scheme 88, bottom). For ortho-substituted substrates, tertiary phosphines were the major products, whereas the phosphonium salts were mainly obtained with less encumbered substrates (e.g. meta- or para-substituted aryl iodides). Steric tuning of the aryl-iodides allowed for the selective formation of secondary or primary phoswhen mesityl (2,4,6-trimethylphenyl) and phines 2.6dimesitylphenyl-iodide was used, respectively. Also, electronwithdrawing groups on the arene ring seemed to disfavor the formation of the quaternary phosphonium salt. Mechanistic experiments (³¹P{¹H} NMR scale) revealed the sequential formation of PhPH₂, Ph₂PH and Ph₃P as reaction intermediates for the model substrate phenyl iodide. The hydrogen atoms of the intermediate species are probably the result of hydrogen atom abstraction from the reductant, NEt₃. Redox potential measurements indicate that a reductive quenching mechanism, in which the excited state of the photocatalyst is reduced by NEt₃ to produce the neutral complex [337], is feasible. Individual experiments using the successive reaction intermediates instead of white phosphorus, afforded $Ph_{A}P^{+}$. with conversions similar or superior to the model reaction with P₄, and each reaction step appears to be photochemically mediated. Although such experiments gave significant insight into the reaction mechanism, the authors highlighted that further experiments are required to fully understand the initial activation path of the P₄ cage. This system represents a rare example of the direct catalytic carbofunctionalization of white phosphorus [252].



Fig. 73. X-ray structures of selected products of functionalization of [Cp^{'''}Co(η⁴-P₄)] (285): a) 288a, b) 289, c) 290, d) 291, e) 292b, f) 293b, g) 295; adapted from Refs. [119,237,238].

3.3. Group 10 metals

The coordination chemistry or activation of white phosphorus with group 10 metals has been increasingly studied in recent years, with most of the investigations involving nickel complexes [253–263], while palladium is undoubtedly the least represented. The formation of nickel or palladium phosphide nanoparticles from white phosphorus has also been described.

3.3.1. Nickel

The 17 valence electron (VE) nickel radicals [CpNi(NHC)] (NHC = IDipp (1,3-bis(2,6-diisopropylphenyl)imidazolin-2-ylidene, **338a**), IMes [1,3-bis(2,4,6-trimethylphenyl)imidazolin-2-ylidene, **338b**)] activate white phosphorus to generate the bimetallic butterfly-P₄ (P₄²⁻) complexes, {CpNi(NHC)}₂(μ,κ¹:κ¹-P₄)] (**339**) (Scheme 89a) [253,254]. The selective reaction of P₄ with **338a** occurs under mild conditions and quantitatively, and contrasts with the reactivity of the latter with sulfur or selenium, in which at least two products formed [253]. The bond distances of the butterfly-P₄ motif lie within the range of P-P single bonds (see Fig. 83a) for the molecular structure of **339b**), and its ³¹P{¹H} NMR spectrum is similar to, *e.g.*, that of species **178** [190,193,195]. The P₄ scaffold in complex **339b** undergoes reversible insertion of the heteroallene phenyl isothiocyanate (Scheme 89b) [254]. The main products obtained are the two isomeric complexes **340a** and **340b**, featuring the unusual bicyclo [3.1.0]heterohexane framework. The analogous reaction between **339a** and a large excess of PhNCS resulted in a mixture containing



Scheme 78. Synthesis of dinuclear tetraphosphido complexes by activation of P_4 at $[K(OEt_2)_2{Co(^{dipp}BIAN)(cod)}]$ (**296a**).

mainly unreacted **339a** (50%), P₄ (35%) and a product similar to **340b** (15%), according to the ³¹P{¹H} NMR spectroscopic monitoring. Both regioisomers **340** were characterized through XRD (Fig. 83**b-c**), and **340a** was isolated as an analytically pure solid. Products **340** equilibrate slowly with the starting materials **339b** and PhNCS in solution [254].

Driess and co-workers [255] showed that β-diketiminato nickel (I) complexes activate the P₄ tetrahedron affording the dinuclear Ni-complexes, $[(L^R Ni)_2(\mu, \kappa^3: \kappa^3 - P_4)]$ $(L^R = HC[CMeN(2, 6 - R_2C_6H_3)]_2;$ R = *i*Pr (**341a**), Et (**341b**); Scheme 90) [256]. Compounds **341** feature a doubly κ^3 -coordinated P₄ ligand, and were synthesized upon reacting P_4 with the corresponding β -diketiminate nickel(I) complexes $[(L^RNi)_2 \cdot toluene]$ (342) or, for 341a, the dihydride $[(L^{iPr} -$ NiH)₂ (**343**) with subsequent dihydrogen release. The κ^3 coordination of the P₄ scaffold was confirmed via XRD (see Fig. 84 for the structure of 341a). Moreover, two P-P bonds are considerably longer than typical single bonds (2.769 Å and 2.531 Å), but shorter than the sum of the van der Waals radii, suggesting activation of the P₄-cage. Dissolution of complex 341a in toluene gave rise to a paramagnetic species, contrasting with the diamagnetic behavior observed in the solid state. It was proposed that 341a underwent reversible dissociation in solution, to generate the paramagnetic complex $[L^{iPr}Ni(\eta^2-P_4)]$ (344), featuring an intact η^2 -P₄ ligand, and **342**, by dimerization of the remaining [L^{iPr}Ni¹] fragment. The presence of 344 in toluene solution was supported by high-resolution electrospray ionization mass spectrometry (HR ESI-MS) and DFT calculations. The same behavior was not observed for the less-hindered analogue 341b. Furthermore, the calculations also indicate that the activation of the P₄ moiety occurs without reduction to P_4^{2-} , therefore featuring Ni(I) metal centers in the activated product.

The symmetric cleavage of P_4 at a Ni(0) center, to yield a butterfly-Ni₂ P_2 complex, was described by the Radius group in



Fig. 74. X-ray structures of the dinuclear cobalt tetraphosphido complexes [(^{dipp}-BIAN)Co]₂(μ , η^4 : η^4 -P₄)]ⁿ⁻ a) n = 2 (**297**), b) n = 1 (**298a**) and c) n = 0 (**299a**); adapted from Ref. [239].

2013 [257]. The nickel(0) precursor $[Ni_2(iPr_2Im)_4(cod)]$ (**345**) (iPr_2 -Im = 1,3-diisopropylimidazolin-2-ylidene) reacts with P₄ to afford the butterfly-Ni₂P₂ complex $[{Ni_2(iPr_2Im)_2}_2(\mu,\eta^2:\eta^2-P_2)]$ (**346**, see Scheme 91a). The characterization of **346** in solution and in the solid state (Fig. 85), as well as optimization of its structure through DFT calculations, are all in good agreement with each other. Analysis of the frontier orbitals and comparison between the planarized and the bent Ni₂P₂ scaffold revealed that the latter is more stable, with an energy barrier of 99.93 kJ/mol. Such stabilization was assigned to a second order Jahn–Teller distortion [257]. The preparation of related Ni₂P₂ compound was later reported by Wolf, Goicoechea and co-workers, by reacting **338** with a 2-phosphaethynolate salt (Scheme 91b) [258]. The complex



Scheme 79. a) Synthesis of the *cyclo*- P_4 cobaltate anion [(PHDI)Co(η^4 - P_4)]⁻ (**300**) by activation of P_4 and b) functionalization of the P_4 moiety with chlorophosphanes followed by fragmentation/rearrangement of the cyclic P_5 ligand.

 $[{Ni(NHC)(CO)}_2(\mu,\eta^2:\eta^2-P_2)]$ (**347**, NHC = IMes; IDipp) shows a similar structure as **346**. While this complex was not prepared from P₄, its reaction with carbon monoxide (2 bar) yields P₄, presumably by dimerization of unsaturated P₂ units. The use of **346** as a transient P₂ source was demonstrated by trapping experiments with 3,4,8,9-tetramethyl-1,6-diphosphabicyclo(4.4.0)deca-3,8-diene (TMDPDD) and 2,3-dimethyl-1,3-butadiene (DMB).

Subsequently, Wolf and co-workers described the activation of white phosphorus at a Ni(0) center affording metal phosphide frameworks [259]. Following on from the reactivity of the Ni(I) complex **338** with P_4 they were able to synthesize [259] nickelphosphido clusters and the first nickel pentaphosphacyclopentadienyl complexes by reaction of the Ni(0) precursors [Ni(NHC)₂] (NHC = IMes (1,3-bis(2,4,6-trimethylphenyl)imidazolin-2-ylidene), 348a; IDipp (1,3-bis(2,6-diisopropylphenyl)imidazolin-2-ylidene), 348b) or [(IDipp)Ni($\eta^6\text{-toluene})]$ (349) with P_4 (Scheme 92). The steric encumbrance of the NHC used had a profound effect on the product formed. Upon reaction between the smaller 348a and P_4 (0.5 equiv.) in toluene, the electron-deficient trinuclear Ni₃P₄ (**350**, Scheme 92, top) was obtained, while in THF the reaction afforded the same cluster-core **350** along with Ni_3P_6 (**351**) and the dinuclear inverse sandwich Ni₂P₅ complex (352a, Scheme 92, middle). According to the Wade-Mingos rules, compounds **350** and **351** can be described as superhypercloso and hypercloso clusters. When utilizing the bulkier [Ni(IDipp)₂] (**348b**, Scheme 92, bottom) as a metal precursor, the reaction afforded the Ni₃P₈ (353) closo-cluster with a novel homoquadricyclanelike P₈ framework as well as the IDipp analogue of the cyclo-P₅-Ni compound [(IPh)₂Ni₂P₅] (**352b**) although separation of the products from the dissociated IDipp ligand proved to be difficult. The use of **349** as a starting material, in the presence of P_4 (0.5 equiv.) also afforded products 353 and 352b. The molecular structures of compounds 350-353 (Fig. 86) showed the presence of monocarbene nickel moieties on the Ni-P framework in each case. Theoretical calculations on the truncated model $[(IPh)_2Ni_2P_5]$ (352', IPh = 1,3-diphenylimidazolin-2-ylidene) were performed, and their

agreement with the experimental EPR characterization revealed that the spin density in this species is evenly distributed between the Ni-atoms.

