Phosphine- and Isatin-Copper Complexes: Anticancer Activity, Mechanisms of Action and Structure-Activity Relationship Examples from the Last Ten Years

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Abstract

Platinum-based metallodrugs candidates are extensively evaluated as antineoplastic agents against several tumoral cell lines, representing a considerable advance in cancer treatment. On the other hand, the hostile off-site toxicity has been limited their potential uses in clinical step. Thus, the search for novel active and selective antitumor drugs is a leading issue in the medicinal inorganic chemistry. As follow, the research focused on less toxic metals has strategic importance and several copper complexes have revealed promising activity. In addition, copper is an essential metal, which can increase the chances of reduced side effects. This review brings some copper complexes that showed highlighted anticancer activity as drug prototypes from the last ten years (2010-2019), involving phosphine and isatin ligands, different redox states, modes of action and the study of structure-activity relationship (SAR).

Keywords: Copper Complexes; Anticancer Activity; Phosphine; Isatin; Structure-Activity Relationship


Introduction

Copper is an essential element with vital roles in several biological processes in the body. This metal can be found in metalloenzymes acting as cofactor. The two more suitable redox states, Cu(I) and Cu(II), as well as the tetrahedral, square-planar and octahedral geometries around the metal center, make copper an attractive metal for design of novel metallodrugs [1,2]. Even though this metal possesses two accessible redox states, the cell uptake of copper species into eukaryotic cells occurs by members of the Ctr copper transporter family (Figure 1) [3].
Copper transporter (Ctr) proteins comprise the cell copper absorption system in eukaryotes. They are identified in yeast, plants and humans (hCtr1), containing three transmembrane segments with carboxyl and amino terminal located on opposite site of the plasma membrane. As represented in Figure 1, the delivery of copper to the Cu, Zn-superoxide dismutase 1 (SOD1) is by the copper chaperones for superoxide dismutase (CCS). Primarily, in the cytosol, occurs the metalation of SOD1, however, a minor quantity of SOD1 is also restricted to the mitochondrial intermembrane space, when superoxide dismutase is present.

The SOD1 catalyzes the conversion of superoxide ion to molecular oxygen and hydrogen peroxide (Equations 1 and 2). Superoxide dismutase’s (SODs), for example, belong to the family of isoenzymes connected in the scavenging of free O2 radicals occurred in respiratory chain. One of the cellular protection systems from oxidative damages is the SOD1 [4].

Equation 1:

\[
\text{SOD-Cu}^{2+} + \text{O}_2^- \rightarrow \text{SOD-Cu}^+ + \text{O}_2
\]

Equation 2:

\[
\text{SOD-Cu}^+ + \text{O}_2^- + 2\text{H}^+ \rightarrow \text{SOD-Cu}^{2+} + \text{H}_2\text{O}_2
\]

The proper function of important metalloproteins can be affected by redox activity of copper (Cu). The Cu is bound to albumin, ceruloplasmin (glycoprotein produced in liver) and transcuprein (a macroglobulin), all in the blood. The genetic disorders can be reflective of unappropriated Cu homeostasis. In humans, the copper concentration is controlled at the cell and organ levels. The copper distribution occurs in the liver via ceruloplasmin and albumin proteins when it is earlier absorbed in the small intestine and stomach. The human Cu dietary is recommended between 1.5 and 3.0 mg Cu/day [5]. Regarding to the redox states, the Cu(II) ion is typically found in the blood, while the cell uptake mechanism involves the reduced Cu(I) form.

Copper is important for antioxidant activity, collagen synthesis, iron transport (the both metabolisms are tightly linked) and lipid peroxidation. The lack or overload of copper concentration can be responsible by declining hematopoiesis by means of anemia and leukopenia (neutropenia), bone abnormalities, myelopathy, optic neuropathy, growth and progression of cardiovascular diseases, among others. Likewise, the reduction of Cu absorption in the blood causes genetic disease, for instance, Menkes syndrome (mutation in the gene encoding ATP7A). On the other hand, the excess of Cu on liver and brain tissues can origin to Wilson disease (mutation in the gene ATP7B). Including other human diseases associated to no regulation of Cu concentration are Alzheimer, Parkinson (both neurodegenerative disorders), ulcers, diabetes and cancer.

