Enhancing the Activity of Drugs by Conjugation to Organometallic Fragments

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Abstract: Resistance to chemotherapy is a current clinical problem, especially in the treatment of microbial infections and cancer. One strategy to overcome this is to make new derivatives of existing drugs by conjugation to organometallic fragments, either by an appropriate linker, or by direct coordination of the drug to a metal. We illustrate this with examples of conjugated organometallic metallocene sandwich and half-sandwich complexes, RuII and OsII arene, and RhIII and IrIII cyclopentadienyl half-sandwich complexes. Ferrocene conjugates are particularly promising. The ferrocene–chloroquine conjugate ferroquine is in clinical trials for malaria treatment, and a ferrocene–tamoxifen derivative (a ferrocifen) seems likely to enter anticancer trials soon. Several other examples illustrate that organometallic conjugation can restore the activity of drugs to which resistance has developed.

Introduction

Resistance to chemotherapy is a major clinical problem. This arises either from inherent lack of sensitivity to current drugs or induced resistance on repeated treatment. In particular, resistance is a problem for the treatment of cancer and microbial diseases. One approach to overcoming this problem is to design new drugs with novel mechanisms of action. Here we focus on the potential for discovery of novel drugs by incorporating organometallic fragments, so extending designs to the wider periodic table beyond purely organic drugs.[1] In particular we highlight the use of organometallic fragments to re-activate organic drugs towards which resistance has developed.

In general, metal complexes are now finding wider applications in medicine, particularly against cancer, but the basis for use has often been empirical without detailed understanding of structure–activity relationships, or identification of their target sites and mechanisms of action. However, times are changing.

The major development of platinum-based anticancer drugs has helped to establish metal complexes as viable treatments.[2–9] The platinum-containing drug cisplatin (Figure 1) and later generation analogues carboplatin and oxaliplatin are now widely used to treat various cancers.[2] More recently, nedaplatin, lobaplatin and heptaplatin have been approved for chemotherapy in certain Asian countries.[10] This family of drugs mostly target DNA, causing structural changes through Pt–DNA adducts which lead to apoptosis and cancer cell death.[2, 11] However, small changes in structure can dramatically change the target, for example, oxaliplatin targets ribosome biogenesis not DNA.[12] Other metal complexes have long been used in treatment, including arsenic trioxide, marketed as Trisenox, for the blood cancer acute promyelocytic leukaemia,[13–15] melasoprol, an organoarsenical for trypanosome infections in Africa,[16] and BiIII antimicrobial/antiulcer compounds such as colloidal bismuth subcitrate (CBS, De-Nol) and ranitidine bismuth citrate (RBC, Pylorid).[17] Padeliporfin di-potassium (TOOKAD) was recently registered for use as a vascular targeted photodynamic therapeutic agent against prostate cancer.[18] The gold complex, auranofin (Figure 1), is prescribed for rheumatoid arthritis treatment; classified as a disease-modifying antirheumatic which slows progression through suppression of inflammation and encouraging cell-mediated immunity.[19] Recently it has entered clinical trials as a broad-spectrum antiparasitic drug.[20]

There are increasing numbers of novel inorganic and organometallic complexes under development as therapeutic agents with several currently in clinical trials or already approved for clinical use.[2,21] In particular, compounds with metals from the platinum group metals (PGM)—ruthenium, rhodium, palladium, osmium, iodidium, and platinum—have risen to prominence in the search for new inorganic drugs, for example, highly water soluble RuIII chlorido-DMSO complexes like NAMI-A (Figure 2), first studied by Mestroni, Alessio, Sava et al.[22–24]
This drug is an anti-metastatic agent, either inhibiting metastasis formation\cite{25} or reducing the size of metastases.\cite{26} It completed phase I clinical trials.\cite{27–30} However, in a phase I/II combination with gemcitabine in patients with non-small cell lung cancer,\cite{31} it did not show improvement over gemcitabine alone.\cite{32}

The antineoplastic agent KP-1019 developed by Keppler and co-workers has completed phase I trials;\cite{30, 33–39} its sodium salt, NKP-1339 is in further clinical development.\cite{40} Ruthenium(II) arene complexes such as RM175\cite{38, 41} and RAPTA-C\cite{42–44} (Figure 2) have stimulated interest in the design of half-sandwich organometallic anticancer complexes, as well as possible applications for treatment of malaria and tuberculosis.\cite{9, 45–47}

TLD1433 is a polypyridyl ruthenium complex and first example of a RuII-based photosensitizer to enter human clinical trials.\cite{48}

The use of organometallic fragments in the redesign and repurposing of organic drugs shows promise for reactivation, combatting resistance, and increasing potency. Recently the power of ferrocene labelling, for example, has been widely demonstrated. Ferrocene is an organometallic compound containing FeI and two cyclopentadienyl ligands in a rigid sandwich structure. This unit is stable in physiological media and has a lipophilic nature that allows it to traverse cell membranes effectively.\cite{49–51} It can also be derivatised with relative ease to allow incorporation into a range of drug scaffolds.\cite{51–53}

In an early example (1975), the phenyl or heteroaromatic groups of penicillin or cephalosporin were replaced by a ferrocene moiety, giving new highly active antibiotics and potent β-lactamase inhibitors.\cite{54}

We have restricted our discussion to derivatives where the structure of the clinical drug remains largely intact and with applications against a variety of diseases and conditions.