Similarly, Weigand, Wolf and co-workers described later [260] the preparation of (NHC)Ni⁰ complexes, obtained by reacting the previously known compound $[(nacnac)Si(\kappa^2-P_4)]$ (**354**, nacnac = C $H[C(Me)N(Dipp)][C(CH_2)N(Dipp)])$ [261] with a suitable "(NHC) Ni⁰" source. Treatment of the nickel(0) synthon [(NHC)Ni(η^2 $vtms)_2$ (355, NHC = IDipp, IMes; $vtms = Me_3SiCH=CH_2$) with 354 yielded different coordination compounds, depending on the stoichiometry used (ratio 355:354 = 1:1 or 1:2). The complexes [(NHC) Ni{ $(\mu,\kappa^2:\kappa^2-P_4)$ Si(nacnac)}₂] (**356**) were obtained after reaction of the Ni(0) precursor with 2 equiv. of 354 (Scheme 93, left), while analogous reaction in 1:1 ratio afforded compound [(IDipp)Ni(µ, κ^2 : κ^2 -P₄)Si(nacnac)] (**357**, for NHC = IDipp). The structures of these complexes confirm the coordination of 354 to the (NHC)Ni(0) fragment (see Fig. 87a-b). In solution, compound 357 decomposes slowly (over a 3 weeks period), dimerizing to yield the unusual Ni₂Si₂P₈ cluster [(IDipp)Ni₂P₈{Si(nacnac)}₂] (**358**) (see Scheme 93, right). An alternative pathway for the clean formation of 358 was found, in which precursor 355 was heated to 60 °C in the presence of **354**, thus circumventing the isolation of complex **357**. The unusual structure of the cluster was determined through X-ray crystallography (see Fig. 87c), and shows strongly varying P-P bond distances (range: 2.1702(8)-3.4219(8) Å). The two "(nacnac)Si" fragments stabilize the P₈ framework.

To investigate the potential of *cyclo*-P₃ complexes as phosphorus-based building blocks, Scheer and co-workers [262] studied the reactivity of the nickel complex $[Cp'''Ni(\eta^3-P_3)]$ (**359**, $Cp''' = \eta^5$ -1,2,4-*t*Bu₃C₅H₂) towards main group nucleophiles and reducing agents (Scheme 94). Upon reaction with potassium, the anionic dinuclear complex **360**, bearing a realgar-type P₈ ligand, was obtained (Fig. 88**a** for its X-ray structure). A similar P₈-motif was also identified for the dinuclear cobalt complex **291** (see Scheme 77, above), with the main difference being that, in the latter case an additional P-P bond has been cleaved [238].



Fig. 75. Molecular structures of a) adduct 302 and selected products of functionalization of the cyclo-P₄ ligand in 300: b) 303a, c) 303c, d) 304, e) 305 and f) 306b; adapted from Ref. [240].

Treatment of 359 with nucleophiles afforded an anionic tripledecker sandwich complex (**361**), a functionalized η^2 triphosphirene species (362) and a ring expansion product (363). The formation of **361** from the reaction of **359** and NaNH₂ contrasts with the behavior of pentaphosphaferrocene (25) towards the same nucleophile, which leads to dinuclear species 227 (vide supra). Based on the results from CV experiments, the oxidation of **361** with AgOTf (OTf = $CF_3SO_3^-$) led to the formation of the neutral triple-decker compound 364. Crystallographic analysis of both triple-decker species reveals that the three-membered phosphorus ring is anionic for complex 361 and neutral for 364 and in the latter case, distorted (shown in Fig. 88b). When 359 was reacted with LiNMe₂ (Scheme 94, bottom), instead of forming a similar tripledecker complex, the highly sensitive η^2 -triphosphirene **362** was obtained as the major product. The molecular structure reveals that the cyclo-P₃ unit is coordinated to the nickel center in an η^2 fashion, while the non-coordinated P-atom bears the NMe2 substituent (see Fig. 88c for the molecular structure). Analogous reaction with LiPPh₂ led, by contrast, to the formation of the highly

heptaphosphane compound Li[(Cp'''Ni)₂($\mu,\kappa^2:\kappa^2$ sensitive (P₆PPh₂)] (**363**), which could not be crystallized. The existence of the $P_6PPh_2^-$ ligand was confirmed after protonation of **363** with HBF₄, which afforded the more stable neutral species 365 (Fig. 88d shows its X-ray structure). Remarkably, the $P_6PPh_2^-$ ligand is formed by a P₆ bicyclic scaffold substituted by an exocyclic PPh₂ group and acts as a bridge between the two nickel centers in **365**. ³¹P VT NMR studies revealed that when a reaction between the nickel cyclo-P₃ sandwich compound **359** and LiPPh₂ occurs, an intermediate analogous to the triphosphirene species 362 is formed, subsequently incorporating a second equivalent of 359 and rearranging to 363 on a rapid timescale. It was proposed that, similarly, in the reaction between **359** and NMe₂⁻ a reaggregation of the polyphosphorus ligand and addition of the nucleophile leads to the formation of an unstable $P_6NMe_2^-$ -containing complex as a minor product [262].

Yakhvarov and co-workers [263] reported on the interaction between coordinatively unsaturated organonickel σ -complexes and white phosphorus leading to the formation of arylphosphinic



Scheme 80. Cobalt complexes featuring a) rectangular P₄ ligands, by the group of Driess [82] and b) cyclo-P₄ or cyclo-P₃ ligands, by the group of Scheer [241].

acids ArP(O)(OH)H. The complexes [(bpy)Ni(Ar)(Br)] [366; bpy = 2,2'-bipyridine, Ar = 2,4,6-trimethylphenyl (Mes) or 2,4,6triisopropylphenyl (Tipp)] were activated either by reaction with a halide scavenger, such as TIPF₆, or electrochemically. Attempts to activate P₄ directly with **366** resulted to no reaction, evidencing that a vacant coordination at the nickel center site was needed. It was proposed that the P₄-cage should coordinate to the metal center, and the authors found evidence of formation of free Mes₂PH and coordinated MesPH₂ when complex 366a was used. In the chemically activated system, addition of small amounts of water or concentrated HCl (37% w/w) did not alter the type of products obtained, after 6 h. However, prolonged reaction (stirring, 6 days) afforded the arylphosphinic acid MesP(O)(OH)H in moderate yields (32%). The analogous reaction with 366b formed TippP(O)(OH)O (24% yield), without detecting Tipp₂PH, probably due to the higher steric demand of the substituents. Electrochemical activation of complexes 366, followed by treatment with P₄, gave rise to the formation of a black precipitate, attributed to nickel phosphides, which after vigorous stirring and addition of HCl (2 M) returned better yields of the arylphosphinic acids ArP(O)(OH)H (Ar = Mes, 53%; Tipp = 42%).

The use of P_4 as a soluble and highly reactive "P" source to form nickel-phosphide and other metal-phosphide nanoparticles ($M_x P_y$) has been extensively documented by Mézailles and co-workers [221,264–267]. In Section 3.1.1, the synthesis of FeP nanoparticles from iron nanoparticles and white phosphorus, was described [221]. An early example of the investigations by this group, albeit previous to the period under review, is the synthesis of nickel phosphide nanoparticles from white phosphorus [264,265]. Ni₂P (367) was obtained by reaction of Ni(0) complexes (such as [Ni (cod)₂] (**368**), cod = 1,5-cyclooctadiene) or preformed (stabilized) Ni(0) nanoparticles with P_4 when used in a 2:1 ratio (Ni/P) [265]. Ni₂P-Ni core-shell monodispersed nanocomposites were also prepared by the same group, using mild reaction conditions and P_4 as phosphorus source [266]. By reacting well-defined monodisperse Ni nanoparticles with substoichiometric amounts of white phosphorus in solution at low temperatures (<220 °C), nanoparticles composed of two crystallized phases (Ni₂P and Ni fcc) were obtained. The crystallization of Ni₂P occurs after phosphorus insertion into the Ni fcc nanoparticles, hence yielding Ni₂P cores surrounded by magnetic Ni shells. The transformation of the nanoparticles into the core-shell structure proceeds without observing other Ni_xP_y crystallized intermediates. Fine-tuning of the magnetic properties of the core-shell Ni-P system was achieved through adjustment of the amount of P₄ used [266]. Later, by contrast, another investigation revealed an interplay between the P diffusion on the nanoparticles and the crystallization phase [221]. When substoichiometric amounts of white phosphorus were used in the Ni-P system (Ni/P <2) in this case, unreacted P₄ was identified (³¹P NMR). Even by using a large excess of white phosphorus, hollow Ni₂P nanoparticles were preferentially obtained from monodispersed nickel nanoparticles in solution up to 320 °C.

