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ortho-Quinone methide (o-QM): a highly reactive, ephemeral and versatile intermediate in organic synthesis

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Abstract: Since its first observation in 1907, *ortho*-quinone methide (*o*-QM) has occupied a strategic place within the framework of reactive intermediates in organic synthesis. In recent years, *o*-QM has enhanced its importance as a versatile reactive intermediate, and its applications in organic synthesis, material chemistry, fine chemicals, and pharmaceuticals are increasing rapidly. This critical review summarizes the key concepts behind *o*-QMs and provides an overview of current applications in organic synthesis to provide appropriate background for synthetic, medicinal and combinatorial developments. This review cover the literature from its origin to the mid of 2014 (112 references).

1. Introduction

ortho-Quinone methides (o-QMs) are short-lived and highly reactive versatile intermediates in organic synthesis, material chemistry, fine chemicals, and pharmaceuticals. The *ortho*-quinone methide (o-QM) was first suggested by Fries in 1907 and is generally formed by the reaction of phenol with aldehyde in the presence of acid or base. The first direct evidence was given by Gardner in 1963 by trapping it at -100 °C. After 1963, its use as versatile intermediate in organic synthesis increased enormously particularly in many tandem [4+2] cycloaddition reactions with a variety of dienophiles. However, there are abundant indirect evidences for the *in situ* generation of *ortho*-quinone methides. Most indirect evidence comes from the structural identification of the products that result from dimerization, trimerization, intramolecular and intermolecular [2+2] cycloadditions as well as the nucleophilic trapping of o-QMs. The chemical behaviour of o-QMs resembles that of α , β -unsaturated ketones due to the presence of 1,3-cyclohexadiene core substituted with a carbonyl and an exo-methylene group. They react very rapidly with nucleophiles and undergo efficient Diels-Alder reactions with electron-rich olefins. Expansive reactivity of the o-QM can be used in linchpin reaction for the construction of various natural products.

Compared to carbocations, carbanions, radicals, carbenes, nitrenes and so forth, *ortho*-quinone methides are relative newcomers to the family of inherently reactive species. Quinone methides exist in three isomeric forms *o*-, *m*-, and *p*-quinone methides (also known as *o*-, *m*-, and *p*-QMs) (Fig. 1). *meta*-Quinone methide is actually a resonance hybrid of two canonical forms, one of which is zwitterionic form stabilized by aromatic conjugation, increasing its polarity and thus enhancing reactivity. The importance of *o*- and *p*-quinone methides in organic synthesis and their role in biochemistry have been studied in detail. Unlike benzoquinones, *o*- and *p*-quinone methide derivatives are highly polarized compounds, usually observed with difficulty or postulated as reactive intermediates because of facile reactions driven by the formation of aromatized phenol derivatives.

Fig. 1 Isomeric forms of quinone methides.

ortho-Quinone methides are highly versatile intermediates that have been extensively harnessed by nature (Fig. 2). A variety of plants, animals, and insects capitalize upon these types of compounds as a means of defence. However, despite general knowledge of o-QMs for over a century these intermediates still lie outside the synthetic mainstream. Pettus and Water^{1a} reviewed o-QMs particularly its preparation, the benefits and limitations associated with each method as well as the applications in total synthesis.

Fig. 2 *ortho*-Quinone methides.

ortho-Quinone methide exists in E/Z geometric isomers that are fluxional in nature, and behaves as a combination of charged zwitterionic and biradical structures (Fig. 3). The distribution among these geometric isomers is believed to result from the differences between non-bonded interactions. If from a steric point of view R^1 substituent is smaller than oxygen, then E-configuration is preferred. However, increasing the size of R^1 substituent can cause

the Z-configuration to predominate. The E/Z ratio proves important in governing the diastereoselective outcome for Diels-Alder cycloadditions.

E-configured R1 R2 Z-configured (if R1 is smaller than carbonyl)
$$R^1$$
 R^2 R^2 R^3 R^4 R^2 R^4 R^2

Fig. 3 Geometric, zwitterionic and biradical structures of *o*-QM.

Quinone methides (QMs) are common reactive intermediates in chemistry as well as in the photochemistry of phenols, attracting much attention recently owing to their biological activity. Although the partial zwitterionic character of QMs makes them both electrophilic and nucleophilic, their reactivity with nucleophiles is especially important in biological systems. It has been demonstrated that QMs react with amino acids and proteins, and inhibit the action of some enzymes. Moreover, they have been implicated as the ultimate cytotoxins responsible for the function of agents such as antitumor drugs, antibiotics, and DNA alkylators in biological chemistry. Later on, it was shown that QMs are not responsible reagents that cross-link DNA, but they probably involve metabolic formation of formaldehyde. To understand the chemistry of anthracycline antitumor antibiotic, a simple stable *o*-QM has been constructed and fully characterized. Reaction of quinone methide with nucleophiles, including 2',3'-isopropylideneadenosine has been examined.

