8-Hydroxyquinolines: a review of their metal chelating properties and medicinal applications

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Abstract: Metal ions play an important role in biological processes and in metal homeostasis. Metal imbalance is the leading cause for many neurodegenerative diseases such as Alzheimer’s disease, Parkinson’s disease, and multiple sclerosis. 8-Hydroxyquinoline (8HQ) is a small planar molecule with a lipophilic effect and a metal chelating ability. As a result, 8HQ and its derivatives hold medicinal properties such as antineurodegenerative, anticancer, antioxidant, antimicrobial, anti-inflammatory, and antidiabetic activities. Herein, diverse bioactivities of 8HQ and newly synthesized 8HQ-based compounds are discussed together with their mechanisms of actions and structure–activity relationships.

Keywords: metal binding compound, antineurodegenerative, anticancer, antidiabetic, multifunctional actions, structure–activity relationships

Introduction

8-Hydroxyquinoline (8HQ) (Figure 1), a quinoline derivative originating in plants as well as from synthesis, has been used as a fungicide in agriculture and a preservative in the textile, wood, and paper industries.¹ 8HQ possesses potent coordinating ability and good metal recognition properties, which means it is widely used for analytical and separation purposes as well as for metal chelation.²

Metal ions play a very important role in biological processes, and metal homeostasis is required for the maintenance of metal balance.³,⁴ Many diseases arise from the loss of homeostasis including metal overload and deficiency, which are caused by abnormal metal metabolism or metal absorption. Of all the hydroxyquinoline derivatives, 8HQ is the most interesting one to be explored, owing to its multifunctional properties, such as diverse bioactivities and therapeutic potentials.⁵

8HQ is the only one, among seven isomeric monohydroxyquinolines, capable of forming complexes with divalent metal ions through chelation.⁶ Most bioactivities of 8HQ and its derivatives originate from their chelating ability. As previously mentioned, metal imbalance is the leading cause for many diseases, therefore, 8HQ is a potent chelator that may restore metal balance and be useful for the treatment of metal-related diseases. In this review, the bioactivities, mechanisms of actions of newly synthesized 8HQ-based compounds, and their structure–activity relationships (SAR) will be discussed.

Antineurodegenerative activity

Transition metals such as Fe, Zn and Cu are found in the brain at relatively high concentration and are required for the brain’s cellular processes including synaptic neuronal
activity and metalloenzyme function; for instance, Cu/Zn superoxide dismutase (SOD), cytochrome C oxidase, etc. 3 Metal homeostasis dysregulation is generally accepted as a key predisposing factor in many neurodegenerative diseases such as Alzheimer’s disease (AD), Parkinson’s disease, multiple sclerosis and others. 4,7 Increasing levels of redox-reactive metal ions, such as Cu and Fe, in specific brain regions can generate reactive oxygen species (ROS) that cause lipid peroxidation and toxic reactive aldehyde products. These finally lead to damage of cellular components. 7 The proteasome is a cellular system that degrades unwanted or abnormal proteins. 9 It is worth noting that metal ions can interact with proteins in the brain and induce their conformational change, leading to protein misfolding and rendering them resistant to proteasomes. Moreover, metal–protein interaction facilitates aggregation and accumulation of misfolded proteins in regions of the brain, 8 leading to neurotoxicity, neuronal dysfunction, and neuronal cell death. 7,9 It has been suggested that dysregulation of metal homeostasis and metal ion–protein interactions are involved in the pathogenesis of neurodegenerative diseases. 10–12 Therefore, metal chelation therapy has been proposed to be a promising approach in restoring metal balance and reducing neurotoxicity caused by metal–protein interaction. 13

An ideal metal chelator for neurodegenerative treatment had been suggested to be a low molecular weight (MW) and lipophilic (uncharged) compound capable of crossing the blood–brain barrier to reach target sites in the brain. 3 In addition, the selectivity of compounds in chelating certain metal ions but not affecting metalloenzymes would also be required for cellular functions. 3 It is necessary that the compound would be able to chelate metal ions in accumulated proteins 1 in order to reverse proteasome resistance, thereby allowing misfolded proteins to be degraded. 14 Moreover, the drug itself is required to have minimal toxicity and side effects. 3 However, merely controlling misfolded protein levels may not be sufficient to reverse the neurodegenerative progression in the brain. 14 Therefore, a new therapeutic strategy for preventing metal–protein interaction has been recently proposed. 14

A series of 8HQ derivatives, such as 5-chloro-7-iod o-8-hydroxyquinoline, or clioquinol (CQ), 5-((4-(prop-2-ynyl)piperazin-1-yl)methyl)quinolin-8-ol (HLA-20), 5-((methyl(prop-2-ynyl)amino)methyl)quinolin-8-ol (M30), and 5-((4-(2-hydroxyethyl)piperazin-1-yl)methyl)quinolin-8-ol (VK-28) (Figure 2), have been reported to exert potent antineurodegenerative effects. 15 Among these, CQ has reached pilot Phase II of clinical trials in AD patients. 16–20 CQ was originally used as an antimicrobial for amoebic dysentery (traveler’s diarrhea); however, after its neurotoxicity was reported among the Japanese in the late 1960s, this drug was withdrawn from oral usage. 21 The proposed mechanism of toxicity is that CQ decreases vitamin B12 bioavailability, which results in neurological symptoms. 22 However, the neurotoxicity can be reversed by vitamin supplementation and dosage control. 20

CQ is a potent chelator containing two electron donor sites located at the quinoline ring nitrogen atom and phenolate oxygen atom, which give rise to its chelating ability (Figure 3A). Moreover, halogen groups are known to increase its lipophilicity to allow its absorption to target sites in the brain. 9 CQ selectively chelates Cu and Zn ions, 9,23 which play
a vital role in misfolded protein production, aggregation, and accumulation that ultimately leads to neurotoxicity in AD.\textsuperscript{9,24} However, the selectivity of the compound can minimize the chance of developing a depletion in systemic metal ions.\textsuperscript{9}

Recently, a dual mechanism of CQ action based on metal–protein interaction has been proposed.\textsuperscript{14} Firstly, as Cu and Zn chelators, CQ can inhibit misfolded protein production and aggregation. Moreover, the chelation of accumulated Zn in misfolded proteins can reverse proteasome resistance and promote misfolded protein degradation.\textsuperscript{14} However, the affinity of the compound to chelate Zn is not enough to alter Zn metalloenzymes.\textsuperscript{14} Secondly, CQ can function as a metal chaperone in transporting metal ions into cells and promoting redistribution of ions that consequently activate cell signaling involved in neuroprotective cascades.\textsuperscript{25} Thus, the activity of CQ can be attributed to two aspects: the prevention of neurotoxicity initiated by metal–protein interaction and the redistribution of metal ions into cells to promote protective functions.\textsuperscript{14} It has been reported that CQ is a potent antineurodegenerative agent that can improve cognitive functions in AD patients;\textsuperscript{1} however, this compound was not further developed owing to manufacturing difficulties. This was due to the presence of a small amount of the carcinogenic contaminant 5,7-diiodo-8-hydroxyquinoline (Figure 3B) that forms during large-scale chemical synthesis.\textsuperscript{14} Thus, PBT2, a second generation CQ, was developed to solve the problem of CQ and to improve its solubility and its ability to cross the blood–brain barrier.\textsuperscript{14} It was observed that PBT2 could selectively chelate Cu and Zn and form neutral soluble complexes capable of passing through cellular membranes. Due to its moderate affinity to metal ions, after entering cells it can release metal ions from the complex. This leads to bioavailable delivery of Cu and Zn into cells.\textsuperscript{26–28} So far, PBT2 has shown improvement of cognitive function as noted for CQ in Phase IIa of clinical trials in AD patients.\textsuperscript{29,30} Results supported by in vitro and in vivo studies suggested that antineurodegenerative efficacies of both CQ and PBT2 are based on their chelating ability and metal ion delivery into cells.\textsuperscript{26–28} Moreover, metal binding affinity of both compounds is high enough to inhibit misfolded protein production and aggregation but not high enough to alter the actions of metalloenzymes.\textsuperscript{27,31}