Concerning cancer, the Fenton reaction with formation of reactive oxygen species (ROS) is also linked to excess of free Cu ions [6]. Breast and prostate tumoral cells contain higher concentrations of Cu than normal cells. The level amounts to 1.67μg Cu/mL in serum of tumor breast cells line and 0.98μg Cu/mL in healthy breast cells control. However, the molecular mechanisms of Cu increasing of tumor cells are not totally elucidated yet. One hypothesis can be associated with the process of copper in angiogenesis.

**Copper and Production of Reactive Oxygen Species (ROS)**

Living aerobic species have molecular oxygen as fundamental molecule in cell breathing. The breathing sequence affords photo-sensible peroxides, superoxide, singlet oxygen, and hydroxyl radical active species (Fenton reaction). In the presence of UV light, an electron
is promoted from fundamental state ($S_0$) to excited empty state ($S_1$) (also, to an excited triplet state), generating an unstable and energetic chemical compound contain oxygen [7].

The copper can bring oxidative stress in Fenton reaction, since Cu(II) ion can suffer reduction in presence of superoxide radical to form a Cu(I) ion and a molecular oxygen. In sequence, the production of hydroxyl radical species can be catalyzed by Cu(I) ions. Thus, the ROS level can significantly increase. There are critical biological processes that are affected by abnormal production of ROS, mainly the programmed cell death (apoptosis), mutation in the biomolecular structure of DNA (AT-GC and GC-AT), changes in cell motility, and disturbs in cell signaling by irreversible oxidation of key proteins. In addition, endogenous copper complexes play vital role in biological processes, mainly oxygen and electron transfer and oxidative DNA damage, both related with elderly and cancer [7,8].

Cancer
Cancer can be defined as a group of diseases where mutant cells undergo excessive proliferation. After disruption of basal membrane, the tumor cells access the blood system and can extent to other tissues of the body (metastasis) [9]. Cancer has been increasingly investigated for being considered the second cause of death in the world. The WHO reported that occurs at least 14 million of new cases/year being 8.8 million of deaths, which have represented US$ 1.16 trillion of costs in public health and loss of productivity [10,11].

The most important landmark for therapeutic of cancer in the 20th century was the discovery of the neoplastic activity for cisplatin. The free Pt(II) ion is highly toxic for human cells, but when further coordinated to ammine and chloride ligands, [PtCl₂(NH₃)₂], became to an extremely effective metallodrug to treat a diversity of cancers, for instance, testicular cancer. Unfortunately, the administration of this platinum metallodrug is still limited by its aggressive adverse effects, due to its low selectivity index [10-12].

The variety of coordination numbers, redox states, geometries, and intrinsic properties of cationic metal ions are great features for the design of therapeutic agents. Metals, such as Ru(II), Pt(II), Pd(II) and Cu(I) and Cu(II) have been extensively studied for the development of metal-based complexes over last decades [13]. Among them, copper-based complexes have aroused much attention since many of them have exerted their anticancer activities with promising selectivity index.