Several important clinical drugs like cisplatin, psoralens, and mitomycin C are known to induce DNA ISC (interstrand cross-links) formation, which can disrupt cell maintenance and replication. Both reductive and oxidative metabolisms form quinone methides and have become a very important topic in drug design and drug safety. Even though *ortho*-quinone methides were investigated in detail 50 years ago, the last few years have brought about a renaissance in their chemistry. This Review summarizes the recent developments in this field and highlights key historical contributions.

2. Generation of *ortho*-quinone methides

ortho-Quinone methides are highly reactive species, which have attracted much attention because of their intriguing structure and properties. A number of methods for the generation

of o-QMs have been developed so far enabling the straightforward construction of diverse aromatic molecules with successful application to a number of total syntheses of complex natural products. In the past decade, the most popular methods were photochemical initiation of o-(α -phenyl) substituted phenols or thermal initiation of various substituents on the benzene ring of ortho-methyleneacetoxy phenols. Lewis acid, base, and chemical oxidants were also used to generate o-QMs. Thus, the method of generation of the o-QM becomes intimately linked with the manner in which it is to be utilized.

2.1. Thermal generation

Among the used procedures to generate o-QMs, most common are thermal and base initiations in which good to excellent stereoselectivity with excellent yields of the corresponding products have been obtained. Only a limited number of accounts have been reported where acid is used to generate the o-QMs. This is mainly due to the low compatibility of acid with the dienophiles, and the greater ionic character of the reaction conditions results the product with low stereoselectivity. Baldwin and co-workers¹² introduced an efficient method for the generation of o-QM from ortho-methyleneacetoxy phenols that was used for the biomimetic synthesis of (\pm)-Alboatrin (Scheme 1). Alboatrin 1 is a phytotoxic natural product, and its biosynthesis has been realized through hetero Diels-Alder reaction of an orcinol derived o-quinone methide 2 and (R)-4,5-dihydro-2,4-dimethylfuran 3.

Scheme 1 Retrosynthesis of Alboatrin.

Baldwin incipiently assumed that dehydration of the corresponding hydroxymethyl orcinol derivative 4 would give o-QM. Compound 4 was obtained by the reduction of precursor 6b, which was found to be unstable and rapidly decomposed under the reaction conditions. Then a diacetate 5 was prepared, which was reduced to its alcohol and consequently underwent transesterification by itself to give an *ortho*-methyleneacetoxy phenol 6a under thermal conditions (Scheme 2). Subsequently, simple heating of 6a in the presence of dimethylfuran 3 afforded acetylalboatrin 7 as the major product (63%), acetyl *epi*-alboatrin (5%), and an inseparable mixture of diastereoisomers 8 (25%). Deacylation of 7 yielded the desired (±)-

Alboatrin, for which spectral data was found to be identical to the natural (\pm) -Alboatrin. Intramolecular hydrogen bonding might facilitate the elimination of acetic acid upon heating through a six-member transition state furnishing the o-quinone methide.

Scheme 2 Synthesis of acetylated (\pm) -Alboatrin (7).

Christopher D. Bray¹³ reported the generation of *o*-QM from deprotonation of *ortho*-hydroxybenzyl acetate **9** with i PrMgCl under mild anionic reaction conditions, which reacts with *exo*-enol ethers **10** as 2π partners in hetero Diels–Alder (HDA) reactions furnishing *mono*-benzannelated spiroketals **11** (Scheme 3). This rapid and simple strategy is clearly applicable to the synthesis of a wide range of natural products including berkelic acid, chaetoquadrin A and cephalostatin.

$$\begin{array}{c}
 \text{HO} \\
 \text{AcO}
\end{array}$$

$$\begin{array}{c}
 \text{iprMgCl} \\
 \text{O} \\
 \text{I0}
\end{array}$$

$$\begin{array}{c}
 \text{O} \\
 \text{I1}
\end{array}$$

Scheme 3 Generation of *o*-QM using *o*-hydroxybenzyl acetate and ⁱPrMgCl.

Rosellon and co-workers¹⁴ developed a method for the generation of *ortho*-quinone methides **13** *via* fluoride-induced desilylation of silyl derivatives of *o*-hydroxybenzyl iodides **12** under mild experimental conditions (Scheme 4).

Scheme 4 Formation of *o*-QMs *via* fluoride-induced desilylation.

Dolatowska and Wojciechowski¹⁵ reported the thermal extrusion of sulphur dioxide from 1,3-dihydrobenzo[c]thiophene-2,2-dioxides **16a** and 1,3-dihydro-2,1-benzisothiazole-2,2-dioxides **16b** to generate the corresponding quinodimethanes **17a** and quinonemethyleneimines **17b**, which undergo Diels-Alder reaction with alkene **18** to give the compound **19** (Scheme 5).

Scheme 5 Generation of *o*-QMs 17 by thermal extrusion of sulphur dioxide.