The Fe ion is considered to be another redox-reactive metal ion that causes oxidative stress via the Fenton reaction. It is found to be elevated in many neurodegenerative diseases, such as AD,\textsuperscript{3} Parkinson's disease,\textsuperscript{32} and amyotrophic lateral sclerosis.\textsuperscript{33} In view of its multifunctional roles, 8HQ-based compounds have been utilized in the treatment and improvement of neurodegenerative patients. For example, M30 and HLA-20 (Figure 4) are novel multifunctional 8HQ-based drugs synthesized by combining an Fe chelating compound possessing an antioxidant activity (VK-28) with the Parkinson's drug (Ladostigil) containing the N-propargylamine moiety (Figure 5), which affords the neuroprotective property.\textsuperscript{14}

As outlined in Figure 6, VK-28 can chelate excessive Fe ions in the brain, thereby preventing the Fenton reaction that produces reactive hydroxyl radicals (OH). It is capable of directly scavenging OH, which gives rise to antioxidant effects.\textsuperscript{35,36} Ladostigil contains the propargylamine moiety that accounts for its inhibition of the monoamine oxidase enzyme.\textsuperscript{34} This enzyme is involved in dopamine oxidation, which generates hydrogen peroxide (H$_2$O$_2$) that initiates the Fenton reaction in the presence of Fe$^{2+}$, leading to oxidative stress in neurons.\textsuperscript{32} Compounds M30 and HLA-20 have moderate chelating affinity toward Fe, Cu, and, Zn, with the order Fe$^{3+}$ > Cu$^{2+}$ > Zn$^{2+}$, and they strongly inhibit mitochondrial membrane peroxidation in vitro with a half maximal inhibitory concentration (IC$_{50}$) in the micromolar range.\textsuperscript{15,37} In vitro studies indicated that M30 upregulates expression of Fe$^{2+}$/O$_2^-$ regulated hypoxia inducible factor, which is a hypoxia mimetic regulator, resulting in neuronal prosurvival and cytoprotective effects.\textsuperscript{38–40} In addition, M30 has been reported to exhibit neurorescue and neuroprotective activities in animal models.\textsuperscript{41}

Therefore, both HLA-20 and M30 are novel multifunctional drugs that exhibit promising antioxidant and neuroprotective effects as well as antidepressant activity. These bioactivities arise from the ability of the compounds to elevate levels of dopamine, serotonin, and norepinephrine in the brain through the inhibition of the monoamine oxidase enzyme.\textsuperscript{42}

**Anticancer activity**

It has been well recognized that redox-active metal ions do not only play important roles in normal cells but are also
essential in cancer cells. Some transition metal ions, such as Fe and Cu are considered as cancer risk factors.\textsuperscript{43–50}

In normal cells, Fe serves as a prosthetic group in many enzymes that are required for physiological processes, such as oxidase, catalase, and ribonucleotide reductase. In contrast, it generates ROS, leading to lipid peroxidation and damage to cellular components, such as lipids, proteins, and DNA.\textsuperscript{51,52}

Thus, Fe plays essential roles in cancer via the generation of ROS as well as serving as a nutrient for the growth of cancer cells.\textsuperscript{43}

Most Fe that exists in the human body is in the protein-bound form that cannot promote lipid peroxidation or ROS formation.\textsuperscript{51} In addition, free Fe per se is a poor catalyst for reactive oxygen metabolites, but Fe toxicity arises when it binds to a low-MW chelator. Therefore, the formed Fe-chelator complex causes the dissociation of H\textsubscript{2}O\textsubscript{2} into OH.\textsuperscript{53} The chelating ability of 8HQ has been proposed to account for its observed cytotoxic activity as afforded by the Fe-8HQ complex.\textsuperscript{54}

The formed Fe-8HQ lipophilic complex is capable of entering and being distributed within cells,\textsuperscript{55} causing massive breakage of DNA strands. In order to repair damaged DNA, large quantities of adenosine triphosphate are required, which consequently leads to cellular adenosine triphosphate depletion and finally cell death.\textsuperscript{56} As such, possible mechanisms of DNA damage were proposed. The Fe-8HQ complex may be formed at specific sites that break the phosphodiester backbone of DNA, acting as chemical nucleases, causing oxidative degradation at the deoxyribose moiety.\textsuperscript{57} In other words, the Fe-8HQ complex acts as a cytostatic drug.\textsuperscript{58}

Another possible mechanism is that the Fe-chelator complex induces membrane damage, that leads to loss of calcium homeostasis, which triggers endonuclease to cleave DNA in an apoptotic-like manner.\textsuperscript{55} Results from SAR studies demonstrated that 8HQ is a crucial scaffold for anticancer activity.\textsuperscript{59} This relationship is derived from the ability of the compound to form chelate complexes with metal ions, incorporated with essential enzymes for DNA synthesis,\textsuperscript{60} possibly, ribonucleotide reductase.\textsuperscript{61} Moreover, bis-type structure of 8HQ is required for potent anticancer activity.\textsuperscript{62}

In fact, S\textsubscript{1} [bis-N-(8HQ-5-ylmethyl)benzylamine] has been reported to form Fe complexes with higher affinity to exert higher antiproliferative effects as compared to o-trensox (ie, the reference drug). However, o-trensox is a very high affinity Fe chelator in hepatocyte cultures.\textsuperscript{60} The results indicated that S\textsubscript{1} is a promising starting point for anticancer drug development.\textsuperscript{60} In addition, metal complexes of mixed
ligands of 8HQ-uracils (Figure 7) have been reported to provide significant cytotoxicity against human cancer cells (ie, HepG2, A549, HuCCA-1, and MOLT-3).63

Recently, great interest in metal complex compounds has extensively increased due to their wide range of applications.64 The interaction of metal complexes with DNA has been studied for biotechnology and medical applications including their use as anticancer drugs.65 The metal complex binds reversibly to DNA via noncovalent interactions, such as electrostatic binding, groove binding, and intercalative binding.66,67 Intercalation between metal complexes and DNA bases is considered to be the most important binding mode giving rise to antitumor activity.68 This causes DNA conformational changes, which finally leads to DNA strand stress and breakage.69

The intercalating ability of metal complex compounds are dependent on the planarity of the ligands, the coordination geometry, types of ligand donor atoms, and metal ions.70 Sulfonamide-substituted 8HQ metal complexes have been reported to exhibit higher DNA binding affinity than that of free ligands.69 The highest binding efficiency among metal complexes that are formed using the same ligands was found to be that of Cu complexes.69 It was suggested that pharmacological activities of metal complexes are dependent on the nature of both the ligands and the metal ions.71 This notion was demonstrated for metal complexes synthesized from different types of metal ions using the same ligand; such metal complexes were found to exert different bioactivities.72,73

Cu ions are a risk factor predisposing to cancer, and they also serve as an essential cofactor for tumor angiogenesis, that is crucial for tumor growth and metastasis.44–47 High levels of Cu in tissue or serum has been found in many cancer patients including those with breast, prostate, colon, lung, and brain cancer.74–78 It was suggested that Cu could be used as one of the selective targets for cancer treatments.79

The anticancer effects of 8HQ derivatives, such as CQ, are related to Cu and Zn ions. As a Cu chelator, CQ exerts selective antiangiogenesis activity toward breast cancer81, prostate cancer,79 leukemia, and myeloma,82 with less effect on normal cells. In addition, the antitumor activity of CQ has been proposed to be tightly associated with proteasome inhibitory ability,79 which is elicited through ionophore actions.