Copper Complexes as Anticancer Agents
Copper-based complexes have been investigated because endogenous metals may promote less side effects in comparison with cisplatin. The mechanisms of action of copper complexes can occur by degradation of DNA, topoisomerase and/or proteasome inhibition. The biological activities of these complexes have been attributed to the synergism between the Cu(I) or Cu(II) metal ions and their ligands. Thus, several ligands for various copper complexes can be explored to find candidates for cancer treatment. As follows, an important feature concerns the oxidation state of copper metal center. Even if Cu(I) species are found in Ctr proteins to active internalization of biological copper in mammalian cells, this same Cu(I) as antitumor agent is rarely found because it is difficulty to stabilize, especially in aqueous environment. So, Cu(I) is easily oxidized to Cu(II) in most part of applications. Therefore, when Cu(I) ions are handled for design of new active complexes, where ligands able to establish robust metal-ligand bond with Cu(I) are mandatory. In this context, the phosphines have been used to prevent the hydrolysis and degradation of the complexes. In addition, soft donor base, such as S-, P-, C-atoms and aromatic amines have preference to bond soft Pearson acids, such as Cu(I). These complexes are frequently four-fold coordinated assuming generally tetrahedral geometry. This spatial orientation is preferred by means of its [Ar] 3d¹⁰ electronic configuration.

Instead, there are much more Cu(II)-based complexes as anticancer agents in literature, because Cu(II) is chemically more stable in aqueous environment. Then, in different Cu(I)-based complexes, a high number of ligands can be exploited [14]. Others types of Cu(II) complexes, with general formula Cu[(N-N)(A-A)]NO₃, where N-N represents a diamine donor-like (phen or bipy) and A-A stands for amino acid, have showed both in vitro and in vivo antineoplastic activity [15]. Moreover, Cu(II)-based complexes containing aromatic heterocyclic ligands have aroused particular attention for the detection of their genotoxic activity. Herein, the ion complex [Cu(phen)₂]²⁺, with planar geometry ligand, exhibited strong interactions to DNA double-helix by intercalation [16].

In view of that, this review explores some copper complexes in two different oxidations states, including...
phosphine and isatin moieties, as examples of promising ligands to reach anticancer activity.

**Phosphine (Derived) Ligands**

Taking into account the \(\pi\)-acid profile, the phosphine ligands strongly bind to electron-rich \(d^{10}\) metal ions, such as Zn(II), Ag(I) and Cu(I). As follows, regarding to the drug response in tumor cells and normal ones, the structure-activity relationship studies have shown direct association between hydrophilic/lipophilic properties of phosphine ligands and the magnitude of selectivity index (SI) for these complexes. Accordingly, different range of solubility can be acquired by the choice of an appropriated phosphine ligand [1,17]. Among a plethora of suitable ligands that are able to stabilize a specific redox state of metal ions, phosphines are the most attractive category of ligands. As follows, these P-donor ligands may stabilize Cu(I) complexes, which have been explored as antitumor agents as well as to form other hydrophilic soluble complexes [18]. The use of phosphine-derived ligands in medicinal inorganic chemistry has given metal-based complexes with promising anti-arthritis, anti-inflammatory and anticancer activities [19].

*In vitro* cytotoxic activities of Cu(I) and Cu(II) complexes containing phosphine fluoroquinolone derivatives as ligands were performed by Bykowska and Komarnicka [20]. In this study, the Cu(I) cations are four-fold coordinated in a tetrahedral geometry by one P atom from phosphine ligand, one iodide anion and two N-atoms from dmp ligand, resulting in a five-membered metallaring, Figure 2 (1,2). Instead, the Cu(II) cations are coordinated by the quinolone fragment in OPCp or OPNr ligands, that are acting as \(k^2\)\(O\)-\(O\)-donors, forming six-membered metallarings and two N-atoms from phenanthroline (phen). Each Cu(II) cation is four-fold coordinated in a square-planar environment, Figure 3 (3, 4).