The sultone of 2-hydroxytoluene- α -sulfonic acid **20** after extrusion of SO₂ upon photochemical irradiation gave o-quinone methide **21**, which upon reaction with N-phenyl maleimides **22** afforded the respective adducts **23** (Scheme 6).

$$X$$

$$X$$

$$Y$$

$$Z0$$

$$X = Y = H \text{ or } CI$$

$$X$$

$$X$$

$$Z1$$

$$X$$

$$Z1$$

$$X$$

$$Z2$$

$$R = Ph, Me$$

$$R' = H, Me$$

$$Y$$

$$Z3$$

$$Z3$$

Scheme 6 Generation of *o*-QM from benzosultones.

Ohwada and co-workers¹⁶ reported retro-Diels-Alder reaction of 4*H*-1,2-benzoxazines **24** to generate *o*-QMs **25** under the thermal conditions, which undergo Diels-Alder reaction with vinyloxy cyclohexane **26** to afford the chroman derivative **27** (Scheme 7). They also presented a mechanistic study of the retro Diels-Alder reaction of benzoxazines **24** with various substituted benzene ring. In general, *o*-QMs are too unstable to be isolated and observed spectroscopically, unless they are trapped with metal complexes such as Cp*Ir. In

all the reported methods, *o*-QMs were generated *in situ*, and underwent 1,4-conjugate addition with nucleophiles or cycloaddition reaction with suitable dienophiles.

Scheme 7 Formation of o-QM via retro-Diels-Alder reaction of 4H-1,2-benzoxazines 24.

o-QMs may be generated through a diverse range of initiation methodologies (Scheme 8). Perhaps the most commonly encountered method for the generation of o-QM is the elimination of a stable molecule, typically a benzylic substituent to generate the methylene and α -carbonyl functionalities with concomitant dearomatisation. Various precursors have been utilized for the thermolytic generation of *ortho*-quinone methides. It should be noted that all thermal generation techniques preclude the application of nucleophiles that are thermally unstable. With any given precursor, there is substantial temperature range for initiation that depends upon the substituents. In general, if the process involves significant non-bonded interactions, then the temperature requirements are higher, while extended conjugation or other stabilizing factors lower the overall temperature requirements (Scheme 8).

Scheme 8 Thermal generation of *o*-QMs.

2.2. Photochemical generation

Apart from the thermal generation of *o*-QM intermediates in presence of different basic or acidic reagents, photochemical energy sources have also been extensively used to generate *o*-QMs. In general, *o*-QM intermediate **29** is generated by light-induced water elimination from *ortho*-hydroxybenzyl alcohols **28** or excited state intramolecular proton transfer (ESIPT) in *ortho*-hydroxystyrenes **30** (Scheme 9).

Scheme 9 Photochemical generation of *o*-QMs.

Majorty of β -cyclodextrin inclusion complexes undergoes photohydration of the olefin preventing the formation of o-QM by insertion of a methylene group between the phenol and the olefin moieties. Miranda *et al.*¹⁷ reported the formation of an o-QM via C–C fragmentation of a zwitterion formed by excited state intramolecular proton transfer (ESIPT) from an *ortho*-allylphenol derivative **31** for the first time (Scheme 10). The mechanism has been explained in terms of the formation of the zwitterionic intermediate **32A** by the initial ESIPT from the phenolic subunit to the double bond. Thus, subsequent C–C bond formation with concomitant ring opening elucidates the generation of the o-QM **32B**, which would be trapped by solvent methanol to give compound **33**. The direct involvement of an o-QM intermediate has been confirmed by laser flash photolysis. Use of 266 nm excitation wavelengths in acetonitrile solvent gave a well defined signal with two maxima around 300 and 400 nm.

Scheme 10 Generation of *o*-QM *via* C–C fragmentation of zwitterions.

Generation of *o*-QM by flash photolysis of *ortho*-hydroxybenzyl substrates **34** in aqueous solution has also been reported (Scheme 11). Hydrogen-ion catalyzed hydration of *o*-QM furnished back to the *ortho*-hydroxybenzyl alcohol, which was anticipated to ensue through initial fast equilibrium protonation of the carbonyl oxygen followed by trapping with water. The slower uncatalyzed hydration proceeds through nucleophilic attack of water on the QM methylene group.¹⁸

Scheme 11 Flash photolysis of *o*-hydroxybenzyl substrate **34** to give *o*-QM.

o-Quinone α-phenylmethide was produced as a short-lived transient species in aqueous solution by flash photolysis of *ortho*-hydroxy-α-phenylbenzyl alcohol, and its rate of decay was measured in HClO₄ and NaOH solutions as well as in CH₃CO₂H, H₂PO₄, and HCO₃ buffers. Results showed that the hydration of quinone methide back to its benzyl alcohol precursor occurs by acid-, base-, and uncatalyzed routes. Kresge and co-workers¹⁹ reported the generation of o-QMs 35 and 36 by photodehydration of the corresponding *ortho*-hydroxybenzyl alcohols 37 and 38. *ortho*-Hydroxybenzyl-p-cyanophenyl ether 39 after photocleavage gave o-QMs 36 (Fig. 4). Flash photolytic techniques have been used to monitor the progress of the reactions because the reactions of quinone methides are very fast.

Fig. 4 Structures of o-hydroxybenzyl alcohols and their photodehydrated o-QMs.