Ionophore is a subset of metal-binding drugs that are capable of transferring multiple metal ions across biological membranes, either in or out of cells.83–85 Two properties are required for metal-binding compounds to act as ionophore,
which are described as follows. First, compounds should have low to moderate metal affinity, allowing them to bind metal ions in higher concentration areas and release them in lower concentration areas. Second, a suitable logarithmic measure of acid dissociation constant value (pKa) is necessary for compounds to be protonated upon entering cell compartments, which induces the release of metal ions from the complex. If the extracellular pH is higher than the pKa of the ionophore, the compound will form a complex with the metal ion. Once the metal complex passes into the cell, where the pH is lower than the pKa, the metal ions will be released. Both properties have been noted for CQ thereby allowing it to act as an ionophore with the capability to transport Cu and Zn ions into cells.

As a Cu ionophore, CQ has been reported to be able to deliver metal ions into cells, where it exerts its activity. CQ has been found to interact with Cu ions in tumor cells to form active Cu complexes which target the proteasome. Either Cu or Cu interact with electron donors, such as thiol and amino groups that are located outside the active site of proteasome thereby causing its conformational changes. These effects finally lead to proteasome inhibition and apoptosis of tumor cells. Some organic Cu complexes including CQ-Cu complexes have been reported to exhibit potent proteasome inhibitory effects on tumor cells but not on normal cells. Moreover, in vivo study of the effect on prostate cancer cells and xenografts by CQ was reported. The results showed that CQ alone can form an active metal complex with Cu of tumor cells, leading to androgen receptor repression, angiogenesis reduction, and apoptosis induction. Particularly, androgen receptor overexpression was found in all stages of prostate cancer, indicating that CQ may serve as an excellent antiandrogen receptor agent for prostate cancer treatment and prevention.

Besides acting as the Cu ionophore, CQ also provides anticancer activity as the Zn ionophore. The CQ-Zn complex is a known proteasome inhibitor; however, its growth inhibitory effect is weaker than that exerted by the CQ-Cu complex. In addition, CQ relays Zn ions to lysosomes and inhibits nuclear factor kappa-light-chain-enhancer of activated B cells (NF-kB) activity thereby leading to lysosomal disruption and cell apoptosis. The ionophore property of CQ was confirmed by adding metal ions, such as Cu and Zn, which can potentiate its cytotoxic activity instead of reversing the effects, as can be expected in the case of metal chelators. This indicated that CQ can transport metal ions into cells and exhibits cytotoxic activity. Furthermore, the anticancer activity of 8HQ derivatives on human cancer cells indicated that the effect is enhanced by Cu ions, a redox-reactive metal ion, as it leads to an elevation of ROS. However, such effects were not observed for Zn ions, which are non-redox reactive metal ions.

It has been further demonstrated that nitro containing 8HQ derivatives such as nitroxoline (8-hydroxy-5-nitroquinoline; NQ) exerted more potent anticancer activity, with a IC of 5–10-fold less than that of CQ (halogenated 8HQ derivative), and may be less neurotoxic. Unlike CQ, the antitumor effect of NQ is mainly exhibited via an increasing level of ROS in cells. The nitro moiety of NQ is a nitrogen radical source that initiates redox reactions, that consequently alters intracellular signaling thereby leading to antiangiogenesis and inhibition of tumor cell growth. These effects are enhanced by Cu but not Zn ions. This hypothesis was supported by studies that demonstrated that NQ acted as an antiangiogenic agent both in vitro and in vivo.

It is notable that Cu ions enhanced the cytotoxic activity of NQ. While Zn ions are known to enhance cytotoxic activity, the activity is found only in association with compounds containing an iodine moiety on the C-7 position of quinoline rings, such as CQ. However, the mechanism by which Zn enhances the cytotoxic activity has not been fully elucidated. The neurotoxicity of CQ has been reported to be involved with the Zn transporting activity. CQ that contains iodine at the C-7 position is capable of acting as a Zn ionophore, while NQ does not. Such an observation explains why NQ is less neurotoxic than CQ. Moreover, neurological diseases have not been reported in patients treated with NQ, suggesting that NQ is a novel compound with less neurotoxicity and should be further developed as an anticancer drug.

Recently, glucoconjugates of 8HQ derivatives were developed as anticancer prodrugs in order to improve the selectivity and to avoid chelation of systemic metal ions. It was reported that glucose avidity, increased glycolysis rate, and overexpression of glucose transporters were found in cancer cells. The study indicated that glucoconjugates could enhance drug delivery owing to the presence of the glucose moiety in drug molecules. The molecular structures of conjugated glucose can mask the chelating properties of compounds until they reach their target sites. Moreover, the presence of glucose in the drug structure promotes a more selective action by exploiting glucose transporters, which were found to be overexpressed in cancer cells. Therefore, glucoconjugated drugs are more selective to cancer cells and can cause less systemic side effects. After glucoconjugates are trapped in target cells, glucose moieties are hydrolyzed by specific β-glucosidases, which allows the compounds
to display chelation and exert antiproliferative effects.\textsuperscript{101} It has been demonstrated that 8HQ-glucoconjugates are novel compounds with potential for further development as selective anticancer treatment.\textsuperscript{99}

### Antimicrobial activity

Antimicrobial effects of 8HQ and its derivatives encompassing antibacterial,\textsuperscript{102-104} antimalarial,\textsuperscript{105-107} antiviral,\textsuperscript{108} antitubercular,\textsuperscript{109} and antidental plaque activities\textsuperscript{110,111} have been previously reported.

#### Antibacterial activity

**Antitubercular activity**

Nonreplicating *Mycobacterium tuberculosis* (TB) or latent TB is more tolerant to most antituberculosis drugs than the replicating type of TB and requires a more prolonged treatment.\textsuperscript{109} In fact, more than 200 8HQ-like compounds were identified to have an inhibitory effect against replicating TB.\textsuperscript{112} Results showed that 8HQ itself exerted the most potent activity among other compounds in its class.\textsuperscript{113} 8HQ can kill both replicating and nonreplicating TB in vitro, with a more potent effect noted for the nonreplicating type.\textsuperscript{109} Moreover, toxicity toward mammalian cells was not observed within the tested range of concentrations (0.1–10 μM), suggesting its safety in humans.\textsuperscript{109} Insight into its mechanism of action was not fully elucidated; however, the bidentate chelating property of 8HQ was probably not the primary mechanism. This was supported by the finding that the minimum inhibitory concentration (MIC) value was not changed by the addition of metal ions such as Fe, Cu, Mn, Zn, or Ni.\textsuperscript{109} Therefore, it was suggested that 8HQ interacts with several molecular targets to evade TB resistance.\textsuperscript{109}

