Abstract: The synthesis of cyclometalated osmium complexes is usually more complicated than of other transition metals such as Ni, Pd, Pt, Rh, where cyclometalation reactions readily occur via direct activation of C–H bonds. It differs also from their ruthenium analogs. Cyclometalation for osmium usually occurs under more severe conditions, in polar solvents, using specific precursors, stronger acids, or bases. Such requirements expand reaction mechanisms to electrophilic activation, transmetalation, and oxidative addition, often involving C–H bond activations. Osmacycles exhibit specific applications in homogeneous catalysis, photophysics, bioelectrocatalysis and are studied as anticancer agents. This review describes major synthetic pathways to osmacycles and related compounds and discusses their practical applications.

Keywords: osmium; cyclometalation; osmium cyclometalated complex; osmacycle; pincer complex; catalysis; photophysics; anticancer activity

1. Introduction

Osmium is the heaviest and rarest stable element in the earth’s crust, with a concentration of approximately 0.05 part per billion. Perhaps the latter accounts for why the coordination and organometallic chemistry of osmium is less advanced than that of other transition metals. Common precursors, such as OsCl$_3$ or M$_2$[OsCl$_6$], are significantly more expensive than their respective ruthenium precursors. Osmium is often associated with highly toxic OsO$_4$, widely used as an oxidizing agent in organic chemistry and as a staining agent in transmission and scanning electron microscopies [1–3]. Fortunately, osmium chemistry is not limited to its oxides, and numerous coordination, organometallic, and cluster compounds have been prepared, though their chemistry is not as deeply explored as that of ruthenium [4]. The osmium chemistry was not extensively reviewed in the literature. The first review was written by Griffith in 1987 [5]. Some osmium compounds were also compiled in *Gmelin Handbuch der Anorganischen Chemie* in the 1980s [6]. In the 2011 edition of the *Encyclopedia of Inorganic Chemistry*, two articles were devoted to coordination and organometallic chemistry of osmium [7,8]. To our knowledge, the chemistry of cyclometalated osmium species was not specifically reviewed, in contrast with the extended coverage of the chemistry of other metalacycles in the last several decades [9–23].

This review attempts to fill the existing gap in covering the cyclometalation chemistry of osmium compounds as they are presented in Figure 1 with M = Os. The metal is usually σ-bonded to a carbon atom and coordinated to E, where E is generally N, P, O, S, As, Se. Undoubtedly, the impact of Michel Pfeffer’s work on the development of cyclometalation reactions was fundamental, and many of his publications, first on palladium then on ruthenium metalacycles, set the path for countless publications [24–27].
2. Historical Background

Ruthenium and osmium belong to group 8 metals, and they frequently form similar complexes. The oxidation states vary from 0 to VIII, though more common are oxidation states between II to IV [4]. Osmium forms stronger metal-metal and metal-carbon bonds, and hence osmium complexes are generally less active catalysts than analogous ruthenium species. Many osmium complexes are made via different or more difficult routes than their ruthenium analogs. Publications on osmacycles started to grow after 2000, prior to which they were nearly nonexistent. Their number is considerably lower than the number of reports on ruthenacycles (Figure 2).

The first osmacycle prepared by Bruce in 1973 was made from \([\text{Os}_3(\text{CO})_{12}]\) and benzo[h]quinoline. A mixture of mononuclear isomers 1 was obtained in low yields after 230 h at reflux in light petroleum [28]. In 1981, Jameson described cyclometalated formazan derivatives of osmium 2 [29]. Desrosiers prepared a hydride osmium–phosphine complex 3 by intramolecular C–H activation in 1986 [30]. Cyclometalated piano-stool derivative of benzoic acid 4 was reported by Kisenyl in 1987 [31]. Later, mononuclear and dinuclear
pincer osmium complexes 5, 6, dinuclear Ru-Os cyclometalated derivative 7 [32,33], cationic mononuclear osma(II)cycle with PPh3 8 [34], and osmacycle 9 from polyquinoline [35] were obtained. Structures of compound 5–11 are shown in Table 1.

**Table 1.** First osmacycles prepared between 1973 and 1997.

| Precursor                                      | Cyclometalated Ligand | Osmacycle                                      | Ref.   |
|------------------------------------------------|-----------------------|------------------------------------------------|--------|
| [Os$_3$(CO)$_{12}$]                            |                       | ![Image](image1.png)                             | [28]   |
| [OsH(O$_2$CCF$_3$)(CO)(PPh$_3$)$_2$]           | ![Image](image2.png)  | ![Image](image3.png)                             | [29]   |
| cis-[Os(PMe$_3$)$_4$H(CH$_2$C(Me)$_3$)]        | PMe$_3$               | ![Image](image4.png)                             | [30]   |
| [Os(p-cymene)Me(Cl)(Me$_2$SO)] prepared from [Os(p-cymene)Cl$_2$]$_2$ | ![Image](image5.png)  | ![Image](image6.png)                             | [31]   |
| [Os(tterpy)(O)$_2$(OH)(NO$_3$)] prepared from K$_2$[Os(O)$_2$(OH)$_4$] | ![Image](image7.png)  | ![Image](image8.png)                             | [32]   |
Molecules 2021, 26, x FOR PEER REVIEW 4 of 52

Table 1. Cont.

| Precursor | Cyclometalated Ligand | Osmacycle | Ref. |
|-----------|-----------------------|-----------|------|
| [Os(tterpy)(O)₂(OH)(NO₃)] prepared from K₂[Os(O)₂(OH)₄] | ![Cyclometalated Ligand](image1) | ![Osmacycle](image2) | [32] |
| [Os(tterpy)Cl₃] prepared from [OsCl₃] | ![Cyclometalated Ligand](image3) | ![Osmacycle](image4) | [33] |
| [Os(Cp)(PPh₃)₂]OTf | ![Cyclometalated Ligand](image5) | ![Osmacycle](image6) | [34] |
| [Os(N=N=N)Cl₃] prepared from [OsCl₃] | ![Cyclometalated Ligand](image7) | ![Osmacycle](image8) | [35] |

* tterpy = 4′-(4-tolyl)-2,2′,6′,2″-terpyridine.

3. Main Precursors

3.1. [OsX₂(PR₃)₃]

Complexes [OsX₂(PR₃)₃], a family of 16 electron osmium(II) precursors, are widely used for the synthesis of osmacycles. Among them, [OsBr₂(PPh₃)₃] is particularly valuable.

3.1.1. mono-Osmacycles

Chakravorty synthesized phenolate osmium complex 11 from 2,6-diformyl-4-methylphenol and [OsBr₂(PPh₃)₃] [36]. The authors proposed the oxidative addition of the aldehyde, followed by decarbonylation, to form an unstable OsIV intermediate. Subsequent reductive elimination of HX yields the cyclometalated complex [36]. Related [OsBr(C=O)(PPh₃)₂(CO)] complexes 12 were similarly prepared, and the substitution of bromide generated cationic species 13 (Figure 3) [37].
The mechanism of cyclometalation of 2-vinylpyridine by \([\text{MCl}_2(\text{PPh}_3)_3]\) (\(\text{M} = \text{Ru, Os}\)) was explored theoretically for the ruthenium case \[38\]. The lower reactivity of osmium allowed to isolate several intermediates proposed. The reaction of \([\text{OsCl}_2(\text{PPh}_3)_3]\) with an excess of 2-vinylpyridine in the presence of \(\text{Cs}_2\text{CO}_3\) and \(\text{NaBF}_4\) results in cyclometalation of the ligand with liberation of \(\text{PPh}_3\) to produce 14. The cationic complex 15 is formed via substitution of the chloride by a second equivalent of 2-vinylpyridine in presence of \(\text{NaBF}_4\). In the absence of \(\text{NaBF}_4\), the neutral cyclometalated complex 14 generates isomers 16 and 17 (Figure 4) \[38\].

Wen and co-workers prepared cyclometalated \([\text{Os}(\text{P~C~P})\text{Cl}(\text{PPh}_3)]\) complexes 25 and 26 (Figure 6) \[40\]. They react with \(\text{H}_2\) to afford dihydrido species. The corresponding carbonyl derivatives can also be prepared in the presence of \(\text{CO}\) \[41\].

Figure 3. Synthesis of C–O cyclometalated complexes.

Figure 4. Cyclometalated complexes derived from 2-vinylpyridine.
3.1.2. Pincer and bis-Cyclometalated Complexes

[OsBr₂(PPh₃)₃] reacts with 2-(2',6'-dimethylphenylazo)-4-methylphenol in refluxing 2-methoxiethanol in the presence of NEt₃ to produce osmium(III) complex 24. The authors consider a coordination of the ligand to osmium(II) in a tridentate C–N–O fashion, followed by the oxidation to osmium(III) in 18 (Figure 5). Thereafter, the loss of HX affords a carbene species 19. In the presence of water, the carbene intermediate could transform into the OsH–C=O fragment (22) after elimination of H₂ in intermediate 21. A posterior reductive elimination of a half equivalent of hydrogen from 22, followed by the migration of a CO fragment (23), would produce the final cyclometalated complex [Os(C–N–O)(CO)] (24) [39].

![Figure 5. Synthesis of C–N–O pincer complexes.](image)

Wen and co-workers prepared cyclometalated [Os(P–C–P)Cl(PPh₃)] complexes 25 and 26 (Figure 6) [40]. They react with H₂ to afford dihydrido species. The corresponding carbonyl derivatives can also be prepared in the presence of CO [41].