Lukeman and Wan²⁰ observed predominate D₂O exchange at the 2' position irrespective of D₂O (MeOD) content during the study of the photochemical incorporation of deuterium on phenylphenol **40** in D₂O (CH₃OD)-CH₃CN solutions with varying D₂O (CH₃OD) content.

Exchange at the 2'-position (but not at the 4'-position) was observed when crystalline samples of compound **40** (deuterated OH i.e. OD) were irradiated. (Scheme 12)

Scheme 12 Photochemical incorporation of deuterium on phenylphenol.

No deuterium exchange was observed in case of compounds **49-52**, while 2,2'-biphenol **48** underwent similar photo exchange reaction as compound **40** (Fig. 5).

Fig. 5 Structure of compounds which do not undergo deuterium exchange.

Wan and co-workers²¹ reported the generation of quinone methide **54** by ESIPT in which naphthyl group acts as the proton acceptor (Scheme 13). The reaction has been carried out in a variety of solvents as well as in the solid state illustrating that the proton transfer is intrinsic (proton transfer do not requires water).

Scheme 13 Generation of *o*-QM by ESIPT.

Zhan and co-workers²² have reported the generation of o-thioquinone methide **59** by flash photolysis of benzothiete **58** in aqueous medium (Scheme 14). The hydration rates of the quinone methide **59** to ortho-mercaptobenzyl alcohol **60** were measured in perchloric acid solution using H_2O and D_2O as the solvent in acetic acid in the presence of tris(hydroxymethyl)methylammonium ion buffers.

Scheme 14 Generation of *o*-thioquinone methide **59**.

Solvent isotope effects on hydronium-ion catalysis of hydration for the two substrates are different. Solvent isotopic effect for the oxygen quinone methide **61** is $k_{\rm H}/k_{\rm D} = 0.42$, while for the sulfur substrate **59** is $k_{\rm H}/k_{\rm D} = 1.66$. The inverse isotope effect ($k_{\rm H}/k_{\rm D} < 1$) in the oxygen containing system specifies pre-equilibrium proton-transfer reaction mechanism by which protonation of the substrate on its oxygen atom being fast and reversible, and capture of the benzyl-type carbocationic intermediate. On the other hand the normal isotope effect ($k_{\rm H}/k_{\rm D} > 1$) in the sulfur system implies the protonation of the substrate on its sulfur atom being rate-determining and carbocation capture is fast (Scheme 15).

Scheme 15 Solvent isotope effects on *o*-QM and *o*-thioquinone methide.

Matsumoto and co-workers²³ reported the thermal and photochemical generation of *o*-QM that was accelerated by the presence of water molecule in reaction solvents via the formation of anionic micelle and vesicle. According to their report amino phenanthrol **65** and their naphthalene analogues act as precursors and transforms to QM **68** in presence of light (Scheme 16).

Scheme 16 Generation of QMs by the formation of anionic micelle and vesicle.

Popik and Arumugam²⁴ reported the efficient generation of quinone methide from 2-hydroxymethyl phenol **69**. Isomeric 3-hydroxy-2-naphthalenemethanol **70** and 2-hydroxy-1-naphthalenemethanol **72** resulted the respective naphthaquinone methides, 2,3-naphthoquinone-3-methide **71** and 1,2-naphthoquinone-1-methide **73** (Scheme 17). During the generation of *ortho*-naphthaquinone methides (*o*-NQMs) **71** and **73** in the flash photolysis of ammonium precursors **70c** and **72c** ensues within the duration of a laser pulse, irradiation of **70a** and **72a** as well as their ethoxy derivatives **70b** and **71b** provides evidence for the detection of a kinetic precursor to *o*-NQMs. The rate of formation of *o*-NQMs does not depend on pH of the solution. Further, the generation of cyclohexadienone **75** and **76** by the photochemical irradiation of three different precursors **74a-c** and **76a-c** have been reported (Scheme 17).

Scheme 17 Generation of o-QM by using phenolic derivatives.

Wan and co-workers²⁵ tactically substituted the hydroxymethyl phenols with 2-hydroxy-2-adamantyl moiety and explored their photochemical reactivity. Upon excitation of AdPh derivative **78** to its singlet excited state, at first it undergoes intramolecular proton transfer followed by the loss of H_2O to give quinone methide **80**, which has been detected by laser flash photolysis (CH₃CN-H₂O 1:1, $\tau = 0.55$ s) and UV-vis spectroscopy (Scheme 18). The quantum yield of QM formation has been increased by three fold and lifetime is enhanced by the introduction of the adamantyl substituent on the *ortho*-hydroxymethyl phenol moiety.

$$\begin{array}{c} \text{OH} & \text{OH} \\ \text{OH} \\ \text{OH} & \text{OH} \\ \text{OH} & \text{OH} \\ \text{OH} \\ \text{OH} & \text{OH} \\ \text{OH} \\ \text{OH} & \text{OH} \\ \text{OH} \\$$

Scheme 18 Formation of adamantyl substituted o-QM 80.