Most of the clinically used antituberculosis drugs are known to be more active against replicating TB and are able to kill both types of TB. The more potent actions were exhibited against the nonreplicating type as was observed in 8HQ, which has not been reported in the literature.\textsuperscript{109} Thus, 8HQ could be a promising compound for the improvement of TB treatment.\textsuperscript{109}

The antitubercular activity of CQ-metal complexes were also reported.\textsuperscript{113} In particular, a series of mixed ligand metal complexes using CQ and 1,10-phenanthroline as ligands to coordinate with transition metal ions were synthesized and tested for antitubercular and antifungal activities.\textsuperscript{113} It was found that the Mn(II) complex was active against TB (MTCC200) with comparable MIC to the standard drug rifampicin with MIC values of 45 µg/mL and 40 µg/mL, respectively,\textsuperscript{113} whereas the Co(II) complex showed more potent activity with MIC 6.4-fold less than that of rifampicin.\textsuperscript{113} This study showed that free ligands and metal complexes exerted higher antitubercular activity than that of metal salts.\textsuperscript{113}

#### Inhibitory effect on *Escherichia coli*

A previous study reported the antimicrobial effects of 8HQ and its derivatives including 2-hydroxyquinoline (2HQ), 4-hydroxyquinoline (4HQ), and 6-hydroxyquinoline (6HQ) as well as 2-methyl-8HQ against human intestinal bacteria (*Bifidobacterium longum*, *Clostridium difficile*, *Clostridium perfringens*, *E. coli*, *Lactobacillus acidophilus*, and *Lactobacillus casei*).\textsuperscript{103} The results from the paper disc agar diffusion method demonstrated that only 8HQ could exhibit anti-intestinal bacterial activity. Strong inhibition was also observed for *E. coli* and *C. difficile* at a concentration of 0.5 mg/disc and for *C. perfringens* at a concentration of 0.1 mg/disc. The SAR study indicated that different positions of the hydroxyl group but not the methyl group on the quinoline ring gave rise to growth inhibitory activity against intestinal bacteria. Particularly, compounds with an OH group at the C-8 position displayed effective *E. coli*, *C. difficile*, and *C. perfringens* inhibitions.\textsuperscript{103} On the other hand, 2HQ, 4HQ, and 6HQ showed no growth inhibition against all of the tested intestinal bacteria.

#### Inhibitory effect on *Staphylococcus aureus*

Aqueous formulation of 8HQ (0.5% 8HQ), or commercially available as AQ\textsuperscript{+}, was reported to strongly
inhibit the growth of *S. aureus* including methicillin-resistant *S. aureus* (MRSA), methicillin-susceptible *S. aureus*, and vancomycin-intermediate *S. aureus* and displayed median MIC of 0.25%, which is equal to an active ingredient concentration of 12.5 μg/mL at optimum pH of 9.2.\(^1\) Lowering of the pH value caused a reduction in its efficacy: pH 7.5 yielded 4-fold reduction, and pH 5.5 resulted in 8-fold reduction.\(^1\) Interestingly, MRSA and vancomycin-intermediate *S. aureus* were equally susceptible to AQ\(^+\) as was observed for methicillin-susceptible *S. aureus*. It was suggested that the susceptibility to AQ\(^+\) was not influenced by antibiotic resistance determinants of the microbe.\(^1\) Data from electron microscopy indicated that AQ\(^+\) actively disrupts bacterial cell walls thereby leading to cell lysis.\(^1\) A time-killing study showed that AQ\(^+\) killed 99.9% of all bacterial cells from tested isolates within 6 hours. The time-killing curve of AQ\(^+\) was similar to that of gentamicin. Moreover, at higher concentrations of AQ\(^+\), a more rapid killing effect was observed.\(^1\) MRSA is carried in the anterior nares, and it should be noted that mupirocin-containing nasal ointment is currently being used to prevent transmission; however, antibiotic resistance had been reported to increase.\(^1\) Owing to the lipophilicity of 8HQ and its potency against various *S. aureus* strains as well as its rapid killing nature, it has been suggested that this compound could be used as topical hand cleansing agent to prevent MRSA transmission.\(^1\)

The efficacy of 8HQ in inhibiting *S. aureus* is dependent on its chelating ability and is enhanced in the presence of Cu. In addition, 8HQ derivatives were found to inhibit *S. aureus* strains.\(^1\) Quantitative structure-activity relationships (QSAR) study performed on 24 substituted 8HQ derivatives showed that ten three-dimensional descriptors such as molecular refractivity (MR), partition coefficient (logP), total energy (E), standard Gibbs free energy (G), lowest unoccupied molecular orbital (LUMO), highest occupied molecular orbital (HOMO), total molecular energy (ToE), Wien index (WInd), Balaban index (Blnd), and octanol-water ClogP were significant for the development of highly predictive \((R^2 = 0.988)\) model.\(^1\) Moreover, potent activity was found in ester derivatives rather than styril derivatives.\(^1\)

**Antidental plaque activity**

Antidental plaque activity of 8HQ derivatives has been reported.\(^1\) Dental plaque is a combination of oral microorganisms colonized on oral surfaces in which a microbial consortium or oral biofilm is formed.\(^1\) *Mutans streptococci* and *Porphyromonas gingivalis* are the most important among such oral microorganisms since they are pathogens of dental caries and periodontal diseases, respectively.\(^1\) At equilibrium, oral biofilms are beneficial for the prevention of exogenous and potentially pathogenic species colonization. However, unfavorable disruption of dynamic balance between host and microbial community at local sites eventually leads to an overgrowth of virulent or pathogenic species causing diseases.\(^1\) *Mutans streptococci* comprises two species, *Streptococcus mutans* and *S. sobrinus*, in which *S. mutans* is highly prevalent in dental plaque and is considered as an etiological pathogen for dental caries.\(^1\) An in vitro study of antidental plaque activity of three 8HQ derivatives, namely 8HQ sulfate, 5-chloro-7-iodo-8HQ (or CQ), and 5,7-dichloro-8HQ, against *S. mutans*, *Streptococcus sanguis*, *Actinomyces viscosus*, and *Actinomyces naeslundii* was reported. The result showed that all 8HQ derivatives differentially inhibited each of the tested organisms.\(^1\) These compounds were prepared in percentage concentrations in polyethylene glycol owing to their sparingly water-soluble nature.\(^1\) CQ and 5,7-dichloro-8HQ exerted a bactericidal effect at 0.05% concentration on *S. mutans* (cariogenic) and *A. viscosus* (periodontogenic), whereas 8HQ sulfate showed a bacteriostatic effect against both pathogens at a higher (0.3%) concentration.\(^1\) It was demonstrated that halogenated 8HQs were more potent dental plaque inhibitors.\(^1\) The low water solubility is a limitation in using these bioactive compounds via an aqueous vehicle such as mouth rinse.\(^1\) However, such compounds could be used via a polyethylene glycol vehicle as an ointment or additive ingredient in dentifrice to control dental diseases.\(^1\)