### Derivatives of Cancer Drugs

Tamoxifen and its active metabolite hydroxytamoxifen (Figure 3) are breast cancer chemotherapeutics. Ferrocenyl derivatives of these drugs, aptly named ferrocifens, have been extensively studied by Jaouen, Vessières, Top and co-workers since the mid-1990s, starting with the ferroenyl complexes shown in Figure 3.\cite{59–61} These early complexes, are selective estrogen receptor modulators (SERMs), and exhibit good inhibitory effects on both MCF-7 (hormone dependent) and MDA-MB-231 (hormone independent) breast cancer cells in contrast to their organic precursor, hydroxytamoxifen, which is active only against the MCF-7 cell line. The discovery of this dual effect prompted the synthesis of several different libraries of ferrocifen-type complexes (Figure 4).\cite{57}

Each series has been extensively studied for their effects on MCF-7 breast cancer cells.\cite{57} In vitro antiproliferative studies on the breast cancer cell lines MDA-MB-231 and MCF-7, the human prostate PC-3 cancer cell line, and healthy cell lines,

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**Figure 2.** Early RuII and RuI complexes studied for anticancer activity. RuII complexes are thought to be activated by in vivo reduction to RuI.

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showed that these compounds have higher selectivity for cancer cells versus normal cells. Ferrocene itself is less active compared to the ferrocifen derivatives and the presence of the ferrocenyl-double bond-phenol motif (Series C, where \( R', R'' = H, \text{OH}, \text{OH} \), Figure 4) is essential for high antiproliferative effects. When the ferrocene unit is incorporated at the end of the sidechains as in Series E, a reduction in activity is observed relative to Series C. Series D, ansa ferrocenyl analogues, do not contain a dimethylaminoalkoxy group like the original ferrocifen complexes (Series A) and were more active than these open-chain homologues.

Comparison of Series C (acyclic) and Series D (ansa) also showed that the closed-ring analogues were more active; the complex with \( R' = \text{OH} \) exhibited an IC\(_{50}\) value of 0.09 μM for resistant MDA-MB-231 cells. It was seven times more potent than its acyclic derivative. A similar increase in activity against the resistant breast cancer cell line was obtained for the ansa complex where \( R' = \text{NH}_2 \) and \( R'' = \text{OH} \). Series F, containing suberoylanilide hydroxamic acid (SAHA) or phenylsuberamide (PSA), was more active (IC\(_{50}\) values between 0.5–0.7 μM) compared to ferrocifen-\( \text{OH} \) (IC\(_{50}\) = 2.6 μM), tamoxifen SAHA (IC\(_{50}\) = 8.6 μM) and tamoxifen PSA (IC\(_{50}\) = 25.2 μM) derivatives. SAHA inhibits histone deacetylases, enzymes involved in gene expression.

The osmium and ruthenium metallocene derivatives of hydroxytamoxifen (Series B) are not as active as their iron homologues (Series A). A detailed account of the synthesis, inhibitory activities and mechanistic studies on ferrocifens has been published. Oxidation of ferrocifens in cells to active quinone methide metabolites appears to play a key role in their mechanism of action.

In Series G, where the ethyl group was replaced with a longer ester or hydroxy-modified alkyl chain, the complexes exhibited low micromolar activity against MDA-MB-231 cancer cells.