Freccero and co-workers²⁶ reported the photogeneration of new Binol quinone methides 83 and 84. They also explored mono- and bisalkylation of free nucleophiles by product

distribution analysis and laser flash photolysis in water solution using Binol quaternary ammonium derivatives **81** and **82** as photo activated precursors (Scheme 19).

Scheme 19 Photochemical generation of binol quinone methide.

Excitation of anthracene derivative **85** to S_1 initiates photodehydration giving the corresponding quinone methide **86** that was detected by laser flash photolysis (LFP) in 2,2,2-trifluoroethanol.²⁷ The QM decays by protonation giving a cation **87**, which subsequently reacts with nucleophiles to give compound **88** (Scheme 20). QMs of the anthrol series is important for potential use in biological systems since the chromophore absorbs at wavelengths > 400 nm. Anti-proliferative investigations conducted with 2-anthrol derivative **85** on three human cancer cell lines showed higher activity for irradiated cells.

Scheme 20 Formation of QM by photodehydration of hydroxylated anthracene.

The vinylidene-quinone methides **92** and **93** were generated by irradiation of 2-alkynylphenols **89** and **90**, respectively that were detected by laser flash photolysis in organic solvents and aqueous acetonitrile (Scheme 21). It was also shown that the UV-vis properties and the electrophilicity are enhanced by a β -silicon effect and solvent polarity.

Scheme 21 Photochemical generation of vinylidene-QMs.

3. Reactions of ortho-quinone methides

The transient nature of *ortho*-quinone methides is due to their propensity to undergo rapid rearomatisation either by Michael addition of nucleophiles or often more usefully by cycloaddition with 2π partners or via oxa- 6π -electrocyclisation. Surprisingly, this rapid reactivity was historically seen as a deterent to synthetic development in the field with the potential for dimerisation and trimerisation sometimes a major stumbling block. To avert this, it is common to employ an excess of the nucleophilic or dieneophilic components and the localized o-QM concentration is often kept low.

o-QMs being a labile intermediates have been stabilized by metal coordination. The examples of simple isolated quinone methides are scarce. In fact in condensed phases, the parent compound has only been characterized spectroscopically at lower temperatures because it is extremely reactive. Spectroscopic evidence of the zwitterionic form of the o-QM derived from vitamin E has only been possible at –78 °C through stabilization by interaction with N-methylmorpholine N-oxide. These results show how difficult is to isolate quinone methides. para-Quinone methides have been discussed as intermediates in the chemistry of lignins and have been used in organic synthesis as electrophiles or electron acceptors. Because the parent o-QM molecule is unstable, NMR spectroscopic data are not available. Therefore, metalated parent ortho-quinone methide complex was examined using two-dimensional NMR spectroscopic techniques to understand its structure and reactivity. o-QMs are also believed to be key intermediates in the action of several antitumor and antibiotic drugs.

3.1. Electrocyclisation and cycloadditions

3.1.1. [4+2] Electrocyclisation and cycloadditions

Baldwin and Adler²⁹ has reported a proficient process for the synthesis of 2,2-dimethyl-2*H*-chromenes in a single step from the corresponding phenol and 3-methyl-2-butenal *via* microwave irradiation in CDCl₃. These reactions generated a transient *o*-QM intermediate **99**, which upon *in situ* electrocyclization gave the final bicyclic chromenes **100** in good yields (Scheme 22).

Scheme 22 Synthesis of chromenes via electrocyclization of *o*-QMs.

ortho-Quinone methides are versatile intermediates involving a minimum of seven carbon atoms, which are mainly involved in 1,4-Michael type additions as well as aza-Michael reactions with various nucleophiles. o-QMs also give Diels-Alder and hetero-Diels-Alder cycloaddition products (Scheme 23). o-QM especially derived from 4-hydroxycoumarin undergo [4+2] cycloaddition reaction with pentafulvenes to afford pyranocoumarin and pyranopyrone.³⁰

Scheme 23 Reactivity of *ortho*-quinone methide with alkenes.

Ploypradith *et al.*³¹ reported the generation of o-QMs by the reaction of compound **103** with p-TsOH on silica. Hetero Diels-Alder (HDA) reaction of *in situ* generated o-QMs **104** with styrenes **105** and **106** gave the corresponding arylchromans **107** and **108** in good yields (Scheme 24).

Scheme 24 Acid-catalysed generation of *o*-QMs and their HDA reaction with styrenes.

Sajiki and co-workers³² established a FeCl₃-catalyzed method for the synthesis of 1-naphthoquinone-2-methides 112a from dihydronaphthalenes 109 and further transformation of the products in an annulation reaction with various allyl silanes to afford biologically useful dihydronaphthopyran derivatives. The annulation can proceed via an *o*-NQM intermediate. The site-selective cleavage of one C=O bond is promoted by the coordination of the two oxygen atoms of 109 to the Lewis acid to give a five-member ring, which is then cleaved to give the zwitterionic intermediate 110. Subsequent migration of the siloxymethyl group (-CH₂OTBS) and aromatization provides the 2-siloxymethyl-1-naphthol 112. Further, Lewis acid induced elimination of the silanol (TBSOH) leads to an *o*-NQM 112a, which undergoes annulation with allyl-TMS through a hetero-Diels–Alder reaction to furnish 113 (Scheme 25).