It is widely known that the antibacterial activity of 8HQ is closely related with its chelating ability, therefore, Fe or Cu chelation is required for the activity.\(^1\) Previous SAR studies indicated that substitution near the nitrogen atom or the phenolic group could alter the chelating ability, which led to a reduction of the antimicrobial activity.\(^1\) This study demonstrated that the hydrophobic logP parameter alone is not adequate for accurate computational prediction. Thus, electronic parameters such as pKa and steric parameters including MW and molecular refractivity are required for antidental plaque prediction.\(^1\) The QSAR study showed that two main factors are involved in antidental plaque activity of 8HQ against *S. mutans*.\(^1\) The results revealed that compounds with increased lipophilicity and electron-withdrawing substituents at the 5-position led to improvements in the activity, whereas bulky substituent groups afforded a decrease in the activity.\(^1\) According to these findings, the most potent *S. mutans* inhibitor should contain small C-5 substituents with lipophilicity and electron-withdrawing properties.\(^1\)
Notably, CQ contains a 5-chloro group with lipophilicity and an electron-withdrawing nature, making this compound a good *S. mutans* inhibitor.\(^\text{110}\)

### Antimicrobial activity of metal complexes and novel compounds

#### Metal-8HQ complexes

The antimicrobial activity of divalent metal-8HQ complexes and their mechanisms of action have been proposed.\(^\text{120}\) It was assumed that 8HQ uses its high lipophilicity to penetrate bacterial cell membranes in order to reach its target site of action, which could possibly be a metal-binding site of bacterial enzymes. The metal-8HQ complex will dissociate into a 1:1 ratio of 8HQ-metal charged complex and 8HQ free ligand.\(^\text{120}\) The charged 8HQ metal complex can bind and block the metal-binding sites on bacterial enzymes, which gives rise to the antimicrobial effect.\(^\text{119}\) Therefore, the lipophilicity, as indicated by the logP, is considered to be an important factor for antimicrobial activity of the investigated compounds.\(^\text{120}\) In addition, the dissociated free ligand of 8HQ possesses high chelating ability that could bind metallic prosthetic groups of microbial enzymes thereby leading to the inhibition of enzymatic activity.\(^\text{5,120}\)

Recently, 8HQ-uracil metal complexes bearing antimicrobial activity (Figure 7) have been reported.\(^\text{121}\) The complexes exhibited growth inhibition against many strains of Gram-positive and Gram-negative bacteria including resistant pathogens, such as *S. aureus*, *Enterococcus faecalis*, and *Candida albicans*.\(^\text{121}\)

Previously, 4-benzenesulfonamide (HQMABS), shown in Figure 10, is a hybrid of 8HQ and sulfanilamide and was reported to be a ligand for metal complexes.\(^\text{5}\) This study showed that HQMABS exhibited more potent antimicrobial activity with higher sensitivity against Gram-positive bacteria as compared to their individual parent compounds (ie, 8HQ and sulfanilamide).\(^\text{5}\) This demonstrates that there is a synergistic effect of 8HQ and sulfanilamide that facilitates the penetration of HQMABS into the site of action in bacterial cells.\(^\text{5}\) Therefore, HQMABS exhibited antimicrobial effects through a similar mechanism to that of 8HQ, as a membrane active agent via metal ion chelation.\(^\text{122}\) On the other hand, all metal complexes of HQMABS displayed weak to moderate activity as compared to their respective free ligand, HQMABS. Moreover, the antimicrobial activity of these compounds is dependent on the nature of the ligands, concentration and lipophilicity of the compound, nature of metal ions, geometry of the complex, and coordinate sites.\(^\text{5}\)

8HQ-based quaternary cationic surfactant

Quaternary cationic surfactants (Figure 11) were synthesized from the reaction of 8HQ and long chain alkyl halides.\(^\text{104}\) The study showed that cationic amphiphilic structures of quaternary salts allowed the compounds to interact with the bacterial lipid bilayer membrane.\(^\text{122}\) The effect may alter the membrane itself or cause toxicity to the membrane thereby leading to bacterial cell death.\(^\text{123}\) The activity of these 8-hydroxyquinolium derivatives is dependent on both the polar heads (ie, size and electronic charge distribution) and the hydrocarbon chain length.\(^\text{104}\) It was found that the activity increased from C-12 to C-14 carbon atoms and decreased in the case of C-16.\(^\text{104}\) This suggested that cationic and long chain hydrocarbons of an appropriate length facilitate bacterial killing via membrane attack.

### Antiviral activity

It is well recognized that nucleic acid binding ability is important for RNA-dependent-DNA polymerase inhibition, which is essential for antiviral activity.\(^\text{124}\) Among the groups of tested metal-binding compounds, 8HQ exhibited high antiviral activity with approximately 50-fold higher activity.\(^\text{124}\) Moreover, the binding activity of the Cu complexes of 8HQ and its derivatives were significantly higher than their respective free ligand forms.\(^\text{124}\) Interestingly, this activity was enhanced markedly when an equimolar concentration of Cu was added.\(^\text{124}\) According to Albert et al,\(^\text{119}\) the ratio of metal complex and their free ligand was shown to affect their antibacterial and antifungal activities; particularly only the 1:1 ratio provided the activity. It was found that increasing the amount of ligands resulted in the formation of more inactive complexes, thereby resulting in decreased activity – which is known as concentration quenching. This phenomenon was observed when high concentrations of drugs gave rise to lower inhibition of DNA synthesis in comparison to using a low drug concentration.\(^\text{124}\) Despite binding to viral nucleic acid, another possible mechanism of antiviral activity is its binding to Zn in enzymes, thereby leading to the inactivation of viral enzymes.\(^\text{125,126}\) However, the stability of the Cu complex is much greater than that of the Zn complex.\(^\text{127}\) This was supported by the study demonstrating that the 8HQ-Cu complex inhibited RNA-dependent-DNA polymerase as well as inactivating Rous sarcoma virus and herpes simplex virus with comparable activity as to that of 8HQ free ligand.\(^\text{112}\) Therefore, antiviral activity as exerted by ligand binding to Zn metalloenzymes may not be possible.\(^\text{112}\)

Macro cyclic polyamines such as AMD3100 (Figure 12) has reached Phase II of clinical trials and is considered to
be a prototype for an antihuman immunodeficiency virus compound.\textsuperscript{108} AMD3100 is known to block host cell entry via blocking cell surface G-protein-coupled receptors, such as CCR5 and CXCR4, which are chemokine receptors.\textsuperscript{128–130} Hydroxyquinoline-polyamine conjugates (Figure 13) were synthesized using hydroxyquinoline conjugation with polyamine backbones or polyazamacrocycles in order to mimic chemokine receptor antagonists.\textsuperscript{108} The results showed that the conjugated compounds elicited antihuman immunodeficiency virus activity against two viral strains, \textit{Human immunodeficiency virus} (HIV), 1 LAV and HIV-1 BaL, whereas CQ and polyazamacrocycle were shown to be inactive.\textsuperscript{108} Interestingly, AMD3100 (the reference compound) is only active against HIV-1 LAV thereby suggesting that the quinoline moiety is necessary for conjugated polyamine compounds as anti-HIV agents against both viral strains.\textsuperscript{108}

