Introduction

Since the Barnett Rosenberg’s discovery of cisplatin in 1965, the use of metal-based compounds in medicine, particularly in the treatment of cancer, has been revolutionized by further development of other platinum drugs such as carboplatin and oxaliplatin. The study of metal-based complexes with potential biological applications has expanded over the course of the years. There are numerous records from the history of metal-based treatments, ranging from the use of mercury(II) sulfide (cinnabar) by the ancient Greeks to the use of arsenic trioxide (trisenox) by traditional Chinese medical practitioners and the use of zinc(II), gold, and copper chelating agents in ancient Egypt to treat various diseases. In nature, various metals found in metalloenzymes and biocatalysts are essential for normal cellular functions. Most of the metals involved in biological processes belong in the transition metal group, occurring in the d-block of the periodic table (groups 3–12). These metal ions are positively charged and exhibit variable oxidation states in aqueous solution, making them very ideal for interaction with negatively charged biological molecules, such as nucleic acid or amino acids. These metal complexes are able to form various three-dimensional configurations due to the range of coordination geometries, leading to various metal–ligand interactions. One of the most common metal complexes in human biology is vitamin B12 (cobalamin) (►Fig. 1), which is required for deoxyribonucleic acid (DNA) and ribonucleic acid (RNA) synthesis and acts as an intracellular superoxide scavenger. The transition metal complexes are typically distinguished by metal ions with more than one coordination
number, which can lead to the formation of a large number of stereochemically and geometrically diverse and stable scaffolds. Metal complexation leads to dramatic changes in the biological activity of ligands, such as improved potency of drug molecules (ligands) that are complexed to metal ions and/or reductions in adverse side effects. The discovery of ferrocene by Kealy and Pauson at Duquesne University, and the subsequent incorporation of this molecule in chloroquine (CQ), suggested that the combination of organometallic compounds with known antimalarial drugs could result in potent antimalarials. Ferroquine (FQ; SSR 97193) is a 4-aminoquinoline CQ analogue which is functionalized with the organometallic ferrocenyl moiety. The metal-modified FQ has been shown to be active against CQ-resistant and CQ-sensitive *Plasmodium falciparum* strains and against *Plasmodium vivax*. FQ has also been shown to potently inhibit autophagy, perturb lysosomal function, and impair prostate tumor growth in vivo.

*N*-heterocyclic carbenes (NHCs) are one of the most widely used ligands in the formation of metal–ligand complex because of their strong α-donating and adaptable π-accepting abilities. NHCs have a very modular structure that allows for the selective modification of their steric and electronic properties. The electronic structure of NHCs can be tuned almost at will by changing the nature of the heterocyclic ring or its substituents, while the steric requirements can be strongly tuned by the nature of the *N*-substituents. In addition to NHCs, Schiff base ligands are widely used in metal complexes due to their low cost, ease of access, simplicity of synthesis, and chemical and thermal stability.

In general, the use of metal-based drugs gives rise to toxic side effects due to the in vivo development of free radicals and reactive oxidant species. The metals normally found in the body that are essential in biological processes include iron, which is involved in redox reactions of oxygen and copper to form heme, a catalytic cofactor in enzymes that can assist in overcoming these side effects. With similar properties to calcium ions, lanthanide(III) ions (classified as rare earth metal ions) can also be used in metal complexes because they possess pharmacological properties such as anticoagulation, anti-inflammatory, antibacterial, antiallergic, and anticancer properties, and exhibit paramagnetic properties at meanwhile. Lanthanides present in pyrrolquinoline quinone-dependent alcohol dehydrogenases in methylotrophic bacteria, where the highly selective Ln(III)-binding protein lammodulin plays a role in intracellular transport.

**Mechanisms of Action**

Endogenous and environmental damage is constantly being inflicted on cellular DNA, resulting in cellular dysfunction and cell death. Nucleic acids remain a key player in the central dogma of molecular biology, where the genetic code in DNA can be transcribed onto messenger RNA and translated into a protein form. DNA damage response (DDR) signaling networks have evolved in cells to ensure genomic stability and to sustain continuous cellular progression and growth. DDR defects play a role in the development of cancer and drug resistance in microbes. One of the mechanisms of action of metal complexes is to target nucleic acid (DNA/RNA), contributing to the disruption of transcription, translation, and other processes, which may ultimately cause death of the cell. In addition to targeting the nucleic acids, the metal complexes can also interact with proteins such as key DNA repair enzymes in DDR signaling pathways. Roughly 30 to 40% of known proteins require transition metals for their normal human biological activity. These metals play a role in proteins by acting as Lewis acids, thus providing reactivity in biocatalytic processes.
These metal ions facilitate redox chemistry through their multiple oxidation states, and they are able to act as cross-linking agents by coordinating at multiple sites in proteins. The action of metal complexes is not, however, selective, in that they do not differentiate between normal cells and tumor cells, and therefore severe side effects would occur due to normal tissue damage. By substituting the metal, changing its oxidation state, and/or coordinating geometry and coordination number, it is possible to fine-tune the metal-complex chemical stability, ligand exchange rate, metal–ligand bond strength, redox potential, ligand conformation, and outer-sphere interactions. Therefore, through the rational design of ligands and metal complexes, it is possible to prepare a variety of metal complexes with potential biological application. Depending on the design of the metal complex, the mechanism of action can proceed via several very different binding interactions with DNA.

**Structure–Activity Relationship**

The structure–activity relationship (SAR) describes the correlation between the chemical structure of a drug molecule and its biological activity. The SAR can be a powerful tool to permit the selection and optimization of potential drug candidates in drug discovery research. SAR studies favor the discovery of structural elements, such as metal components, coordination mode variations, labile water molecules, chelated ligands, etc., that may have an important cooperative effect on biological activity. The SAR can be influenced through the choice of ligand type (e.g., arene), metal, coordination mode, rate of hydrolysis, charge, and pKa. The choice of the ligand (lipophilicity) can influence the cytotoxic activity and coordination mode (where S,O, S,N, C,N, and N,N-bidentate donor systems [excluding bipyridines and bipyrimidines] generally are reported to yield high biological activity with more potency toward metallo drugs tethered to bioactive ligands such as indoloquinolines, paulilones, and flavonoids) (Fig. 2). Thus, both ligand structure and organometallic fragment are beneficial in bioactivity, where the latter entity modulates the solubility of the bioactive ligands.

A key feature of metal complexes acting as a prodrug is the labile ligand which dissociates from the metal to give the active species. The presence of the leaving group can exert subtle effects, where chlorido and iodido complexes are more potent than the bromido analogues. The rate of hydrolysis is critical, because if the complexes hydrolyze too quickly, they may not reach the target site, thus, the rate of hydrolysis is a key factor for metal complexes. A study on a series of charged (promoting interaction with oppositely charged biomolecules) ruthenium polypyridyl complexes revealed good antibacterial activity, owing to the target of this class of compounds being the highly negatively charged bacterial surface, resulting in damaged and deformed cell walls. Many properties of molecules, such as solubility, lipophilicity, permeability, and protein-binding ability, are influenced by pKa values, which have an impact on pharmacokinetic processes such as absorption, distribution, metabolism, and excretion. According to a study on Ru(II) coordination complexes as antiproliferative agents, the pKa values of the aquated species control the reactivity of the active complex and allow the drug to be active in specific cells.

**Binding to Nucleic Acids**

DNA is a key therapeutic target for transition metal complexes and has a wide range of intracellular interactions. Nucleobases follow Chargaff’s rules whereby a 1:1 (purine: pyrimidine) relationship exists between guanine (G) and cytosine (C), and adenine (A) and thymine (T) (Fig. 3). The B-form of duplex DNA, which is the most common in biology, is a right-handed double helix with two antiparallel sugar–phosphate chains. The minor (4.8 Å) and major (10.5 Å) grooves in the helical structure are the result of the angle between the glycosidic bonds and...
hydrogen bonds. The dehydrated form of B-DNA, termed A-DNA, has a similar but more stable and compact structure, with 11 base pairs per helical turn and a 2.55 Å axial rise between base pairs. Z-DNA (linked to cancer, autoimmune and neurological diseases) is produced in vivo during the transcription process as a result of torsional strains generated by RNA polymerase moving along the sequence of the DNA double helix. This provides a range of possible intermolecular interactions, including irreversible covalent binding, reversible groove interactions, or intercalation between metal complexes and DNA, due to the structural complexity and polymorphism of DNA. Studies on the effect of DNA metal complexes have shown that metal coordination geometry and the ligand configuration can influence binding behavior: square planar metal complexes were observed to have a greater capacity for deeper insertion as an intercalator than complexes with octahedral or tetrahedral geometry.

In contrast to intercalation, different DNA adducts can form as a result of covalent bonding, which is the most common type of interaction for metal-based anticancer drugs (Fig. 4). These adducts range from monoadducts (one bond is formed with DNA and the other coordination site of the ligand remains equated or protein-bound), to intrastrand crosslinks (1,2-intrastrand or 1,3-intrastrand where two bonds are formed on the same strand between consecutive base pairs, or bonds are formed with base pairs that are one base apart, respectively) and interstrand crosslinks (bonds are formed on opposite strands of the double helix) between the metal complexes and DNA bases. These interactions can lead to various changes to the DNA structure such as bending of the DNA double helix and unwinding of the DNA, leading to cell death. Under certain conditions, binding with DNA can inhibit winding or ensure that no bending occurs, thus affecting replication and transcription. Studies have also shown that different metal ions can
interact with the DNA in different ways. The alkali and alkaline earth metals mainly interact with the DNA phosphate groups, whereas the metal ions, such as Cd\(^{2+}\), Pb\(^{2+}\), and trivalent lanthanides can interact with both phosphates and bases.\(^{35}\)

**Metal-Ion-Mediated Base Pairs**

Studies on the binding of metals to nucleosides have extended to metal-ion-mediated base pair formation. The incorporation of metal complexes into DNA via metal-mediated base pairing has been established as having the potential to expand the genetic alphabet (incorporating nonstandard bases into DNA) and to afford new DNA structures (noncanonical), and functionalities in which the metal properties can be shared with the DNA structure.\(^{36}\) The use of metal-ion base pairs can lead to the design of noncomplementary base pairs of natural pyrimidine bases such as thymine–thymine (T–T) and cytosine–cytosine (C–C) pairs. The most common are metal-ion-bridged T–Hg\(^{2+}\)–T and C–Ag\(^{+}\)–C. The applications of metal-mediated base pairing range from metal-ion sensors (detection of Hg\(^{2+}\)) that are capable of forming T–Hg–T base pairs, to redox sensors and biomolecule sensors for detecting single-nucleotide polymorphisms.\(^{37}\) It has been proposed that metal-ion base pairs could either activate or inhibit the activity of enzymes such as DNA polymerases, endonucleases, ligases, and exonucleases. This could lead to the development of a DNA aptamer capable of inhibiting DNA polymerase.\(^{38}\) The metal ions can also be incorporated in the design of DNA molecular switches as observed on divalent metal ions such as Mg\(^{2+}\), Ca\(^{2+}\), and Mn\(^{2+}\), which are able to stabilize DNA duplexes and form stable complexes (hairpin, triplex, G-quadruplex, and i-motif) with ethylenediamine-stabilized DNA phosphate groups, whereas the metal ions, such as Cd\(^{2+}\), Pb\(^{2+}\), and trivalent lanthanides can interact with both phosphates and bases.