Mézailles, Carenco and co-workers also described the synthesis of crystalline bimetallic phosphide Ni–Cu–P nanoparticles, which were obtained by reacting core–shell copper–nickel nanoparticles with white phosphorus [267]. The authors synthesized core–shell



Fig. 76. X-ray structures of selected β-diketiminato cobalt(cyclo-P_n) complexes: a) 307b, b) 309c, c) 310 and 307c; adapted from Refs. [82,241].

Cu_{0.25}Ni_{0.75} nanoparticles and reacted them with P₄, yielding crystalline Ni–Cu–P nanoparticles. This investigation afforded hollow monocrystalline (Ni,Cu)₂P nanoparticles together with Cu nanoparticles, which crystallized in a mixed phase isostructural to Ni₂P, using a stoichiometry aiming for full phosphidation of Ni into Ni₂P. For this system, it was possible to gain insights into the mechanism of formation of the nanoparticles. According to the proposal by this group, a metal preference of phosphorus towards nickel instead of copper was observed, yielding hollow Ni-rich phosphide nanoparticles and Cu nanoparticles. The insertion of phosphorus into the bimetallic nanoparticles triggered their amorphization and recrystallization, thus promoting an outward migration of copper to leave a void in the core. In the course of the copper migration, an amorphous intermediate of (Ni,Cu)₂P could initially form, to crystallize later into mixed phosphide. Alternatively, the formation of pure Ni₂P with subsequent evolution towards (Ni,Cu)₂P could also occur. Finally, the migrated copper crystallizes into Cu fcc nanoparticles [267].

3.3.2. Palladium

Very few reports on the chemistry of palladium towards white phosphorus were published in the period 2011–2020. The synthesis of Pd-secondary phosphine oxide compounds from P₄-derived H₃PO has been recently published in a collaboration between the groups of Yakhvarov and Hey-Hawkins [49]. However, the transformation of P₄ into H₃PO was carried out electrochemically. Subsequent reaction of the produced H₃PO with cyclic ketones afforded the secondary phosphine oxides, which were used as ligands towards palladium(II) [49].


Scheme 81. Aromatic cyclo-P₄ ligands at cobalt obtained after activation of coordinated P₄-tetrahedra.



Fig. 77. X-ray structures of a) 313 and b) 314; adapted from Ref. [81].

Nevertheless, and despite the fact that the contribution dates from 2010 and strictly speaking falls outside the scope of this review, it is worth mentioning that the group of Yakhvarov described a catalytic system for the transformation of P_4 into

 H_3PO_3 [268]. By reacting $[Pd(PPh_3)_2Br_2]$ (**369**) with P_4 , in the presence of a stoichiometric amount of NaBPh₄ in THF/water, labile cationic Pd(II)-solvato complexes were generated, and phosphorous acid was obtained. The formation of H_3PO_3 , along with the



Scheme 82. Insertion of P₄ into the framework of a cobalt-tin (Co₂Sn₂) cluster.



Fig. 78. Molecular structure of the cluster $[Ar'_2Sn_2Co_2P_4]$ (316); adapted from Ref. [242].



Scheme 83. Formation of P_4 ligands *via* coupling of P_2^{2-} ligands on **281b**.

minor products H_3PO_4 and $H_4P_2O_7$, was confirmed *via* NMR spectroscopy. The products were converted to H_3PO_4 after addition of small amounts of aqueous HCl (2 N). Mechanistic studies (NMR, MALDI-MAS) revealed that a ligand exchange of the phosphine ligands at Pd(II) by the P₄-cage (Scheme 95), followed by hydrolysis of the coordinated P₄ tetrahedron led to the major product H_3PO_3 . Furthermore, spectroscopic evidence indicated that the species $[Pd(PPh_3)_3]^{2+}$ and $[Pd(PPh_3)_2]$ were present during the reaction. Consequently, it was proposed that the oxidation of white phosphorus occurs with partial reduction of $Pd(II) \rightarrow Pd(0)$, which is maintained in solution by recoordination of the PPh₃ ligands liberated at an earlier stage in the reaction. No further evidence for the mechanistic proposal was given.

The formation of palladium phosphide nanoparticles was described by Mézailles and co-workers [221,269]. By reacting Pd nanoparticles synthesized in oleylamine with exact amounts of P_4 (0.5 equiv. or 0.1 equiv. of P_4 , respectively), both PdP_2 (370) and Pd_5P_2 (**371**) nanoparticles were selectively obtained [221]. For the crystallization of the two different phases, high temperatures were required. For nanoparticles with lower P-content, even at the high temperature used (310 °C), it was not possible to crystallize the amorphous Pd₃P (372) obtained. Hence, the authors described the amorphous plain Pd₃P nanoparticles as having a particularly high stability [221]. Further investigations revealed that the formation of crystallized palladium phosphide nanoparticles is a two-step process [269]. In a first step, amorphous Pd–P nanoparticles are formed at moderate temperatures, while crystallization occurs at higher temperatures (270-310 °C). Moreover, the crystallization temperature was found to be strongly dependent on the Pd/P ratio. The Pd-P system proved to be complex at the nanoscale, and only three crystalline phases were identified, as a result: PdP₂, Pd₅P₂, and Pd₃P. Quantitative phase-to-phase transformation from P-poor phosphides (Pd_3P and Pd_5P_2) to the P-rich PdP_2 was also demonstrated. The use of a defined molecular precursor such as P₄ resulted key for the study of a nanoscale phase diagram for this system, given that it reacts stoichiometrically to generate P (0) species under mild conditions [269].

3.3.3. Platinum

The platinum-promoted opening of a coordinated P₄-ligand was reported by Peruzzini and co-workers, as described in Section 3.1.2 (Scheme 71**a**, *vide supra*) [178,222]. The cationic complexes *trans*-[Ru(PP)₂(H)(η^{1} -P₄)]⁺ (PP = dppm or dppe, **246**), featuring intact P₄ tetrahedra, were reacted with [(C₂H₄)Pt(PPh₃)₂] (**247**) in CH₂Cl₂, to yield the heterobimetallic compounds [{Ru(PP)₂(H)}(μ , κ^{1} , κ^{2} -P₄) {Pt(PPh₃)₂]BF₄ (PP = dppm or dppe, **248**). Here the P₄ moiety has undergone activation of one P-P bond, with insertion of the platinum unit (Scheme 71**a**) [178,222]. Similar behavior was observed when the piano-stool complex was reacted with the Pt(0) precursor, yielding the P₄-butterfly compound [{CpRu(PPh₃)₂})(μ , κ^{1} , κ^{2} -



Fig. 79. X-ray structures of selected products of P-P coupling at cobalt: a) 317a, b) 320 and c) 321; adapted from Ref. [244,245].



Fig. 80. X-ray structures of a) 322b, b) 323a, c) 324 and d) 325; adapted from Ref. [209]. 75



Scheme 84. Synthesis of the *catena*- P_4 heterodinuclear complex **322b** and functionalization of the P_4 -ligand with chlorophosphanes.

P₄){Pt(PPh₃)₂]⁺ (**251**; Scheme 71**b**, *vide supra*). For the three bimetallic species, dynamic processes in solution were identified [178,222] through variable temperature ³¹P{¹H} NMR and ³¹P{¹H} EXSY experiments. The fluxional process was described as a combination of two independent motions. Complete scrambling of all of the phosphorus atoms in the P₄-cage was observed as the result of chemical exchange at the Ru-center, specifically between the Rubonded and the free P atoms of the tetrahedron (swinging or pendulum-like), without rupture of the Ru-P bond, thus passing through an η^2 -P₄{Pt(PPh₃)₂} motif. In a second process, the plat-



Fig. 81. X-ray structure of $(\eta^3$ -P₃)Co(CNAr^{Mes2})₃ (327); adapted from Ref. [248].



Scheme 86. Mechanism proposed by Schoeller for the activation of P_4 at [Rh $(dppm)_2]^*$ (329).

inum center exchanges between the three P-atoms not coordinated to ruthenium [178]. For **248a**, such a proposal was confirmed through DFT calculations on the model compound **248**', featuring $H_2PCH_2PH_2$ instead of dppm [222]. The activation parameters for the dynamic processes, calculated from the VT NMR experiments, were reported [178,222].



Scheme 85. Synthesis of a cyclo-P₃ cobalt complex by Figueroa and co-workers [248].



Scheme 87. Synthesis of the diphosphorus compound $[Rh_2(CO)_2(\mu-dppm)_2(\mu,\eta^2-P_2)]$ (**335**) by Caporali, Peruzzini and co-workers [251].



Fig. 82. X-ray structure of $[Rh_2(CO)_2(\mu\text{-}dppm)_2(\mu,\eta^2\text{-}P_2)]$ (335), adapted from Ref. [251].