Paclitaxel (Figure 5) is a diterpene that belongs to the group of antimitic agents known as taxanes. Marketed as Taxol or Onxal, it is an intravenous anticancer drug used in treatment of ovarian and breast carcinomas and Kaposi’s sarcoma. Paclitaxel exerts its chemotherapeutic effect through promotion of tubulin polymerisation and stabilisation and impairment of microtubules causing mitotic arrest and apoptosis of cancer cells. Ferrocenyl derivatives of paclitaxel have been studied for antiproliferative activity in the human tumour cell lines A549 (alveolar basal epithelial cell adenocarcinoma), Colo 205 (colorectal adenocarcinoma), Hep G2 (hepatocellular carcinoma), MCF7 (breast adenocarcinoma) and SW620 (colorectal adenocarcinoma) by Rychlik and co-workers. Efficacy was dependent on the position of...
the ferrocene moiety, and the N-benzoyl substituted derivatives (ferrocene-paclitaxel derivatives type I, Figure 5) showed the most promising activity. The spatial positioning of ferrocene greatly affected the activity; for the type I derivatives the best activity was observed when there was no aromatic spacer (IC_{50} = 0.005–0.0015 \mu M), against W620, A549, Colo 205, HCT116, Hep G2 and MCF7 cells. Lower activities were noted for complexes containing the ferrocene substituted N-benzoyl group in the order m-substituted > p-substituted > o-substituted. Substitution of the 3'-phenyl group in the phenylisoserine side chain of paclitaxel with ferrocenyl groups (ferrocene-paclitaxel derivatives type II) did not yield any notable activity. These complexes were also assayed for activity in a panel of five multidrug-resistant cell lines derived from SW620 and characterized by overexpression of various ATP-Binding Cassette (ABC) proteins, namely ABCG2 (SW620C line), ABCB1 (SW620M and SW620E lines) and ABCB1 (SW620D, SW620E, and SW620V lines). ABC proteins are a group of transporter proteins responsible for drug resistance and reduced bioavailability of drugs by efflux of the drug from cells. In particular, the expression of the ABCB1 efflux protein adversely affects the activity of taxanes in cancer cells. Complexes of type I showed good inhibition of the ABCB1-expressing cell lines, SW620D, SW620E and SW620V. Determination of the induction of tubulin polymerization and ROS generation by these compounds led to the conclusion that there is no general mechanism for these ferrocenyl-paclitaxel complexes. Compounds showing robust pro-oxidative properties exhibited lower anticancer activities and complexes that were good polymerization inducers showed either high or low anticancer activity. Nevertheless, these ferrocenyl-paclitaxel complexes are a good starting point for further development of ferrocene taxane anticancer drugs and it is likely that one such complex will reach clinical trials.

5-Fluourouracil is administered intravenously to treat a variety of cancers including, colon, stomach and breast, and is used topically for some skin cancers. Moderate inhibition was observed for ferrocene-uracil derivatives (Figure 5) in the MCF-7 breast cancer cell line. Structure-activity analysis showed that introduction of methyl or fluoro substituents on the uracil ring in Ferrocenyl-uracil-A activated the antiproliferative effects, with Ferrocenyl-uracil-B being the most potent (IC_{50} 23.8 \mu M). The ruthenocenyl-uracil complex did not show any appreciable activity and measurement of ruthenium levels in HT-29 cells after exposure to a 100 \mu M concentration of the complex revealed that Ru is taken up by the cells over time but poorly, which may account for the low activity observed. These complexes were not screened in a resistant cancer cell line so their selectivity cannot be assessed.

Antibacterial screening of ferrocene-uracils C-E on Gram-positive Staphylococcus aureus (SA) and Staphylococcus epidermidis (SE) bacterial strains revealed moderate to good activity for these complexes. Their MIC values were the same for methicillin-sensitive (MSSA), methicillin-resistant (MRSA), and vancomycin-resistant (VRSA) strains, suggesting no selectivity towards resistant bacteria. Ferrocenyl-uracil-D was the most promising, with MIC values of 28 \mu g\,mL^{-1} on the SA strains and 16 \mu g\,mL^{-1} on the SE strain. 5-Fluourouracil was not tested at the same time as the ferrocenyl complexes.

### Derivatives of Antibiotics

Fluoroquinolones (e.g., ciprofloxacin, CP, Figure 6) are a class of antibiotics, used especially for treatment of respiratory and urinary tract infections. CP is a broad spectrum agent also used as a prophylactic for neutropenia and in veterinary medicine. CP is active against both chloroquine-sensitive and chloroquine-resistant Plasmodium falciparum strains, which led to studies of its effect on patients with acute falciparum malaria. However, the drug was less potent than chloroquine; displaying delayed inhibitory effects, thus limiting its potential use as an antimalarial. As a result of success with ferroquine (vide infra), Biot and co-workers extended their search for organometallic conjugates to ferrocenyl-CP derivatives. An enhancement of efficacy was reported when compared to CP and the antimalarial drugs, chloroquine, quinine and artemunate, displaying IC_{50} values between 0.8–3.9 \mu M in CQ-resistant (W2) and CQ-sensitive (3D7) parasite strains. All the complexes were active against both W2 and 3D7, suggesting that these ferrocenyl-CP derivatives are either not susceptible to the same resistance mechanisms as CQ, or that they have different molecular targets. Complex Fc-CP-A was the most potent, with IC_{50} values of 1.0 and 0.8 \mu M against 3D7 and W2, respec-
tively, 20 times greater (3D7) and 460 times greater (W2) than CQ, and comparable to artesunate.\textsuperscript{[79]}