![Figure 2: Chemical structures of Cu(I) complexes: [Cu(PCp)dmpI] and [Cu(PNp)dmpI]. (1) Modified with cyclopropane substituent. (2) Modified with ethyl substituent. Structure-activity relationship (SAR) of Cu(I) complexes, with logP and IC\(_{50}\) values.](image-url)
The tumor cell lines A549 (human lung) and CT26 (mouse colon) were used to screen the ability of all compounds to inhibit cell viability via MTT assay. Also, the DNA-affinity of them was assessed by means of fluorescence staining, flow cytometry and gel electrophoresis methods. As a result, the Cu(I) complexes were more active than Cu(II) derivatives in both tumor cell lines. These complexes induced apoptosis, in prevalence. In addition, all cooper complexes appear as DNA-binding metallo-intercalators into two different modes. The Cu(II) complexes damaged the plasmid DNA in the presence of ROS, while the Cu(I) derivatives triggered single-strand breaks on DNA backbone chain. Remarkably, the free ligands did not present significant DNA-affinity. The calculated \( \log P \) values revealed that \([Cu^{II}(OPCp)phen]\) (3) and \([Cu^{II}(OPNr)phen]\) (4) possess more hydrophilic profile when compared with more hydrophobic complexes: \([Cu^{I}(PCp)dmpI]\) (1) and \([Cu^{I}(PNr)dmpI]\) (2) (Figure 2&3). It is hypothesized that Cu(I) complexes had efficient uptake through lipid cell membrane than Cu(II) derivatives [20]. Therefore, the structure-activity relationship (SAR) analysis disclosed that cytotoxic profile of the Cu(I) compounds is dependent on the lipophilic property of the fluoroquinolone ligand [21].

Still concerning the SAR analysis, the absence of ethyl group and fluorine atoms on piperidone and fluoroquinolone rings, respectively, can be the reason of increasing activity. The ethyl group has established prospective metabolic availability due to oxidative N-demethylation pathway [22]. Likewise, cyclopropyl group on piperidone ring appear as an auxophore moiety, as observed in 1 and 3. The enhanced π-characteristic regarding to the coplanarity, decreased C-C bonds distances, and increased polarity of C-H bonds rather than in alkane derivatives (2 and 4), result on increased bioavailability. The fluorine atoms have strong electron-
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withdrawing properties, so they exert influence on special orientation, pKa, membrane permeability, and pharmacokinetic properties [23]. In addition, each hydrogen bonding-donor/acceptor O-atom supports the water solubility and increases the molecule hydrophilicity [24,25].

The Cu(II) complexes, 3 (IC50 = 8.3 ± 1.7 μM) and 4 (IC50 = 15.9 ± 2.1 μM), are less active than Cu(I) complexes 1 (IC50 = 2.9 ± 0.3 μM) and 2 (IC50 = 5.0 ± 0.3 μM). For the Cu(I)-based complexes, the mechanism is related with breaks of the backbone and DNA interactions, through redox reactivity involving ROS generation. In Cu(II) complexes, the mechanism can be associated with ROS amount inside the cells.

Villarreal and collaborators synthesized a Cu(I)-phosphine polypyridyl metal complex, where the Cu(I)-metal center is four-fold coordinated by two P-atoms from PPh3 ligand and two N-atoms of the dppz ligand. The complex display a general formula [Cu(PPh3)2(dppz)]NO3 (5) (Figure 4) [26].

![Figure 4: Chemical structure of Cu(I) tetrahedral complex with IC50 value in human prostate cell line (DU-145).](image)

In this work, the binding constant (k0) of complex 9 with *Calf thymus* DNA (ct-DNA) measured 1.80 ± 0.17 (x 10⁴mol L⁻¹). This data supported a reversible interaction concerning the complex 9 and ct-DNA allied to intercalative mode. This similar intermolecular contact was observed during crystal packing analysis for dppz ligand (blue-square in Figure 4). Moreover, the PPh3 groups (red-square in Figure 4) were chosen to avoid the kinetic instability [27].

Thus, the cytotoxicity of the complexes was evaluated by MTT assays in different types of tumoral cells, such as A549 (human lung), DU-145 (human prostate), MCF-7 (human breast) and the non-tumor cells lines V79-4 (Chinese hamster lung), as well as MRC-5 (human lung). The complex 9 showed IC50 (μM) values: 0.36 ± 0.01 (A549); 2.63 ± 0.25 (MCF7); 0.78 ± 0.04 (DU-145); 0.34 ± 0.04 (V79-4) and 0.52 ± 0.02 (MRC-5), which supported its powerful cytotoxicity. In all tested cell lines, the ligands showed no drug response at IC50 >50μM, therefore considered inactives.