The dinuclear complex $[({(dppp)Pt}_2)_2(\mu,\eta^2:\eta^2-P_2)]$ (dppp = 1, 3-bis(diphenylphosphanyl)propane, 373) was reported in 2010 by Mézailles and co-workers [270]. The formation of 373 (Scheme 96) was achieved using the Pt(0) precursor, [(dppp)Pt (C_2Ph_2) (374) in the presence of white phosphorus (0.5 equiv.). Fig. 89 shows the molecular structure of the complex, determined using X-ray diffraction. DFT calculations showed that the P₂ bridging scaffold in **373** can be considered as a P_2^{4-} ligand and, thus the metal centers have been oxidized to Pt(II). The observation of a red precipitate along with complex 373, and the theoretical studies supported the proposal that a "P₂" fragment was eliminated from the P₄ cage as a highly reduced P₂^{4–} fragment [270]. In 2013, Radius and co-workers also concluded that the P2 ligand in 373 corresponded to a P₂⁴⁻ unit, while revisiting such investigations in the frame of their own study on the related Ni₂P₂ complex [{Ni₂(*i*Pr₂- $Im_{2}_{2}(\mu,\eta^{2}:\eta^{2}-P_{2})$] (**346**), discussed above [257]. Furthermore, this group performed calculations on the platinum analogue [{Pt₂(*i*Pr₂- $Im_{2}_{2}(\mu,\eta^{2}:\eta^{2}-P_{2})$] (**375**) to determine the influence of the metal center (Ni vs. Pt) and of the ancillary ligand (NHC vs. diphosphine) on the nature of the P₂ bridge. The results indicated that the ligand does not exert a significant influence on the distribution of the frontier orbitals of the platinum complexes. Moreover, the structural features calculated for **375** are in good agreement with those found for 346, with the main difference being a longer M-P distance, attributed to the increase in atomic radius for the platinum complex. Nevertheless, it was found that for the platinum analogue, the planarized structure would be significantly more destabilized than in the case of the nickel compound. The stronger destabilization would also be caused by a second-order Jahn-Teller distortion, and by unfavorable bonding interactions between the platinum atoms and the P_2 ligand [257].

3.4. Group 11 metals

The chemistry of white phosphorus towards coinage metals has been of interest in the period under review [181–187,195,271–27 9], with most of the examples involving either copper or gold species.

3.4.1. Copper

The coordination of white phosphorus towards coinage metal salts was investigated by the groups of Green, Russell, Slattery and co-workers [271]. Suitable cationic coinage metal precursors were generated from simple inorganic salts in the presence of Lewis acids, and then reacted with P₄. Equimolar amounts of MX (for M = Cu, Au, X = Cl; for M = Ag, X = OTf) and $GaCl_3$ in dichloromethane, followed by addition of P₄, led to the formation of ioncontacted coordination polymers for copper and silver, of the general formula $(P_4MGaCl_4)_n$ [M = Cu (**376**); M = Ag (**377**)]. The molecular structure of **376** (see Fig. 90) features chains of CuGaCl₄ units linked by P₄ molecules. Each P₄ tetrahedron is coordinated through P₂ edges to a different copper center, connecting two metal centers and aggregating to form the coordination polymer. The bond lengths on the coordinated edge of the P₄ molecule feature elongated distances (P-P bond = 2.3744 Å) with respect to the free P₄ tetrahedron (2.21 Å), while the distances of uncoordinated edges (2.1742–2.1787 Å) are in the range of the distances in the latter. The product is thermally unstable and sensitive to oxygen [271].

Russell and co-workers also investigated on the coordination of the intact P₄ tetrahedron towards coinage metal-phosphine/–NHC compounds (Scheme 97a) [272]. The series of complexes [(PtBu₂-Ar)M(η^2 -P₄)]⁺ (M = Cu (**378**), Au (**379**); PtBu₂Ar = JohnPhos [PtBu₂(*o*-biphenyl)], tBuXPhos [PtBu₂(2',4',6'-triisopropylbiphe nyl)] were obtained upon treatment of a suitable (PtBu₂Ar)MX (MX = CuI, AuCl) precursor with a halide scavenger, followed by addition of one equivalent of white phosphorus. Both copper complexes present similar structural features, according to XRD (see Fig. 91 for the structure of [(JohnPhos)Cu(η^2 -P₄)]⁺, **378a**). As previously observed for analogous compounds [271,273–275], a dynamic behavior was observed on the NMR timescale for the P₄ ligand, featuring a unique singlet on ³¹P{¹H} for the tetrahedron in each example of the series of complexes.

In 2016, Lammertsma and co-workers reported on the coordination chemistry of P₄ towards coinage metal cations such as Cu⁺ and Au⁺ [276]. The commercially available *N*-heterocyclic carbene (NHC) complexes (IDipp)MCl ([M = Cu (380), Au (381); IDipp = 1,3bis(diisopropylphenyl)imidazolin-2-ylidene)] were suitable precursors, in the presence of a chloride scavenger, for the synthesis of the M-P₄ cationic adducts, $[(IDipp)M(\eta^2-P_4)]^+$ (M = Cu (**382**), Au (383); Scheme 97b). Almost simultaneously, Russell and coworkers also reported on the synthesis of complex 383 by a different synthetic approach (see Scheme 97) [272]. ³¹P{¹H} NMR spectra of the complexes revealed broad singlets (δ = -483.1 ppm for **382** and $\delta = -464.4$ ppm for **383**) indicative of fluxional behavior, similar to other η^2 -P₄ complexes. VT NMR measurements (-90 °C) confirmed the dynamic behavior in solution, for both complexes [271–273]. Computational calculations on the bonding situation for $[(IDipp)M(\eta^2-P_4)]^+$ agree with the spectroscopic observations, suggesting that the P₄ ligands remain intact, and that oxidative addition by P-P bond cleavage is disfavored. Additionally, the stability of the complexes is in agreement with the theoretical



Scheme 88. Photocatalytic direct functionalization of white phosphorus with aryl iodides using [Ir(dtbbpy)(ppy)₂]PF₆ ([**337**]PF₆). Conversions were determined by quantitative ³¹P NMR experiments using Ph₃P=O as an internal standard [53].



Scheme 89. a) Synthesis of $\{CpNi(NHC)\}_2(\mu,\kappa^1:\kappa^1-P_4)\}$ (**339**, NHC = IDipp, IMes) and b) insertion of phenyl isothiocyanate into a P-P bond (NHC = IMes).

data, with the gold complex being more stable than the copper analogue [276].

Neutral copper complexes containing intact E₄ tetrahedra (E = P, As) as ligands were reported in 2015 by the group of Scheer [277]. The neutral dicopper compounds [{(nacnac)Cu}₂(μ , η^{2} : η^{2} -E₄)]

 $[nacnac = [{N(C_6H_3iPr_2-2,6)C(Me)}_2CH]^-; E = P (384a), As (384b)]$ featuring intact E₄ tetrahedra as bridging η^2 : η^2 -ligands were synthesized and fully characterized. 384a is obtained upon reaction between [(nacnac)Cu(MeCN)] (385) and P₄ in a 2:1 ratio (Scheme 98, and Fig. 92 for the molecular structures). The complex is stable under argon in the solid state for long periods of time but decomposes slowly in solution and when exposed to light. The molecule features an almost square-planar copper center, while the P-P bonds within the tetrahedron are elongated with respect to the free white phosphorus, as revealed by crystallography. Moreover, reacting **384a** with a stronger Lewis base such as pyridine, leads to a displacement of the bridging $\eta^2: \eta^2-P_4$ ligand, releasing free P₄. By reacting **384a** with P₄, the mononuclear species $[(nacnac)Cu(\eta^2-P_4)]$ (**386**), bearing an intact side-on coordinated P₄ tetrahedron, was obtained in solution. Such a mononuclear complex can also be obtained from the copper precursor 385 and an excess of P₄. In solution, a dynamic behavior on a NMR timescale was observed for complex **386**. VT ³¹P{¹H} along with ³¹P{¹H} EXSY NMR experiments indicated that, at low temperatures (up to 213 K), the main fluxional process is the tumbling (as previously mentioned, the $\eta^1 \rightarrow \eta^2 \rightarrow \eta^1$ walk down an edge of the tetrahedron) of the coordinated P₄. At higher temperatures, dissociation of the P₄ ligand takes place. Such release results in a shifted equilibrium between the mononuclear and dinuclear complexes, 386 and 384, respectively (Scheme 98). For both complexes, DFT calculations confirm the coordination of the intact P₄ tetrahedron.

The chemistry of the chelating metalloligand $[{Cp'''Fe(CO)_2}_2(\mu, \kappa^1:\kappa^1-P_4)]$ (178b) towards copper precursors described by the



Fig. 83. X-ray structures of a) **339b** and b) and c) the bicyclo[3.1.0]heterohexane products (**340**) of insertion of phenyl isocyanate into the butterfly- P_4 motif; adapted from Ref. [254].

group of Scheer (see Section 3.1.1), demonstrated the potential of such complexes to form heterometallic complexes containing P_n ligands [195]. The same group also studied the interaction between the mixed P_nAs_m ligand complex, [{Cp'''Fe}₂($\mu,\eta^4:\eta^4-P_nAs_{4-n}$]] (**179c**, Section 3.1.1) and CuCl [192]. Upon reacting **179c** with CuCl in a 1:2 ratio, the 1D coordination polymer [{Cp'''Fe}₂($\mu,\kappa^4:\kappa^4:\kappa^1:\kappa^1-E_4$]{Cu(μ ,Cl)}₂(MeCN)]_{∞}·(CH₂Cl₂)₂ (**387**, E₄ = P_nAs_{4-n}) was obtained. According to the occupancy pattern of the E₄ moiety and the XRD molecular structure (see Fig. 93), the undulated chain of the coordination polymer is formed through interaction of the phosphorus atoms with the copper centers.