![Figure 6. Synthesis of P–C–P pincer complex from 1,3-(R₂PCH₂)₂C₆H₄.](image)

The diphosphine [1,3,5-(CH₃)₃-2,6-(tPr₂-PCH₂)₂C₆H] was also reacted with [OsCl₂(PPh₃)₃]. The metal promotes the cleavage of one C–CH₃ bond with liberation of methane (Figure 7). In the presence of H₂, dihydride heptacoordinated complex 27 is formed, while [OsH(Cl)(PPh₃)₃] generates a pentacoordinated complex 28 [41].
The diphosphine {1,3,5-(CH$_3$)$_3$-2,6-(iPr$_2$-PCH$_2$)$_2$C$_6$H} was also reacted with [OsCl$_2$(PPh$_3$)$_3$]. The metal promotes the cleavage of one C−CH$_3$ bond with liberation of methane (Figure 7). In the presence of H$_2$, dihydride heptacoordinated complex 27 is formed, while [OsH(Cl)(PPh$_3$)$_3$] generates a pentacoordinated complex 28 [41].

Figure 8. Synthetic route for a series of P~C~P pincer complexes by a double C−H activation.

Other examples involving C−H activation are shown in Figure 9. Majumder et al. synthesized osmacycles 36 and 37 from 2-(arylazo)phenols and [OsBr$_2$(PPh$_3$)$_3$] [43], and Das found an unusual cyclometalation of N-arylbenzohydroxamic acids leading to 38 [44].

Bis-pincer complex 39 was obtained from [OsCl$_2$(PPh$_3$)$_3$] and diphosphine 2,6-(CH$_2$PPh$_2$)$_2$C$_6$H$_3$ at a ligand-to-complex ratio of 3:1 (Route 1 in Figure 10). On the other hand, the ratio of 1:1 provides mono-pincer complex 25. An exchange of chloride by triflate affords 40 and a subsequent reaction with the second equivalent of the ligand leads to bis-pincer complex 39 (Route 2 in Figure 10) [45].
Other examples involving C–H activation are shown in Figure 9. Majumder et al. synthesized osmacycles from 2-(arylazo)phenols and [OsBr₂(PPh₃)₃] [43], and N-N-arylbenzohydroxamic acids leading to similar [48] (Figure 11). Osmafuran (Route 2 in Figure 10) [45]. The efficient cyclometalation to give osmabenzenes were prepared in a similar way [47]. The η²-alkyne-coordinated alcohol complex 41 can be transformed under acidic conditions to a η²-coordinated α,β-unsaturated ketone 42. Additionally, 41 reacted with triphenylphosphine to give 43 in presence of Bu₄NX (X = Cl, Br) with elimination of OH. The efficient cyclometalation to give 45 with η²-coordinated allyl alcohol was also performed. Upon thermal treatment, 45 converts to a mixture of four conjugated osmacycles (42, 46–48) [48]. Osmafuran 49 was prepared similarly [48] (Figure 11).

3.1.3. Others Osmacycles

Gong studied the interaction between HCC==C(H)CH=CH₂ and [OsCl₂(PPh₃)₃] in THF leading to η²-allyl alcohol osmacycle complexes, which could further be converted to osmabenzenes, cyclic osmium η²-allene, osmafuran and α,β-unsaturated ketone complexes in a one-pot reaction [46] (Figure 11). It is worth mentioning that the “donor center” in such osmacycles is a C=C double bond, narrowly fitting into the definition of a metalacycle. Nevertheless, the authors classify the complexes as osmacycles. The cyclometalation takes place through a nucleophilic attack of PPh₃ at the coordinated alkyne [47]. This method results in efficient preparations of conjugated osmacycles in high yields and allows the synthesis of complexes containing a phosphonium moiety incorporated in the metalacycle. Osmabenzenes were prepared in a similar way [47]. The η²-alkyne-coordinated alcohol complex 41 can be transformed under acidic conditions to a η²-coordinated α,β-unsaturated ketone 42. Additionally, 41 reacted with triphenylphosphine to give 43 in presence of Bu₄NX (X = Cl, Br) with elimination of OH. The efficient cyclometalation to give 45 with η²-coordinated allyl alcohol was also performed. Upon thermal treatment, 45 converts to a mixture of four conjugated osmacycles (42, 46–48) [48]. Osmafuran 49 was prepared similarly [48] (Figure 11).
3.2. \([\text{OsX}_2(\text{CO})(\text{PR}_3)_3]\)

Related precursors \([\text{OsX}_2(\text{CO})(\text{PR}_3)_2]\) were also explored. The oxidative addition of the ortho C–H bond of the phenyl group of imine \(\text{Ph}_2\text{C}=\text{NH}\) to \([\text{Os(Cl)}(\text{Ph})(\text{PPr}_3)_2(\text{CO})]\) followed by the reductive elimination of benzene and the coordination of the iminic nitrogen generates osmacycle 50 (Figure 12). The process could be viewed as an unusual \(\sigma\)-bond metathesis [49].

![Figure 11. Unusual osmacycles prepared by activation of alkynes.](image)

![Figure 12. Synthesis of cyclometalated complex by oxidative addition of the phenyl group of an imine and subsequent reductive elimination.](image)

Clark et al. studied by X-ray crystallography four cyclometalated osmium complexes (51–55) obtained via the transmetalation reaction between hydride precursors \([\text{OsH(Cl)}(\text{CO})(\text{PPh}_3)_3]\) and organomercurial derivatives of 2-phenylpyridine. A series of related complexes (56–58) in which the phenylpyridine ligand was modified by electrophilic substitutions was also reported (Figure 13) [50].
3.3. Osmium Hydride Complexes

Esteruelas and coworkers have extensively been working on C–H bond activations mediated by transition metals. Their results represent an important step in the development of new synthetic routes through C–H bond activations by osmium. Notably, osmium hydrides complexes favor high oxidation states (4 and 6) and display a wider range of stoichiometries and structures than ruthenium derivatives. Osmium hydride complexes can also exhibit lability of the hydride ligands under thermal conditions [53]. Particularly, the hexahydride osmium(VI) precursor, [OsH$_6$(P$_3$Pr$_2$)$_2$], was widely used to prepare a variety of osmacycles. For example, this hexahydride osmium complex is capable of

![Figure 13. Osmium(II) complexes with cyclometalated phenylpyridines.](image)

Bennett et al. also showed that the reaction between [Hg(o-C$_6$H$_4$PPh$_2$)$_2$] and [MH(Cl)(CO)(PPh$_3$)$_3$] (M = Ru, Os) form the corresponding four-membered metalacycles 59, elemental mercury and free triphenylphosphine (Figure 14) [51]. It was suggested that the reaction occurs through the oxidative addition of Hg–Ar to a coordinatively unsaturated intermediate generated by the dissociation of PPh$_3$ and the posterior reductive elimination of ArH and Hg [52].

![Figure 14. Use of organomercurial derivative to obtain cyclometalated complexes.](image)
generating a tetrahydride [OsIVH4(P^3Pr3)2] species and one hydrogen molecule by thermal activation [53]. The tetrahydride complex can then coordinate the oxygen atom of the carbonyl group of an aromatic ketone and subsequently activates C−H or C−F bonds for the corresponding cyclometalation step [54].

3.3.1. [OsH6(PR3)2]

Studies by NMR revealed that [OsH6(PR3)2] reacts with cycloalkyl or phenyl methyl ketone to form cyclometalated compounds 60, 61 and 62 (Table 2) [49,50,54,55]. The reactivity of 61 and related complexes will be discussed in Section 4.2. The reaction of [OsH6(PR3)2] with 2-vinylpiridine affords 63, and a secondary C−H bond activation generates complex 64 [56]. The C(sp^3)−H activation occurs when [OsH6(PR3)2] reacts with 8-methylquinoline. Activation of the methyl group and elimination of two H2 molecules gives 65. Experiments with methyl deuterated 8-methylquinoline and DFT calculations confirmed the mechanism [57].

Table 2. Osmacycles obtained from [OsH6(P^3Pr3)2].

| Cyclometalated Ligand | Osmacycle | Yield | Ref. |
|-----------------------|-----------|-------|-----|
| L9                    | ![Image](image1.png) | 62%   | [55]|
| L10                   | ![Image](image2.png) | 61%   | [54]|
| L11                   | ![Image](image3.png) | 62% (X = H, R = Ph) 74% (X = F, R = CH3) 60% (X = F, R = Ph) | [54]|
| L12                   | ![Image](image4.png) | 92%   | [56]|


Table 2. Cont.

| Cyclometalated Ligand | Osmacycle | Yield | Ref. |
|-----------------------|-----------|-------|------|
| ![Image](image1) | ![Image](image2) | 55%   | [56] |
| ![Image](image3) | ![Image](image4) | 97%   | [57] |
| ![Image](image5) | ![Image](image6) | 23% (E = N) | [58] |
| ![Image](image7) | ![Image](image8) | 60% (X = F, R = Ph) | [58] |
| ![Image](image9) | ![Image](image10) | 45% (E = CH) | [58] |
| ![Image](image11) | ![Image](image12) | 50% | [59] |
| ![Image](image13) | ![Image](image14) | 91% | [59] |
| ![Image](image15) | ![Image](image16) | 99% | [59] |
Table 2. Cont.