In another study of Marzano and collaborators, Cu(I)-based complexes (Figure 5) containing tridentate tris(pyrazolyl)borate and tertiary phosphine ligands: [HB(pz)3-Cu(PCN)] (6), and [HB(3,5-Me2pz)3Cu(PCN)] (7), where HB means tris(pyrazol-1-yl) borate, were founded out as promising drug candidates [28]. The poly(pyrazolyl) borates (HB), presented in green square in 6, were chosen for the chemical stabilization of metal complexes [29]. The alkyl chain bonded in phosphine atom (represented in red square in 6) increased lipophilicity, while the CN substituents increased inhibition [30].

![Figure 5: Structure-activity relationship (SAR) of Cu(I) tetrahedral complexes with IC50 values in cervical cells line (A431).](image)

The electron-donating methyl groups in the pyrazolyl moieties of the [HB(3,5-Me2pz)3Cu(PCN)] (green squares in 7) resulted in significant reduction of the *in vitro* antitumor activity. The antiproliferative activity for the
Cu(I) complexes were tested against human tumor cell lines of colon (HCT-15), breast (MCF-7), melanoma (A375), cervical (A431), pancreatic (BxPC3), lung (A549), and neuroblastoma (SHSY5Y). The IC$_{50}$ values were calculated by MTT assay after 72 hours of incubation with the complexes. The values obtained for 6 (in μM) are: 0.73 ± 0.17 (MCF7); 0.41 ± 0.18 (A431); 0.99 ± 0.54 (HCT-15); 0.65 ± 0.23 (A375); 0.62 ± 0.46 (BxPC3); 0.77 ± 0.83 (SHSY5Y) and 0.84 ± 0.25 (A549). In the same context, for 7 the values of IC$_{50}$ are: 8.02 ± 2.36; 9.27 ± 1.35; 10.15 ± 1.57; 5.63 ± 2.15; 5.74 ± 2.15; 5.36 ± 1.19 and 10.08 ± 2.54, respectively. In addition, 6 were more effective against tumor cell lines than cisplatin. A similar behavior is observed for phosphine-copper (I) complexes, where a different modus operandi from cisplatin can be observed. In addition, the use of phosphine donors can stabilize reduced copper (I) species, which are commonly incorporated by eukaryotic cells.

In the work by Santini and collaborators, three sequence of water soluble complexes with tetrahedral geometry (Figures 6 & 7), named [M(thp)$_4$]$_x$ (THP: tris(hydroxypropyl)phosphine), [M(thpp)$_4$]$_x$ (THPP: tris(hydroxypropyl)phosphine) and [M(PTA)$_4$]$_x$ (PTA: 1,3,5-Triaza-7-phosphaadamantane), where M = Cu(I), Ag(I) and Au(I), were tested. The Cu(I) complexes were always found to be the most effective against human breast (MCF-7), lung (A549), colon (HCT-15), cervix (HeLa) and melanoma (A375) cell lines. The IC$_{50}$ (μM) for [Cu(thp)$_4$]PF$_6$ (8) was 9.11 ± 2.71 (A549), [Cu(thpp)$_4$]PF$_6$ (9) was 18.22 ± 2.11 (A549) and [Cu(PTA)$_4$]PF$_6$ (10) was 17.40 ± 1.76 (A549). The best value found to colon cells (HCT-15) for 8 (IC$_{50}$ = 2.00 ± 0.03). Among the tested compounds, metal-THP species present better cytotoxic activity than metal-PTA and metal-THPP. The calculated logP specified that all tried complexes were hydrophilic in nature sorted as 8 (logP = -1.23) > 10 (logP = -0.97) > 9 (logP = -0.62), which can be related with antiproliferative activity of the complexes.