**Drug Resistance**

It is well known that drug resistance (extrinsic and intrinsic) to anticancer and antimicrobial medications is high. Although metal complexes, such as platinum drugs, may be initially effective against cancer, cancer cells develop resistance over time due to factors such as more efficient DNA damage repair, inactivation of drug with glutathione and metallothionein, and drug efflux with various transport systems located in the cell membrane.\(^{40}\) Elevated levels of copper transporters ATP7A and ATP7B have been linked to a decreased accumulation of cisplatin in cells and, as a result, to lower cytotoxicity.\(^{41}\) There are proteins which consist of an abundance of cysteine, such as glutathione and metallothionein, which inactivate cisplatin and other similar drugs within the cell by coordinating to thiol groups. The World Health Organization reports that with regard to microbes, we are now on the verge of a postantibiotic era, and that antibiotic treatment failures will be common over the next few decades.\(^{42}\) Cationic silver (Ag\(^{+}\)) resistance has been recognized for several years, and it has been observed that exposing bacteria to toxic heavy metals such as silver can induce the emergence of antibiotic resistance via the process of co-selection.\(^{43}\) As a result, novel strategies and formulations for anticancer and antimicrobial metal complexes with nonclassical modes of action are being investigated.

**Metal-Catalyzed Cleavage of Nucleic Acids**

**Hydrolysis**

Nucleic acids can undergo hydrolytic (enzymatically reversible), oxidative, and photocatalytic cleavage as a result of interactions with metal complexes. The Lewis acidity of the metal has been shown to determine the hydrolysis rate of the metal-bound phosphoester after the formation of the metal–phosphate intermediate, as demonstrated by the biological activity of natural metallonucleases.\(^{15}\) Since the neighboring 2′-OH group can be deprotonated by metal complexes in the RNA backbone, in comparison to DNA, RNAs are more hydrolysis-prone by nucleophilic attack. The phosphoester bond (mono-, di-, or tri-ester) can be found in proteins, nucleic acids, and lipids. This bond forms the backbone of DNA and RNA by connecting the adjacent nucleotides. In nature, the hydrolysis of this bond is required during DNA repair, posttranslational modification of proteins, and energy metabolism.\(^{44}\) Metal complexes, unlike enzymatic activity, can be used as cost-effective therapeutics with tunable functionality. The catalysis of the phosphoester bond (P–O) has highlighted the fact that metal ions can act as Lewis acids by activating the phosphoryl group for nucleophilic attack (Fig. 5). The pKa of the leaving group through the coordination of the alcoholic oxygen of the phosphodiester can also be reduced and metal-coordinated water molecules can be deprotonated.\(^{45}\) Using the metal complex based on copper and cobalt, the rate of DNA cleavage in comparison with the non-meta-catalyzed reaction was demonstrated to increase by a factor of 10 to 100 million.\(^{46}\) Several artificial nucleases that promote hydrolysis of the nucleic acid phosphate backbone have been synthesized, including Cu(II), Cr(III), Zn(II), Ce(IV), Zr(IV), La(III), Fe(III), and Co(III) complexes.\(^{35}\)

**Oxidation**

Another mechanism for nucleic acid degradation is irreversible oxidative cleavage, which requires redox-active metals. In this process, reactive oxygen species (ROS) are required for the hydrogen abstraction of the deoxyribose/ribose ring, which is followed by spontaneous cleavage of C–C and C–O bonds.\(^{47}\) The discovery of [Cu(phen)\(_2\)]\(^{2+}\) (phen = 1,10-phenanthroline) provided the first synthetic chemical nuclease for the development of new artificial metallonucleases.\(^{48}\) Colibactin, a human intestinal bacterial genotoxin (colorectal cancer) metabolite, has been shown to cause DNA damage (double strand breakage) and genomic instability following the formation of a copper complex that releases ROS such as superoxide and singlet oxygen species.\(^{49}\) Similarly, Cu(II) complexes with either phenanthroline or hydrazine ligands have been identified as self-activating metallonucleases where DNA can be damaged or cleaved by ROS generation in the absence of a reducing agent.\(^{50}\)

Copper derivatives generally interact noncovalently with DNA through intercalation, electrostatic interaction, and/or major or minor groove-binding; and in this they differ from
platinum drugs, such as cisplatin, which interact covalently with DNA nucleobases.\textsuperscript{51} Several naturally occurring antimicrobial peptides (AMPs) require metal ions such as Zn\textsuperscript{2+}, Na\textsuperscript{+}, and Mn\textsuperscript{2+} for antimicrobial activity. Upon formation of the AMP–metal complex, through redox chemistry they are able to inhibit DNA and RNA replication, retard protein synthesis, permeabilize the cell membrane, as well as disrupt proton and ion transmembrane gradients, and suppress cell-wall biosynthesis.\textsuperscript{52} The nucleobase guanine has been observed to be the most sensitive toward oxidation due to hydroxyl radicals (Fenton reaction), singlet dioxygen (Diels–Alder cycloaddition), or electron transfer as depicted in \textbf{Fig. 6}. The hydroxyl radicals attack the carbon atom C8 of guanine (and also the ribose sugar at C5′ > C4′ > C3′ > C2′ > C1′), while the singlet dioxygen undergoes a Diels–Alder cycloaddition with guanine across the imidazole ring; during electron-transfer pathways this leads to abstraction of one electron from guanine to produce the guanine radical cation.\textsuperscript{53} 

Metal complexes with redox properties are also capable of oxidizing the sugar unit via the production of radical species. The shape of the DNA duplex and the C–H bond-energy differences between the distinct carbon atoms play a key role in influencing the accessibility of the Csp3–H bonds of the deoxyribose within the major and minor grooves. The C1′ position is buried within the minor groove of B-DNA, thus making it inaccessible to some free radicals. The 2′-position of deoxyribose, in comparison, has high C–H bond strength and low accessibility of the corresponding hydrogen atoms, which means that the position can be changed only by γ-irradiation of DNA. Modification of the C3′ position, which is located in the major groove of DNA, makes a minor contribution to DNA strand break in B-DNA. The C4′ is situated on the outer edge of the minor groove of the DNA. The C4′–H4′ bond is relatively weak, which makes the 4′-position of deoxyribose a major target for hydroxyl radicals and other minor groove-binding oxidants, accessible to metal complexes such as iron bleomycin, [Cu(phen)]\textsuperscript{2+} and Fe(II)-EDTA. Both hydrogen atoms at the 5′-position of deoxyribose are highly accessible in the minor groove of B-DNA. Computational studies have indicated that H4′ abstraction can disrupt the deoxyribose moiety, while H5′ abstraction can lead to DNA cleavage at the 5′-position.\textsuperscript{54}
Photocatalysis

In photocatalysis, the inorganic or organic chromophores, once irradiated, induce an energy or electron transfer to the substrate. Both respond to ultraviolet light, leading to photocatalysis, photocaging, and photoswitching. Photocatalysis using metal complexes has been well studied in the context of modification of amino acids, peptides, and proteins. In terms of nucleic acid cleavage, visible light-based photosensitizers (PSs), relying on metal complexes such as ruthenium(II) and iridium(III) with riboflavin as a ligand, have resulted in the design of more efficient and site-selective DNA photocleavage induced by the generation of ROS. In photodynamic therapy, the drug TOOKAD Soluble, which utilizes a palladium-based PS, has been approved for use in treating low-risk prostate cancer, while the ruthenium-based PS TLD-1433 is currently undergoing clinical trials.

There are different mechanisms of photocatalytic action of metal complexes which can lead to cell death. During irradiation, which can lead to the formation of toxic singlet oxygen species, local heat production and dissociation (oxidation state remains unchanged) are responsible for radicals binding to biomolecules. Reduction of the metal leads to biomolecule interaction and bidentate ligand cleavage, resulting in ligand dissociation and subsequent binding to a biomolecule.

Approved drugs include Xcytrin (Gd(III)), Antrin (Lu(III)), TOOKAD (Pd(II)), Photrex (Sn(IV)), and Photosens (Al(III)).

Anticancer Activity

Cancer is defined as a malignant disease caused by unusual cell growth due to carcinomas (epithelial cell-derived), sarcomas (connective tissue), germ cell seminoma and dysgerminoma (testes and ovary), blastomas (embryonic tissue) and lymphoma, and leukemia (hematopoietic [blood-forming] cells). Global statistics indicate that lung cancer had the highest mortality rate (18.4%), followed by breast cancer (6.6%), colon cancer (5.8%), stomach cancer (8.2%), esophagus (5.3%), pancreas (4.5%), liver cancer (8.2%), and prostate cancer (3.8%). DNA remains the main target for metal-based anticancer drugs. Over the years, DNA metallointercalation has been extensively explored because this process can be used in potential anticancer drugs. Several metal complexes, ranging from ruthenium, osmium, iron, and titanium to copper, have been found to exhibit both anticancer and antibacterial properties.