The rhenium carbonyl derivative of CP (Figure 6) has activity similar to free CP towards Gram positive and Gram negative bacteria.\textsuperscript{[76]} The Ru\textsuperscript{a} arene complex p-cy-Ru-CP containing chelated CP retained moderate anti-microbial activity in a clinical E. coli isolate that is resistant to first, second and third generation β-lactam antibiotics.\textsuperscript{[77]}

Organometallic derivatives of the anti-TB drug isoniazid\textsuperscript{[78, 79]} (INZ) have been widely studied, with its pyridyl ring being an attractive metal coordination site. Tirkey et al. reported the synthesis of cymantrenyl hydrazone (cymantrenyl-INZ) and ferrocenyl hydrazone (ferrocene-INZ-A-D) analogues of INZ (Figure 7).\textsuperscript{[80]} Selected complexes were tested for activity in bacterial strains B. subtilis, E. coli, S. aureus, K. pneumoniae and P. aeruginosa. Cymantrenyl-INZ was generally less active (MIC values 125–250 \(\mu\)gm L\(^{-1}\)) compared to the ferrocenyl analogue (ferrocene-INZ-A; MIC values: 31.25–125 \(\mu\)g mL\(^{-1}\)). No data for INZ were reported. The activity of these complexes was also compared to reported MIC values for organic derivatives or the ferrocenyl and cymantrenyl complexes; the organometallic complexes were found to have superior antimicrobial activity.\textsuperscript{[80]} The authors speculated that the presence of the ferrocenyl and cymantrenyl groups may increase cell permeability and lipophilicity of these molecules thus increasing activity. They also suggested that \(\pi\)-electron delocalisation and hindrance of metal binding sites in crucial enzymes in the microorganisms could also play a role. However, no experimental data confirming this were reported.

Stringer and co-workers carried out structure–activity studies of ferroceny (ferrocene-INZ-E-G) and heterobimetallic complexes (ferrocene-INZ-H-J) in malarial, trichomonal and mycobacterial strains.\textsuperscript{[81]} Most of the complexes showed higher antimycobacterial activity towards \(M.\) tuberculosis (H37Rv strain) in the glycerol-based GAST-Fe growth medium (MIC\(_{90}\) or MIC\(_{99}\) < 1 \(\mu\)M). Ferrocene-INZ-E (containing no aromatic spacer between INZ and ferrocene) and the heterobimetallic complexes showed activities comparable to the free parent drug, INZ. None of the complexes were as active as INZ against \(Mtb\). As antiparasitics, low to moderate inhibition was observed against the \(T.\) vaginalis strain G3. Interesting trends were noted in the NF54 chloroquine-sensitive strain of \(P.\) falciparum. In contrast to the antimycobacterial activity, INZ was found to be inactive against the NF54 PF strain. The ferrocene-INZ-E was not active (>100 \(\mu\)M) while ferrocene-INZ-F and G containing the aromatic spacer were remarkably more active; ca. 3 times greater for ferrocene-INZ-F (31.52 \(\mu\)M) and 65 times greater for ferrocene-INZ-G (1.58 \(\mu\)M). The heterobimetallic complexes ferrocene-INZ-H-J also displayed low IC\(_{50}\) values of 2.99–7.82 \(\mu\)M. Screening against the normal Chinese Hamster Ovarian (CHO) cell line revealed low toxicity, indicative of good selectively for microbial strains.

Ruthenium-arene complexes (plumbagin-INZ Ruthenium arene A and B) of a hybrid ligand prepared from the natural plant product plumbagin, a yellow naphthoquinone dye, and INZ have been reported by Spoerlein-Guettler et al.\textsuperscript{[82]} Due to the relative insolubility of plumbagin-INZ Ruthenium arene B, only plumbagin-INZ Ruthenium arene A was tested for anti-cancer activity on two cell lines that are sensitive to plumbagin (518A2 melanoma and HCT-116 colon carcinoma cells), the multi-drug resistant KBV1/Vbl cervix carcinoma cell line and non-malignant CHF chicken heart fibroblast cell line. Compared to free plumbagin and cisplatin, the ruthenium complex showed higher activity in all tumour cell lines and was not toxic to the CHF cell line up to 50 \(\mu\)M. Notably, in the multidrug resistant cell line (KBV1/Vbl) plumbagin-INZ-Ru-arene A displayed an IC\(_{50}\) of 3.80 \(\mu\)M while cisplatin and plumbagin were less active (IC\(_{50}\) 11.4 and 26.2 \(\mu\)M, respectively), promising for further work on drugs that can overcome resistance.