Scheme 25 FeCl₃-catalyzed synthesis of 1-naphthoquinone-2-methides.

Lindsley *et al.*³³ has reported the total synthesis of polemannones B **114** and C **115**, which are highly oxygenated benzoxanthenones derived from *Polemannia montana* by employing catalytic Cu(II)/sparteine oxidant system for β , β -phenolic couplings and cascade Diels–Alder reaction by tandem inverse-electron demand in 75-90% yield (Fig. 6).

Fig. 6 Structure of polemannones.

The polemannones are unique because there is an extra electron donating ether moiety. Compounds containing electron-donating ether moieties *para* to the phenol such as **116** stabilizes the *o*-QM intermediate and provide the 'push' in the inverse-electron demand in Diels–Alder reaction during the synthesis of **118** *via o*-QM **117** (Scheme 26). If two electron-donating ether moieties are present, the second one is always located at *meta* to the phenol.

Scheme 26 Synthesis of carpanone *via* electrocyclization of *o*-QM 117.

But exceptionally, it is wonderful that *o*-methoxy group as in **119** equally stabilize the *o*-QM intermediate and provide the 'push' in the inverse electron demand Diels-Alder reaction to provide unnatural benzoxanthenone **121** (Scheme 27).

Scheme 27 Synthesis of benzoxanthenone 121.

Jha and co-workers³⁴ reported the synthesis of naphthopyrans **133** from 2-naphthol **129**, a secondary amine **124**, and 3-hydroxy-2,2-dialkylpropanal **122** in the presence of a catalytic amount of *p*-toluenesulfonic acid. This reaction is a one-pot retro-aldol disintegration of 3-hydroxy-2,2-dialkylpropanal followed by formation of a Mannich base intermediate **130** from 2-naphthol, a secondary amine, and formaldehyde (retro-aldol product). The Mannich base undergoes disproportionnation to give a QM intermediate **132** and a secondary amine. Compound **125** reacts with amine **124** to generate enamine **128**. QM intermediate undergoes electrocyclization with enamines to generate the final product **133** (Scheme 28).

Scheme 28 Synthesis of naphtha pyrans 133 *via* electrocyclization of *o*-QM with enamines.

3,3-Dialkyl-2-morpholino-3,4-dihydro-2*H*-naphtho[1,2-b]pyrans **137** were synthesized by the reaction of 1-naphthol **134**, morpholine **135** and compound **136** in the presence of catalytic amount of *p*-TSA (Scheme 29).

Scheme 29 Synthesis of naphtho fused pyrans via *o*-QM.

Lee *et al.*³⁵ have reported an efficient one-pot approach for the synthesis of benzopyranobenzo pyrans and naphthopyranobenzo pyrans. Resorcinol and naphthols undergo domino aldol type reaction followed by hetero Diels-Alder reaction with the benzaldehyde. Reaction of compound 139 with resorcinol 140 in the presence of ethylenediamine diacetate (EDDA)/TEA first gives intermediate 141 as an aldol-type product. TEA promoted the dehydration of intermediate 141 to form quinone methides 142a and 142b. Because of sp²-geminal effect the endo-transition structure 142b must have been more favourable than the *exo*-transition structure 142a due to the 1,3-allylic strain. The regiospecificity is attributed to the products 143a and 143b due to possibility of hydrogen bonding between the hydroxyl group and carbonyl group (Scheme 30).

Scheme 30 Synthesis of benzopyranobenzo pyrans and naphthopyranobenzo pyrans.

ortho-Quinone methide **147** arising from a formal [2+2] cycloaddition between aryne **144** and DMF was found to undergo a [4+2] cycloaddition with ester enolate **148** or ketenimine anion to produce diverse coumarins **150** (Scheme 31).³⁶

Scheme 31 [4+2] Cycloaddition of *o*-QM with ester enolate.

Silver oxide-mediated oxidation of phenol substrates **151** for the synthesis of o-QMs **152** has been developed by Lei *et al.*³⁷ for the biomimetic syntheses of novel trimeric natural products, (\pm)-schefflone and tocopherol trimers **153** (Scheme 32).

OH
$$R^4$$
 Ag_2O $benzene$ R^2 R^3 R^4 R

Scheme 32 Silver oxide-mediated generation of *o*-QM for the synthesis of tocopherol trimers.

Bharate and Singh³⁸ have reported the synthesis of anti-malarial robustadials **A** and **B 159** starting from commercially accessible phloroglucinol. Biomimetic three-component reaction that involves the *in situ* generation of an *o*-QM *via* Knoevenagel condensation followed by Diels-Alder cycloaddition with (-)- β -pinene **157** gave robustadials **A** and **B**. A retrosynthetic pathway is shown in Scheme 33.