However, to date only platinum-based drugs have been used extensively to treat cancer. Cisplatin is the first-generation platinum-based drug, which subsequently led to the development of carboplatin and oxaliplatin, both of which have been approved worldwide. Later generations of platinum drugs such as lobaplatin, nedaplatin, and heptaplatin are used in China, Japan, and South Korea, respectively. The mechanism of action of these drugs is based on binding to DNA in cancer cells through the formation of adducts with the cellular DNA, which lead to distortions that cannot be recognized by DNA repair mechanisms. The cisplatin analogues of titanium, vanadium, and iron have been shown to react with DNA specifically in tumor cells. Platinum-based anticancer agents, which are clinically used to treat 50% of malignant cancers, have several disadvantages, such as low bioavailability, severe side effects, poor stability, and inherent or acquired resistance. Pt(IV) complexes can be designed as prodrugs with kinetic inertness and a low spin d6 octahedral geometry, making them more stable for oral administration, while axial ligand modification can improve pharmacological properties and reduce side effects and drug resistance.
Inactive platinum(IV) prodrug complexes, such as Satraplatin, need to undergo reductive elimination by endogenous reductants to release the active square-planar platinum(II) core pharmacophore with concomitant dissociation of the axial ligands (to improve lipophilicity and solubility). Under (pseudo)physiological conditions, anticancer transition metal complexes containing biologically active ligands can behave in a variety of ways, including no release of a bioactive ligand, release of a bioactive ligand and cytotoxic metal-containing species from the initial multi-targeted complex (prodrug), and release of a bioactive ligand from an inactive metal-containing species in which the initial complex represents a drug carrier. To overcome the intrinsic resistance as well as prevent the development of acquired resistance, combining the mechanisms of action appears to be beneficial for cancer therapy. Complexes with dual anticancer and antibacterial properties can also be beneficial for cancer therapy because cancer patients have a weakened ability to fight infections. The siderophore deferoxamine B was successfully derivatized to form mono- and bi-dentate complexes with ruthenium, thus generating the dual anticancer and antibacterial agents. Siderophores are essential in order for bacteria to scavenge iron(III) from their environment. The release of biologically active ligand from the complex has been shown to be essential for potent activity to occur, in which cleavable linkers triggered by phosphatase, esterase, or reductase enzymes can be used.

**Antimicrobial Activity**

It was estimated that in 2018, approximately 2.4 million people in Europe, North America, and Australia would die from drug-resistant microbes over the following 30 years, at a cost of up to 3.5 billion U.S. dollars annually, due to global increase in microbial resistance. Over the years, the study of metal complexes as potential antimicrobial agents has attracted significant interest. Complexes of antibiotics with metal ions can affect the geometry of the antibiotic, leading to improved biological activity due to the presence of the metal ion within the organic compound structure. The drug Arsphenamine (Salvarsan), which contains an arsenic ion, was developed in 1909 and observed to be an effective treatment for syphilis and African trypanosomiasis. Other Food and Drug Administration (FDA)-approved metal-based complexes, termed metalloantibiotics, for antimicrobial applications include silverdine (silver), auranozin (gold), ganite (gallium), and pylera (bismuth). DNA interaction via an intercalative binding mode for photo and oxidative cleavage of bacterial DNA through an ROS and OH radical mechanism was well demonstrated in Cu(II), Ni(II), and Co(III) complexes of 2-furymethylamine Schiff base ligands. Peptide antibiotics have also been studied, and when complexed to metal ions, they will gain a higher positive charge, allowing them to interact more strongly with polyanionic DNA and RNA molecules. Bleomycin, for example, is able to form stable complexes with redox metal ions, and generate free radicals to cleave the nucleic acid chain.

**Antibacterial Activity**

Resistance arises from the misuse and overuse of antibiotics. Currently, there has been an alarming increase in bacterial resistance to several antibiotics available. Historically, clay was used to prevent wound infection. The study of the clay from the Amazon region (kaolin and smectite) or volcanogenic hydrothermal clay (illite-smectite) has been shown to inhibit even drug-resistant bacteria. The clay consists of aqueous Fe(II) in synergy with Al(III) to generate ROS, which is able to inhibit bacteria though disruption of the cell membrane and DNA. Currently bismuth (bismuth...
tribromophenate) and silver-based (silver sulphadiazine) antimicrobials are two metal-based therapeutics in clinical use. Over 906 metal-containing compounds have been submitted to the Community for Open Antimicrobial Drug Discovery for evaluation, of which 246 compounds have been observed to have antibacterial and antifungal activity. From investigations of antibacterial activity of metal complexes, several factors have been defined as significant, including (1) the chelate effect of ligands; (2) the structure of the N-donor ligands; (3) the total charge of the complexes; (4) the presence and nature of the complex counter ions; and (5) the nuclearity of the metal complex.

**Antiviral Activity**
The use of metal-based therapeutics, including gold, bismuth, arsenic, antimony, and mercury-based compounds to combat diseases such as tuberculosis and syphilis, caused cytotoxic problems, and lead to discontinued use. Over the years, the world has witnessed the (re)emergence of infectious diseases of viral origin, such as those caused by the dengue virus, Zika virus, chikungunya viruses (CHIKV), human immunodeficiency virus (HIV), yellow fever, measles, influenza A, severe acute respiratory syndrome coronavirus (SARS-CoV), Middle-East respiratory syndrome coronavirus (MERS-CoV), SARS-CoV2/Covid-19, as well as Ebola. Various metal complexes have been observed to have activity against: (1) HIV (nickel, copper, magnesium, lanthanum, ruthenium, and vanadium); (2) herpes simplex virus (cobalt, silver, iridium, gold, palladium, and platinum); (3) influenza (magnesium and copper); (4) hepatitis C (copper); (5) Sindbis virus (cobalt); (6) SARS-CoV (gold and bismuth); (7) cytomegalovirus (platinum); (8) chikungunya virus (gold); (9) poliovirus (platinum); (10) Zaire virus (gold), and (11) West Nile virus (lanthanum, cerium, neodymium, and praseodymium). Zinc ions exhibit antiviral properties against influenza and herpes in that they suppress the viral life cycle; they also play a major role in innate and adaptive immune signaling pathways. Like the antimicrobial properties of metallothioneins (small, cysteine-rich proteins) which bind to either zinc or copper to induce antiviral activity, synthetic metal-based complexes could also be a tool against viral infections. Phototherapy can induce antiviral activity from metal complexes in addition to antiviral activity caused by oxidative stress. Chelating the metal ion(s) within the active site is another method of antiviral inhibition.

**Natural Products**
Natural-based compounds such as curcumin have been shown to exhibit various therapeutic properties such as anti-inflammatory, antimicrobial (antivirus, antibacterial, and antifungal), antioxidant, and anticancer properties. These compounds are also capable of forming metallocomplexes once they chelate various metal ions. Quinone and polyphenol metabolites are produced by microorganisms, plants, and some animals. These metabolites have been widely used in medicine to combat bacterial infections. These natural products have been shown to exhibit metal-binding activity which would be useful in the formation of metal complexes for therapeutic use. Several widely used anticancer therapeutics are originated from natural sources, examples being irinotecan, vincristine, etoposide, and paclitaxel from plants (Fig. 9); as well as actinomycin D and mitomycin C from bacteria and marine-derived bleomycin.

Fig. 9 Widely used anticancer natural products.
lipophilicity due to electron shift from donor atoms of the ligand to the positively charged metal ion.

**Sensing and Imaging**

Over the years, several fluorophores (e.g., hoechst and 4′-6-diamidino-2-phenylindole, known as DAPI) that target DNA have been used for sensing and imaging applications for genetic diseases, biological process, and tumorigenesis (theragnostic application). The metal complex Fe-EDTA (C₁₀H₁₂FeN₂O₈) is one of the widely used molecular probes which provides information on biological activities associated with nucleic acids, including protein–DNA/RNA interactions, structure of DNA/RNA, and footprinting of nucleic acids.¹⁵ The use of luminescent transition and lanthanide metal complexes has been shown to be an alternative, as these complexes are able to achieve long wavelengths of luminescence and longer emission lifetimes, which are ideal for real-life application due to deep tissue penetration in the tissue optical window. It has been reported that the complexes are able to bind to DNA via π–π interactions between polyaromatic systems on the metal complex and nucleobases, resulting in intercalation and insertion between base pairs of duplex DNA.⁸⁶

Binding into the grooves of DNA or direct coordination has also been observed. Phosphorescent transition metal complexes generally display large absorption–emission Stokes shifts, which can also be advantageous for potential biological applications. Through the use of strong σ-donor groups and multidentate ligands, the observed cleavage of the metal–ligand bonds can be avoided, leading to the reduction of the phosphorescence metal complex fluorophores. A variety of luminescent metal complexes, based on rhenium, ruthenium, osmium, iridium, rhodium, platinum, europium, and terbium, have been designed for DNA sensing and imaging application.⁸⁷ These compounds can be used in photodynamic therapy due to their light-responsive properties. In addition, the photoluminescence properties of metal–organic frameworks have also been applied in drug delivery and bioimaging. The presence of the conjugated π-electron system allows for binding single-stranded DNA molecules.⁸⁸

The approved magnetic resonance imaging (MRI) contrast agents include Gd(III)-based complexes such as Eovist, Ablavar, and Dotarem (→ Fig. 10); for Mn(II) they include Lumenance and Mangafodipir, and for Fe(III) they include Ferumoxsil and Feridex I.V.(iron oxide nanoparticles).⁶⁰

Radiometals are becoming more widely available and are routinely used in the development of radiotracers for diagnostic and therapeutic purposes. With more than half of the cancer patients requiring radiotherapy to target localized solid tumors, the combination of multiple variants of immunotherapy with different forms of radiotherapy has given rise to radioimmunotherapy. Unlike the conventional radiotherapy treatment technique, radioimmunotherapy helps to overcome organs exposure to direct or scattered irradiation. Nuclear properties of radiometals, ranging from γ-ray and β⁺ particle emission (imaging) to Auger electron and α-particle emission (treatment) in combination with long half-lives, are excellently suited with the comparatively long biological half-life of monoclonal antibodies in vivo.⁹⁰ In radiotherapeutics, cancer cells are tracked and eliminated using monoclonal antibodies/peptide that have been radiolabeled with a β-emitting radionuclide. The FDA has approved only one radioimmunoconjugate (RIC)-[90Y]-Ibritumomab tiuxetan (Zevalin) for the treatment of indolent CD20-positive B cell lymphoma.⁹⁰ While the Novartis’ peptide-based Lutetium-177 (177Lu) prostate-specific membrane antigen (177Lu-PSMA) was granted a breakthrough therapy designation for patients with metastatic castration-resistant prostate cancer (CRPC).⁹¹ The α-particle emitter ²²⁳RaCl₂ (Xofigo) was also successfully launched by Bayer for the treatment of osseous metastases in CRPC.⁹² A recent review of therapeutic radio-pharmaceutical concepts and concerns was published elsewhere.⁹³