The SAR (Figures 6 and 7) revealed that the presence hydroxyl groups in 8 and 9 increased hydrophilicity. In a more acid cancer tissue, the most active aqua-form of PTA would be predominant through introduction of electron-withdrawing groups [31]. Herein the absence of these in PTA complexes could explain no significant IC$_{50}$ values. Thus, the decreasing lipophilicity of a molecule is an effective approach to design active Cu (I) complexes with hydrophilic tertiary phosphine ligands. In this context, the copper complexes are promising when compared with other metal complexes with phosphine derived ligands.

Furthermore, the cytotoxic effects of these copper complexes have proven to induce damage in ubiquitin-proteasome system (UPS). The UPS is a key mechanism in which eukaryotic cells degrades proteins to preserve the protein homeostasis. In normal cells, the natural load of degraded substrates is in equilibrium with the cell UPS. Instead, regarding the cancer cells this load is amplified for the reason of the overexpression of modified proteins [32-34].
Isatin Derived Ligands

Isatin (2,3-dioxindole) is an important heterocyclic moiety, structurally related with the indole privileged structure. In addition, isatin is also a privileged substructure for the design of novel molecules [35]. Some biological activities for derived compounds found in literature are antitumor, antimicrobial, anti-inflammatory, analgesic, anti- mycobacterial, anticonvulsant, antiviral, anthelmintic, antioxidant, and CNS depressant effects [36]. Among these, undoubtedly the antitumor activity has been defined as more evident, mainly for halogenated isatin-derivatives [37]. Then, complexes containing these worthy biologically active scaffolds cover a remarkable class of compounds for design of novel target-specific drugs. In fact, many copper complexes of isatin Schiff-base ligands exhibit a wide range of pharmacologic properties, such as antimicrobial, antibacterial, antifungal and antitumor [38-42]. Complexes formed by N1-substituted-isatin-3-thiosemicarbazone and Cu(I) halides were reported by Sharma and collaborators, as represented in Figure 8 [43]. Isatin-thiosemicarbazone ligands are described as k2N,S- or k2O,N,S-donors. The isatin-β-thiosemicarbazone has been recognized to be more selective to promote selective cell death to overexpressing P-glycoprotein cells. The P-glycoprotein is a key carrier in multidrug resistance cells (MDR) [44].

In the complex 11, Cu(I) ion is four-fold coordinated in a distorted tetrahedral environment. The metal center is connected by one bromide, one P atom from triphenylphosphine and one neutral isatin-N1-methyl-thiosemicarbazone that act as k2N,S-donor. The complex 12 differs from the former only by the halide ligand. These Cu(I)-compounds were screened in human tumor cell lines L123 (lung) and HepG2 (liver) to check their cytotoxicity profile. The IC50 values for complexes 11 and 12 ranged at 8.31-25.50 μM (L123) and 18.65-25.50 μM (HepG2), respectively. These data showed the influence of the electron-withdrawing effect from halide ligand, i.e., the higher electron-withdrawing properties of bromine atom than chloride one in cytotoxic efficiency.

The Cu (I)-complexes [CuBr(N3,S-H2jtsc-N1-Me)(Ph3P)] (11) and [CuCl(N3,S-H2jtsc-N1-Me)(Ph3P)] (12) are shown in Figure 8. In addition, chlorine atom presents an electron-withdrawing property higher than bromine atom, resulting in changes of pKa, membrane permeability, preferential orientation, potency and pharmacokinetic properties, as well as in better inhibition, with the lowest IC50 value.

Molecular modeling (MM) studies are extremely useful for better understanding of the correlation between drug candidates and their biological interactions [45]. The docking studies of the complexes to MFE-23 (anticarcinoembryonic antigen, PDB ID 1QOK) indicated the promising anticancer activity for them. The MFE-23 is an important antibody used as biomarker of carcinoembryonic antigen (CEA). Thus, several research groups have put efforts to design antitumor drugs with MFE-23-affinity. Based on MM studies, it was observed that H-donor group NH (green squares in Figure 8) forms hydrogen bonding interactions with O atom from side chains of glutamine (Q249) and tyrosine (Y247) residues. Likewise, NH group, showed hydrogen bonding contacts with S atom of cysteines (C248 and C184) and N atom of tryptophan (W195) residue.