Scheme 33 Retro-synthesis of Robustadials A and B.

Robustadials **A** and **B** have been anticipated to be formed biogenetically from the Diels–Alder cycloaddition of o-QM **158** with (-)- β -pinene **157**. The key intermediate o-QM **158** could be generated by the oxidation of diformylated synthon **154** or by coupling of phenol **155** and aldehyde **156** *via* a Knoevenagel condensation (Scheme 34).

Scheme 34 Synthesis of Robustadials A and B.

The treatment of 155 with 156 and (-)-β-pinene 157 in the presence of sodium acetate in acetic acid under microwave irradiation (1000 W) for 4 min resulted in the formation of desired products in a combined yield of 68% (Scheme 34). Similar results were obtained when the reaction was carried out under conventional heating at 80 °C for 2 h. The *o*-QM 158 is produced *in situ* by treatment of addition of phloroglucinol 155 with isovaleraldehyde 156 followed by dehydration (Knoevenagel-like condensation). Then 157 can undergo the cycloaddition reaction with *o*-QM 158a and 158b via two different orientations and each can lead to two pairs of diastereoisomers 159a and 159b. The formation of two isomers attributed due to the attack of oxygen of the QM only to the more substituted terminus of the pinene double bond (Scheme 34).

The biosynthesis of the meroterpenoid guajadial **163** has been achieved *via* hetero-Diels-Alder reaction between caryophyllene **160** and an *o*-QM. Biomimetic synthesis of guajadial

163 and psidial A 164 involved three-component coupling reaction between caryophyllene 160, benzaldehyde 161, and diformylphloroglucinol 162 in an aqueous medium (Scheme 35). It has been envisaged that the relative stereochemistry of guajadial 163 would be partly controlled by the conformation of caryophyllene 160 during the hetero-Diels-Alder reaction. The essential *o*-QM 165 could be derived retrosynthetically from a simple Knoevenagel condensation between diformylphloroglucinol 162 and benzaldehyde 161. Consequently a domino three-component one-pot reaction between diformylphloroglucinol 162, benzaldehyde 161, and caryophyllene 160 would constitute a simple biomimetic synthesis of guajadial 163.

Scheme 35 Synthesis of the guajadial and psidial A.

The intermolecular Diels-Alder reactions between o-QMs and double bonds that are not electron-rich are known to be difficult to achieve due to the easy dimerization of the o-QMs, which in turn accounts for the novelty and efficiency of the protocol. Caryophyllene **160** being certain degree of conformational mobility and it was not certain how this flexibility would influence the facial bias in the *in vitro* hetero-Diels-Alder reaction. And from ¹³C NMR studies and molecular mechanics calculations it has come to known that caryophyllene has four possible conformations ($\alpha\alpha$, $\alpha\beta$, $\beta\alpha$, $\beta\beta$), which vary according to the relative nature of the exocyclic methylene and olefinic methyl groups. It has been considered the proposed biogenetic hetero-Diels-Alder reaction must involve the $\beta\alpha$ conformer (the most stable conformer) of caryophyllene **160** with proposed transition states (Scheme 36). The stereo

chemical upshot of the hetero- Diels-Alder reaction was being mainly controlled by the local conformation and intrinsic chirality of caryophyllene 160.

$$\begin{array}{c} \text{Guajadial} \end{array} \Longrightarrow \begin{array}{c} \text{HO} \quad \text{CHO} \\ \text{Ph} \quad \text{OH} \\ \text{($\beta\alpha$, trans)} \end{array} \end{array} \\ \begin{array}{c} \text{Ph} \quad \text{OHO} \\ \text{OHC} \quad \text{CHO} \\ \text{OH} \quad \text{CHO} \\ \text{OHC} \quad \text{OHO} \\ \text{OHC} \quad \text{CHO} \\ \text{CHO} \quad \text{CHO$$

Scheme 36 Possible conformations of guajadial and psidial A.

An enantioselective cycloaddition of an *o*-QM **167** with a chiral enol ether **168** was reported for the first time by Selenski and Pettus, ⁴⁰ which has been portrayed along with the formal synthesis of (+)-tolterodine **171** and total synthesis of (+)-mimosifoliol **172** as shown in scheme 37. These syntheses demonstrate a three-component one-pot benzopyran approach for the construction of chiral benzylic junctions. A low-temperature anionic method for *o*-QM generation was achieved in high *endo/exo* ratios for ensuing cycloadditions. ⁴¹

Scheme 37 Synthesis of (+)-mimosifoliol.

o-QM generation results from a series of events called the o-QM cascade. The cascade began with the nucleophilic addition of an organometallic reagent to an OBoc salicylaldehyde **166a**. The consequential benzyloxy anion **166b** attacks the OBoc carbonate to form a cyclic intermediate **166c**, which then disintegrates to form a more stable phenoxide **166d**. In particular conditions, β -elimination of the OBoc residue generates an o-QM **166e** as mentioned in scheme 38. The cascade invasion is both metal and temperature dependent and presumably represents the strength of the oxygen-metal bond and the electrophilicity of the metal cation like an aluminium reagent proceeds to intermediate **166b**, while a lithium reagent prolongs to intermediate **166d** and Mg reagent assists the ultimate conversion to **166e**. The consequential geometry of the o-QM olefin is E except a large group is situated at the 6-position of the salicylaldehyde.