![Fig. 10](image-url) Approved photodynamic therapy agents and widely used molecular probes.
Molecular Techniques for DNA Interaction Characterization

The electron-rich phosphate backbone, donor heteroatoms in the nucleobases, and intricate secondary and tertiary structures create potential binding environments for both “free” metal ions and discrete complexes. Metal complexes bind to DNA through various modes, for example (1) noncovalent intercalation (insertion of a planar, usually aromatic, ligand [or part thereof] between the stacked base pairs of DNAs); (2) insertion (incorporation of a ligand into the base pair stack); (3) groove binding; and (4) coordination by discrete phosphate, such as the phosphate clamp, with shape and charge of the complex determining the mode of interaction. The transition metal complexes are able to bind to DNA via these two interactions, which can be elucidated by using electronic absorption spectroscopy (fixed concentration of metal complexes with incremental addition of calf thymus DNA), fluorescence quenching (ethidium bromide [EB] competitive binding), cyclic voltammetry, X-ray crystallography, circular dichroism, or viscosity measurements (Table 1). Gel electrophoresis experiments can be conducted to characterize DNA cleavage by means of the interaction between plasmid DNA and the complex.

The melphalan protection assay (DNA/RNA ladders crucial for accurate sizing), which also requires an agarose gel, can aid in DNA alkylation (monoaalkylated covalent DNA adduct) inhibition studies where preferred drug–DNA binding into the minor groove is established. When using crystallography to study drug interactions with DNA or oligonucleotides, the main difficulty lies in growing a high-quality crystal of reasonable size. This requires a significant quantity of homogeneous and pure materials. To conduct DNA–metal complex interaction via X-ray crystallography, 3G synchrotron facilities/microsource X-ray diffractometer techniques are required. This method allows definitive visualization of metalloprotein–DNA binding interactions. In the past, X-ray crystallography has been an effective choice for atomic-level visualization of metalloproteins, despite some drawbacks such as the fact that X-ray-induced structural and electronic changes often occur at the site of greatest interest.

To avoid radiation damage, techniques including serial synchrotron crystallography and/or X-ray free-electron lasers (XFELs) can be used. Serial synchrotron and XFEL crystallography have been used in studying metalloproteins through the collection of metal redox and ligation state data at room temperature. Mass spectrometry data are able to give information on covalent metalloprotein–DNA binding interactions/binding stoichiometry/noncovalent binding

| Technique                | Advantage                                      | Limitations                                                   |
|-------------------------|-----------------------------------------------|---------------------------------------------------------------|
| X-ray crystallography   | Detailed structural data                       | Solid-state interactions only                                 |
| NMR                     | Modification of electronic properties of complex | Active nuclei and diamagnetic compounds                        |
| Mass spectrometry       | Very small quantities                          | Gas-phase results may not always translate to solution        |
| Viscosity               | • Ease of use  
• Low cost  
• Suitable for bulk analysis | Metal ions themselves can compact DNA (charge neutralization) |
| Circular dichroism      | • Excellent sensitivity and reproducibility  
• Provides structural information related to specific binding interactions  
• Small sample size | • Caution must be taken with chiral metallo drugs  
• Experimental conditions must carefully maintain DNA structure  
• Limited solvent choice (UV activity) |
| UV thermal melting      | • Simple experimental setup  
• High sensitivity and reproducibility  
• Suitable for comparing/ranking binding affinity (SAR)  
• Small sample size | • Metallo drugs stability/optical transparency at elevated temperatures  
• Nonphysiological temperatures (>37°C) |
| Fluorescence            | High-throughput analysis possible when combined with 96-well plates | Reliance on signal from a fluorogenic reporter                  |
| Agarose gel electrophoresis | • Simple experimental setup  
• Cost effective  
• Small sample size  
• Fast analysis time | Quantitation of DNA damage/modification                       |
| On-chip microfluidics   | Quantitative method with high sizing resolution | Expense, and scaling between DNA sizes requires change of microfluidic chip, and results in analytically challenging. |
| UV-Vis                  | Direct binding analysis                        | Solvent choice                                                 |

Table 1 Characterization methods for metalloprotein–DNA interactions (adapted from Kellett et al94)

Abbreviations: NMR, nuclear magnetic resonance; UV, ultraviolet; Vis, visible.
Heteronuclear Bimetallic Complexes

Heteronuclear bimetallic complexes are designed for their enhanced bioactivity when compared with that of monometallic species expected in combining two distinct metal centers within the same molecule. With the limitations of single metal complexes such as auranoitin and cisplatin, various other metal ions can be added to the coordination sphere to give a bimetallic complex with introduced properties. The combination of the classical transition metals such as Pt, Ru, and Au with other metal moieties may produce complexes with improved pharmacokinetic and pharmacodynamic properties. The multiple metal ions can influence the overall redox properties of the complexes, hydrolysis behavior, as well as interactions with biomolecules. In cancer or microbes that have developed a resistance mechanism that renders one of the metals redundant, the second or third metal may show some activity. Evidence suggested that a heterometallic titanocene–gold chemotherapeutic compound was successfully synthesized and characterized and found to be effective against renal cancer by inhibiting migration, invasion, and angiogenesis. Mechanistic studies of the gold–titanocene complex revealed that the anticancer activity of the complex is partly attributable to the interaction of gold ions with ubiquitin, suggesting that the titanocene moieties are cleaved off under biological conditions. The complex was observed to have no interaction with DNA.

Computational Studies

Quantum Mechanical Modeling

Various computational tools are available for the characterization of metal complexes and investigation of electronic properties and reactivity. Computational methods can be used to determine molecular structures, including bond length, bond angle parameters, molecular orbital analysis, quantum chemical electronegativity (χ), absolute hardness (η), chemical potentials (μ), absolute softness (ω), global softness (σ), global electrophilicity (S), and electronic charge (ΔNmax). These calculated molecular and electronic properties provide key insights into the rationalization of the observed reactivity. Density functional theory (DFT) is a useful tool in computational chemistry. Various software programs, such as Gaussian, Vienna Ab initio Simulation Package, Amsterdam Density Functional, and ORCA, have implemented this theory and are highly cited. The merit and accuracy of the metal complex data generated by DFT depends on the level of theory employed in the study.

In recent years, with the addition of novel functional groups (such as carbohydrates to the NHC ring), the application of NHC ligands in medicinal chemistry has been expanding. Carbohydrates have the potential to form metal complex due to their selective uptake into tumor cells, multihydroxy functionality and well-defined stereochemistry suitable for metal coordination. They also have properties of biocompatibility, chirality, water solubility, and low toxicity.
Increased DNA repair is the main mechanism of cisplatin resistance in A2780 ovarian cancer cell lines. According to DFT studies using relativistic reactivity descriptors, DNA repair is increased because cis-planaramineplatinum(II) complexes (e.g., cisplatin) primarily form bifunctional intrastrand Pt(GG) adducts.\textsuperscript{112}

**Molecular Docking**

Various DNA structures such as A-, B-, and Z-DNA from PDB files 1VJ4, 1BNA, and 2DCG can be used in docking studies. Several docking techniques have been found to be effective for investigating molecular docking of metal complexes. To conduct docking studies, the target DNA structure is first imported into the software, and then hydrogen atoms are added followed by energy optimization. For the metal–complex ligand, it is ensured that hydrogen atoms have been added to the compound, and a conformational search is conducted in the case of all the compounds to identify the best conformers; then the MMFF94 force field was used to minimize energy minimization. In some docking softwares, the DNA remains rigid, while the torsional bonds of the metal complexes are free to rotate. Studies on Co(II), Cu(II), Ni(II), and Zn(II) complexes of naphthoquinone have shown strong binding interaction between these compounds (Cu(II) > Zn (II) > Ni(II) > Co(II)) and DNA through an intercalative mode of binding stabilized by intermolecular hydrogen bonding.\textsuperscript{113}

More than 60 different docking tools and programs have been developed for both academic and commercial use (\textsuperscript{114}Table 2). Examples of the applications of DNA binding using these programs include: (1) AutoDock for copper (II), cobalt (II), Ni(II), Ru(II), and Zn(II) complexes\textsuperscript{114–117}; (2) AutoDock Vina for Zn(II) metal complex\textsuperscript{118}; (3) Molecular Operating Environment (MOE) software for Cu(II), Co(II), Ni (II), Fe(II), VO(II), Cr(III), Mn(II), Zn(II) complexes, Pd (II), and Ag (I)\textsuperscript{119–121}; and (4) GOLD for Cu(II), Co(II), Zn(II), and Cr (II).\textsuperscript{122} Docking can take two forms: flexible (flexible side chain) and rigid (rigid receptor). Rigid docking is an ideal choice when standard computational systems were used to screen databases (large numbers of ligands), while flexible docking is an ideal choice when high-end computational power is available, and pocket shape changes are required during docking.\textsuperscript{123} Based on their treatment of ligand flexibility, search algorithms range from systematic conformational searches, which allow ligands (metal complexes) of interest to rotate in all directions, which allows more docking interactions but is time consuming (e.g., Dock and FlexX), to stochastic algorithms, which randomly change the structure or the position of the allowed ligands (metal complex) of interest (e.g., GOLD and AutoDock) and simulation algorithms, which require high computational cost to make the potential protein–ligand complex structure ideal for molecular dynamics (MD) and energy minimization (e.g., CHARMM, Amber, and GROMACS).\textsuperscript{124}