| Cyclometalated Ligand | Osmacycle | Yield | Ref. |
|-----------------------|-----------|-------|------|
| ![L20](image)          | ![71](image) | 90%   | [59] |
| ![L21](image)          | ![72](image) | 60%   | [60] |
| ![L22](image) ![L23](image) ![L24](image) | ![73](image) | 71% (L22) | [61] |
| ![L25](image)          | ![74](image) | 64%   | [62] |
| ![L26](image)          | ![75](image) | 80%   | [62] |
| ![L27](image)          | ![76](image) | 53%   | [63] |
| ![L27](image)          | ![77](image) | 60%   | [63] |
The same hexahydride osmium precursor in the presence of 0.5 eq of 2,6-bis[1-[(4-methylphenyl)imino]ethyl]pyridine is involved in a double C−H activation of the pyridine ring producing a dimetallaticyclic system 66 (Table 2). The product was characterized by an X-ray study and the presence of three fused rings was confirmed [58]. Other bimetallic osmacycles were prepared from [OsH₆(PPr₃)₂] by activation of the ortho C−H bond of N,N’-di- and tetr phenyl bipyridines. Starting from 2-phenylpyrididine, compound 67 was first obtained (Table 2). Using 2,2’-diphenyl-4,4’-bipyridine in 1:3 and 3:1 ratios with respect to osmium allowed to isolate mononuclear 68 and dinuclear 69 osmacycles, respectively (Table 2). The corresponding phenyl and anthracenyl derivatives of bipyridine also form dinuclear osmacycles (70, 71) [59]. Complex [OsH₆(PPr₃)₂] and 2-azetidinones substituted at 4-position by an N-heterocycle and in 3-position by a phenoxy group afford corresponding cyclometalated complexes (72, 73). This reaction is associated with the metal-mediated degradation of the ligand which involves two C−H bond activations and a parallel C−N and C−C rupture within the four-membered rings [60,61] (Table 2). Double cyclometalation takes place when [OsH₆(PPr₃)₂] reacts with 2,6-diphenylpyridine to give tridentate C−N−C pincer compound 74. In the presence of HBF₄ the latter can suffer the addition of one proton at the metalated carbon to form the cationic mono-cyclometalated complex, which eliminates 2,6-diphenylpyridine in the presence of acetonitrile to generate the corresponding coordination derivative. When the ligand is 2-phenoxy-6-phenylpyridine, a double C−H activation also occurs. Five- and six-membered rings 75 are produced (Table 2) [62]. Other reactions with [OsH₆(PPr₃)₂] involve the cyclometalation of azole ligands by selective C−H bond activation which can discern the coordination of N over “N−H side” in the ligand (76, 77). Such behavior is also observed with 2-phenylthiazole and 2-phenylbenzothiazole, where the cyclometalation occurs at the “C−N side” [63]. The reaction of the hexahydride precursor with benzonitrile in toluene promotes the C−H activation and the reduction of the triple C≡N bond forming the corresponding imine. The resulting complex 78 is produced in a 75% yield. The substitution of PPr₃ by PPh₃ in such complexes was also studied [64]. Complex [OsH₆(PPr₃)₂] can also activate directly
the 8-position in 2-, 3-, 6- and 7-methylquinolines to obtain corresponding [OsH₅κ²-C₈₅N-(quinilinyl)](P₂Pr₃)₂ and [OsH₅κ²-C₈₅N-(quinilinyl-η-Me)](P₂Pr₃)₂ (79, n = 2, 3, 6, 7) complexes. It is interesting to note that a four-membered cyclometalated fragment is formed in all cases (Table 2) [65]. Complex [OsH₄(P₂Pr₃)₂] reacts with 2-vinylpyridine to form an osmacycle 80 where C(sp²)-H activation of the vinyl is observed (Table 2) [56].

A curious reductive cyclometalation occurs when the dihydride 16e OsIV complex [OsH₂Cl₂(P₂Pr₃)₂] reacts with an excess of allylamine. Cyclometalated OsII species 81 is formed alongside with the [HP₂Pr₃]Cl salt (Figure 15). The final product is coordinated by two molecules of allyl amine. One of them is cyclometalated through the insertion of the alkene fragment into the Os–H bond. The mechanism proposed by the authors suggests a C–H activation to justify the [HP₂Pr₃]Cl formation. This cyclometalated complex 81 is the starting material to form a variety of other complexes by either ligand metathesis or CO and allene insertions [66].

Figure 15. Reaction between [OsH₂Cl₂(P₂Pr₃)₂] and allylamine.

The cyclometalation of helicenes occurs similarly (Figure 16). [OsH₂Cl₂(P₂Pr₃)₂] reacts with the ligand and the cyclometalation to form 82 is accompanied by the reduction of the metal center to OsIV in refluxing toluene [67].

Figure 16. Cyclometalation via by C–H activation assisted by reductive elimination of hydride ligands.

3.3.2. *bis*- and *tris*-Osmacycles

Several osmacycles that contain two osmium–carbon σ-bonds (*bis*-cyclometalated), as well as some rare examples with three osmium–carbon σ-bonds (*tris*-cyclometalated) were prepared from [OsH₄(P₂Pr₃)₂]. The first examples were classified by Esteruelas et al. as "multiple C–H bond activations". Complex [OsH₄(P₂Pr₃)₂] reacts with phenyl-substituted pyrimidines or triazines to form the corresponding [Os(C–N–C)] pincer derivatives. The reaction with phenyl pyrimidines leads to mono-cyclometalated complexes along with the pincer species. The phenyl triazine ligand forms pincer complex 85 but can also generate a dinuclear *bis*-cyclometalated osmium complex 86 (Figure 17) [68].
More recently, Esteruelas et al. described tris-cyclometalated osmium complex 87 produced from 3 eq of \(N,N\)-diphenylbenzimidazolium chloride [Ph\(_2\)BImH]Cl and [OsH\(_6\)(P\(_3\)Pr\(_3\))\(_2\)] in the presence of a base. In acidic medium (HBF\(_4\)), one cyclometalated phenyl group of 87 is protonated and a reductive elimination of hydride with another phenyl group forms a cationic octaedral complex 88 stabilized by two agostic interactions between osmium and C–H bonds (Figure 18) [69]. The replacement of chloride by BF\(_4^-\) in the \(N,N\)-diphenylbenzimidazolium salt promoted the tridentate coordination to form 89. Monocyclometalated complex 90 was made using of one equivalent of [Ph\(_2\)BImH]Cl. However, complex 87 could be obtained through the subsequent reaction with another equivalent of the ligand [69].

![Figure 17. Synthesis of bis-cyclometalated complexes from [OsH\(_6\)(P\(_3\)Pr\(_3\))\(_2\)].](image1)

![Figure 18. Synthetic routes to tris-cyclometalated complexes from monocyclometalated derivatives.](image2)
3.4. $[\text{Os}(\eta^6\text{-arene})\text{Cl}_2]_2$

The $[\text{Os}(\eta^6\text{-arene})\text{Cl}_2]_2$ dimers usually display similar reactivity as their widely used ruthenium analogues. The bridging chloride bonds are readily broken in polar solvents or in the presence of coordinating ligands. Monomers formed were used to prepare a variety of osmacycles [70].

3.4.1. mono-Osmacycles

In 2003, piano-stool neutral 92 and cationic 93 osmacycles were prepared from the dimeric $[\text{Os}(\eta^6\text{-C}_6\text{H}_6)(\mu\text{-Cl})\text{Cl}_2]$ precursor (Figure 19) [71]. We are particularly proud that the first experiments were carried out in the lab of Michel Pfeffer in Strasbourg. Needless to say that background for this work was previously created by Michel through the thorough investigation of the chemistry of matching ruthenium complexes [72–75]. The thorough investigation of the chemistry of matching ruthenium complexes [72–75]. Coordinated benzene and acetonitrile in 93 are easily substituted by 2,2’-bipyridines or 1,10-phenanthroline to form octahedral cationic complexes (94, 95, 96) [71]. Complex 92 can also be synethetized through a transmetalation reaction with organomercurial derivatives and the $[\text{Os}(\eta^6\text{-C}_6\text{H}_6)(\mu\text{-Cl})\text{Cl}_2]$ dimer [71]. Similarly, series of complexes bearing cyclometalated 2-phenylpyridine and $N,N$-dimethylbenzylamines with different electron-donating groups such as 4-OMe (97b) and 3,5-(OMe)$_2$ (97c) were prepared with the aim to modulate the electron density at the osmium center [76].

![Figure 19. Cyclometalated osmium complexes obtained from $[\text{Os}(\eta^6\text{-C}_6\text{H}_6)(\mu\text{-Cl})\text{Cl}_2]$.](image)

Transmetalation between $[\text{Os}(\eta^6\text{-C}_6\text{H}_6)(\mu\text{-Cl})\text{Cl}_2]$ and silver(I) compounds is exemplified by the reaction of the 1-phenyl-3-methyl-1H-benzimidazolium $[\text{AgPhMeBIm}]$ complex. Intermediate 98 forms first followed by the metalation of the phenyl group in the presence of $\text{Al}_2\text{O}_3$. Piano-stool mono-cyclometalated complex 99 is the final product (Figure 20) [77]. It takes two days for triazolium silver salts to react with $[\text{Os}(\eta^6\text{-C}_6\text{H}_6)(\mu\text{-Cl})\text{Cl}_2]$ to give structurally similar complex 100 after treatment with KPF$_6$ [78]. This methodology was also successfully employed to prepare cyclometalated complexes of iridium and ruthenium [78].
3.4.2. *bis*- and *tris*-Osmacycles

A few *bis*- and *tris*-cyclometalated osmium complexes have been described [79,80]. Such compounds differ from pincer complexes, as they contain two or three cyclometalated ligands, respectively. Complexes \[\text{[Os}(\text{phpy})_2(\text{bpy})]\text{PF}_6 (102)\] and \[\text{[Os}(\text{phpy})_3] (101)\] were prepared from organomercurial Hg(phpy)_2 and *mono*-cyclometalated complex 92. Their electrochemical properties were compared with those of the corresponding coordination complex \[\text{[Os}(\text{bpy})_3](\text{PF}_6)_2 (103)\] and *mono*-cyclometalated complex \[\text{[Os}(\text{phpy})(\text{bpy})_2]PF_6 (94)\], (Figure 19). Each Os–C bond decreases the OsII/OsIII reduction potential by ca. 500 mV ranging from 888 mV for coordination compound 103 to −958 mV (vs. Ag/AgCl) for *tris*-cyclometalated complex 101 [79,80] (Figure 21).