Supramolecular Cu-(P_n) (n = 2) or Cu-*cyclo*- P_n (n = 3, 5, 6) structures have also been described by the group of Scheer [144,148,181–187,278] (see Section 2.4.2). The interaction between "naked" copper ions and chromium or molybdenum complexes featuring P_2 or *cyclo*- P_n ligands [148,278], or pentaphosphaferrocene (**25**) [181–187,278] proves the potential of such compounds to act as metalloligands towards Lewis acids, thus assembling supramolecular structures. "Inorganic fullerenes" have been obtained by interaction of **25** with Cu(I) halides [181–187].



Scheme 90. Reversible activation of P₄ at Ni(I) by Driess and co-workers [256].



Fig. 84. X-ray structure of 341a; adapted from Ref. [256].

Likewise, mononuclear or dinuclear Cu(I) complexes resulted from reacting P_2Mo_2 or *cyclo*- P_3 chromium or molybdenum metalloligands with "naked" copper cations, while the interaction with **25** afforded a one-dimensional coordination polymer featuring a paddle-wheel-shaped building block [144,278].

Mézailles and co-workers described the formation of copper phosphide nanoparticles from copper nanoparticles and white phosphorus [221]. The quantitative reduction of $Cu(acac)_2$ (acac = 2,4-pentanedionate) with oleylamine yielded copper nanoparticles which were reacted with P₄, using the stoichiometric amount required (1/12 equiv. of P₄) to yield Cu₃P (**388**) nanoparticles. Under very mild conditions (100 °C, 30 min), it was possible to crystallize the nanoparticles, yielding an original core-shell Cu-Cu₃P structure. No amorphous intermediate could be observed. The mild conditions used for the crystallization of the copper nanoparticles contrast with those required for the iron (Section 3.1.1), nickel (Section 3.3.1) or palladium (Section 3.3.2) nanoparticles.

3.4.2. Silver

As previously discussed (see Section 3.4.1), upon reacting P_4 with CuCl or AgOTf in the presence of GaCl₃, coordination polymers



Scheme 91. Butterfly- Ni_2P_2 complexes by a) Radius and co-workers [257] and b) Wolf, Goicoechea and co-workers [258].



Fig. 85. X-ray structure of $[{Ni_2(iPr_2Im)_2}_2(\mu,\eta^2:\eta^2-P_2)]$ (346); adapted from Ref. [257].

of general formula (P₄MGaCl₄)_n [M = Cu (**376**); M = Ag (**377**)] were obtained [271]. The silver polymer **377** (Fig. 94), however, presents structural differences with respect to the copper analogue **376**. In this case, the P₄ tetrahedron is not part of the polymer core skeleton but coordinates to the silver metal centers in an η^2 -mode outside the main polymer net. The latter is composed of a AgGaCl₄ ladder. Coordination of the P₄ molecule in this polymer creates a distortion on the bond lengths within the tetrahedron, with the bond distance of the coordinated edge (2.3403 Å) being elongated in comparison to the rest of the P-P bonds (2.1670 to 2.2161 Å).

Scheer and co-workers have extensively studied the chemistry of P_n metalloligands towards Lewis acidic coinage metal cations, which leads to interesting supramolecular structures [144,148]. For example, organometallic-organic hybrid polymers were obtained by the interaction of $[Ag_2(110)_4]^{2+}$, [see Section 2.4.2, $[Cp_2Mo_2(CO)_4(\eta^2-P_2)]$ (110)], with the pyridine linker, di (pyridine-4-yl)ethene. Flexible coordination of the ligands, monomer–dimer equilibria, and dynamic behavior in solution were evidenced, giving rise to the formation of rectangular or 1D hybrid networks [144].



Scheme 92. Ni(0)-mediated degradation and aggregation of P_4 by Wolf and co-workers [259].

3.4.3. Gold

The interaction of P₄ with coinage metal salts in the presence of GaCl₃ afforded copper(I) and silver(I) coordination polymers **376** and **377**, as discussed in sections 3.4.1 and 3.4.2 [271]. By contrast, analogous reaction with AuCl yielded the until then unknown homoleptic cationic complex $[Au(\eta^2-P_4)_2]^+$ (**389**, Scheme 99) [273], which completed the series of coinage metal homoleptic complexes of the P₄ tetrahedron [273–275], $[M(\eta^2-P_4)_2]^+$ (M = Cu, Ag, Au). The molecular structure of **389** (Fig. 95a) reveals that, similar to the previously known examples [273-275], the cationic complex features two η^2 -coordinated distorted P₄ tetrahedra, and is ion-separated. The bond distance of coordinated P-P edge is also elongated, as in the polymers 376 and 377. Through gas-phase quantum chemical studies and a thermodynamic cycle it was possible to rationalize the differences in the formation of complex 389 in contrast with polymers 376 and 377. Weak M-P₄ interactions were determined in all cases. However, the formation of $[M(\eta^2 - P_4)_2]^+$ (M = Cu, Ag) is slightly endothermic, affording the coordination polymers 376 and 377, which are built through M-Cl linkage.

In 2016, the selective functionalization of white phosphorus by addition of ArLi to a gold complex the N-heterocyclic carbene (NHC) gold(I) complex 383 was achieved by the group of Lammertsma [276]. The synthesis of 383 (Scheme 97, vide supra), featuring an intact η^2 -coordinated P₄ tetrahedron, was already described in Section 3.4.1. above. The complex was characterized through X-ray diffraction (Fig. 95b). The Au(I) center resides in a distorted trigonal planar molecular geometry, with very similar Au-P bond lengths (2.4043 and 2.4286 Å) and an acute P1-Au1-P2 angle (57.79°). The stability of 383 allowed for additional studies regarding its reactivity. At low temperatures, 383 reacts with aryl lithium compounds (Scheme 100), yielding an asymmetrically substituted butterfly-P₄ ligand after P-C bond formation and cleavage of one P-P bond, as shown in intermediate **390**. ¹H NMR spectral analysis, as well as molecular structure determination through XRD, revealed that the reaction led to a doubly substituted RP₄ ligand. The latter coordinates two [(NHC)Au] fragments, the second of which formed after displacement of P₄ from a second equivalent



Fig. 86. Molecular structures of (NHC)Ni-phosphido clusters and inverse sandwich compound obtained by activation of P₄: a) 350, b) 351, c) 353 and d) 352b; adapted from Ref. [259].



 $\label{eq:scheme 93. Synthesis of Ni(0) complexes with bridging butterfly-P_4 ligands, and thermolysis to yield an unusual Ni_2Si_2P_8 cluster.$

of **383**. The resulting bimetallic cationic complex **391** (see Fig. 96 for the molecular structure) was obtained in high yield. The ability of the butterfly- P_4 ligand to coordinate a second [(NHC)Au] fragment was found to be comparable with the reactivity observed with metalloligand **178b** upon interaction with Cu(I), yielding

complexes **183** and **184** [195], described in Section 3.1.1. above. As mentioned before, compound **383** was reported, almost simultaneously, also by the group of Russell (see Scheme 97) [272].

Additionally, the formation of the complexes [$(PtBu_2Ar)Au(\eta^2 - P_4)$]⁺ [$PtBu_2Ar = JohnPhos$ (**379a**), tBuXPhos (**379b**)] was described



Fig. 87. X-ray structures of a) 356a, b) 357 and c) the Ni₂Si₂P₈ cluster 358; adapted from Ref. [260].

(Scheme 97b). Single-crystal characterization revealed that the coordination of the P₄ tetrahedron is affected by the steric hindrance on the phosphine co-ligand (see Fig. 95c-d). In complex 379b, the more sterically encumbered tBuXPhos ligand causes a rotation of approximately 90° on the P₄ molecule with respect to 379a, and that the P-P bond in 379a is also shorter than in 379b (2.341(3) Å versus 2.4182(8) Å, respectively). Thus, the geometry around the gold center in complex 379b was initially described as T-shaped. However, DFT calculations, along with structural data, indicate that the differences observed might be attributed to a displacement in the binding of the P₄ tetrahedron between the limits of the Au^I(η^2 -P₄) and Au^{III}(η^2 -P₄) descriptions. In the latter case, a P_4^{2-} ligand would result from the oxidative addition of a P-P bond to the gold center. It was suggested that a marked shift towards Au^{III} character and a preference to adopt planar structures could be possible for the gold complexes with respect to the copper analogues 378, which tend to be tetrahedral. In both cases the arene from the PtBu₂Ar ligands would occupy a fourth coordination site. Such studies accounted also for the fluxionality observed on the NMR timescale, suggesting free rotation of the coordinated P₄



Scheme 94. Reactivity of the neutral cyclo-P₃ complex $[Cp^{\prime\prime\prime}Ni(\eta^3-P_3)]$ (359, $Cp^{\prime\prime\prime}=\eta^5-1,2,4-tBu_3C_5H_2)$ with nucleophiles or potassium.

molecule, which should occur *via* an η^1 -coordinated M¹ intermediate with a deactivated P₄ unit.