**HEX software is widely used for in silico docking studies of metal complexes (Cu(II), Co(II), Ni(II), Zn(II), and Os(IV)), as it maintains the structural conformation during docking.\textsuperscript{125,126} Docking of metallodrugs has posed difficulties when dealing with the formation of coordination bonds between the metal and a donor of an amino acid side chain. Many docking programs (e.g., GOLD and AutoDock) have limitations, as the covalent docking approach is used. By including coordination scoring parameters in GOLD, Sciortino and colleagues were able to predict metal complex–protein interactions with good agreement with the experimental structures (root mean square deviation is < 1.0 Å).\textsuperscript{126} Studies of Pt(II)–p-phenylenediamine complex interacting with double-stranded and G-quadruplex DNA revealed that computational studies can correlate with in vitro studies, in which evidence of planar aromatic cationic metal complexes and double-stranded DNA was observed by intercalation.\textsuperscript{127}

**Molecular Dynamics**

Current docking approaches cannot fully account for ligand reorganizations and exchange reactions in the metal-centered coordination sphere, particularly in the case of bulky

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|}
\hline
Software & Search algorithm & License \\
\hline
AutoDock 4.2 & Lamarckian genetic algorithm & Academic \\
AutoDock Vina & Local optimization & Academic \\
DOCK & Shape matching & Academic \\
BIOVIA Discovery Studio & Shape matching & Commercial/academic \\
FlexX & Shape matching & Commercial \\
Glide-Schrodinger & Hybrid & Commercial \\
GOLD & Genetic algorithm & Commercial/academic \\
ICM & Hybrid & Commercial \\
LIGANDFIT & Shape matching & Commercial \\
MOE & Hybrid & Commercial \\
Surflex-Dock & Shape matching & Commercial \\
SWISSDOCK & Evolutionary optimization & Academic \\
\hline
\end{tabular}
\caption{List of available docking software (adapted from Ferreira et al\textsuperscript{128} and Prieto-Martínez et al\textsuperscript{129})}
\end{table}

Abbreviations: ICM, internal coordinate mechanics software; MOE, molecular operating environment.
Metal Complexes as DNA Synthesis and/or Repair Inhibitors

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Metallo-drugs. MD-based docking has been proposed as an alternative method, which considers the plasticity of metal-centered complexes during recognition and binding, while ligand reorganizations and exchange reactions will require the assessment of ad hoc methodological developments. MD simulations can predict how each atom in a system moves over time based on a general model of the physics governing interatomic interactions, and highlighting important dynamic molecular processes such as conformational transition, ligand binding, and protein folding. However, experimental techniques, such as X-ray crystallography, cryo-EM, NMR spectroscopy, and mass spectrometry, can only provide static snapshots of metal complex binding and interactions. The hydration shells of biomolecules may have an enormous impact on their structure and function. Mixed quantum mechanics/molecular mechanics (QM/MM) is the most commonly adopted approach to studying metal-containing biomolecules. Since QM computations are computationally expensive, a timescale of 100s picoseconds to 1 nanosecond is commonly used, which is insufficient for chemical reactions, conformational transitions, and drug binding/dissociation study. In 2019, Van Rixel et al used the QM/MM technique to describe aromatic Pt-compound intercalation-binding mode to a noncanonical site created by two DNA tetrads, as well as the drug-induced deformation. The studies on Au(I)-compounds binding to telomeric DNA G-quadruplex were experimentally confirmed using ESI-MS and X-ray diffraction, while QM/MM simulations confirmed its binding mode within a G-quadruplex. Apart from biomolecule interactions, QM/MM was also successfully used to investigate the emission spectrum of platinum(II) bipyridine complex, which was due to the excited state of metal–metal to ligand charge transfer triplet (³MMLCT). Numerous molecular simulation packages, such as CHARMM, AMBER, NAMD, OpenMM, Gaussian, Q-Chem, Turbomole, Psi4, ORCA, and Gromacs, have QM/MM functionalities with built-in QM modules or interfaces.

Drug-Like Properties

In addition to carrying out docking studies, it is also possible to evaluate the drug-like properties of metal complexes such as absorption, distribution, metabolism, elimination, and toxicity (ADMET). These properties are crucial, since when a drug is administered via oral ingestion, it must cross membranes (influenced by lipophilicity [hydrophobicity]) of numerous cells to reach its site of action (absorption); it is distributed to its site of action through the circulatory systems (distribution); it is metabolized by a variety of enzymes into other products called metabolites (metabolism); it is excreted via the bowel in the feces, and kidneys through the urine or in the biliary excretion, and can lead to risk of toxicity if accumulated in the body (excretion). According to Lipinski’s rule, the molecule with Log p < 5 should have drug-like characteristics, and thus have a higher tendency to penetrate the biological membrane. The total polar surface area parameter helps us predict the way of drug transport (less than or equal to 140 Å) inside the various parts of the body, including gastrointestinal tract, blood-brain barrier, and cell membrane, as well as oral bioavailability. Active compounds are also expected to have a molecular weight of less than 500 g/mol, no more than 5 hydrogen bond donors, no more than 10 hydrogen bond acceptor sites, and less than 10 rotatable bonds for flexibility. However, compounds such as antibiotics have been shown to violate some of the rules, but still have good pharmacokinetic and pharmacodynamic properties. Jeyaraman et al. have used SWISS ADME to determine the ADMET properties of Cu(II), Ni(II), Co(II), and Zn(II) complexes.

Nanotechnology

The use of nanomaterials as enzyme mimics capable of degradation of nucleic acids has received increasing attention. The “nanozymes” mimic the redox activity of enzymes, and are capable of cleaving both RNA and DNA. Similar to endonuclease, cadmium telluride quantum dots, cerium oxide nanoparticles, and single-layered graphene oxide have been shown to be able to cleave DNA though hydrolytic, oxidative, or photocatalytic mechanisms. Through the use of metal-amphiphilic molecules, they have become a source of metal ions and serve as protective agents (surfactant) for the synthesis of stable metallic/metallic oxide nanostructure fabrication, thus providing an advantage over the conventional nanoparticle synthesis procedures. Over the years, specific interactions between nanoparticles and DNA or protein as well as other biomolecules have been observed. Silver clusters templated by DNA (Ag-DNA) highlight the potential nanoparticle synthesis procedures. DNA duplex is used as a nanoscale rigid molecular arm in DNA nanotechnology to spatially locate addressable transition metals, which are normally coordinated by organic molecular pockets and form vertices. Metallic nanostructures with electrical, magnetic, and optical properties can be synthesized in a well-controlled, flexible, and highly addressable motif for diagnostics and drug delivery systems. Because of the chemical properties of DNA, it has been possible to regulate the nucleation and growth of nanocolloids in which metals or metal ions bind to specific sites on the DNA backbone, resulting in the development of silver, platinum, and gold nanowires and nanoparticles. Biofunctionalization of metal nanoparticles using aptamer-appended DNA tetrahedron nanostructures has also been shown to be useful in medicine, as it allows selective targeting of the cancer cells and acts as a MRI contrast agent. The interaction of thymine and mercury makes it possible to develop ultrasensitive detection methods to detect mercury at subnanomolar concentrations, due to the formation of metallo-DNA duplexes (dT-Hg-dT)n. The use of nanoparticles is very suitable for mediating endocytosis to promote intracellular accumulation of drugs in cells, resulting in targeted delivery to reduce side effects and improve bioavailability of poorly soluble drugs. By adding Chlorin e6 (PS) to the surface of mesoporous silica nanoparticles and encapsulating cisplatin, nanotechnology was observed to provide
simultaneous chemotherapy and phototherapy in cancer eradication that can overcome cisplatin resistance.\textsuperscript{150}

**Toxicity**

Clinical use of the platinum (Pt) metal-based cisplatin drug has been observed to cause numerous side effects (nephrotoxicity, ototoxicity, neurotoxicity, and inborn or acquired drug resistance), which remains a challenge to overcome in the preparation of efficient anticancer drugs.\textsuperscript{151} Due to the lack of tumor selectivity of cisplatin, intravenous administration of the drug leads to interaction with human serum albumin in the presence of cysteine residue. Therefore, severe nausea, vomiting, loss of hearing, and kidney damage often occur when cisplatin is used for chemotherapy. Long-term effects may be different, requiring dialysis, kidney transplant, or medication cocktails to support other systems in the body.\textsuperscript{152} This means that a lower dose should be administered to avoid side effects; however, this leads to cancers rapidly developing resistance due to suboptimum therapeutic levels in comparison with cisplatin and the related drugs carboplatin, in which bidentate dicarboxylate ligands replace the more labile chlorides of cisplatin, and oxaliplatin, in which the stable 1,2-diaminocyclohexane ligand leads to more tolerable compounds.\textsuperscript{153} This highlights the evidence that ligand design can help ensure the biocompatible metal complex for medicinal application. Novel strategies for rational metallotherapeutic design have been proposed to help overcome these issues. According to the analogy principle, the essential metal ion can be used as a reference for the biological behavior of a nonessential metal ion in relation to similarity in charge, ionic radius, coordination number, coordination stability, redox potential, and so on.\textsuperscript{154} As a result, if a nonessential metal ion mimics or promotes intrinsic metal signaling, it will almost certainly be tolerated. This can help in selection of metals that will have minor or no metal toxicity.

**Conclusion**

It is clear that the unique properties of metal ions for the design of new drugs can be beneficial in medical inorganic chemistry. The role of transition metal complexes as therapeutic compounds has grown in importance in the field of inorganic chemistry, particularly in DNA interaction. In this review, metal-based therapeutics have made advances in the treatment of cancer and infectious diseases. In the case of metallotherapeutics, the metal ions are identified as enhancing the biological activity of known drugs and improving their mechanism of action by acting as metal-binding sites. With an expanding number of computer programs being used in computational chemistry and computer-aided drug discovery, these tools can now be used for \textit{in silico} metallotherapeutic studies. Although metallotherapeutics are difficult to develop, their unique properties provide valuable opportunity for discovering novel drugs with special mechanisms of action. This would assist in overcoming complex medical obstacles encountered in the world.

**Funding**

This work was supported by the South African National Research Foundation (Grant No. 129549) and the University of South Africa.