**Figure 20.** Synthesis of *mono*-cyclometalated complexes through transmetalation reactions with silver(I) derivatives.

**Figure 21.** Synthesis of *bis*- and *tris*-cyclometalated complexes (top) and cyclic voltammograms showing the influence of the number of Os–C bonds on the OsII/OsIII reduction potentials. Bottom: cyclic voltammograms of 103, 94, 101 and 102 (1 mM) in MeCN: 25 °C, scan rate 0.1 V s\(^{-1}\), 0.1 M \(n\text{Bu}_4\text{NPF}_6\), glassy carbon electrode. Inset shows a plot of the formal reduction potential vs. the number of C donors.
Imidazolium complex 99 can be converted to a complex with four acetonitrile ligands 105, which undergoes a second transmetalation with the organosilver compound to give 106. The latter affords tris-cyclometalated complex 107 as shown in Figure 22 [77].

3.5. Other Precursors

Complexes [cation]2[OsX₆] and [OsX₃(terpy)] (terpy = 4′-(4-tolyl)-2,2′:6′,2″-terpyridine) were also used as osmacycle precursors. Complex [Et₄N]2[OsCl₆] reacts with 1,3-(CH₂P₂Bu₂)₂C₆H₄ in the presence of NEt₃ affording 16e pincer complexes [OsCl(P~C~P)(CO)] (108) and [OsCl(H₂)(P~C~P)] (109) [81]. It is suggested that Os⁴⁺ is first reduced to Os²⁺ by methanol to produce CO. When the osmium precursor and the ligand are heated in 2-propanol, the solvent is not decarbonylated and dihydride complex 109 is obtained (Figure 23) [81].
Complex [Et4N]2[OsCl6] reacts also with 1,5-bis(di-terbutylphosphino)pentane in the presence of NEt3 and H2 in 2-pentanol to give P–C–P alkyl complex 110. A posterior thermal treatment initiates second C–H bond activation via α-elimination to give carbene complex 111 (Figure 24) [82].

An interesting cycloosmation occurs within alkenyl–allenylidene–acetonitrile complex 109 which can form an osmacyclopentapyrrole at reflux in acetonitrile (Figure 25). The proposed mechanism is presented in Figure 25 [83].
Various Os\textsuperscript{III} bis(N-methylbenzimidazolyl)benzene or 1,3-di(pyridin-2-yl)benzene complexes were prepared from [OsCl\textsubscript{3}(Mebip)] (118) (Mebip = bis(N-methylbenzimidazolyl)pyridine) or [Os(ttpy)Cl\textsubscript{3}] (121) (ttpy = 4′-(4-tolyl)-2,2′:6′,2″-terpyridine) as shown in Figure 26 [84].

Figure 26. Synthesis of bis-pincer complexes.

4. Representative Reactions of Osmacycles
4.1. Reactivity of Chelate and Pincer Complexes

Ligand substitution in 59a (see also Figure 14) yields neutral mono-osmacycle 124 [51]. On the other hand, ruthenium analogue 59b displayed a different behavior since a parallel insertion of CO into the Ru-C bond in addition to ligand substitution accounts for the formation of 126 (Figure 27). In order to force the CO insertion into the Os-C bond, complex 59a was treated with AgSbF\textsubscript{6}. However, the cationic species 125, product of the substitution of the chloride ligand, was obtained [51].

Figure 27. Reactions of four-membered metalacycles.
Reactions of 2-phenylpyridine osma(II)cycle 51 are summarized in Figure 28 [50]. The cyclometalated ligand is activated by OsII towards electrophilic substitution allowing the introduction of diverse functional groups. Nitration occurs at both the 4- and 6-position of the phenyl ring, whereas bromination takes place at the 4-position only. Complex 51 reacts also with CO2/H+ or Bu3SnCl as shown in Figure 28 [50].

Gusev et al. studied the reactivity of P~C~P pincer compounds (Figure 29) [81]. Complex 108 reacts with NaBH4 and the mixture of two hydrides 128 and 129 is obtained. A chloride for iodide exchange takes place in 109 in the presence of MeI. The reaction of 109 with NaBH4 gives trihydride 131. Complex 132 with two H2 molecules is also obtained in the presence of H2 at −80 °C; at higher temperatures one of the coordinated H2 undergoes oxidative addition to yield complex 133 [81].

Reactions of P~C~P pincer complex 25 with a series of alkynes afford carbene and carbyne osmacycles (Figure 30) [40]. Complex 25 is also convertible to monohydride 141 and trihydride 142 in the presence of NaH (Figure 30). In an excess of phenylacetylene, 142 gives 143 with three incorporated phenylacetylene ligands. Remarkably, the coordination of all three is different including acetylide- and vinylidene-binding motifs. Vinylidene complex 135 reacts with Tl(OAc) to form 144. Chloride is being replaced by acetate which acts as a bidentate ligand [85].
Figure 29. Reactivity of P~C~P pincer complexes 108 and 109.

Figure 30. Reactions between osmium P~C~P pincer complex 25 with alkynes.

Figure 31. Reactions of P~C~P pincer complex 26 with H2 and CO.
Related P–C–P pincer complex 26 reacts with H₂ giving rise to dihydride species 145. Carbonyl derivatives 146 and 147 can also be prepared in the presence of CO (Figure 31) [41].

Figure 31. Reactions of P–C–P pincer complex 26 with H₂ and CO.

Dihydride [OsH₂(P²Pr₃)₂(C=C′~N)] (73) reacts with HBF₄ to yield [OsH₂(P³Pr₃)₂(C=C″~N)]BF₄ (148) where the carbon in α position to the central carbon of the pincer ligand was protonated by the acidic medium [60]. Similarly, C~N~C′ pincer complexes 74 and 75 accept one proton at a metalated carbon to form cationic monocy- cyclometalated complexes, which further react with nitriles as shown in Figure 32 [62].

Figure 32. Protonation of the cyclometalated fragment by HBF₄.
4.2. Reactions of Hydride Complexes

Hydrides in osmacycles promote reactions of adjacent ligands. For example, complex 80 reacts with HBF₄ to give 63 with regenerated 2-vinylpyridine, the Os-trihydride unit being transformed into the Os-hydride-dihydrogen functionality. Subsequent treatment of 63 at 50 °C in the presence of Ph₂CO induces the reduction of the double bond and formation of 64. The elimination of benzophenone in acetonitrile at room temperature gives 153 [56]. In the case of trihydride complex 61 (Figure 33), addition of HBF₄, however, affords dihydrogen derivative 154, which is converted to cationic complexes (155, 156) in the presence of NaCl or CsF, the cyclometalated fragment being unaffected [86].

![Figure 33. Reactions of hydride complexes with HBF₄.](image)

4.3. Ligand Substitution in Osmacycles

Piano-stool osmacycle 93 is a versatile precursor for a series of octahedral 2,2'-bipyridine, 1,10-phenanthroline and MeCN complexes (Figure 34). The substitutions are due to the lability of the coordinated benzene in polar solvents such as MeCN and MeOH. Cyclometalated N,N-dimethylbenzylamine derivatives 97 behave similarly affording complexes 159–162 [71,76,79,80].

Complex 44 reacts with PMe₃ and benzonitrile leading to cationic osmacycles. New species such as 167 with a η²-allene ligand are formed. Osmabenzene derivatives are also produced in many instances [87]. The reaction pathways of 44 are summarized in Figure 35.

Interesting ring expansions of the cyclometalated fragment take place when 49 reacts with HC≡CCH(OH)Ph or PhC≡CH to give complexes 170–174 (Figure 36) [48].
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Figure 34. Synthetic routes to octahedral cyclometalated derivatives by ligand substitution starting from piano-stool complex 93.

Complex 44 reacts with PMe3 and benzonitrile leading to cationic osmacycles. New species such as 167 with a $\eta^2$-allene ligand are formed. Osmabenzene derivatives are also produced in many instances[87]. The reaction pathways of 44 are summarized in Figure 35.

Interesting ring expansions of the cyclometalated fragment take place when 49 reacts with HC≡CCH(OH)Ph or PhC≡CH to give complexes 170–174 (Figure 36)[48].

Figure 35. Studies of reactivity of complex 44.
5. Applications of Osmacycles

Osmacycles find applications in many areas including homogeneous catalysts; they are used as chemical sensors, luminescent materials, and anticancer agents. Relevant examples are summarized below.

5.1. Catalysis

Although ruthenium and palladium cyclometalated complexes have been widely investigated in homogeneous catalysis [19–21,88–90], osmium analogues are not as widely used. Perhaps they have not yet demonstrated their potential. However, there are a few examples where osmacycles show good catalytic activity and look like promising candidates for new applications. In particular, binding of H₂ to 25 affords 175 with η²-bound H₂ (Figure 37) which might be a key intermediate in catalytic hydrogenation [91].

Figure 36. Ring-expansion reaction from a five-membered to a nine-membered osmacycle.

Hexamerization of phenylacetylene at pincer complex 25 is a promising reaction which gives a mixture of isomers of diosmium complex 176 with a µ-1,2-bis(η²-cyclopentadienyl)-1,2-diphenylethane bridging ligand (Figure 38) [92].

Figure 37. Coordination of dihydrogen to a pincer complex.

Figure 38. Hexamerization of phenylacetylene at osmium pincer complex.
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Figure 37. Coordination of dihydrogen to a pincer complex.