On the other hand, the Cu(II) complexes can be used to cleave the DNA via oxidative pathway. In this case, in the work of Teoh and contributors, Cu(II) complexes of isatin thiosemicarbazone derivatives demonstrated this capacity [46]. Each Cu(II) metal center is bonded by one tridentate k3O,N,S-donor ligand, one chloride anion and
the metals are linked each other through two sulfide bridges comprising a binuclear complex. Each metal center is embedded in a distorted square pyramidal environment. Their activity in the human colorectal carcinoma cell line (HCT116) is represented in Figure 9.

**Figure 9:** Binuclear Cu(II) complexes of isatin thiosemicarbazone derivatives. The IC₅₀ values were obtained out by MTT assay, as represented for colorectal carcinoma cell line (HCT116).

These binuclear Cu(II) complexes demonstrate the biological importance of indole moiety for activity, highlighted in green squares (Figure 9). In addition, as evidenced in red squares in 14, the fluorine atom, with strong electron-withdrawing properties, resulted in high inhibition of tumor cells growth. In the case of complex 15, the presence of alkyl group as substituent resulted in decreased potency [47].

Furthermore, the cleavage DNA via oxidative pathway is observed. The complexes were incubated with H₂O₂ and DNA was degraded. These results support that Cu(II) complexes can cleavage plasmid DNA (pBR322 DNA) via ROS formation. The reduction of H₂O₂ into hydroxyl radical by the oxidation of transition metal complexes can damage DNA structure by strand-chain breakage. In this case, this process occurs through the redox conversion into Cu(II)/Cu(I) species [48-51].

In different analysis about targets, Bulatov and collaborators, reported the pharmacological evaluation of isatin-Schiff base analogues and its Cu(II) complex in different tumor cell lines by assessing their effects on cellular metabolism, real-time cell proliferation and toward p53-positive in MCF7 cells. Also, it was promoted p53-dependent gene expression inducing apoptosis. The tumor suppressor of p53 normally reduces the efficiency to arrest damaged cells from perpetrating their genetic errors into future generations. In fact, p53 is the most altered gene in cancer. The inactivation of tumors suppressors occurs in most types of human cancers [52,53].

Thus, the work, as represented in Figure 10, explored potential antitumor properties of Cu(II) complex 17 in p53-positive tumors, such as breast cancer (MCF7) cell line. To validate p53 associated effects of Cu(II) complex on tumor cells, a comparison between cells lines in different p53 status was performed, (p53 positive versus p53 negative).
In this context, the treatment of MCF7 p53<sup>WT</sup> cells with 17 increased the p53-protein levels. However, ligand 16 did not have effect on p53 activation. These findings suggested the higher cytotoxicity and more reduced proliferation of tumor cells from the complexes than free ligands [54].

The studies with p53 are extremely important in cancer research because this protein family works as tumors suppressor. Since inactivation of tumors suppressors occurs, the apoptosis mechanism is affected. Mutant p53 are expressed by p53-positive and p53-negative. They are performed to act into several important cell pathway, such as regulating cell division and metabolism. Thus, disturbs related with p53 can result in cancer promoting. Because of the loss function of p53, the over-expression of mutant p53 is generally considered as a bad prognosis factor, associated with nodal metastasis [51,55,56]. The Cu(II) complex 22 exhibited improved cytotoxicity toward p53-positive and p53-negative MCF7 cells and induced apoptosis signaling in both p53 MCF7 cells.