Isoniazid has also been incorporated into the Ru\textsuperscript{a} complex bpy-Ru-INZ (Figure 7). This complex was screened against Gram-positive Bacillus subtilis, Gram-negative E. coli, and Mycobacterium smegmatis, which possess the three major classes of cell envelope. It was inactive in the dark, but rapidly (<500 ps) released isoniazid on irradiation of the complex with visible light.\textsuperscript{[83]} The complex was 6 times more potent towards \(M.\) smegmatis compared to isoniazid alone and relatively nontoxic towards normal human cells. This strategy for combating AMR could be applied to a range of Ru\textsuperscript{a} and Ir\textsuperscript{a} photoactivatable organometallic complexes.

Biguanides can exhibit antimicrobial and antiviral activity, and recently have been incorporated as chelating ligands into Cp\(^*\), Cpy\(^{ph}\) and Cpy\(^{phph}\) complexes.\textsuperscript{[84]} Several such complexes, containing inactive biguanides, exhibit potent activity against Gram-negative and Gram-positive bacteria (including methicillin-resistant Staphylococcus aureus (MRSA), and high antifungal potency towards C. albicans and C. neoformans, with minimum inhibitory concentrations (MICs) in the nanomolar range, and low cytotoxicity towards mammalian cells. They can restore
the activity of vancomycin against vancomycin-resistant Enterococci (VRE) and disrupt and eradicate bacteria in mature biofilms.

**Derivatives of Antiparasitic Drugs**

The aminoquinoline pharmacophore has been incorporated into many antiparasitic drugs, particularly antimalarials.\[^{85,86}\] Although resistance to several of the current quinoline drugs is emerging, they still remain attractive for derivatisation.\[^{85}\] Organometallic conjugates of such drugs also show potential for overcoming resistance, Ferroquine being an important example.\[^{87–90}\] Chloroquine (CQ, Figure 8) belongs in the 4-aminoquinoline structural group and is currently used in regions affected by CQ-sensitive strains of *P. vivax*, *P. ovale*, and *P. malariae*. It is no longer recommended for treatment of malaria caused by *P. falciparum*, due to extensive resistance.\[^{91}\]

CQ is administered orally and has also been prescribed for other illnesses such as rheumatoid arthritis and lupus.\[^{92}\] The mononuclear Rh\(^{1}\) complex (CQ-Rh) proved active against *P. falciparum* strains in vitro.\[^{93}\] It exhibits activity similar to CQ diphosphate, and in vivo studies against rodent malaria parasite *P. berghei* revealed that at the same concentration of CQ diphosphate required to kill 50% of parasitemia, CQ-Rh reduced parasitemia by 74%. Ferroquine has a ferrocene moiety incorporated into the side-chain of CQ and has impressive antiplasmodial activity in both CQ-sensitive and CQ-resistant strains.\[^{6,88–90}\] It has completed Phase I and is expected to complete Phase II clinical trials against uncomplicated malaria soon. Ruthenium-arene CQ conjugates (CQ-Ru-A–D) have been evaluated for in vitro activity against three chloroquine resistant (W2, Dd2 and K1) and four chloroquine sensitive (FcB1, 3D7, PFB and F32) *P. falciparum* strains.\[^{94}\] They were moderate inhibitors of the chloroquine-sensitive *P. falciparum* strains with activities lower than CQ diphosphate. However, against CQ-resistant strains, these complexes displayed consistently better activity compared to CQ diphosphate. The most potent inhibitor was CQ-Ru-C (Figure 9), which was 5x more potent than CQ diphosphate against the Dd2 and K1 strains. These results illustrate clearly the concept of enhanced activity of metal-chloroquine conjugates against resistant malarial strains.

The antimalarial primaquine (PQ, Figure 9) is a prescribed oral treatment for malaria caused by *P. vivax* and *P. ovale*, and *Pneumocystis pneumonia*.\[^{95}\] Matos et al have studied the antimicrobial efficacy towards plasmodial, leishmanial and pneumocystal strains of a library of ferrocene-primaquine conjugates (Fc-PQ-A–M).\[^{92,95,96}\] Against the CQ-resistant *P. falciparum* strain W2, only complex FC-PQ-F was active, whilst all of them were active against *Pneumocystis carinii*.\(^{95}\) The complexes containing amino acid type linkages between PQ and ferrocene showed anti-Pneumocystis activity with FC-PQ-F and FC-PQ-G being the most potent (inhibitory concentrations; 2.69 and 1.15 \(\mu\)g/mL, respectively). It appears that the degree of lipophilicity of the amino acid linkers in these complexes may play a role; higher activity was observed for the complexes that contained the more lipophilic groups (iso-butyl and benzyl). In vivo studies of the complexes in the liver stage model *P. berghei ANKA-GFP*-infected BALB/c mice and *Anopheles stephensi* mosquitoes were carried out to confirm if these complexes could inhibit the sporogonic cycle of plasmodia.\[^{96}\] Complexes FC-PQ-B and FC-PQ-G displayed high dose-dependent inhibition with the former able to completely impair the parasite’s sporogonic cycle at 50 \(\mu\)mol kg\(^{-1}\) (95% inhibition); a significantly higher activity than free PQ (26.9% inhibition). In vitro *P. berghei* inhibi-
tion assays found that most of the complexes show low micromolar IC\textsubscript{50} values with five of the FC-PQ complexes performing better (IC\textsubscript{50} = 0.17–2.82 μM) than PQ (IC\textsubscript{50} = 7.50 μM). As a result of the appreciable antimalarial activity, they were also screened against the promastigote and intramacrophagic amastigote forms of L. infantum.\textsuperscript{100} All complexes inhibited growth of L. infantum to some degree and complex Fc-PQ-C was identified as a potential new antileishmanial lead. At 60 and 40 μM concentrations, it caused >96% reduction in amastigotes per 100 macrophages in the intramacrophagic amastigote form of the parasite.