Hexamerization of phenylacetylene at pincer complex 25 is a promising reaction which gives a mixture of isomers of diosmium complex 176 with a \( \mu \)-1,2-bis(\text{η}_5\text{cyclopentadienyl})-1,2-diphenylethane bridging ligand (Figure 38) [92].

Figure 38. Hexamerization of phenylacetylene at osmium pincer complex.

Baratta et al. contributed significantly to the homogeneous catalysis by cyclometalated Ru and Os complexes. Benzo[\( h \)]quinoline C–N–N osmium pincers were tested as hydrogenation catalysts. Three osmium complexes [OsX(C–N–N)(dppb)] (177) (\( X = \text{Cl}, \text{H}, \text{OR} \); dppb = 1,4-bis(diphenylphosphino)butane) were studied in catalytic hydrogen transfer from isopropanol to asymmetric ketones. Using a catalyst charge of 0.005% mol, conversions above 94% in less than 30 min with TOFs of \( 10^5 – 10^6 \) h\(^{-1} \) were achieved. The ruthenium analogues of 177 were slightly more active for the same process, with conversions above 97% within 10 min, presenting TOFs values in the order of \( 10^6 \) h\(^{-1} \) [93]. Chiral ligands in 178 were tested for the asymmetric hydrogenation of prochiral ketones and conversions up to 92% with 74–94% enantiomeric excess (\( ee \)) were reported [94]. Structural modifications of the benzo[\( h \)]quinoline ligands gave two new series of chiral osmium complexes 179. The complexes with a Josiphos ligand were used in the hydrogenation of acetophenone with conversions up to 97% with TOFs around \( 10^4 \) h\(^{-1} \) and \( ee \) up to 86%. In the same work, the ruthenium analogue of 179a was also studied. Conversions up to 95% with TOFs around \( 10^4 \) h\(^{-1} \) and \( ee \) between 90 and 99% were obtained. However, in the case of ruthenium the reaction was performed at 40 °C instead of 70 °C for the osmium catalyst [95]. Different C–N–N pincer ligands were also used, keeping the Josiphos ligand system in complexes 180. These were used for catalytic ketone reduction. Conversions were above 93% with TOFs \( 10^4 – 10^5 \) h\(^{-1} \) and \( ee \) between 90 and 99%. The ruthenium analogues were also obtained, and the catalytic activity for both metals was comparable [96]. The results are summarized in Figure 39.
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Figure 39. Hydrogenation reactions catalyzed by osmium pincer complexes. Osmium C~N~N pincer complexes 177 (X = Cl) and 178a were employed in dehydrogenation of alcohols to form the corresponding ketones and H₂. The dehydrogenation of α-tetralol using a catalyst charge of 0.4% mol was run with 36–44% conversions in 24 h. Their ruthenium analogues were also tested and were found to be more active, with conversions between 90–93% at the same reaction time [97]. The osmium C~N~N pincer complexes 181 and 182 were used for the racemization of alcohols with good results. In this case, a similar catalytic activity was observed for both ruthenium and osmium analogues (Figure 40) [98].

Osmium C~N~N pincer complexes 177 (X = Cl) and 178a were employed in dehydrogenation of alcohols to form the corresponding ketones and H₂. The dehydrogenation of α-tetralol using a catalyst charge of 0.4% mol was run with 36–44% conversions in 24 h. Their ruthenium analogues were also tested and were found to be more active, with conversions between 90–93% at the same reaction time [97]. The osmium C~N~N pincer complexes 181 and 182 were used for the racemization of alcohols with good results. In this case, a similar catalytic activity was observed for both ruthenium and osmium analogues (Figure 40) [98].
Fluorinated pincer complex [OsH(CF₃P-C-P)(cod)] (183) (Figure 41) obtained from [Os(cod)(η³-2-methylallyl)₂] was tested in alkane dehydrogenation. The catalyst showed a longer lifetime than its ruthenium analogue. The dehydrogenation of cyclooctane in the presence of tert-butylethene as a hydrogen donor was carried out. A significant production of cyclooctene was detected within a few minutes at 200 °C [99].

Esteruelas’ group reported on the metalation of imidazolium salts using [OsH(PPr₃)₂]. Stable compounds 185 and 187, like possible intermediates in catalytic hydrogenation, were isolated (with yields from 83 to 89%) through the osmium reduction in the presence of HBF₄. The authors also explored the substitution of the n²-bonded molecular hydrogen by a coordinating molecule like acetonitrile [100].

The metalation of 1-(2-methoxy-2-oxoethyl)-3-methylimidazolium chloride afforded cyclometalated acyl complex 188 which coordinates dioxygen, dihydrogen, and carbon monoxide (Figure 42). Complex 188 was used as a catalyst for alcoholysis and hydrolysis of pinacolborane. Using a 2% mol catalyst charge for the alcoholysis reaction, TOFs between 62 and 3644 h⁻¹ were obtained at 50% conversion. For the hydrolysis carried out with the same amount of catalyst, TOFs were 473–1648 h⁻¹ at 50% [101]. The 2-(aminomethyl)pyridine-based complexes 190 were used in the imine-to-amine hydrogenation (Figure 43). Depending on the reaction conditions, conversions between 26 and 99% were observed. In this work, the ruthenium equivalents of 190 were also obtained. In general, under the same reaction conditions, the osmium complexes were found to be less active than the corresponding ruthenium derivatives [102].
Esteruelas et al. described the heterobinuclear iridium–osmium compound $[(P\text{r}3)2(H)2Ir\{\mu-(\kappa^2-N\text{py},N\text{imine}-B\text{MePI-}\kappa^2-N\text{imine},C_4\text{iso})\}OsH3(P\text{r}3)2]$ (191) incorporating 1,3-bis(6′-methylpyridyl-2′-imino)isoindoline (HBMePI) as a bridging ligand. This complex was tested in the base-free dehydrogenation of secondary alcohols using 7% mol of catalyst. Conversions up to 84% were obtained. Interestingly, heterobimetallic complex 191 proved to be more active than the corresponding monometallic components taken independently (Figure 44) [103].

Figure 42. Examples of hydride complexes used in the alcoholysis and hydrolysis of pinacolborane.

\[
\begin{align*}
&\text{H}_2\text{O} + \text{HBPin} \rightarrow \text{HOBPin} \\
&\text{HOBPin} \rightarrow \text{HBPin} - \text{H}_2 \\
&\text{HBPin} \rightarrow \text{(BPin)}_2\text{O}
\end{align*}
\]

Figure 43. Hydrogenation of imines catalyzed by C~N~N pincer complexes.
Esteruelas et al. described the heterobinuclear iridium–osmium compound [(P^3Pr_3)Ir_2(μ-κ^2-N_{py},N_{imine}-BMePI-κ^2-N_{imine},C^4_{iso})OsH_3(P^3Pr_3)_2] (191) incorporating 1,3-bis(6'-methylpyridyl-2'-imino)isoindoline (HBMPI) as a bridging ligand. This complex was tested in the base-free dehydrogenation of secondary alcohols using 7% mol of catalyst. Conversions up to 84% were obtained. Interestingly, heterobimetallic complex 191 proved to be more active than the corresponding monometallic components taken independently (Figure 44) [103].

![Figure 44](image_url)

**Figure 44.** Heterobimetallic complex used in dehydrogenation of secondary alcohols.

The resolution of a helical chiral octahedral osmium complex afforded optically pure Λ (192) and Δ (193) enantiomers (Figure 45). The Δ enantiomer was tested as a catalyst in two processes, viz. C(sp^3)–H aminations of sulfonylazide and azidoformate. For the former, a product yield of 96% was observed using 2% mol of catalyst, with an enantiomeric ratio of 92:8 (Figure 45). Under similar conditions, a yield of 86% was registered in the case of azidoformate with the enantiomeric ratio of 89:11. This is the first example of an osmium complex with a central (helicoidal) chirality successfully used in asymmetric catalysis [104]. It is important to mention that Meggers’ group also developed highly effective enantioselective catalytic processes with metal-centered chiral octahedral ruthenium and rhodium complexes [105–107].

![Figure 45](image_url)

**Figure 45.** Asymmetric catalysis by cyclometalated osmium complexes with central chirality.
Octahedral cyclometalated 2-phenylpyridine Ru\textsuperscript{II} and Os\textsuperscript{II} complexes were investigated as catalysts for the atom transfer radical polymerization (ATRP). Polymerization of styrene was catalyzed by ruthenium complexes though the osmium counterparts were not able to mediate the polymerization and just traces of the polymer were observed [108].