As a way to exemplify other applications of copper complexes on inorganic medicinal chemistry, Sreekanth and collaborators, studied 5-methoxyisatin thiosemicarbazones with different N-terminal substituents and their Cu(II) complexes, as represented in Figure 11 [57]. The isatin-derivative ligand acts as a tridentate <i>k</i><sup>3</sup>O,N,S-donor to Cu(II). Chloride anion completes a square-planar geometry around the metal center. Three different terminal substituents were chosen: pyrrolidine (18), morpholine (19) and cyclohexyl ammine (20).
The complexes 18 and 19 were evaluated for their antifungal, antibacterial, and antitumor activities in *S. aureus*, *B. subtilis* and *S. epidermidis* (Gram-positive), *E. Coli*, *P. aeruginosa* and *P. Vulgaris* (Gram-negative), *A. niger*, *A. flavus*, *F. solani*, *C. albicans* and *C. lunata* and HeLa (cervical), A549 (lungs), MCF-7 (breast) and HEK293 (normal embryonic kidney).

The Cu(II) complex 18 was the most active against tumors cells. It exhibited wide range of wide range of activity in MCF-7, A549 and HeLa cells, presenting IC$_{50}$ = 14.83 ± 0.45 µM, 17.88 ± 0.16 µM and 6.89 ± 0.42 µM, respectively. Doxorubicin was used as control. For Cu(II) complex 19 the IC$_{50}$ values obtained was 49.38 ± 0.16 µM, 55.52 ± 0.63 µM and 28.79 ± 0.32 µM, respectively in MCF-7, A549 and HeLa. As highlighted in red square for complex 18, the pyrrolidine substituent is more hydrophobic than cyclohexyl substituent from complex 20 (MCF-7, A549 and HeLa, with IC$_{50}$ values 25.02 ± 0.51 µM, 64.97 ± 1.06 µM and 10.86 ± 0.65 µM, respectively), due to the absence of the protic hydrogen (NH). Additionally, the authors concluded that the highest activity against tumor cells of complex 18 can be explained because of its greater hydrophobic nature in comparison with complexes 19 and 10. The compound 18 can interact with the hydrophobic center of DNA more effectively.46 On the other hand, the complex 20 showed promising potency as antimicrobial drug against *B. subtilis* (MIC: 10.35 µg/mL), *S. aureus* (MIC: 8.56 µg/mL), *P. aeruginosa* (MIC: 21.36 µg/mL), *S. epidermidis* (MIC: 6.50 µg/mL), and *P. vulgaris* (MIC: 17.64 µg/mL) in comparison with ciprofloxacin. The presence of H atom at N4-position makes the complex less lipophilic. Additionally, possible H-bonding contacts from this moiety (red square, in 20) can be the reason of activity against bacteria. Regarding the antifungal activity data, the complex 20 had better results against *A. niger* (MIC: 11.67 µg/mL), average activity in *C. albicans, A. flavus, and C. lunata* (MIC: 17.56 µg/mL, 8.14 µg/mL, and 14.38 µg/mL, respectively), compared to standard Nystatin and the other complexes 18 and 19. It is also important to highlight that the methoxy group (in green square, Figure 11), as an electron-donating group, can increase the electron density on aromatic ring, resulting on important properties for biological activity [58].

**Conclusion**

Some copper complexes with phosphine and isatin derivates as ligands, studied through 2010-2019 were reported in this review. We have to consider that copper allows different oxidation states, ligands and geometries of the metal center, leading to drug design versatility. In addition, as copper is an essential metal, it increases the chances for obtaining complexes with less side effects.

Copper complexes have different mechanisms of action to induce apoptosis in tumoral cells. For Cu(I)-based complexes the mechanism is related with the break of DNA backbone without redox activity. On the other hand, for the Cu(II)-based complexes, the mechanism can be linked to ROS production inside the cells. In addition, Cu(II)-complex can also bound to DNA by intercalation when planar ligands are used. Some copper complexes can show their cytotoxic effects through ubiquitin-proteasome system (UPS) and inducing mutant p53 for apoptosis. These aspects, combined with SAR information were explored in this review, furnishing suitable features for the design of novel bioactive copper complexes targeting anticancer activity.

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