### Antiparasitic activity

#### Antimalarial activity

Malaria is considered to be a life-threatening infectious disease worldwide.\textsuperscript{131} Quinoline-containing compounds have been used for malarial treatment, such as quinine.\textsuperscript{132} Unfortunately, drug-resistance has been continuously reported\textsuperscript{107}, thus, the search for novel quinolone-based compounds is a demanding issue.\textsuperscript{133} Studies have shown that high sensitivity of human malaria to such compounds is mainly due to high lipid-water logP and metal-binding constants.\textsuperscript{134–139} It was noted that the inhibition of \textit{Plasmodium falciparum} multiplication and the chelating ability of the compounds were directly correlated.\textsuperscript{106} Chelators are known to interact with parasitic enzymes in different ways, such as by interacting with sulfhydryl groups, with amino groups, and with certain metal ions of enzymes.\textsuperscript{137} 8HQ as a potent chelator with high lipophilicity, and is known to possess an antimalarial effect against the intracellular stage of malaria in red blood cells by inhibiting a variety of metalloprotein oxidase enzymes, thereby resulting in the inhibition of glycolysis and parasitic growth.\textsuperscript{135} In vivo toxicity of 8HQ derivatives as diabetogenic agents (Figure 14) has previously been reported. However, a small group of substituents at the C-5 and C-7 positions on the quinoline ring can markedly lower the toxicity in higher animals.\textsuperscript{106} Substitution with a chlorine group at the C-5 or at both the C-5 and C-7 positions of 8HQ (Figure 14A and B, respectively) was shown to increase the lipid solubility and chelating ability of the compounds. These increases are expected to be an effect of the phenolic group that leads to improvement in metal chelation.\textsuperscript{106} Surprisingly, in this case, an improvement of the antimalarial activity was not observed.\textsuperscript{106} Unlike to be found in bacterial systems, the addition of an extra aromatic ring at the C-5 and C-6 positions (Figure 14C and 14D, respectively) was shown to not only increase its lipophilicity but also led to a decrease in its antimalarial activity. This could be attributed to the bulky
structures of the aromatic ring, that may cause steric effects and thereby hinder the compound from interacting with macromolecules on plasmodial enzymes or receptor sites.\textsuperscript{106} It was suggested that the substitution of C-5 or C-7 positions with electron-withdrawing groups or aromatic rings could improve the lipophilicity of the compounds and is likewise beneficial for drug delivery to target an intracellular site of action. Thus, improved chelating ability and reduced in vivo toxicity were observed, but antimalarial activity showed no improvement. This indicated that the antimalarial effect as afforded by these compounds may be contributed to other factors.\textsuperscript{106}

**Linkages of antiparasitic and anticancer activities**

Metabolic pathways in intracellular parasites (ie, *T. gondii* and *P. falciparum*) and in cancer cells are more sensitive to oxidative stress than normal cells and are dependent upon glycolysis in order to produce energy. In addition, some anticancer drugs were found to have antimalarial activity.\textsuperscript{138–140} Results from computational screening of quinoline-based antitumor compounds showed that nitrogen substitution at the C-5 position of the quinoline ring is required for antiprotozoal activity. Apparently, 5-nitroso-8HQ (NSC3852) (Figure 15) exerted the most potent predicted activity and is therefore relevant to the experimentally observed activity against *T. gondii*, with a half maximal effective concentration of 78.6 nM. But, its activity against *P. falciparum* was found to be eight times less active than that for *T. gondii*.\textsuperscript{105} So far, NSC3852 was reported to exert anticancer activity as elicited through the production of superoxide anion and nitric oxide (NO) against breast cancer cells.\textsuperscript{141} In contrast, antiprotozoal activity of NSC3852 was attributed to different mechanisms.\textsuperscript{105} Particularly, it was hypothesized that NSC3852 increased the intracellular oxidative stress via indirect mechanisms. This involves arylation of protein sulhydryls and the depletion of intracellular glutathione,\textsuperscript{142} which resulted in the metabolism of NSC3852 to non-redox cycling naphthoquinone and NO release from the nitroso group of the compound.\textsuperscript{105}

Interestingly, antiparasitic and anticancer activities of 8HQ derivatives are potentiated upon complexation with metal ions such as Cu and Zn.\textsuperscript{84} A series of antimony-8HQ complexes were synthesized and found to exhibit more potent antitrypanosomal and cytotoxic activities when compared to Sb salt (SbCl$_3$) and their free ligands, which are 8HQ, 5-chloro-8HQ, and 5-chloro-7-iodo-8HQ (or CQ).\textsuperscript{143}

**Antioxidant activity**

Oxidative damage is frequently found in many diseases such as aging, atherosclerosis, cancer, diabetes,\textsuperscript{144} and neurodegenerative diseases.\textsuperscript{7} Free radicals are continuously produced in cells through a wide range of biological processes.\textsuperscript{144} For example, the changing oxidation stage of Cu, which is a cofactor of SOD, results in the generation of ROS.\textsuperscript{145} Therefore, antioxidant defenses, such as those afforded by tocopherol, ascorbic acid, SOD enzyme, and catalases, are necessary in the maintenance of homeostasis.\textsuperscript{146} Many phenolic compounds, derived from either natural sources or synthetic methods, have been reported as potent antioxidants.\textsuperscript{147–149} An effective antioxidant activity of phenolic compounds is dependent on the stability of the phenoxyl radical formed in the reaction\textsuperscript{150} as well as the position of a substituent which affects the phenoxyl radical.\textsuperscript{151}
8HQ derivatives have been reported as potent antioxidants, which arises from their chelating ability. It is widely known that mixed ligand metal complexes can commonly occur in biological fluids from various bioactive ligands with metal ions. Interest in the area of metal complexation has steadily increased. SOD is one of the most useful antioxidant enzymes, known to convert superoxide into $\text{H}_2\text{O}_2$ and oxygen. However, this enzyme has certain limitations such as short shelf-life, low lipid solubility, low penetration into the cell, and high MW. The SOD structure has a central metal atom surrounded by the protein structure, thus, accordingly, great focus has been reported on the synthesis of small-sized lipophilic metal complexes in efforts to mimic the SOD activity.

A series of mixed ligand metal complexes using 8HQ, 5-iodouracil, and 5-nitouracil as ligands were synthesized and studied for their antioxidant activity using SOD assay. The results showed that amongst the different tested metal complexes, the 5-iodouracil-Mn-8HQ complex was shown to exert the highest activity, with an IC$_{50}$ of about 3-fold less than that of the free ligand 8HQ. This indicated that the coordination of metal ion into the free ligand can lead to enhancement of SOD activity. However, complexation of different types of metal ions with the same combination of ligands resulted in different bioactivities.

**Anti-inflammatory activity**

Nitric oxide is a short-lived free radical product generated by the conversion of L-arginine to L-citrulline, which is facilitated by nitric oxide synthase (NOS). There are three isoforms of NOS comprised of endothelial NOS, inducible NOS, and neural NOS. Small amounts of NO produced by endothelial NOS and neural NOS were shown to play important roles in the maintenance of homeostasis. In contrast, large amounts of NO from iNOS is generated in pathological conditions and inflammation. NO production is regulated by the expression of the iNOS gene, which is mainly modulated at the transcriptional level via the binding of transcription factors to sites such as NF-$\kappa$B sites, activator protein-1, interferon regulatory factor 1, and CCAAT/enhancer-binding protein $\beta$ (C/EBP$\beta$). The NF-$\kappa$B site is essential for lipopolysaccharide-mediated NO production in response to inflammation. Macrophage-derived NO acts as a host-defense mechanism against microbes and tumors, as well as a regulator of proinflammatory genes in vivo. It has been found that 8HQ inhibits lipopolysaccharide-induced NO production. Particularly, it suppresses iNOS mRNA expression and iNOS promoter activity by inhibition of NF-$\kappa$B activation and C/EBP$\beta$ DNA-binding activity. The study demonstrated that 8HQ possesses anti-inflammatory activity and could be further developed for the treatment of inflammatory diseases.