5.2. Chemical Sensors and Biosensors

Oxidoreductases are enzymes that catalyze oxidative and reductive reactions. They are used in various amperometric biosensors, including glucometers [109]. Active sites of the enzymes do not usually exchange electrons with an electrode and therefore low-molecular-weight compounds are commonly used to move the electrons. Known as electron shuttles or mediators, they are often transition-metal complexes. In particular, cyclometalated ruthenium derivatives were successfully coupled with a number of oxidoreductases such as glucose oxidase, glucose and alcohol dehydrogenases, or peroxidases [110–112]. The M\textsuperscript{II}/III reduction potential is an essential feature for optimal performance of biosensors [113], and since the potentials are tunable as it is illustrated in Inset to Figure 21 [80], the corresponding osmium-based mediators were exploited. In particular, a series of octahedral osmium(II) complexes of the general formula \([\text{Os}(\text{N~C})(\text{N~N})(\text{MeCN})_2]\)PF\(_6\) [80], the corresponding osmium-based mediators were exploited. In particular, a series of \([\text{Os}(\text{phpy})_2(\text{bpy})]\)PF\(_6\) (159, 161) and \([\text{Os}(\text{N~C})(\text{N~N})_2]\)PF\(_6\) (94, 95, 162, 163) were investigated, where C~N is cyclometalated 2-phenylpyridine or N,N-dimethylbenzylamine and the N~N ligands are 2,2’-bipyridines or 1,10-phenanthrolines. The complexes were evaluated as electron shuttles with glucose oxidase (GO) for potential applications in amperometric enzymatic glucose sensors. All complexes showed extremely fast electron transfer with the enzyme, with the second-order rate constants \(k_2\) for bimolecular oxidation of reduced glucose oxidase in the range of \((0.67 – 2.90) \times 10^6 \text{ M}^{-1} \text{s}^{-1}\) (Figure 46 and Table 3) [76].

\[
\begin{align*}
\text{GO}_{\text{ox}} + \text{D-glucose} & \quad \text{GO}_{\text{red}} + \text{D-glucuronolactone} + 2\text{H}^+ \\
\text{GO}_{\text{red}} + 2\text{Os(III)} & \quad \text{GO}_{\text{ox}} + 2\text{Os(II)} \quad (k_2) \\
2\text{Os(II)} - 2\text{e} & \quad 2\text{Os(III)} \quad (\text{electrode}) 
\end{align*}
\]

**Figure 46.** Electron transfer pathway between osmium shuttle and glucose oxidase (GO).

| Complex | \(E^\circ\) (mV) | \(k_2\) (M\(^{-1}\)s\(^{-1}\)) |
|---------|----------------|-----------------|
| 162     | -51            | 0.67 \times 10^6 |
| 163     | 13             | 4.80 \times 10^6 |
| 161     | 32             | 2.00 \times 10^6 |
| 94      | 84             | 2.90 \times 10^6 |
| 95      | 31             | 1.80 \times 10^6 |
| 159     | 109            | 2.90 \times 10^6 |

Discussed above mono- \([\text{Os}(\text{phpy})(\text{bpy})_2]\)PF\(_6\) (94), bis- \([\text{Os}(\text{phpy}_2)(\text{bpy})]\)PF\(_6\) (102) and tris-cyclometalated \([\text{Os}(\text{phpy})_3]\) (101) derivatives turned out to be attractive objects for electrochemical studies. Using related coordination complex \([\text{Os}(\text{bpy})_3]\)PF\(_6\)_2 (103), it was shown that the Os\textsuperscript{II}/Os\textsuperscript{III} reduction potential significantly decreases as the number of Os~C bonds increases. Rates of electron transfer between the osmium center and the active sites of oxidoreductases including horseradish peroxidase (HRP) were investigated (Figure 47). Cyclic voltammetry was used to estimate the rate constants. In the case of HRP, the results showed that the rate constants \(k_2\) are higher than \(k_3\) for all compounds, i.e., electrons move faster from Os\textsuperscript{II} to compound I than to compound II (see Figure 47). The value of both rate constants increases in the series \([\text{Os}(\text{bpy})_3]\)PF\(_6\)_2 < \([\text{Os}(\text{phpy})(\text{bpy})_2]\)PF\(_6\)
(94) < [Os(phpy)₂(bpy)]PF₆ (102) and then decline for the tris-cyclometalated complex (101). It should be mentioned that all complexes are very reactive. The Monte Carlo docking simulations helped to evaluate how the complexes approach the active site of HRP [80].

**Figure 47.** Mono-, bis-, and tris-cyclometalated complexes used as electron shuttles and electron transfer pathway with horseradish peroxidase (HRP).

Bis-cyclometalated complex [Os(phpy)₂(bpy)]PF₆ (102) was tested as a mediator in a prototype of an amperometric biosensor to quantify hydrogen peroxide (Figure 48) An iron(III)-TAML (TAML = tetraamido macrocyclic ligand) catalyst used as an alternative to peroxidase enzymes was immobilized on an electrode surface with the osmium mediator. The activity and sensitivity of such a device were similar to those of a biosensor based on HRP enzyme [114].

**Figure 48.** Conceptual principle for a sensor using an iron(III)-TAML catalyst and the bis-osmacycle as a mediator.

5.3. Electronic Properties and Photophysics

Due to a unique combination of spectroscopic, photophysical, photochemical and electrochemical properties, octahedral Ru°II and Os°II polypyridyl complexes have been extensively studied in various fields such as photochemical conversion of solar energy,
photocatalysis and in molecular electronic devices. Their properties can be modulated by modifying the ligand structures and by introducing ancillary ligands [115,116]. The major goal is to enhance excited state lifetimes of the compounds at room temperature by increasing the energy gap between the radiative 3MLCT and quenching 3MC states by destabilizing the 3MC state using cyclometalated ligands [117,118]. However, the enhanced spin-orbit coupling of the third-row elements when compared to the second-row elements usually makes optoelectronic properties of osmium complexes notably different from those of ruthenium derivatives [119]. Nevertheless, as we discuss in this section, switching from ruthenium to osmium has proven to be successful in generating promising materials.

Sierra et al. synthesized dinuclear cyclometalated trihydride osmium(IV) complexes (69–71, 194, 195) (Figure 49) which were studied by cyclic voltammetry. Two quasi-reversible processes corresponding to the OsIV/OsV and OsV/OsVI redox features were observed. The OsIV complexes showed emission around 360 nm, with quantum yields being between 0.005 and 0.020. The excitation spectra suggested that a ligand-centered transition is at the origin of the emission. Moreover, spectrochemical studies showed that the emission spectra remained very similar upon OsIV to OsV oxidation at 0.05 V vs. Fc+/Fc, since only a slight increase in intensity was observed, with quantum yields from 0.013 to 0.026. Nevertheless, when a potential of 1.10 V was applied to carry out the oxidation of OsV to OsVI, a bathochromic shift, from around 360 to around 400 nm, was observed along with an increase in the luminescence intensity with quantum yields around 0.030–0.110. DFT calculations suggest that the photophysical properties vary due to the conversion of trihydride species to monohydride-dihydrogen derivatives during sequential oxidation processes [59].

Kapturkiewicz et al. prepared the series of cationic osmium complexes 196 of the general formula [OsCl(N–C)(PPh3)2(CO)] (N–C is a cyclometalated substituted 2-phenylpyrididine) and measured their UV-visible absorption and emission spectra (Figure 50). The complexes showed emission at 455–532 nm attributed to 3MLCT transitions. Importantly, only the solids emitted light at room temperature; the compounds in solution were emissive only at a low temperature (77 K). The emission lifetimes are relatively long, between 10 and 20 μs [120].
Bis-(P–C–P) pincer osmium complex 39 (Figure 10) has a strong absorption at 250 nm attributed to the ligand π→π* transitions. The emission at 546 nm in a dilute solution of 2-MeTHF at 77 K with a quantum yield of 0.6 and a lifetime of 8.0 µs suggests phosphorescence. The solid complex is also luminescent at room temperature at 556 nm, with a quantum yield of 0.03 and a lifetime of 0.3 µs. The electronic structure of the complex and related transitions were explored by DFT [45].

Carbene pincer ruthenium and iridium compounds display remarkable luminescent properties [121]. Related osmium C–C~C complexes 197 and 198 were obtained by reacting [OsCl4(N–N)] precursors (N–N = 2,2′-bypiridine, 1,10-phenanthroline, 4,4′-diphenyl-2,2′-bypiridine) with bis-imidazolium or bis-benzimidazolium hexafluorophosphates in refluxing ethylene glycol as a source of the CO ligand (Figure 51). The UV-visible absorption data showed intense and high-energy absorption at λ lower than 330 nm and a less intense band at λ > 330 nm. The complexes are also emissive upon photoexcitation with emission maxima in the red region (674–731 nm). The quantum yields reported were between 10−4 and 10−2 and emission lifetimes in the order of 10−1 µs. The photophysical parameters are sensitive to the modification of N–N and C–C~C parts, suggesting that the emissive-excited state should involve both fragments and the emission occurs due to the energy dissipation of d(Os)→π*[N–N] MLCT transitions [122].

The photophysical studies of the C–C′–C mono- and bis-pincer compounds (199–204) in Figure 51 revealed an intense absorption between 341 and 363 nm along with a less intense band between 396 and 405 nm. TD-DFT calculations allowed to attribute the high energy bands to interligand charge-transfer transitions and the low energy bands to metal-to-ligand charge-transfer processes. Particularly, the bis-pincer complexes 202–204 exhibited emission in the blue-green region (475–578 nm), both in the solid state at room temperature and in toluene solution (at room temperature and 77 K) with lifetimes in the range of 10–29 µs. The best quantum yield is observed for complex 202 (R = CF3) with a value of 0.62. This last complex 202 (R = CF3), with emission in the blue region, was chosen for the fabrication of a model OLED device, obtaining promising results [123].

Figure 50. Luminescent [OsCl(N–C)(PPh3)2(CO)] complexes.
phosphino ligand) charge-transfer transitions. In addition, the absorption spectra contained an intense band at 330 nm and a less intense band at 404 nm attributed to metal-to-ligand charge-transfer transitions [124].

Figure 51. Luminescent carbene-based pincer complexes.

The osmium complexes 205 and 206 with a tetradeinate C–C–C–C ligand (C$_{phenyl}$–C$_{carbene}$–C$_{carbene}$–C$_{phenyl}$) in Figure 51 can be viewed as bis-cyclometalated compounds. The complexes with two coordinated DMSO ligands provide options for their replacement by bidentate diphosphines or dienes. Complexes bearing 1,2-bis(diphenylphosphino)benzene emit light upon photoexcitation in the solid state at room temperature and in solution of 2-MeTHF at 77 K. The bands between 630 and 549 nm were attributed to the $\pi$-$\pi^*$ HOMO (Os and NHC fragment) to LUMO (diphenylphosphino) and metal-to-ligand (osmium to phosphino ligand) charge-transfer transitions. In addition, the
absorption spectra contained an intense band at 330 nm and a less intense band at 404 nm attributed to metal-to-ligand charge-transfer transitions [124].