Ragogna and co-workers [279], described gold complexes of the previously reported P₄-derivatized low-valent germanium compound [(2,6-Mes₂C₆H₃)₂Ge(κ^2 -P₄)] (**392**) [280]. The metalation of **392** yielded either monosubstituted or disubstituted species, with the coordination occurring selectively at the wing-tip P-atoms (Scheme 101). The selective coordination at the P-atoms adjacent to germanium, was confirmed by DFT calculations and single-crystal X ray analysis for the monosubstituted complex [{(2,6-Mes₂C₆H₃)₂Ge(κ^2 : κ^1 -P₄)AuCl]] (**393**, Fig. 97). Based on the theoretical analysis, the authors suggested [279] that the activation of P₄ at the germanium fragment results in a relative concentration of electron density at the wing-tip P-atoms rendering them better donors and explaining the selective coordination towards gold fragments. ³¹P{¹H} VT NMR studies revealed that **393** is more stable than the related digold compound **394**.

Nanoscaled gold phosphides were also reported by the group of Mézailles and Sanchez, using white phosphorus as a soluble phosphorus donor [281]. By reacting gold nanoparticles with P₄, in an analogous manner as previously described for Ni₂P and Cu₃P (see sections 3.3.1 and 3.4.1) [221], instead of forming nanoparticles of the metastable gold phosphide (Au₂P₃, **395**), other nanostructures containing Au₂P₃, such as nanowires, were obtained. In a first attempt to produce such nanoparticles, moderate to intermediate temperatures (up to 250 °C) were used, observing that the reaction of gold nanoparticles with P₄ was limited to their surface. The authors concluded that gold nanoparticles do not convert into gold phosphide under such conditions. However, it was possible to prepare Au-P passivated gold nanoparticles by optimizing the initial reaction parameters. Furthermore, when harsher conditions (320 °C) were used, the formation of Au_2P_3 nanowires as well as aggregation into Au₂P₃-Au nanostructures, was observed. In the latter case, the gold phosphide domains were systematically larger than the unreacted gold nanoparticles.

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Fig. 88. X-ray structures of selected products of functionalization of $[Cp'''Ni(\eta^3-P_3)]$ (**359**): a) dinuclear complex bearing a realgar-type P₈ ligand (**360**), b) neutral tripledecker **364**, c) η^2 -triphosphirene compound **362** and d) dinuclear complex featuring a protonated P₆PPh₂⁻ ligand **365** (hydrogen atoms omitted for clarity); adapted from Ref. [262].



Scheme 95. Mechanistic proposal for the Pd(II) catalyzed hydrolysis of P₄ [268].



Scheme 96. Synthesis of $[(\{(dppp)Pt\}_2)_2(\mu,\eta^2;\eta^2-P_2)]$ (373) by Mézailles and co-workers [270].

4. Surface coordination chemistry of exfoliated black phosphorus

The presence of a formal lone pair of electrons on each phosphorus atom in black phosphorus (BP) is reminiscent of the elec-



Fig. 89. X-ray structure of $[(\{(dppp)Pt\}_2)_2(\mu,\eta^2;\eta^2-P_2)]$ (373); adapted from Ref. [270].

tronic structure of the white allotrope of phosphorus, P_4 . This suggests the possibility of immobilizing metal fragments onto the BP surface. Recently, experimental studies have demonstrated that BP is able to form coordinative bonds of covalent character with the surface of metal nanoparticles grown *in situ* [282]. However, this has received limited attention both experimentally and theoretically. The very few examples reported testify that major difficulties must be overcome when converting P_4 into BP. Theoretical calculations [283] have shown that, like P_4 , BP is a much weaker Lewis base than phosphines. This is mainly attributed to



Fig. 90. X-ray crystal structure of the coordination polymer $(P_4CuGaCl_4)_n$ (376); adapted from Ref. [271].



Scheme 97. Synthesis of coinage-metals adducts of white phosphorus (M = Cu, Au) by the groups of a) Russell [272] and b) Lammertsma [276].



Fig. 91. X-ray structure of $[(JohnPhos)Cu(\eta^2-P_4)]^+$ (378a); adapted from Ref. [272].



Scheme 98. Fixation and release of intact P₄ tetrahedra at copper(I), by Scheer and co-workers [277].

the partial delocalization of the lone pairs of electrons on the phosphorus centers. Furthermore, the chair conformation of each P_{6^-} unit in the skeleton of BP implies a tilting of the lone pairs at approximately 30° from the plane of the layer (see Fig. 98).

The η^1 coordination mode to a metal fragment limits the choice of ancillary ligands on the metal center, as larger substituents will quickly increase the steric repulsion from the neighboring P atoms. Besides η^1 , η^2 and η^3 -coordination modes were identified on the BP surface (see Fig. 99). However, as lone pairs on distinct P-atoms are not perfectly convergent, these geometries do not possess an optimal orbital overlap. This could be overcome at the cost of some local distortion of the BP surface. However, this is difficult due to its rigid structure.

Taking into account both the low donating ability and its steric requirements, the task of developing the coordination chemistry of BP seems very challenging. Only small and labile coordination complexes should be considered, to make the displacement of a co-ligand by BP in the coordination sphere of a metal more facile.

4.1. Black phosphorus functionalization with naked adsorbed metal atoms (adatoms)

One particular approach for metal surface decoration that avoids the need for ligands displacement is the use of naked metal atoms. One such technique is physical vapor deposition (PVD), whereby a metal is brought to the vapor phase and neutral atoms are deposited onto a target substrate (i.e. BP). If the interaction with the target material is strong enough to prevent aggregation, isolated atoms may be immobilized on its surface or can even diffuse into the lattice. The effect of adsorbed metal atoms (called adatoms) on the BP layers has been addressed theoretically and can span from the tuning of electronic and transport properties of BP to an increased affinity for selected gas molecules. Kulish and co-workers [284] carried out a first principle study on 20 different adatoms on monolayer BP. All the investigated metals preferred an η^3 coordination mode on BP with Ni, Co, Pd and Pt having the strongest adsorption energies, with Pt reaching a maximum value of - 4.71 eV. Notably, it has been shown experimentally (XAS, XPS) that M(0) systems, where M = Pd, Pt, in the form of nanoparticles, can interact strongly via a covalent bond with the BP surface [282,285]. Further studies on M_{adatom}/BP from other groups corroborated these trends [286-288]. S.P. Koenig et al. [289] used direct current sputtering to deposit Cu adatoms on a thin BP flake exfoliated mechanically. The latter was used to build a field effect transistor (FET). Ab initio calculations were carried out



Fig. 92. X-ray structures of copper-diketiminato complexes featuring η²-coordinated intact P₄ ligands; adapted from Ref. [277].



Fig. 93. Molecular structure of the 1D coordination polymer $[{Cp'''Fe}_2(\mu, \kappa^4: \kappa^4: k^1:k^1-E_4)[Cu(\mu, Cl)]_2(MeCN)]_{\infty} (CH_2Cl_2)_2$ (**387**, $E_4 = P_n As_{4-E_4} = P_n As_{4-R_4}$; adapted from Ref. [192].



Fig. 94. X-ray crystal structure of the coordination polymer (P₄AgGaCl₄)_n (377); adapted from Ref. [271].

on a Cu/four-layer-BP model system, considering both adsorbed (η^3) and intercalated Cu atoms (see Fig. 100).

In both these configurations, Cu behaves as an electron donor, transferring one electron to BP and leaving a stable closed shell Cu⁺ cation. This turns BP into an n-type semiconductor, with a remarkable electron mobility of 2140 cm²V⁻¹s⁻¹, measured at low temperature. Z. Guo and co-workers [290] tackled the limited durability of BP-based FETs, caused by surface oxidation, by func-



Scheme 99. Synthesis of the homoleptic complex $[Au(\eta^2-P_4)_2]GaCl_4$ (**389**).

tionalizing the BP surface with silver. However, instead of using Ag (0) atoms as with PVD, Ag⁺ cations were anchored to the surface of BP by a chemical procedure. First, BP nanosheets (lateral dimension ~300 nm) were exfoliated mechanically and transferred onto a Si/SiO₂ wafer. Then, BP was exposed to Ag⁺ ions by dipping the silicon wafer into a solution of AgNO₃ in NMP (N-Methyl-2-pyrrolidone). XPS measurement revealed that Ag⁺ had been successfully immobilized on the BP surface (see Fig. 101**a**). The region around 133 eV in the P 2p XPS spectrum of BP is quite diagnostic for direct metal-phosphorus bonding (see also references [282] and [285]). Thus, a deshielded peak at 133.0 eV was assigned to the P-Ag bonds (Fig. 101**b**).

First principle calculations carried out on this system showed that Ag^+ adsorbs to the BP surface by the η^3 coordination mode, interacting with three distinct lone pairs of electrons (see Fig. 102), as was the case of isoelectronic Cu⁺ described previously. However, Ag^+ ions behave as a p-type dopant for BP, drawing electron density away from the layer. The coordination of silver ions not only enhanced the transistor stability, leaving the flake morphology unaltered after several days of air exposure, but it also improved the transistor performance, doubling the hole mobility.



Scheme 100. Functionalization of P_4 by reaction of **383** with aryl lithium compounds.