**Pb transportation across erythrocyte membranes**

Long-term exposure to Pb from the environment causes lead poisoning. Pb accumulation is found in bone tissue and is also distributed to other organs. Moreover, Pb accumulation in bone tissue is correlated with an increased risk of cardiovascular mortality, cognitive changes, and neurodegenerative diseases. The majority of Pb in whole blood is not found in the plasma but is sequestered in erythrocytes, which take up Pb via anion exchangers. The accumulation of Pb in erythrocytes affects erythrocyte enzymes, especially $\Delta$-ALA dehydratase and 5-nucleotidase. Such accumulation also accelerates the vascular clearance of red blood cells and thrombin generation, which may lead to an increased risk of developing cardiovascular disease. Most of the chelators that are employed for chelation therapy are not capable of passing across the erythrocyte membrane. Thus, the effective

![Figure 14 Substituted 8-hydroxyquinoline derivatives.](image)

![Figure 15 Structure of NSC3852. Abbreviation: NSC3852, 5-nitroso-8-hydroxyquinoline.](image)
reduction of the total body Pb level is not achieved and gives only a short period of therapeutic benefit.\textsuperscript{83,175} Therefore, a combination of Pb ionophore and Pb chelating agent was recommended as a new strategy for lowering total body Pb accumulation.\textsuperscript{176}

\(\Delta\)-ALA dehydratase is well recognized as a Zn-containing erythrocyte enzyme.\textsuperscript{83} Pb is believed to displace Zn from the enzyme, thereby leading to enzyme inactivation. However, the addition of Zn has been reported to restore \(\Delta\)-ALA dehydratase activity in vivo.\textsuperscript{177,178} Since Pb ions enter erythrocytes through anion channels, if they were not trapped in extracellular areas they could reenter cells.\textsuperscript{83} Thus, the increase of intracellular Zn may reverse Pb binding and facilitate the release of Pb from the enzyme. This potentiates intracellular Pb ions to exit from erythrocytes into the extracellular space, where they are effectively eliminated by Pb chelators.\textsuperscript{83}

CQ is a hydrophobic halogenated 8HQ derivative known to act as a Zn ionophore. It is capable of transporting Zn across the cellular membrane.\textsuperscript{80} An in vitro study demonstrated that CQ acting as a Zn ionophore could facilitate Pb escape from erythrocytes into the extracellular space. An increased intracellular Zn level thereby allows more effective chelation.\textsuperscript{83} This suggested that 8HQ derivatives and other classes of Zn ionophore could be further developed as a combined agent, acting as Pb chelators by lowering the total body Pb accumulation.\textsuperscript{83}

**Building block for artificial carbohydrate receptors**

Artificial carbohydrate receptors exploiting carbohydrate-based molecular recognition processes via noncovalent interactions of sugar binding\textsuperscript{179–181} have been developed (Figure 16) for diagnostic and therapeutic purposes.\textsuperscript{182} Binding preference in carbohydrate recognition is an important factor for effective systemic outcome.\textsuperscript{183} An in vitro study demonstrated that 8HQ-based receptors elicited higher affinity to \(\beta\)-galactoside as compared to that of quinoline-based receptors.\textsuperscript{183} Moreover, the addition of more 8HQ moieties into the receptor can potentiate higher affinity.\textsuperscript{183} It was suggested that the quinoline hydroxyl group plays an important role in complex formation and molecular recognition of carbohydrates. Therefore, 8HQ-based compounds could potentially be used as a building block for artificial carbohydrate receptors.\textsuperscript{183}

**Potential antidiabetic activity**

Insulin/insulin-like growth factor-1 signaling pathway (IIS) is an evolutionarily conserved pathway, which regulates lifespan and longevity in various species from nonvertebrates to humans.\textsuperscript{184} In nonhuman organisms, decreased IIS has been found to be associated with extended lifespan and protection against oxidative stress damage.\textsuperscript{185,186} In addition, experiments on mouse models also demonstrated a significant role of down-regulated IIS in the maintenance of metabolic homeostasis and oxidative defense.\textsuperscript{184} In humans, individuals with decreased IIS as found in those with Laron syndrome were shown to exhibit a lower rate in the development of diseases of civilization including acne, cancer, and diabetes mellitus (DM).\textsuperscript{184}

Forkhead box proteins (FOXO) are crucial regulators against oxidative stress conditions. Activation of FOXO in the

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**Figure 16** 8-Hydroxyquinoline is used as a building block for artificial carbohydrate receptors.

**Figure 17** FOXO1 functions are related to glucose homeostasis and providing protection against oxidative stress.

**Abbreviations:** FOXO1, forkhead box protein O1; SOD, superoxide dismutase.
nucleus results in an expression of many downstream effectors including antioxidant enzyme genes such as superoxide dismutase (MnSOD) and catalase. At the promoter level, FOXO1 induces the expression of the Hmox1 gene (heme oxygenase-1) thereby leading to the reduction of mitochondrial respiration and ROS formation.\textsuperscript{185} Moreover, in pancreatic \( \beta \)-cell, FOXO1 functions to inhibit \( \beta \)-cell proliferation and prevent \( \beta \)-cell apoptosis\textsuperscript{187} as shown in Figure 17. Therefore, the level of nuclear FOXO is important for oxidative stress resistance and cytoprotection of cells.\textsuperscript{187,188}

Overstimulated IIS is closely associated with the pathogenesis of DM. Overconsumption of hyperglycemic diet can cause high blood glucose levels, which induces glucose/FOXO1-mediated \( \beta \)-cell proliferation in order to produce insulin for controlling blood glucose.\textsuperscript{188} This long-term phenomenon generates insulin resistance and hyperinsulinemia, which are the hallmarks of DM type 2.\textsuperscript{188} IIS is found to regulate nuclear distribution of FOXO proteins.\textsuperscript{185} The induction of IIS, caused by hyperglycemic diet, induces Zn-dependent phosphorylation of nuclear

**Figure 18** The role of CQ as Zn ionophore in controlling blood glucose level via inhibition of FOXO1.

**Abbreviations:** IIS, insulin/insulin-like growth factor-1 signaling pathway; CQ, clioquinol; FOXO1, forkhead box protein O1; G6Pase, glucose-6-phosphatase; SOD, superoxide dismutase; PEPCK, phosphoenolpyruvate carboxykinase; Pi, phosphorylation.
FOXO1, that subsequently triggers their exclusion from the nucleus.

Reduction of nuclear FOXO1 levels as a result of phosphorylation leads to three events. First, an impairment of oxidative stress resistance of β-cells as caused by downregulation of MnSOD and catalase gene expressions. Second, an increased proliferation and apoptosis of β-cells. Third, the inhibition of FOXO1 serves as key regulators for the inhibition of hepatic gluconeogenesis. Therefore, FOXO1 inhibition suppresses hepatic enzymes necessary for glucose production, such as phosphoenolpyruvate carboxykinase and glucose-6-phosphatase, at the transcriptional level. Obviously the first two events promote β-cell apoptosis thereby leading to impaired insulin synthesis and altered blood glucose level, while the last event facilitates the lowering of blood glucose as shown in Figure 18.

8HQ has been reported as a diabetogenic agent owing to its ability to harm β-cells. As a Zn ionophore, 8HQ carries Zn into cells in the form of lipophilic, uncharged complexes and releases Zn inside the cells to promote Zn-dependent FOXO1 phosphorylation thereby leading to oxidative damage and apoptosis of β-cell. In addition, acidic proton (H+) released from the –OH group of 8HQ causes damage to β-cells.