Bis-tridentate cyclometalated complexes [Os(N~N~N)(N~C~N)]PF$_6$ (207, 208) have been tested as molecular wires, and compounds 208, in which the N~C~N ligand with a triarylamine substituent is separated from the metal center by an oligophenyl bridge, were of special interest (Figure 52). This is due to the presence of two electrochemically active centers, the metal and the triarylamine substituent which can undergo a N$^{**}$/N$^0$ process (neutral nitrogen to positive radical nitrogen). These molecular wires show two anodic redox features between +0.1 and +0.9 V vs. Ag/AgCl, the first one assigned to the Os$^{II}$/Os$^{III}$ process and the second one to the N$^{**}$/N$^0$ process, and an electron transfer from the neutral amine to the Os$^{III}$ center was observed in the one-electron oxidized form [Os$^{III}$,N] [125].

![Figure 52. Bis-tridentate cyclometalated osmium complexes studied as molecular wires.](image)

Esteruelas et al. studied the luminescent properties of C~C~N pincer complexes derived from 2-azetidones (see Table 2). At room temperature, complex (73-L22) showed yellow emission in the solid state at 571 nm (lifetime 1.1 µs) and at 562 nm in a toluene solution (lifetime 2.0 µs), as well as green emission at 77K at 540 nm (lifetime 6.6 µs) [60]. Complexes 74 and 75 in Table 2 are phosphorescent upon photoexcitation in a poly(methyl methacrylate) (PMMA) film and in 2-MeTHF at room temperature and 77 K. Both compounds emit in the yellow region. The 2,6-diphenylpyridine complex 74 is more efficient, the quantum yields being 0.28 (film) and 0.56 (solution). The introduction of oxygen between the pyridine and phenyl ring in 75 lowered the emission (quantum yields below 0.10). Curiously, the iridium analogue of 75 showed the opposite trend [126].

The absorption spectra of C~N~N and C~N~C pincer complexes 209–212 in Figure 53 are similar. They present two main bands between 279 and 317 nm and between 364 and 394 nm. TD-DFT calculations allowed to assign the high energy absorptions to ligand-centered transitions while the low energy absorptions were assigned to metal-to-ligand charge-transfer processes. The compounds were emissive upon photoexcitation in the solid state at room temperature and in 2-MeTHF solution both at room temperature and at 77 K. The use of pincer ligands allowed to obtain more rigid structures reducing both the energy dissipation and the difference between the excited state and the ground state structures. As a consequence, enhanced quantum yields in the range of 0.08–0.59 were obtained, lifetimes being 1.5–5.0 µs in the solid state and 0.9–5.8 µs in solution [127].

The electronic structure of osmium bis-pincer complexes 213 with N~C~N cyclometalated pyrene ligands (Figure 54) was explored by DFT and TD-DFT. It was found that HOMO orbitals are mainly osmium in character. The LUMO orbitals were largely associated with ligand contributions. There are two intense bands absorption spectra between 282 and 372 nm assigned to intraligand transition from the pyrene and terpyridine fragments. Two other intense bands between 410 and 447 nm were associated with intraligand charge-transfer (ILCT) transitions. The bands observed between 513 and 556 nm were due to MLCT transitions [128]. These experimental results launched a theoretical study which helped to optimize photophysical features of the compounds. It was concluded that compounds 214 in Figure 54 should display optimal properties [129].
As a consequence, enhanced quantum yields in the range of 0.08–0.59 were obtained, lifetimes being 1.5 μs. Curiously, the iridium analogue of non-cyclometalated complexes. The cytotoxic activity in HeLa cancer cells was determined first by the evaluation of the half-maximum effective concentration (EC50) at which the HOMO orbital is contributed by pyrene and metal centers. The LUMO orbital has charge-transfer (ILCT) transitions. The bands observed between 513 and 556 nm were due to MLCT transitions [128]. These experimental results launched a theoretical study which showed the opposite trend [126].

DFT calculations performed for bimetallic complex 215 shown in Figure 54 suggested that the HOMO orbital is contributed by pyrene and metal centers. The LUMO orbital has contributions from the pyrene moiety and pyridine rings. Two intense bands observed between 457 and 507 nm were assigned to ligand-to-ligand charge-transfer transitions. An electronic coupling between the neighboring osmium centers affects cyclic voltammograms of osmium compounds and brings about two reversible OsII/OsIII redox features overlapping at −0.15 V (vs. Fc/Fc+). A similar behavior is observed for ruthenium but not for iridium complex [130].

### 5.4. Anticancer and Biological Properties

Transition metal derivatives have been extensively studied in medicinal chemistry, often with emphasis on their anticancer activity. Organometallic complexes are currently receiving special attention [131–137]. The leadership belongs to ruthenium compounds including the clinical studies of metal-based anticancer drugs [138–141]. Omae published a

![Figure 53](image-url) Phosphorescent C-N and C-N-C osmium pincer complexes.

![Figure 54](image-url) Luminescent bis-pincer complexes with cyclometalated pyrene-type ligands.
comprehensive review on anticancer properties of cyclometalated complexes with a few osmium examples [132]. Pfeffer and Gaiddon dedicated a review to ruthenacyles [142]. Examples of a few osmacycles were included in the recent review by Zhang devoted to biological properties of osmium compounds [131]. Here, we show the most recent examples of osmium cyclometalated complexes used in biological assays.

Meggers and Xia compared the anticancer activity of a series of osmium derivatives, including complexes 43, 49, 164, 174, a new η²-allene compound (216, Figure 55), and other non-cyclometalated complexes. The cytotoxic activity in HeLa cancer cells was determined first by the evaluation of the half-maximum effective concentration (EC\textsubscript{50}) at which viability of the cells is reduced to 50% after 24 h incubation. Observed values for EC\textsubscript{50} were between 1 and 30 µM. The best results were obtained with allene complex 216 which exhibited cytotoxic activity with EC\textsubscript{50} = 1 µM. Because of this and its high stability, studies of this complex were prioritized. The compound was tested against Burkitt-like lymphoma (BJAB), showing an inhibition of the proliferation at concentrations lower than 1 µM. Another assay revealed that the compound induces apoptosis in BJAB cells by DNA fragmentation. The results showed that even a 0.3 µM concentration, 216 induced DNA fragmentation in 25% of the cells, with the highest DNA fragmentation observed in 75% of the cells at 5 µM. In addition, other experiments showed that apoptosis occurs through the intrinsic mitochondrial pathway. The compound was also tested in leukemia cells (Nalm6), Vcr-resistant NAIm6 cells and Dau-resistant NAIm6 cells (the last two are drug-resistant leukemia cells) showing a lower activity, with EC\textsubscript{50} higher than 1 µM, but with effective activity at 5µM [143].

![Figure 55. η²-Allene complex with promising cytotoxic activity.](image)

The successful use of ruthenacyles and osmacycles as efficient electron shuttles for oxidoreductase enzymes described in Section 5.2 prompted the investigation of the potential use of these compounds for biological applications. Gaiddon and Pfeffer evaluated the cytotoxic properties of such compounds on series of human cancer cell lines. An exhaustive work was published where a series of osmium cyclometalated compounds was tested in in vitro cell growth inhibition in A172 glioblastoma cell line. The structures of the compounds studied are summarized in Figure 56. First results with the piano-stool complexes (92, 93, 217–219) showed moderate cytotoxic activity with IC\textsubscript{50} values of 10–100 µM. The octahedral complexes with bidentate cyclometalated ligands (94, 158, 159, 220–224) exhibited very promising results with IC\textsubscript{50} values in the nanomolar range. Pincer complexes (225, 226) showed very poor cytotoxic activity, with IC\textsubscript{50} values 21–375 µM, which could be attributed to the presence of three labile acetonitrile ligands. Interestingly, the pincer complexes with polypyridine ligands (227, 228) showed very good results with IC\textsubscript{50} values between 0.3 and 4.1 µM. There was no correlation between the cytotoxicity and structure. However, the reduction potentials were measured for all complexes and the results showed that the compounds with the potentials between 0.3 and 0.6 V (vs. SCE) displayed the...
lowest IC\textsubscript{50} values. The lipophilicity was also determined and the complexes with the highest activity showed log(P\textsubscript{o/w}) values around 2. These results allowed to hypothesize that the compounds could strongly modify the metabolism of the cells by interactions with oxidoreductases and this would partially explain the relationship between the activity and the reduction potentials [144]. Another important target of such studies is to discover molecules that can overcome drug resistance induced by cisplatin. Recently, the mechanisms governing the variability in the cytotoxicity of two ruthenium cyclometalated compounds and their osmium equivalents were studied (Figure 34, complexes 95 and 159). Their anticancer properties in vitro and in vivo were first evaluated and genes involved in their sensibility/resistance were identified by correlating their cytotoxicity with transcriptomic data of 60 cancer cell lines. Docking and functional studies demonstrated that inhibition of known resistance mechanisms, ABCB1 export and EGFR expression, allowed to improve the activity of cyclometalated complexes. Interestingly, switching from ruthenium to osmium favored the cytotoxicity while reducing the sensibility to the ABCB1 export mechanism [145].