4.2. Black phosphorus functionalization with a titanium complex

In 2016, Y. Zhao and co-workers [291] were the first to study the functionalization of exfoliated BP with a transition metal fragment, namely TiL₄ (L = p-toluenesulfonate). Ultrasmall BP nanosheets were prepared following a solvent-assisted procedure, carried out in NMP, previously described by the same group [292].



Fig. 95. X-ray crystal structures of cationic gold complexes featuring η^2 -coordinated intact P₄ ligands: a) [Au(η^2 -P₄)_2]⁺ (**389**), b) [(IDipp)Au(η^2 -P₄)]⁺ (**383**), c) [(JohnPhos)Au(η^2 -P₄)]⁺ (**379b**); adapted from Refs. [271,272,276].



Fig. 96. X-ray crystal structure of the functionalized butterfly-P₄ gold complex **391**; adapted from Ref. [276].

By this method, BP quantum dots (BPQDs) with a lateral size of 2. 6 ± 1.8 nm and a thickness of 1.5 ± 0.6 nm were obtained. The exfoliated material was then stirred with TiL₄ in NMP for 15 h to obtain TiL₄@BP. XPS measurements of the isolated material confirmed the presence of titanium and the broad peak at binding energy 132.4 eV in the P 2p XPS spectrum was assigned to direct P–Ti bonds (see Fig. 103**d**).



Scheme 101. Functionalization of an activated- P_4 germanium compound by coordination to Au(I).



 $\label{eq:Fig. 97. X-ray crystal structure of [{(2,6-Mes_2C_6H_3)_2Ge(\kappa^2:\kappa^1-P_4)AuCl}] \mbox{ (393); adapted from Ref. [279].}$



Fig. 98. a) Top and b) side view of the phosphorene lone pairs. Image b) clearly shows the tilting of the lone pairs with respect to the plane of the layer and the misalignment of two adjacent lone pairs due to P-P bond torsion of 60°; adapted from Ref. [283].

However, no further studies were carried out to shed light on the structure of this adduct and it is unclear how BP is interacting with the metal (i.e. if it has displaced one of the sulfonate ligand to give a charged L₃Ti⁺–P adduct). NMR measurements performed on a TiL₄@BP suspension, although corroborating the persistency of sulfonate coligands in the collected material, did not help to clarify this point. A remarkable effect of the functionalization with TiL₄ was the increased stability of BPQDs towards ambient conditions. To compare the behavior of pristine BP with TiL₄@BP, dilute water suspension of the two were prepared. The suspensions were then exposed to air and their absorbance in the UV/VIS range was monitored over time. As shown in Fig. 104, the absorbance of bare BP at 450 nm decreased by 55% of the original value after 72 h, compared to the 5% for TiL₄@BP. It should be noted that when organic solvent solutions of BP and TiL₄@BP were exposed to air, they showed similar stability (see Fig. 104d).

A follow-up to this work [293] has been the biomedical application of TiL₄@BP, driven by the increasing attention devoted to BP in the life sciences. In 2015, M. Pumera and co-workers [294] demonstrated the low cytotoxicity of 2D BP and found it to be concentration-dependent and between that of graphene oxide and transition metals dichalcogenides. Moreover, BP is biodegradable, since in physiological medium, i.e. in presence of water and oxygen, it oxidizes forming phosphates [295,296], which are naturally present in the human body. Additionally, BP has been shown to be an efficient photosensitizer with superior quantum yields and a pronounced ability to generate singlet oxygen $({}^{1}O_{2})$ [297], opening it up to applications in photothermal and photoacoustic (PA) therapy. TiL₄@BP has been used as a contrast agent for *in vivo* PA imaging of cancer. The method is based on the adsorption of near-IR light from a contrast agent to image a tumor location. Combining the enhanced stability imparted to BP by the functionalization with TiL₄ and its high near-IR extinction coefficient, TiL₄@BP outperformed previously reported nanostructured contrast agents such as bare Au nanorods [298]. In vivo experiments were performed with TiL₄@BP as a PA contrast agent to image tumor area in nude mice (Balb/C nude) previously inoculated with MCF-7 cancer cells. Remarkably, after injecting TiL₄@BP, the mice showed larger PA signals in the tumor than pre-injection, the maximum intensity being reached after 4 h (see Fig. 105). This further demonstrated the high potential of TiL₄@BP in clinical applications as a PA contrast agent, characterized by both a selective accumulation at tumor sites and a remarkable spatial resolution.

4.3. Black phosphorus functionalization with cisplatin derivatives

Since the seminal work of B. Rosenberg in 1965 [299], the cisplatin (*cis*-diamminedichloroplatinum, **396**) family has been lar-



Fig. 99. The three potential bonding modes of BP towards a metal fragment. a) η^1 coordination with formation of single P-M bond, b) η^2 coordination and c) η^3 coordination; adapted from Ref. [283].



Fig. 100. a) Schematic drawing of the two possible location of Cu: adsorbed on the surface (Cu_{ad}) and interstitial (Cu_{int}) . b) Charge density difference of the Cu_{ad} . The red and blue regions refer to electron depletion and accumulation, respectively. c) Calculated band structure of pristine four-layer BP, four-layer BP with adsorbed Cu (Cu_{ad}) , and four-layer BP with interstitial Cu (Cu_{int}) . The dashed line indicates the Femi level for each case. Reprinted with permission from Fig. 1 in Ref. [289]. Copyright 2016 American Chemical Society.



Fig. 101. a) P 2p and b) Ag 3d XPS spectra of bare BP and Ag*/BP, recorded after 3 days of air exposure. Reprinted with permission from Fig. 1 in Ref. [290]. Copyright 2017 John Wiley and Sons.

gely studied for its antitumor activity and it is still used today to treat several cancers, including testicular, ovarian, cervical and lung cancer, either alone or in combination with other antitumor agents.

Searching for a metal fragment with powerful anticancer activity, M. Fojtů *et al.* [300] used cisplatin (CP, **396**) and oxaliplatin (OP, **397**; see Fig. 106) to modify BP and use it for drug delivery.

A well-characterized human ovarian cell line A2780, derived from tumor tissue of an untreated patient, was used as a target in *in vitro* studies. As both **396** and **397** are known to be effective towards this type of cancer, the goal was to highlight a potential enhancement in cytotoxicity induced by the 2D BP carrier. BP nanosheets were prepared by liquid exfoliation in NMP. The exfoliated material, having lateral dimensions in the range 100–500 nm and a thickness of few tenths of nm, was afterwards dispersed in water, **396** or **397** were added, and the mixture was incubated for 24 h. The functionalized BP nanosheets, BPCP (BP modified with **396**) and BPOP (BP modified with **397**), were isolated and characterized. The P 2p XPS spectra, beside the characteristic signal of pristine BP, displayed a smaller broad peak slightly above



Fig. 102. a) Scheme illustrating adsorption of Ag⁺ on BP. b) Three different views of the BP/Ag⁺ structure. Adapted from Ref. [290].

~133.5 eV possibly due to Pt–P bonds. Both BPCP and BPOP featured a double contribution in their Pt 4f spectra, suggesting the presence of two distinct Pt(II) species, the minor component being slightly more electron-depleted (see Fig. 107**b**). In the case of BPOP, this contribution amounted to 30% of total Pt in the sample. The nature of these species however was not further investigated.

In vitro studies revealed that while cisplatin activity was almost unaffected by the presence of BP, the cytotoxicity of oxaliplatin towards ovarian cells was significantly increased, reaching a maximum enhancement of 21% when the BPOP concentration used was 30 μ g mL⁻¹. As in all the experiments performed the total concentration of **397** was the same, only the amount of BP being varied, the superior suppression of cancer cell growth is ascribed solely to the BP. Though no detailed investigations of the mechanism were carried out, the authors attributed the cytotoxicity enhancement to a major cellular uptake of **397** once bound to the BP shuttle and the different behavior of **396** is due to its supposed weaker interaction with BP.

J. Zhang *et al.* [301], in disagreement with the findings from M. Fojtů and co-workers described above [300], also tried to immobilize cisplatin on 2D BP but no Pt incorporation could be detected by XPS. Thus they switched to a cisplatin derivative, namely [Pt $(NH_3)_2(NO_3)_2$] (**398**). Given the non-coordinating behavior of NO_3^- , the charged moiety $[Pt(NH_3)_2]^{2+}$, being highly electrophilic,

should then interact better with the surface of the BP nanosheets. BP was reacted with Pt-NO₃ in water/NMP under an inert atmosphere for 18 h. High resolution Pt 4f XPS confirmed the successful incorporation of platinum, featuring Pt 4f_{7/2} and 4f_{5/2} peaks at 73.0 and 76.3 eV respectively. However, no shifts in peak positions were observed compared to the precursor **398**. Since d⁸ Pt²⁺ favors a square planar coordination, an η^2 coordination of BP would be expected with the two NO₃ as counteranions. Remarkably, along-side an increased environmental stability compared to pristine BP, BP nanosheets functionalized with **398** showed a significantly greater cytotoxicity and apoptotic-promoting capability towards cisplatin resistant cell-lines A2780 and HepG2. This was proposed to be the synergetic effect of BP, which previous works demonstrated to be efficient in damaging cancer even when it is unfunctionalized [302].