Artemisinin is a natural product that has been used for decades to treat malaria. Its derivatives, artesunate and artemether (Figure 10) are also part of the arsenal of WHO-recommended drugs for malarial infections caused by P. falciparum.\textsuperscript{97} The mechanism of action of artemisinin (ART) and its derivatives has yet to be clearly defined, but it is accepted that these drugs selectively target parasites because of their enhanced uptake by parasitized erythrocytes compared to healthy erythrocytes.\textsuperscript{98} The endoperoxide bridge present in ART likely interacts with the iron in haem to generate radical species that affect the redox balance of the malaria parasite.\textsuperscript{99}

Reiter and co-workers synthesised a series of ferrocene-artemisinin (Fc-ART) derivatives (Figure 10) that contained one or two ART moieties.\textsuperscript{100} Their activities were assessed against two leukaemia cell lines (human acute lymphoblastic CCRF-CEM and the multidrug resistant CEM/ADR5000), human cytomegalovirus (HCMV) and Pf strain 3D7. All the complexes (Fc-ART-A–D) were highly potent against CCRF-CEM cells (IC\textsubscript{50} ≤ 0.13 μM) and CEM/ADR5000 (IC\textsubscript{50} ≤ 1.96 μM). The complexes containing two ART moieties (Fc-ART-B and Fc-ART-D) are almost twice as active against CCRF-CEM cells as the complexes with just one ART (Fc-ART-A). However, for CEM/ADR5000, the opposite effect was observed. With the exception of Fc-ART-C, all the complexes exhibited activities of <0.50 μM towards the HCMV cell line. Their inhibitory effect on the CQ-sensitive P. falciparum strain was also significant (IC\textsubscript{50} values 7.20–30.2 μM).

de Lange et al recently reported a series of amino-artemisinin-1,2-disubstituted ferrocene hybrids which contain a piperazine linker between ART and various ferrocenyl moieties (Fc-ART-E–H, Figure 11).\textsuperscript{101} They were tested on asexual blood stage strains of P. falciparum (CQ-sensitive NF54 and CQ-resistant K1 and W2) as well as early and late blood stage gametes (ESG-NF54 and LSG-NF54). All complexes displayed notable activity, with Fc-ART-G, which contains a morpholino fragment bonded to Fc, being the most potent (IC\textsubscript{50} = 0.86 nM (K1) and 1.4 nM (W2)). Gametocytocidal activity was tested at 1 μM and 100 nM concentrations and for ESG-NF54 complexes Fc-ART-E–G showed >93% inhibition at both concentrations; they were slightly less active (≥84%) in LSG-NF54. Fc-ART-E–H were further screened against the human melanoma cell line A375 and normal human embryonic kidney cell line HEK293. With the exception of Fc-ART-G, all complexes had higher activity against A375 compared to HEK293. It is significant that ferrocenyl-artemisinin derivatives inhibited both sexual and asexual blood stages of P. falciparum, since this may lead to the preparation of new drugs that can block transmission.

Complexes Fc-ART-I and Fc-ART-J also have high antiproliferative activity towards CCRF-CEM lymphoblastic leukaemia cells.\textsuperscript{102} Fc-ART-J contains a ferrocenyl diester bridge between the ART moiety and the bioactive natural benzofuran eugonol and was less potent than Fc-ART-I towards drug-resistant CEM/ADR5000. Overall, Fc-ART-I is a promising new lead with good efficacy in both leukaemia cell lines.