**Conflicts of Interest**

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

**References**

1. Rosenberg B. Chapter 2 - Cisplatin: its history and possible mechanisms of action. In: Prestayko AW, Crooke ST, Carter SK, Carter SK, eds. Cisplatin: Current Status and New Developments. New York, NY: Academic Press: 1980:9–20
2. Ndagi U, Mlhongo N, Soliman ME. Metal complexes in cancer therapy - an update from drug design perspective. Drug Des Devel Ther 2017;11:599–616
3. Jory J, Cobalamin, microbially and epigenetics. In: Patel V, Preedy V, eds. Handbook of Nutrition, Diet, and Epigenetics. Cham: Springer; 2017:1707–1725
4. Karaca Ö, Meier-Menches SM, Casini A, Kühn FE. On the binding modes of metal NHC complexes with DNA secondary structures: implications for therapy and imaging. Chem Commun (Camb) 2017;53(59):8249–8260
5. Shakeri A, Panahi Y, Johnston TP, Sahebkar A. Biological properties of metal complexes of curcumin. Biofactors 2019;45(03):304–317
6. Peter S, Aderibigbe BA. Ferrocene-based compounds with antimalaria/anticancer activity. Molecules 2019;24(19):1–27
7. Eyase FL, Akala HM, Johnson JD, Walsh DS. Inhibitory activity of ferroquine, versus chloroquine, against western Kenya Plasmodium falciparum field isolates determined by a SYBR Green I in vitro assay. Am J Trop Med Hyg 2011;85(06):984–988
8. Kondratksy A, Kondratska K, Vanden Abeelee F, et al. Ferroquine, the next generation antimalarial drug, has antitumor activity. Sci Rep 2017;7(01):1–15
9. Doddi A, Peters M, Tamm M. N-heterocyclic carbene adducts of main group elements and their use as ligands in transition metal chemistry. Chem Rev 2019;119(12):6994–7112
10. Falivene L, Cavallo L. Theoretical NMR spectroscopy of N-heterocyclic carbenes and their metal complexes. Coord Chem Rev 2017;344:101–114
11. Kumar S. Recent advances in the Schiff bases and N-heterocyclic carbenes as ligands in the cross-coupling reactions: a comprehensive review. J Heterocycl Chem 2019;56(04):1168–1230
12. Jia P, Ouyang R, Cao P, et al. Recent advances and future development of metal complexes as anticancer agents. J Coord Chem 2017;70(13):2175–2201
13. Kaczmarek MT, Zabiszak M, Nowak M, Jastrzab R. Lanthanides: Schiff base complexes, applications in cancer diagnosis, therapy, and antibacterial activity. Coord Chem Rev 2018;370:42–54
14. Cotruvo JA Jr, Featherston ER, Mattocks JA, Ho JV, Laremore TN. Lanmodulin: a highly selective lanthanide-binding protein from a lanthanide-utilizing bacterium. J Am Chem Soc 2018;140(44):15056–15061
15. Yu Z, Cowan JA. Metal complexes promoting catalytic cleavage of nucleic acids-biochemical tools and therapeutics. Curr Opin Chem Biol 2018;43:37–42
16. Yusoh NA, Ahmad H, Gill MR. Combining PARP inhibition with platinum, ruthenium or gold complexes for cancer therapy. ChemMedChem 2020;15(22):2121–2135
17. Crittenden CM, Novelli ET, Meaffey MR, et al. Structural evaluation of protein/metal complexes via native electrospray
ultraviolet photodissociation mass spectrometry. J Am Soc Mass Spectrom 2020;31(05):1140–1150
18 Cao Q, Li Y, Freisinger E, Qin PZ, Sigel RK, Mao ZW. G-quadruplex DNA targeted metal complexes acting as potential anticancer drugs. Inorg Chem Front 2017;4(01):10–32
19 Zhang P, Sadler PJ. Redox-active metal complexes for anticancer therapy. Eur J Inorg Chem 2017;2017(12):1541–1548
20 Ni L, Zhao H, Tao L, et al. Synthesis, in vitro cytotoxicity, and structure-activity relationships (SAR) of multidentate oxidoanadium(iv) complexes as anticancer agents. Dalton Trans 2018; 47(30):10035–10045
21 Schmidlehner M, Flocke LS, Roller A, et al. Cytotoxicity and preliminary mode of action studies of novel 2-aryl-4-thiopyr- one-based organometallics. Dalton Trans 2016;45(02):724–733
22 Arshad J, Hanif M, Movassaghi S, et al. Anticancer Ru II complexes. Chemistry 2019; 25(61):4868–4871
23 Riedel CA, Flocke LS, Hejli M, et al. Introducing the 4-phenyl-1,2,3-triazole moiety as a versatile scaffold for the development of cytotoxic ruthenium(II) and osmium(II) arene cyclometalates. Inorg Chem 2017;56(01):528–541
24 Meier-Menches SM, Gerner C, Berger W, Hartinger CG, Keppler BK. Structure-activity relationships for ruthenium and osmium anticancer agents – towards clinical development. Chem Soc Rev 2018;47(03):909–928
25 Zaki M, Hairat S, Aazam ES. Scope of organometallic compounds based on transition metal-arene systems as anticancer agents: starting from the classical paradigm to targeting multiple strategies. RSC Advances 2019;9(06):3239–3278
26 Feng Y, Sun WZ, Wang XS, Zhou QX. Selective photoinactivation of methicillin-resistant Staphylococcus aureus by highly positively charged Ru III complexes. Chemistry 2019;25(61):13879–13884
27 Hachey AC, Havrylyuk D, Glazer EC. Biological activities of polypyridyl-type ligands: implications for bioinorganic chemistry and light-activated metal complexes. Curr Opin Chem Biol 2021;61:191–202
28 Pal M, Nandi U, Mukherjee D. Detailed account on activation mechanisms of ruthenium coordination complexes and their role as antineoplastic agents. Eur J Med Chem 2018; 150:419–445
29 Kominami H, Kobayashi K, Yamada H. Molecular-scale visualization and surface charge density measurement of Z-DNA in aqueous solution. Sci Rep 2019;9(01):1–7
30 Kulkarni M, Mukherjee A. Understanding B-DNA to A-DNA transition in the right-handed DNA helix: perspective from a local to global transition. Prog Biophys Mol Biol 2017;128:63–73
31 Chakraborty D, Wales DJ. Probing helical transitions in a DNA duplex. Phys Chem Chem Phys 2016;19(01):878–892
32 Zavarykina TM, Atkarskaya MV, Zhizhina GP. The structural and functional properties of Z-DNA. Biophysics (Oxf) 2019;64(05):445–461
33 Pages BJ, Ang DL, Wright EP, Aldrich-Wright JR. Metal complex interactions with DNA. Dalton Trans 2015;44(08):3505–3526
34 Ghosh S. Cisplatin: the first metal based anticancer drug. Bioorg Chem 2019;88:102925
35 Zhou W, Saran R, Liu J. Metal sensing by DNA. Chem Rev 2017;117(12):8272–8325
36 Mandal S, Müller J. Metal-mediated DNA assembly with ligand-based nucleosides. Curr Opin Chem Biol 2017;37:71–79
37 Scharf P, Müller J. Nucleic acids with metal-mediated base pairs and their applications. ChemPlusChem 2013;78(01):20–34
38 Park KS, Lee CY, Park HG. Metal ion triggers for reversible switching of DNA polymerase. Chem Commun (Camb) 2016; 52(27):4868–4871
39 Wan L, Lam SL, Lee HK, Guo P. Rational design of a reversible MgII-EDTA-controlled molecular switch based on a DNA mini-dumbbell. Chem Commun (Camb) 2020;56(70):10127–10130
40 Makovec T. Cisplatin and beyond: molecular mechanisms of action and drug resistance development in cancer chemotherapy. Radiol Oncol 2019;53(02):148–158
41 Ferreira CR, Gahl WA. Disorders of metal metabolism. Transl Sci Rare Dis 2017;2(3–4):101–139
42 Li LG, Xia Y, Zhang T. Co-occurrence of antibiotic and metal resistance genes revealed in complete genome collection. ISME J 2017;11(03):651–662
43 Hu Q, Jayasinghe-Arachchige VM, Zuchniarz J, Prabhapar R. Effects of the metal ion on the mechanism of phosphodiester hydrolysis catalyzed by metal-cyclen complexes. Front Chem 2019;7:195
44 Diez-Castellnou M, Martinez A, Mancin F. Chapter Four - Phosphate ester hydrolysis: the path from mechanistic investigation to the realization of artificial enzymes. In: Williams IH, Williams NH, eds. Advances in Physical Organic Chemistry. New York, NY: Academic Press; 2017:129–186
45 Kettenmann SD, Louka FR, Marine E, et al. Efficient artificial nucleases for mediated DNA cleavage based on tuning the steric effect in the pyridyl derivatives of tripod tetraamine-cobalt(II) complexes. Eur J Inorg Chem 2018;2018(20–21):2322–2338
46 Jastrzęb R, Nowak M, Skrobalska M, et al. DNA as a target for lanthanide(III) complexes influence. Coord Chem Rev 2019; 382:145–159
47 Łüdtke C, Sobottka S, Heinrich J, et al. Forty years after the discovery of its nucleolytic activity: [Cu(phen)2] II shows unattended DNA cleavage activity upon fluorination. Chemistry 2021;27(10):3273–3277
48 Wernke KM, Xue M, Tirala A, Kim CS, Crawford JM, Herzon SB. Structure and bioactivity of colibactin. Bioclin Med Chem Lett 2020;30(15):1–11
49 Heinrich J, Stubbe J, Kulak N. Cu(II) complexes with hydrazonemetalized phenantridines as self-activating metalonucleases. Inorg Chim Acta 2018;481:79–86
50 McGilver TJP, Afsarpoor S, Marmion CJ. Copper complexes as artificial DNA metalonucleases: from Sigman’s reagent to next generation anti-cancer agent? Inorg Chim Acta 2018;472:12–39
51 Alexander JL, Thompson Z, Cowan JA. Antimicrobial metallopeptides. ACS Chem Biol 2018;13(04):844–853
52 Brissos RF, Caubet A, Gamez P. Possible DNA-interacting pathways for metal-based compounds exemplified with copper coordination compounds. Eur J Inorg Chem 2015;2015(16):2633–2645
53 Biswas PK, Chakraborty S. Targeted DNA oxidation and trajectory of radical DNA using DFT based QM/MM dynamics. Nucleic Acids Res 2019;47(06):2757–2765
54 Kuhlmann A, Herrmann S, Weinberger M, Penner A, Wagenknecht HA. Photocatalysis with nucleic acids and peptides. Phys Sci Rev 2018;3(11):1–12
55 King TA, Mandrup Kandemir J, Walsh SJ. Spring DR. Photocatalytic methods for amino acid modification. Chem Soc Rev 2021;50(01):39–57
56 Koo B, Yao H, Choi HJ, Kim M, Kim C, Kim KT. Visible light photochemical reactions for nucleic acid-based technologies. Molecules 2021;26(03):1–24
57 McFarland SA, Mandel A, Dumoulin-White R, Gasser G. Metal-based photosensitizers for photodynamic therapy: the future of multimodal oncology? Curr Opin Chem Biol 2020;56:23–27
58 Reesing F, Szymanski W. Beyond photodynamic therapy: light-activated cancer chemotherapy. Curr Med Chem 2017;24(42):4905–4950
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Miranda VM. Medicinal inorganic chemistry: an updated review on the status of metallotherapeutics and prominent metalloprotein candidates. Rev Inorg Chem 2021. Doi: 10.1515/reivc-2020-0030