Kandioller developed the coordination of a series of 4-phenyl-1,2,3-triazole-based ligands to ruthenium and osmium to prepare complexes 229 in Figure 57. Their cytotoxic activity was evaluated on three cancer cell lines: non-small-cell lung cancer (A549), colon adenocarcinoma (SW480) and human ovarian carcinoma (CHI/PA-1). The free ligands did not show cytotoxic activity within the studied concentrations range. However, the ruthenium derivatives bearing ligands a and b showed significant cytotoxicity within the studied concentrations range. However, the ruthenium derivatives bearing ligands a and b showed significant cytotoxicity with IC\textsubscript{50} values around 13 µM in A549, 7 µM in SW480 and 4 µM in CHI/PA-1. Osmium counterparts displayed an even higher activity, ligands a and b being also the most active, with IC\textsubscript{50} around 6 µM for A549, 3.6 µM for SW480, and 1 µM for CHI/PA-1 cells. The topoisomerase IIα inhibition was investigated but no compound was capable of inhibiting its function, indicating that the mechanism of action does not follow this pathway. The cell cycle distribution in SW480 in response to treatment with the complexes was determined and only the osmium derivatives exhibited elimination of the S phase fraction, information relevant in order to understand the mechanism of action [146].

Using a triazole-based ligand, Makhubela prepared a series of half-sandwich ruthenium, rhodium, iridium and osmium complexes (230), Figure 57. The complexes were tested on different cancer cell lines (leukemia MT4, cervical cancer HeLa, kidney adenocarcinoma HEK293 and lung cancer A549), and the CC\textsubscript{50} (cytotoxic concentration with 50% of the maximum decrease in cell viability) was calculated. The ruthenium derivative (230a) exhibited a moderate activity against A549, but the osmium compound (230b) was much more active with a CC\textsubscript{50} value of 6.4 µM, comparable to cisplatin used as reference. However, its activity was very modest against the other cancer cell lines. Interestingly, the rhodium and iridium derivatives were more effective against all cancer cell lines when compared to their ruthenium and osmium analogues [147].

Benzimidazole-based ligands were used by Ruiz to prepare the series of new osmium complexes 231, Figure 57. Their antiproliferative activity was tested in six different human cancer cell lines (including cisplatin-resistant cell lines), as well as non-tumorigenic human endothelial hybrid cells and Buffalo green monkey cells. All osmium compounds showed a higher cytotoxic activity than cisplatin in all cancer cell lines, with IC\textsubscript{50} values lower than 10 µM. The cytotoxicity against the non-cancer cells was very similar to cisplatin. The cellular concentrations of both metals were measured in A2780 cells and the results showed that the osmium cellular uptake is ten times higher than for platinum. Other studies revealed that the compounds caused a decrease of the ROS levels and, as a result, an affectation in the G0/G1 phase in the cell cycle is observed [148].

In a very recent work, Gómez-Gallego, Sierra and Esteruelas explored the reaction between osmium precursor [OsH\textsubscript{6}(P\textsubscript{iPr\textsubscript{3}}\textsubscript{2})] and nucleosides. Cyclometalated complexes 232–235 were obtained in good yields (Figure 58). The authors mentioned that this methodology could be applied to the functionalization of oligonucleotides with promising potential biological applications [149].
to osmium favored the cytotoxicity while reducing the sensibility to the ABCB1 export mechanism [145].

Figure 56. Series of cyclometalated osmium complexes tested in the in vitro cell growth inhibition in A172 glioblastoma cell line.

Kandioller developed the coordination of a series of 4-phenyl-1,2,3-triazole-based ligands to ruthenium and osmium to prepare complexes [145]. Their cytotoxic activity was evaluated on three cancer cell lines: non-small-cell lung cancer (A549), colon adenocarcinoma (SW480) and human ovarian carcinoma (CHI/PA-1). The free ligands did not show cytotoxic activity within the studied concentrations range. However, the ruthenium derivatives bearing ligands \(a\) and \(b\) showed significant cytotoxicity with IC\(_{50}\) values around 13 \(\mu\)M in A549, 7 \(\mu\)M in SW480 and 4 \(\mu\)M in CHI/PA-1. Osmium counterparts displayed an even higher activity, ligands \(a\) and \(b\) being also the most active, with IC\(_{50}\) around 6 \(\mu\)M for A549, 3.6 \(\mu\)M for SW480, and 1 \(\mu\)M for CHI/PA-1 cells. The topoisomerase II \(\alpha\) inhibition was investigated but no compound was capable of inhibiting its function, indicating that the mechanism of action does not follow this pathway. The cell cycle distribution in SW480 in response to treatment with the complexes was determined and only the osmium derivatives exhibited elimination of the S phase fraction, information relevant in order to understand the mechanism of action [146].

Figure 57. Complexes tested against various human cancer cell lines.

Using a triazole-based ligand, Makhubela prepared a series of half-sandwich ruthenium, rhodium, iridium and osmium complexes [147], Figure 57. The complexes were tested on different cancer cell lines (leukemia MT4, cervical cancer HeLa, kidney adenocarcinoma HEK293 and lung cancer A549), and the CC\(_{50}\) (cytotoxic concentration with 50% of the maximum decrease in cell viability) was calculated. The ruthenium derivative \(230a\) exhibited a moderate activity against A549, but the osmium compound \(230b\) was much more active with a CC\(_{50}\) value of 6.4 \(\mu\)M, comparable to cisplatin used as reference. However, its activity was very modest against the other cancer cell lines. Interestingly, the rhodium and iridium derivatives were more effective against all cancer cell lines when compared to their ruthenium and osmium analogues [147].

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Many more biological applications of osmium-based complexes remain to be discovered and exploited. For instance, the unique photophysical properties of osmacycles (see Section 5.3) will undoubtedly open the field for further research on their use as PDT agents. On the other hand, deeper in vivo studies are needed to confirm properties such as cellular uptake ability, cellular distribution, as well as to determine essential pharmacokinetic data.

6. Conclusions

As this review demonstrates, even though osmacycles chemistry is still in a growing stage, cyclometalated osmium compounds are essential in many aspects. The broadest chemistry of osmium and the low reactivity of osmium compounds grants determination and studies of a variety of mechanisms, which in turn provide tools to design novel complexes with a wide variety of relevant properties and applications. Due to the enormous number of molecules, known or readily accessible, that can lead to cyclometalation reactions, unique applications of osmacycles will see the light in the near future. Many cyclometalated osmium derivatives have shown remarkable catalytic activity in various reactions, principally hydrogenations and dehydrogenations, often similar and in some cases superior to those reported for analogous ruthenium systems. Such results exhibit the potential of osmacycles for the design of new highly effective and robust catalytic systems for chiral and non-chiral processes. On the other hand, new osmium-containing cyclometalated materials will continue to be studied for their activity in biological systems, and they will likely be involved in the development of innovative chemical sensors, biosensors and improved drugs.

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Abbreviations

2-MeTHF 2-methyl-tetrahydrofuran
A2780 Human ovarian cancer cell line
A549 Human lung cancer cell line
AND/OR Boolean operator for browsing in databases
ATRP Atom Transfer Radical Polymerization
B3LYP Becke three-parameter exchange and Lee–Yang–Parr correlation
BJAB Burkitt-like lymphoma
Bn Benzyl substituent
bpy 2,2′-bipyridine
C~C~N Pincer ligand with Carbon-Carbon-Nitrogen, as donor atoms
C~C~C Pincer ligand with Carbon-Carbon-Carbon, as donor atoms
C~C~C~C Tetradentate Ligand with Carbon-Carbon-Carbon-Carbon as donor atoms
C~E Bidentate ligand with Carbon and E as donor atoms
C~N Bidentate ligand with Carbon and Nitrogen as donor atoms
C~N~C Pincer ligand with Carbon-Nitrogen-Carbon as donor atoms
C~N~N Pincer ligand with Carbon-Nitrogen-Nitrogen as donor atoms
C~N~O Pincer ligand with Carbon-Nitrogen-Oxygen as donor atoms
C~O Bidentate ligand with Carbon and Oxygen as donor atoms
CC~50 50% Cytotoxic concentration
CHI/PA-1 Human ovarian carcinoma cells
cod 1,4-cyclo-octadiene
DFT Density Functional Theory
DMSO Dimethyl sulphoxide
DNA Deoxyribonucleic acid
dpbH 1,3-di(2-pyridyl)benzene
dppb 1,4-bis(diphenylphosphino)butane
E Donor atom such as: N, P, O, S, As, Se
EC~50 50% Effective concentration
GO Glucose oxidase
HBMePI 1,3-bis(6′-methylpyridyl-2′-imino)isoindoline
HEK293 Kidney adenocarcinoma cell line
HeLa Cervical cancer cell line
HOMO Highest Occupied Molecular Orbital
HRP Horseradish peroxidase
IC~50 50% Inhibitory concentration
ILCT Intraligand Charge-Transfer
log(Po/w) N-Octanol-water partition coefficient
LUMO Lowest Unoccupied Molecular Orbital
Mebib 2-deprotonated form of 1,3-bis(N-methylbenzimidazolyl)benzene
Mebip bis(N-methylbenzimidazolyl)pyridine
MLCT Metal-to-Ligand Charge Transfer
MT4 Leukemia cell line
N~C~N Pincer ligand with Nitrogen-Carbon-Nitrogen as donor atoms
N~N Bidentate ligand with Nitrogen-Nitrogen as donor atoms
N~N~N Terdentate ligand where the atom donors are Nitrogen
Nalm6 Leukemia cell line
NHC N-Heterocyclic Carbene
NHE Normal Hydrogen Electrode
NMR Nuclear Magnetic Resonance
OLED Organic Light-Emitting Diode
Otf Triflate
P~C(H)~P Ligand with only phosphorous as donor atoms
P~C~P Pincer ligand with phosphorous–carbon–phosphorous as donor atoms
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