Recently, Gasser et al. have investigated the design of organometallic derivatives of various anthelmintic drugs as anti-par-
Anthelmintics are a class of drugs used to treat diseases caused by parasitic worms. One of their earliest reports focused on ferrocenyl derivatives of Praziquantel (PZQ).\[103\] PZQ, marketed as Biltricide, is an effective treatment for urogenital and intestinal schistosomiasis.\[108\] However, low cure rates have been reported in some countries, thus prompting study of new drug leads.

An extensive library of Fc-PZQ complexes (Figure 12) was tested for activity on Schistosoma mansoni (S. mansoni), cervical cancer cell line HeLa, and the healthy lung cell line MRC-5. Most of the complexes were either inactive or mildly active against S. mansoni. The Fc-PZQ derivatives that showed some inhibitory effects on HeLa cells were less active in healthy MRC-5 cells.\[103\] Half-sandwich chromium carbonyl complexes of PZQ were also prepared.\[104, 107\] PZQ has a stereogenic centre (labelled * in Figure 12), and thus the complexes were initially tested as racemic mixtures (Cr-PZQ-A and Cr-PZQ-B).\[107\] Compared to their ferrocenyl analogues, Cr-PZQ-A and Cr-PZQ-B were better inhibitors of S. mansoni, displaying IC$_{50}$ values of 0.25 and 0.27 μM respectively. Additionally, they were inactive up to a concentration of 100 μM against MRC-5 cells and showed low activity in HeLa cells. The complexes were also relatively stable in D$_2$O and in human plasma. These promising results prompted investigation into preparation and testing of the pure optically active diastereomers of Cr-PZQ-A and Cr-PZQ-B and the three major metabolites identified from in vitro metabolism studies (PZQ, Cr-PZQ-C and cis-4-PZQ-OH, Figure 11).\[104\] The data showed that only the (R)-isomers, complexes (R,Rp)-Cr-PZQ-A and (R,Sp)-Cr-PZQ-B, were highly active with IC$_{50}$ values similar to free PZQ. They were also better inhibitors compared to Cr-PZQ-C and cis-4-PZQ-OH. In vivo studies in mice infected with chronic S. mansoni revealed that, in contrast to the in vitro data, the racemates Cr-PZQ-A and Cr-PZQ-B were toxic and only reduced the worm burden by 24 and 29%, respectively, at a single dose of 400 mg kg$^{-1}$. At this dose, PZQ reduces the worm burden by 96%.

Oxamniquine (OXA) is another antischistosomal agent and organometallic derivatives of OXA (Figure 13) were designed with the premise that these complexes may show different binding modes compared to free OXA in the enzyme, S. mansoni sulfotransferase.\[105\] The most active compounds in vitro were also tested in an in vivo mouse model. Of the four ferrocenyl derivatives (Fc-OXA-A-D), only Fc-OXA-A showed 100% reduction in the viability of S. mansoni newly transformed schistosomula (NTS) in vitro. Fc-OXA-B was the second most active complex at 76.2%; higher than free OXA. In the in vivo studies, at a 200 mg kg$^{-1}$ dose, only Fc-OZA-A and free OXA showed 100% clearance of all worms from the host.

Two additional OXA derivatives, Rc-OXA and Bz-OXA were synthesised to understand whether the ferrocene moiety of Fc-OXA-A is essential for activity, or if a purely organic derivative would be more active. In both the in vitro and in vivo models, the ruthenocenyl complex (Rc-OXA) showed superior activity over Fc-OXA-A and Bz-OXA.

Organo-ruthenium, rhodium and iridium derivatives of the sulfad drug sulfadoxine (Figure 14) have been reported by Chellean et al.\[109\] Screening for in vitro activity against CQ-sensitive 3D7, CQ-resistant Dd2 and LSGNF54 parasite strains and Mycobacterium tuberculosis found that the OM-SFD ruthenium derivatives were inactive. The Rh analogues were more potent than their iridium counterparts but overall, the OM-SFD type B iridium complex containing the quinoline-imino functionality and the Cp$^{6}$Ph ligand was the most active in the whole library. More...
over, the parent drug, sulfadoxine, was not active in most of the assays, affirming the hypothesis that organometallic conjugates of drugs can beneficially affect bioactivity. Most of the Rh and Ir complexes displayed inhibitory activity in the sexual LSG assay, while the clinical drugs, pyrimethamine, sulfadoxine and chloroquine were not. This is yet another study that validates the idea of organometallic conjugation of a drug can alter its biological target, as noted by others.[110–122]

Derivatives of Antifungal Drugs

Clotrimazole (CTZ), miconazole (MCZ) and tioconazole (TCZ) (Figure 15) belong to the azole antifungal drug class. Mono-, bis- and tris-azole ruthenium-arene complexes, where the azole drug is bound to the metal via one of the imidazole nitrogen atoms, were assayed for antifungal and antiparasitic activity.[123] As antifungals, none of the complexes were as active as their corresponding free azole drug. However, the complexes showed good reduction in radial fungal growth at both 0.5 and 0.01 mm concentrations. At a dose of 100 mg L−1, all of the complexes containing MCZ as ligand(s) showed moderate antischistosomal activity, and interestingly, the tris-MCZ ruthenium complex also showed gender-specific selectivity. After 72 h exposure to the complex, female worms showed only a marginal decrease in motility, while all of the male worms died. The mono-azole complexes decomposed slightly in DMSO, while the bis- and tris- complexes remained intact.