Nandanwar SK, Kim HJ. Anticancer and antibacterial activity of transition metal complexes. ChemSelect 2019;4(05):1706–1721

Bray F, Ferlay J, Soerjomataram I, Siegel R, Torre L, Jemal A. Erratum: global cancer statistics 2018: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J Clin 2020;70(04):313–313

Bergamo A, Dyson PJ, Sava G. The mechanism of tumour cell death by metal-based anticancer drugs is not only a matter of DNA interactions. Coord Chem Rev 2018;360:17–33

Sodhi RK, Paul S. Metal complexes in medicine: an overview and update from drug design perspective. Cancer Ther Oncol Int J 2019;14(01):25–32

Li X, Liu Y, Tian H. Current developments in Pt(IV) prodrugs conjugated with bioactive ligands. Bioinorg Chem Appl 2018;2018:8276139

Wong DYQ Probing the platinum(IV) prodrug hypothesis. Are platinum(IV) complexes really prodrugs of cisplatin? In: Wong DYQ, ed. Rethinking Platinum Anticancer Drug Design: Towards Targeted and Immuno-chemotherapeutic Approaches. Singapore: Springer; 2018:55–71

Starpa P, Trávníček Z. Non-platinum complexes containing releasable biologically active ligands. Coord Chem Rev 2019;395:130–145

Laurent Q, Batchelor LK, Dyson PJ. Applying a Trojan horse strategy to ruthenium complexes in the pursuit of novel anti-bacterial agents. Organonmetallics 2018;37(06):915–923

Hofer U. The cost of antimicrobial resistance. Nat Rev Microbiol 2019;17(01):1

Ramotowska S, Wysocka M, Brzeski J, Chylewska A, Makowski M. A comprehensive approach to the analysis of antibiotic-metal complexes. Trac-Trend Anal Chem 2020;123:1–9

Swinne DC. Phenotypic drug discovery: history, evolution, future. In: Beverley Isherwood, Augustin A, eds. Phenotypic Drug Discovery. London: Royal Society of Chemistry; 2020:1–19

Frei A. Metal complexes, an untapped source of antibiotic potential? Antibiotics (Basel) 2020;9(02):1–24

Venkateswarlu K, Kumar MP, Rambabu A, et al. Crystal structure, DNA binding, cleavage, antioxidant and antibacterial studies of CuII, Ni(II) and Co(III) complexes with 2-[(furan-2-yl)methyl-(imino)methyl]-6-ethoxyphenol Schiff base. J Mol Struct 2018;1160:198–207

Jeżowska-Bojczuk M, Stokowa-Soltyś K. Peptides having antimicrobial activity and their complexes with transition metal ions. Eur J Med Chem 2018;143:997–1009

Londono SC, Hartnett HE, Williams LB. Antibacterial activity of aluminium in clay from the Colombian Amazon. Environ Sci Technol 2017;51(04):2401–2408

Frei A, Zuege J, Elliott AG, et al. Metal complexes as a promising source for new antibiotics. Chem Sci (Camb) 2020;11(10):2627–2639

Kumaravel G, Ponna Utthra P, Ramam N. Exploiting the biochemical efficacy of benzimidazole based Schiff base complexes with L-Histidine as a co-ligand: combined molecular docking, DNA interaction, antimicrobial and cytotoxic studies. Bioorg Chem 2018;77:269–279

Cirri D, Pratesi A, Marzo T, Messori L. Metallo therapeutics for COVID-19. Exploiting metal-based compounds for the discovery of new antiviral drugs. Expert Opin Drug Discov 2021;16(01):39–46

de Paiva REF, Marçal Neto A, Santos IA, Jardim ACG, Corbi PP, Bergamini FRG. What is holding back the development of antiviral metalloproteins? A literature overview and implications for SARS-CoV-2 therapeutics and future viral outbreaks. Dalton Trans 2020;49(45):16004–16033

Read SA, Obeid S, Ahlenstiel C, Ahlenstiel G. The role of zinc in antiviral immunity. Adv Nutr 2019;10(04):696–710

Wiehe A, O’Brien JM, Senge MO. Trends and targets in antiviral phototherapy. Photochem Photobiol Sci 2019;18(11):2565–2612

Carcelli M, Fisicharo E, Compari C, et al. Metal-chelating properties and antiviral activity of some 2-hydroxyphenyl amides. Polyhedron 2017;129:97–104

Dandawate P, Padhye S, Schober R, Biersack B. Discovery of natural products with metal-binding properties as promising antibacterial agents. Expert Opin Drug Discov 2019;14(06):563–576

Huang M, Lu JJ, Ding J. Natural products in cancer therapy: past, present and future. Nat Prod Bioprospect 2021;11(01):5–13

De A, Ray HP, Jain P, Kaur H, Singh N. Synthesis, characterization, molecular docking and DNA cleavage study of transition metal complexes of o-vanillin and glycine derived Schiff base ligand. J Mol Struct 2020;1199:126801

Liu HK, Sadler PJ. Metal complexes as DNA intercalators. Acc Chem Res 2011;44(05):349–359

Berrones Reyes J, Kuimova MK, Vilar R. Metal complexes as optical probes for DNA sensing and imaging. Curr Opin Chem Biol 2021;61:179–190

Dong J, Zhao D, Lu Y, Sun YY. Photoluminescent metal–organic frameworks and their application for sensing biomolecules. J Mater Chem B Mater Energy Sustain 2019;7(40):27244–27267

Boros E, Holland JP. Chemical aspects of metal ion chelation in the synthesis and application antibody-based radiotheracers. J Labelled Comp Radiopharm 2018;61(09):652–671

Puttemans J, Lahouette T, D’Hayvetter M, Devogest N. Beyond the barrier: targeted radionuclide therapy in brain tumors and metastases. Pharmaceuticals 2019;11(08):1–23

Sun M, Niazi MO, Nelson A, Skafida M, Niaz MJ. Review of 177Lu-PSMA-617 in patients with metastatic castration-resistant prostate cancer. Cureus 2020;12(06):1–8

Müller C, Béthé M, Geistlich S, van der Meulen NP, Schibli R. Targeted radiotherapeutics from bench-to-bedside’. Chimia (Aarau) 2020;74(12):939–945

Sowa Dumond AR, Scott PJ. Concepts and issues for therapeutic radiopharmaceuticals. In: Scott P, Kilbourn M, eds. Handbook of Radiopharmaceuticals. Hoboken, NJ: Wiley; 2020:23–42

Kellett A, Molphy Z, Slater C, McKee V, Farrell NP. Molecular methods for assessment of non-covalent metallodrug-DNA interactions. Chem Soc Rev 2018;48(04):971–988

Hough MA, Owen RL. Serial synchrotron and XFEL crystallography for studies of metalloprotein catalysis. Curr Opin Struct Biol 2021;71:232–238

Ebrahim A, Moreno-Chicocho T, Appleby MV, et al. Dose-resolved serial synchrotron and XFEL structures of radiation-sensitive metalloproteins. IUCrJ 2019;6(Pt 4):534–551

Marchant A, Rosu F, Zenobi R, Gabelica V. Thermal denaturation of DNA G-quadruplexes and their complexes with ligands: thermodynamic analysis of the multiple states revealed by mass spectrometry. J Am Chem Soc 2018;140(39):12553–12565

Eyers CE, Vonderach M, Ferries S, Jeacock K, Eyers PA. Understanding protein-drug interactions using ion mobility-mass spectrometry. Curr Opin Chem Biol 2018;42:167–176

Timerbaev AR. Application of ICP-MS to the development of metal-based drugs and diagnostic agents: where do we stand? J Anal At Spectrom 2021;36(02):254–266

Arnesano S, NMR spectroscopy to study the fate of metallotherapeutics in cells. Curr Opin Chem Biol 2021;61:214–226

Lincoln P, Wilhelmsson LM, Nordén B. Chapter 3: Slow DNA binding. In: Waring MJ, ed. DNA-targeting Molecules as Therapeutic Agents. London: The Royal Society of Chemistry; 2018:45–73