The SAR study of 8HQ and its derivatives indicated that their binding affinity to Zn, charge of complex, and acidity are determinant factors for diabetogenicity of the investigated compounds. As mentioned, FOXO1 phosphorylation is Zn-dependent, therefore, compounds that possess affinity to bind and form uncharged complexes with Zn are also capable of penetrating into cells thereby leading to the induction of FOXO1 phosphorylation and the triggering of its exclusion from the nucleus. Reduced nuclear FOXO1 levels ultimately lead to β-cell destruction and apoptosis.

A series of 8HQ and derivatives containing OH groups at different positions on the quinoline ring (ie, 2HQ and 4HQ) as well as ionizable functional groups (ie, CO₂H and 8HQ)

8HQ derivatives with different substitutions.
Abbreviations: CQ, clioquinol; HQ, hydroxyquinoline.

Figure 19 8HQ derivatives with different substitutions.
Abbreviations: CQ, clioquinol; HQ, hydroxyquinoline.

Figure 20 Liver functions on controlling glucose metabolism and antidiabetic actions of 8-hydroxyquinoline derivatives.
Abbreviations: CQ, clioquinol; FOXO1, forkhead box protein O1; G6Pase, glucose-6-phosphatase; GLUT-1, glucose transporter; HIF-1α, hypoxia inducible factor; HLA-20, 5-((4-(prop-2-ynyl)piperazin-1-yl)methyl)quinolin-8-ol; InsR, insulin receptor; M30, 5-((methyl(prop-2-ynyl)amino)methyl)quinolin-8-ol; PEPCK, phosphoenolpyruvate carboxykinase.
Table I (Continued)

### Potential antidiabetic activity

- FOXO proteins are crucial regulators against oxidative stress conditions.
- Nuclear FOXO1 is important for the expression of many downstream effector genes, which function to maintain glucose homeostasis and protect against oxidative stress condition.
- FOXO1 exclusion from the nucleus activates β-cell proliferation and insulin production; however, this also reduces oxidative stress resistance leading to β-cell apoptosis.
- A compound is considered diabetogenic if it has the ability to harm β-cells, including decreasing antioxidant defenses and activating β-cell proliferation leading to β-cell apoptosis.
- 8HQ has been reported as a diabetogenic agent due to its ability to induce Zn-dependent FOXO1 phosphorylation via action of the Zn ionophore as well as its acidity caused by release of H⁺ from OH groups. These phenomena lead to β-cell apoptosis, which affects insulin production and control of blood glucose.
- It was suggested that the ideal antidiabetic agent should be capable of decreasing blood glucose level via induction of FOXO1 phosphorylation as well as minimizing β-cell damage.
- CQ is considered a potential antidiabetic agent. It has the ability to induce FOXO1 phosphorylation without diabetogenic effect. It was suggested that increased lipophilic, electronic, and steric effects may be involved in the lack of diabetogenicity.
- 8HQ-based antineurodegenerative agents such as M30 and HLA-20 have beneficial effects on controlling blood glucose via Fe chelation. Glucose metabolism is directly controlled by hepatic factors. The decreased hepatic Fe levels can mimic hypoxic conditions that results in lower blood glucose level.

**Abbreviations:** C/EBPβ, CCAAT/enhancer-binding protein beta; CQ, clioquinol; HIV, Human immunodeficiency virus; MRSA, methicillin-resistant Staphylococcus aureus; NF-κB, nuclear factor kappa-light-chain-enhancer of activated B cells; ROS, reactive oxygen species; SOD, superoxide dismutase; HQ, hydroxyquinoline; FOXO, forkhead box protein.

SO₃H) and polar amino groups were investigated in vitro. The results showed that 8HQ and 8-hydroxyquinaldehyde (Figure 19A) displayed diabetogenic effects and strongly induced FOXO1 phosphorylation in the presence of Zn²⁺. However, all 8HQ derivatives with substituents at C-2 (CO₂H) and C-5 (SO₃H) (Figure 19B) positions showed no diabetogenicity, and no induction of FOXO1 phosphorylation. This could be attributed to anionic charge groups (CO₂⁻ and SO₃⁻) that inhibited their penetration into the cell and specific compartments. Furthermore, 2HQ and 4HQ (Figure 19C) displayed no diabetogenic effects and no induction of FOXO1 phosphorylation. This suggested that the OH group in the C-8 position on the quinoline ring was crucial for the observed diabetogenic effect. Interestingly, CQ (Figure 19D), a derivative of 8HQ-bearing substitutions with hydrophobic groups at C-5 (chlorine) and C-7 (iodine) positions induced FOXO1 phosphorylation without diabetogenic effects. The phosphorylation of FOXO1 arises from its CQ property as the
Zn ionophore. It was suggested that increased hydrophobicity alone is unlikely to account for the nondiabetogenicity of CQ; however, other factors such as electronic and steric effects may be involved in the lack of diabetogenic effects.

According to IIS regulatory roles on glucose homeostasis, an ideal antidiabetic agent should be capable of inducing FOXO1 phosphorylation as well as minimizing oxidative damage to β-cells. Recent studies suggested that the scope of drug design may focus on small Zn-binding FOXO1 regulators targeting lipophilic compounds that do not release H⁺ after Zn-binding. Therefore, CQ is a potential candidate for DM treatment.

Antidiabetic activity of other 8HQ derivatives, such as M30 and HLA-20, have been reported to function through different mechanisms. The liver is the center of Fe homeostasis, and hepatic enzymes are found to be key regulators of hepatic gluconeogenesis. Glucose metabolism is directly controlled by hepatic factors that in turn are regulated by Fe levels, as is the case for the glucose transporter and the insulin receptor. As Fe chelators, M30 and HLA-20 generate low levels of Fe, thereby mimicking hypoxic conditions. Such conditions increase the expression of Fe/O² regulated hypoxia inducible factor. The activation of hypoxia inducible factor leads to the expression of downstream effector genes including the glucose transporter and the insulin receptor, which consequently causes lower blood glucose levels as shown in Figure 20.

In addition, both compounds exhibit protective effects against oxidative damage to β-cells. A study on pancreatic β-cell lines indicated that cytoprotective effects of HLA-20 and M30 arise from combined actions in which they act as Fe chelators. M30 and HLA-20 directly decrease ROS formation by inhibiting the Fenton reaction as well as indirectly increasing catalase activity, possibly by maintaining nuclear FOXO1 levels. Furthermore, both compounds promote antiapoptotic effects against H2O2 stress-induced mitochondrial dysfunction. Mitochondria are known to play important roles on cell cycle regulation, and their dysfunction is a critical event leading to cell apoptosis. M30 and HLA-20 exert antiapoptotic effects by maintenance of a mitochondrial factor called Δψm, which is necessary for mitochondria survival.

Due to IIS and the FOXO1 phosphorylation inductive ability and significant cytoprotective effects, 8HQ and its derivatives could be a promising class of compound that merits further development for the treatment of DM.

**Conclusion**

Metal imbalance plays a crucial role in the etiology of many diseases that affect quality of life. The objectives of treatment...
are not only aimed at restoring metal balance but also at minimizing cellular damage. 8HQ and its derivatives possess diverse pharmacological and biological activities, which are a result of their chelating ability (Table 1). Interestingly, such bioactivities originate from multiple mechanisms (Table 2). These mechanisms of actions function in restoring metal homeostasis as well as promoting protective effects. Therefore, 8HQ and its derivatives are considered as promising candidates that should be further developed as therapeutics for many diseases.

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