Sanchez-Delgado and co-workers prepared a small library of ruthenium-arene CTZ complexes (Ru-CTZ-A-D) for application as antileishmanial and antitrypanosomal agents.[124] They noted a synergistic effect on activity of CTZ metalation. All of the complexes displayed LD50 values lower than free CTZ for both promastigotes of L. major and epimastigotes of T. cruzi. The most potent complex was Ru-CTZ-A (LD50 = 0.015 μM (L. major) and 0.10 μM (T. cruzi)); respectively 110 × and 58 × more active than free CTZ. Ru-CTZ-A-D also did not show any appreciable toxicity towards human osteoblast cells.

The cyclopentadienyl Ru complex, Ru-CTZ-E, has more potent antiparasitic and antitumor activities compared to free CTZ.[125] In T. cruzi and T. brucei, Ru-CTZ-E was 7 × (IC50 = 0.25 μM) and 41 × (IC50 = 0.60 μM) more active than CTZ, suggesting that this complex could be developed as a broad spectrum drug for trypanosomatid parasites. It was also highly cytotoxic in ovarian A2780, breast MCF7, and cervical HeLa cancer cell lines, inhibiting proliferation at concentrations between 0.50 and 5.0 μM.

Compared to free CTZ (IC50 = 5.7–20.5 μM) and cisplatin (IC50 = 1.9–28 μM), Ru-CTZ-E was a more potent antitumor agent, suggesting potential for anticancer treatment.

Conclusions

In this review we have described briefly how the conjugation or direct coordination of clinical drugs to organometallic fragments can enhance or restore biological activity of anticancer and antimicrobial drugs. Such conjugation has an effect on drug uptake and delivery and can change target sites, introducing modified mechanisms of action that can combat resistance, and sometimes reduce off-target toxicity.[51–54] The best studied anticancer conjugates are the ferrocene-tamoxifen derivatives of Jaouen et al. which target oestrogen receptors.[57] The addition of the ferrocene fragment introduces new redox chemistry into the mechanism of action through formation of quinone methides as a result of oxidative metabolism. Although structurally analogous, the ruthenocene and...
osmocene derivatives have different redox properties and are less active. Similarly, organometallic conjugation can improve the activity of antibiotics. The most successful example to date is that of ferroquine which as a chloroquine-ferrocene conjugate is active against resistant malarial strains, and relatively non-toxic (at gram doses). Ferroquine is not only in clinical trials for malaria treatment, but also a candidate for repositioning as a cancer therapeutic. Its ability to induce lysosomal dysfunction may be important to the mechanism of anticancer activity. Ferroenyl conjugates of the antiinflammatory drug artemisinin also exhibit antiparasitic activity, and interestingly also exhibit antileukaemic activity. Organometallic half-sandwich fragments can also be effective in restoring activity to antibiotics. For example, some cyclopentadienyl IrIII fragments can restore activity to the antiparasitic drug sulfadoxine.

Fragment-based drug design (FBDD) is now a well-established method for exploring chemical space and discovering new drugs. Several drugs designed using this approach have already entered the clinic. The examples described here show that organometallic fragments can bring a new dimension to FBDD, providing fragments which offer more than just structural (e.g. H-bonding, hydrophobic and electrostatic) interactions with target sites such as proteins. Organometallic fragments can redox centres with potentials tunable by not only the C-bound ligand(s), but also the other ligands on the metal ion. They can also act as transporters of other biologically active molecules. In transition metal complexes, C-bound ligands often have high trans and cis effects which promote the release of other ligands. Organometallic fragments containing carbon monoxide ligands, for example, can be activated by photo-irradiation, a potentially attractive procedure for achieving selective attack on target cells and minimizing side-effects. Bearing in mind the large number of metal ions which form stable organometallic compounds, the range metal oxidation states available, as well as variations in coordination number and geometry, organometallic fragments can introduce a vast range of new pharmacophores with unique interactions with target sites. This strategy holds much promise for the discovery of drugs with novel mechanisms of action which can combat resistance.

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**Conflict of interest**

The authors declare no conflict of interest.

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**Keywords:** anticancer complexes · antimicrobial complexes · bioinorganic chemistry · bioorganometallic chemistry · coordination chemistry

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