Morgan SM, El-Sonbati AZ, Eissa HR. Geometrical structures, thermal properties and spectroscopic studies of Schiff base...
complexes: correlation between ionic radius of metal complexes and DNA binding. J Mol Liq 2017;240:752–776
103 Malik MA, Dar OA, Gull P, Wani MY, Hashmi AA. Heterocyclic Schiff base transition metal complexes in antimicrobial and anticancer chemotherapy. MedChemComm 2017;9(03):409–436
104 Raoofmoghaddam S, Zhou Y-P, Wang Y, Driess M. N-heterocyclic silylenes as powerful steering ligands in catalysis. J Organomet Chem 2017;829:2–10
105 Zhao W, Ferro V, Baker MV. Carbohydrate–N-heterocyclic carbene metal complexes: synthesis, catalysis and biological studies. Coord Chem Rev 2017;339:1–16
106 Afzal M, Al-Loheedan HA, Usman M, Tabassum S. Carbohydrate-based heteronuclear complexes as topoisomerase Ix inhibitor: approach toward anticancer chemotherapeutics. J Biomol Struct Dyn 2019;37(06):1494–1510
107 Lopes J, Alves D, Morais TS, et al. New copper(I) and heteronuclear copper(I)-ruthenium(II) complexes: synthesis, structural characterization and cytotoxicity. J Inorg Biochem 2017;169:68–78
108 van Niekerk A, Chellan P, Mapolie HF. Heterometallic multinuclear complexes as anti-cancer agents-an overview of recent developments. Eur J Inorg Chem 2019;2019(30):3432–3455
109 Elie BT, Fernández-Gallardo J, Curado N, Cornejo MA, Ramos JW, Contet M. Bimetallic titanocene-gold phosphane complexes inhibit invasion, metastasis, and angiogenesis-associated signaling molecules in renal cancer. Eur J Med Chem 2019;161:310–322
110 Kiwaam HA, El-Mowafy AS, El-Bindary AA. Synthesis, spectral characterization, DNA binding, catalytic and in vitro cytotoxicity of some metal complexes. J Mol Liq 2021;326:115381
111 Qi SC, Hayashi JJ, Zhang L. Recent applications of metal complexes based on density functional theory. RSC Advances 2016;6(81):77375–77395
112 Alvarado-Soto L, Ramirez-Tagle R. Relativistic structure-activity relationship of cisplatin (II) complexes. J Struct Chem 2020;61(05):688–693
113 Koshiba A, Parthiban C, Ciattini S, Chelazzi L, Elango KP. Metal complexes of naphthoquinone based ligand: synthesis, characterization, protein binding, DNA binding/cleavage and cytotoxicity studies. J Biomol Struct Dyn 2018;36(16):4170–4181
114 Anupama B. DNA binding interactions, docking and antioxidative studies of ternary copper (II) complexes. J Mol Struct 2020;1216:127988
115 Kondori T, Shahrazi O, Akbarzadeh-T N, Aramesh-Boroujeni Z. Two novel bipyridine-based cobalt (II) complexes: synthesis, characterization, molecular docking, DNA-binding and biological evaluation. J Mol Struct Dyn 2021;39(02):595–609
116 Neelakantan MA, Balaramunug K, Balakrishnan C, Subha L. Interaction of acidic amino acid Schiff base metal complexes with DNA/BSA protein and antibacterial activity: spectral studies, DFT calculations and molecular docking simulations. Appl Organomet Chem 2018;32(04):e4259
117 Zhao J, Li S, Wang X, Xu G, Gou S. Dinuclear organoruthenium complexes exhibiting antiproliferative activity through DNA damage and a reactive-oxygen-species-mediated endoplasmic reticulum stress pathway. Inorg Chem 2019;58(03):2208–2217
118 Mishra DK, Singh UK, Das A, et al. DNA binding, amelioration of oxidative stress, and molecular docking study of Zn(II) metal complex of a new Schiff base ligand. J Coord Chem 2018;71(14):2165–2182
119 Sharfaldin AA, Ewmas AH, Jaremko M, Hussien MA. Transition metal complexes of 6-mercaptopurine: characterization, theoretical calculation, DNA-binding, molecular docking, and anticancer activity. Appl Organomet Chem 2021;35(01):e6041
120 Abdel-Rahman LH, Adam MSS, Abu-Dief AM, et al. Synthesis, theoretical investigations, biocidal screening, DNA binding, in vitro cytotoxicity and molecular docking of novel Cu (II), Pd (II) and Ag (I) complexes of chlorobenzylidene Schiff base: promising antibiotic and anticancer agents. Appl Organomet Chem 2018;32(12):e4527
121 Abdel-Rahman LH, Abu-Dief AM, Aboelezz MO, Hassan Abdel-Mawgoud AA. DNA interaction, antimicrobial, anticancer activities and molecular docking study of some new VO(II), Cr(III), Mn(II) and Ni(II) mononuclear chelates encompassing quinolinate imine ligand. J Photochem Photobiol B 2017;170:271–285
122 Kavitha B, Srawanthi M, Saritha Reddy P. DNA interaction, docking, molecular modelling and biological studies of α-Vanillin derived Schiff base metal complexes. J Mol Struct 2019;1185:153–167
123 Rawal K, Khurana T, Sharma H, et al. An extensive survey of molecular docking tools and their applications using text mining and deep curation strategies. PeerJ 2019;7:1–177
124 Wang Q. Protein-ligand Docking Application and Comparison Using Discovery Studio and AutoDock. Fargo, ND: North Dakota State University; 2017
125 Pursuwani BH, Varma RR, Patel MN. Synthesis, characterization and biological applications of Osmium(IV) complexes. J Pure Appl Sci 2018;26:69–77
126 Scirotino G, Rodríguez-Guerra Pedregal J, Ledós A, Garribba E, Maréchal JD. Prediction of the interaction of metallic moieties with proteins: an update for protein-ligand docking techniques. J Comput Chem 2018;39(01):42–51
127 Ang DL, Kelso C, Beck JL, Raph SF, Harmen DG, Aldrich-Wright JR. A study of Pt(II)-phenanthroline complex interactions with double-stranded and G-quadruplex DNA by ESI-MS, circular dichroism, and computational docking. J Biol Inorg Chem 2020;25(03):429–440
128 Ferreira LG, Dos Santos RN, Oliva G, Andricopulo AD. Molecular docking and structure-based drug design strategies. Molecules 2015;20(07):13834–14321
129 Prieto-Martínez FD, Arciniega M, Medina-Franco JL. Molecular docking: current advances and challenges. TIP Rev Esp Cienc Quim Biol 2018;21:65–87
130 Riccardi L, Genna V, De Vivo M. Metal–ligand interactions in drug design. Nat Rev Chem 2018;2(07):100–112
131 Hollingsworth SA, Dror RO. Molecular dynamics simulation for all. Neuron 2018;99(06):1129–1143
132 Lage D, Elaesser T, Hynes JT. Perspective: structure and ultrafast dynamics of biomolecular hydration shells. Struct Dyn 2017;4(04):044018
133 Janoš P, Spinello A, Magistrato A. All-atom simulations to studying metallo drugs/target interactions. Curr Opin Chem Biol 2021;61:1–8
134 van Rixel VHS, Busemann A, Wissingh MF, et al. Induction of a four-way junction structure in the DNA palindromic hexanucleotide 5′-d(CGTACG)-3′ by a mononuclear platinum complex. Angew Chem Int Ed Engl 2019;58(28):9378–9382
135 Guarra F, Marzo T, Ferraroni M, et al. Interaction of a gold(i) dicarbene anticancer drug with human telomeric DNA G-quadruplex: solution and computationally aided X-ray diffraction analysis. Dalton Trans 2018;47(45):16132–16138
136 Nakagaki M, Aono S, Kato M, Sakaki S. Delocalization of the π-system of some of the bases transition metal complexes in antimicrobial and anticancer activities. Dalton Trans 2018;47(45):16132–16138
137 Grover A, ed. Drug Design: Principles and Applications. Singapore: Springer; 2017:113–133
139 Anthony EJ, Bolitho EM, Bridgewater HE, et al. Metallodrugs are unique: opportunities and challenges of discovery and development. Chem Sci (Camb) 2020;11(48):12888–12917
140 Chagas CM, Moss S, Alisaraie L. Drug metabolites and their effects on the development of adverse reactions: revisiting Lipinski’s Rule of Five. Int J Pharm 2018;549(1–2):133–149
141 Jeyaraman P, Alagarraj A, Natarajan R. In silico and in vitro studies of transition metal complexes derived from curcumin-isoniazid Schiff base. J Biomol Struct Dyn 2020;38(02):488–499
142 Zhang J, Wu S, Ma L, Wu P, Liu J. Graphene oxide as a photocatalytic nuclease mimicking nanozyme for DNA cleavage. Nano Res 2020;13(02):455–460
143 Bhar R, Kaur G, Mehta SK. Experimental validation of DNA interactions with nanoparticles derived from metal coupled amphiphiles. J Biomol Struct Dyn 2018;36(14):3614–3622
144 Swasey SM. On the interactions of silver with DNA: from metal-mediated base pairings to fluorescent clusters. Los Angeles, CA: University of California; 2017:1–202
145 Hu Q, Li H, Wang L, Gu H, Fan C. DNA nanotechnology-enabled drug delivery systems. Chem Rev 2019;119(10):6459–6506
146 Jujin S, Nummelin S, Kostiainen MA, Linko V. DNA nanostructure-directed assembly of metal nanoparticle superlattices. J Nanopart Res 2018;20(05):1–11
147 Li N, Shang Y, Han Z, Wang T, Wang ZG, Ding B. Fabrication of metal nanostructures on DNA templates. ACS Appl Mater Interfaces 2019;11(15):13835–13852
148 Ghosh D, Datta LP, Govindaraju T. Molecular architectonics of DNA for functional nanoarchitectures. Beilstein J Nanotechnol 2020;11:124–140
149 Pandeeswar M, Senanayak SP, Govindaraju T. Nanoarchitectonics of small molecule and DNA for ultrasensitive detection of mercury. ACS Appl Mater Interfaces 2016;8(44):30362–30371
150 Mirzaei S, Hushmandi K, Zabolian A, et al. Elucidating role of reactive oxygen species (ROS) in cisplatin chemotherapy: a focus on molecular pathways and possible therapeutic strategies. Molecules 2021;26(08):2382
151 Kathiresan S, Muges S, Annaraj J, Murugan M. Mixed-ligand copper(ii) Schiff base complexes: the vital role of co-ligands in DNA/protein interactions and cytotoxicity. New J Chem 2017;41 (03):1267–1283
152 Kumar M, Kumar G, Kant A, Masram DT. Role of metallodrugs in medicinal inorganic chemistry. In: Ul-Islam S, Hashmi AA, Khan SA, eds. Advances in Metallodrugs: Preparation and Applications in Medicinal Chemistry. Salem, MA: Scrivener Publishing LLC; 2020:71–113
153 Lazarević T, Rilak A, Bugarčić ŽD. Platinum, palladium, gold and ruthenium complexes as anticancer agents: current clinical uses, cytotoxicity studies and future perspectives. Eur J Med Chem 2017;142:8–31
154 Yang X. Regulating cellular stress responses: an emerging strategy for rational metallodrug design. Future Med Chem 2018;10 (06):611–614