4.4. Black phosphorus functionalization with lanthanide complexes

Following the approach outlined for surface modification of BP with TiL₄ (see Section 4.2), L. Wu et al. [303] attempted the metal functionalization of BP nanosheets using lanthanide complexes. Lanthanides have large atomic radii, which should allow for an easier interaction with the lone pairs of electrons of the phosphorus atoms. As in the case of TiL₄, a sulfonate anion was chosen as an ancillary ligand, since it was thought suitable to enhance the electrophilicity of the metal, promoting the coordination to BP. A variety of LnL_3 species (with L = trifluoromethanesulfonate and Ln = Gd, Y, La, Nd, Sm, Eu, Tb, Er, see Fig. 108) were tested in the surface functionalization of BPQDs [292] having lateral dimension 2.6 ± 1.8 nm and thickness 1.5 ± 0.6 nm. A suspension of BPQDs in NMP was stirred at room temperature under an inert atmosphere with an excess of LnL₃. By this method it was possible to decorate the BP surface with lanthanide pendants, as confirmed by EDS analysis of the isolated LnL₃@BPQDs (see Fig. 108b).

Although theoretical investigations were not carried out on the system, the authors propose that LnL₃ species will coordinate to three phosphorus atoms on the BP surface (η^3 -BP), as shown in Fig. 109. This sounds reasonable considering the large atomic radius of lanthanide metals, which favors a larger hapticity of BP. Such coordination has been shown with Lu³⁺ [67] and Y³⁺ [63] complexes bearing a trihapto coordinated *cyclo*-P₃ ligand, derived



Fig. 103. Fabrication and characterization of TiL₄-decorated BPQDs (named TiL₄@BP). a) Synthesis and structural formula of TiL₄; b) surface coordination of TiL₄ to BP; c) TEM image with inset HRTEM image of TiL₄@BP; d) P 2p and e) Ti 2p XPS spectra of TiL₄@BP, acquired after 3 days of air exposure. Reprinted with permission from Fig. 1 in Ref. [291]. Copyright 2016 John Wiley and Sons.



Fig. 104. Stability of bare BP and TiL₄@BP in water. a) Absorption spectra of bare BP and b) TiL₄@BP dispersed in water after exposure to air for 0, 6, 24, 48, and 72 h; c) variation of the absorption ratios (A/A₀) at 450 nm of BP and TiL₄@BP over time, in water; d) A/A₀ of TiL₄@BP over time in EtOH and DMF. Reproduced with permission from Fig. 2 in Ref. [291]. Copyright 2016 John Wiley and Sons.



Fig. 105. *In vivo* PA performances of TiL₄@BP. a) Time-dependent PA images of the MCF-7 cells in the xenografted tumor in the BALB/c nude mice after intravenous injection of TiL₄@BP. The signal intensity bar is shown on the right side. b) Quantitative analysis of each ROI signal in a). Maximum intensity is recorded after 4 h. c) Typical 3D PA images of the tumor taken at different times pre- and post-injection of TiL₄@BP. Reproduced with permission from Fig. 6 in Ref. [293]. Copyright 2017 John Wiley and Sons.



Fig. 106. Molecular structures of a) cisplatin (396) and b) oxaliplatin (397).

from their interaction with white phosphorus. XPS studies of GdL₃/ BPQDs at the P 2p core level highlighted the presence of an electron-depleted phosphorus species at 132.9 eV, assigned to the P-atoms bonded to the electrophilic GdL_3 fragment (Fig. 108d). Furthermore, an appreciable red-shift is observed in the Gd 3d XPS spectrum compared to free GdL_3 (Fig. 108c), in accordance with the donation of electron density from the σ orbitals of BP to Gd. As with the TiL₄ functionalization, LnL₃@BPQDs showed an enhanced stability towards air and water with respect to bare BPQDs, as revealed by UV–Vis absorption measurements carried out on aqueous suspensions aged over 8 days.

This functionalization strategy was extended from BPQDs to BP nanosheets having an average size of 300 nm. The environmental stability of the obtained LnL₃@BP was evaluated *via* TEM. As reported in Fig. 110, while bare BP nanosheets were fully degraded after 8 days of exposure, GdL₃@BP kept its morphology. Accordingly, recording UV/Vis absorption measurements in aqueous sus-



Fig. 107. a) Core level P 2p XPS spectra of pristine BP, BPCP and BPOP and b) relative Pt 4f core-level. All scans were referenced to adventitious C 1s peak at 284.5 eV. Reproduced with permission from Fig. 6 in Ref. [300]. Copyright 2017 John Wiley and Sons.



Fig. 108. Characterization of GdL₃-coordinated BPQDs (named GdL₃@BPQDs). a) TEM image and HRTEM inset; b) EDS spectra; c) Gd 4d XPS spectra of GdL₃@BPQDs and free GdL₃; d) P 2p XPS spectrum of GdL₃@BPQDs. Reproduced with permission from Fig. 1 in Ref. [303]. Copyright 2018 John Wiley and Sons.

pensions of LnL₃@BP over time, only a minimal loss in absorbance was observed, meaning the stabilization effect is not related to flake dimensions (BPQDs or BP nanosheets). Furthermore, this remarkable stabilizing effect was common to all the lanthanides under study.

Depending on the lanthanide selected, the series $LnL_3@BPQDs$ exhibited different physicochemical properties: samples containing Eu^{3+} , Nd^{3+} and Tb^{3+} possessed fluorescence, while Gd@BPQDs showed magnetic resonance responsiveness. The fluorescence spectra of the free LnL_3 species were almost superimposable to



Fig. 109. Atomistic model of the adduct between BP and LnL₃ fragment, L = OTf = Tri-fluoromethanesulphonate; adapted from Ref. [303].



Fig. 110. Stability evaluation of bare BP nanosheets and $GdL_3@BP$. TEM images of a) $GdL_3@BP$ nanosheets and b) bare BP nanosheets exposed to air for 0, 4, and 8 days. Reproduced with permission from Fig. 3 in Ref. [303]. Copyright 2018 John Wiley and Sons.

those recorded for LnL₃@BPQDs. The absence of any shift upon coordination was proposed to be due to the electronic transitions responsible for fluorescence only concerning the 4f levels of the metal, whereas the coordination to BP is expected to involve only 4d orbitals. Paramagnetic Gd³⁺ is commonly employed as an MRI contrast agent in commercial solutions, such as Magnevist, Dotarem and Primovist. All these formulations contain Gd³⁺ chelates. As they are paramagnetic, when in a physiological medium the Gd³⁺ species interacts with the protons of water molecules, reducing their T₁ NMR relaxation times. Image contrast in MRI is based on differences in the (average) relaxation times of tissues, thus contrast agents enhance resolution. Considering the properties of Gd³⁺ and the biocompatibility of BP, preliminary studies were carried out on aqueous suspensions of GdL₃@BPQDs, highlighting a linear increase in $1/T_1$ intensities with concentration. Further studies are needed to fully exploit the potential of GdL₃@-BPQDs in this field.

5. Conclusion and perspectives

The present survey on the coordination chemistry of elemental phosphorus towards transition metals summarized the impressive results achieved in the last decade in the activation of P₄ at a metal center and gives an initial review of the surface coordination of exfoliated black phosphorus (BP). Sections 2 and 3 described a rich variety of phosphorus-containing structural motifs, such as P₄ activation, degradation, aggregation products, functionalized compounds, and nanostructured phosphides. Stabilization of such structures is achieved through an array of spectator ligands which include cyclopentadienyl derivatives, the extended family of β diketiminato ligands, bulky isocyanides, amides, (chelating) phosphines and N-heterocyclic carbenes (NHCs). In metal complexes, the nuclearity of the polyphosphido ligand obtained from P₄ is generally dictated by the electron count of the metal center. Nevertheless, the electronic and steric properties of the ancillary ligands can play an important role in guiding structure and reactivity (see for example the differently substituted cyclopentadienyl iron complexes in Section 3.1.1, and the substitution of trimethylphosphine with trimethylphosphine as a ligand in ruthenium complexes described in Section 3.1.2). Furthermore, while the coordination and initial activation of P₄ at TMs can already be considered an established and mature field, the functionalization of the activated phosphorus motifs has been much less explored and the release of the resulting P_n moieties remains a highly desirable research objective. This is especially relevant for the further development of phosphorus-based synthesis on an industrial level. Nevertheless, the chemistry presented in these sections provides further, although still rather limited, details into the mechanisms of activation of white phosphorus by transition metals. In recent work, the long-envisioned target of a catalytic functionalization of white phosphorus has been achieved.

Finally, the rediscovery of the black allotrope, whose surface behaves as a polydentate phosphine ligand, brings the reader briefly to the world of materials science. Section 4 summarizes its promising reactivity towards unsaturated transition metal complexes, which represents the starting point not only for new heterogeneous catalysts but also for new 2D phosphorus-based materials endowed with unique properties.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgments

M.C. thanks the Italian Ministry for University and Research (MUR) for funding the project PRIN 2017 KFY7XF FERMAT "Fast ElectRon dynamics in novel hybrid-2D MATerials". Thanks are also expressed to Italmatch Chemicals (Arese, Italy) for financing a doctoral grant to L.G. R.W. would like to thank a number of dedicated former and present co-workers for their vital contributions to the group's research in the field. Their names are in the references. In addition, generous financial support by the European Research Council (CoG 772299) is gratefully acknowledged.

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