Scheme 38 Generation of *o*-QM using anionic method.

o-QM in the presence of chiral enol ether addicted in a diastereoselective [4+2] cycloaddition. The facial selectivities of enol ethers **173** and **174** through cycloaddition are shown in Scheme 39.

Scheme 39 Cycloaddition reaction of *o*-QM with chiral enol ether.

The *syn* affiliation flanked by the phenyl residue of the benzopyran and the oxygen substituent of the acetal results from an *endo* transition state. By taking pseudoallylic strain and Houk's calculation into vindication, the enol ether reacts in the *s-trans* conformation. The *o*-QM endures reaction opposite the methyl residue of enol ether **173** and opposite the phenyl residue of enol ether **174**. The *S*-stereocenter of **173** and the 2*R*-stereocenter of **174** induce the *S*-configuration of the resulting benzyl junction of compound **175** (Scheme 39). The absolute configuration of the chiral enol ether determines the absolute configuration of the resulting benzopyran. The diastereoselectivity of *trans*-2*S*-phenyl-1*R*-cyclohexanol vinyl ether **174** was outstanding for all the *o*-QMs that ultimately created *R*-mimosifoliol and *R*-tolterodine.

The substituted enol ether 177 should also undergo a diastereoselective cycloaddition. Addition of phenyl Grignard reagent to aldehyde 176 in the presence of enol ether 177 affords benzopyran 178 creating three chiral centers with a 60% yield and 95% de (Scheme 40).

Scheme 40 Diastereoselective synthesis of compound **178**.

Singh and co-workers⁴² have reported the biomimetic synthesis *S*-Euglobals and their antileishmanial, antimalarial, and antimicrobial activities. Compound **179** was treated with formaldehyde and different terpenes like (+)-2-carene (**182**), (1R)-(-)-myrtenol (**183**), and (1R)-(-)-nopol (**184**) in the presence of NaOAc in CH₃COOH resulted in the development of respective pairs of regioisomers, **185** and **186** from 2-carene, **187** and **188** from myrtenol, and **189** and **190** from nopol in good yields. Similarly, treatment of **180** and **181** with formaldehyde and different terpenes in the presence of sodium acetate in acetic acid resulted in the formation of desired cycloadducts, **191-193** and **194-196**, respectively (Scheme 41).

Scheme 41 Biomimetic synthesis of *S*-euglobals.

Singh and co-workers⁴³ also described the synthesis of new analogues of euglobals **200-203** by biomimetic approach (Scheme 42). These synthetic compounds differ from natural euglobals in the nature of monoterpene and acyl functionality. Oxidative addition of phloroglucinol-terpene with DDQ (2,3-dichloro-5,6-dicyano-1,4-benzoquinone) in nitromethane gave the corresponding o-QM, which further undergoes intermolecular cycloaddition with monoterpenes such as β -pinene **157**, α -pinene **197**, (+)-3-carene **198** and (–)-camphene **199**.

Reagents and conditions: (a) PhNO₂, DDQ, 50 °C, 1 h

Scheme 42 Biomimetic synthesis of *S*-euglobals analogues.

Kucklaendera *et al.*⁴⁴ reported 1,4,9,10-anthradiquinone as precursor for antitumor compounds according to which the naphtho-condensed indoles **206** were obtained by the reactions of unsubstituted 1,4-anthraquinone **204** with enamines **205** *via* the normal Nenitzescu method. Indoles **206** were converted to Mannich bases followed by dimerization through the Diels–Alder reaction of *o*-QM intermediate **207** to give the compound **208** (Scheme 43). These heterocyclic compounds synthesized here were appraised for their anticancer properties in the NCI's human disease oriented *in vitro* anticancer screen.

Scheme 43 Synthesis of naphtho-condensed indoles.

Generation of *o*-QMs by the photolysis of carboxylate derivatives **209** in aqueous acetonitrile was reported by Steinmetz and Chen.⁴⁵ The carboxylate leaving groups are swiftly eliminated by photolysis of the 1,4-benzoquinone **209** to produce an *o*-QM intermediate **211** that is trapped by [4+2] cycloaddition with unreacted starting material **209** and generated photoadduct **212**. The carboxylate leaving groups are easily expelled upon photocyclisation to benzoxazolines **210** under aqueous conditions (Scheme 44).

Scheme 44 Synthesis of compound **212** *via* photo generated *o*-QM.

Photolysis of 0.02 M solution of compound **209** in a mixture of 30% D₂O and 70% CD₃CN with a 120 W sunlamp formed benzoic acid in 58% yield with 100% conversion according to ¹H NMR spectroscopy. The other photo product **212** was formed in 36% yield, which was isolated by chromatography and identified as 1:1 mixture of diastereomers by ¹H and ¹³C NMR spectroscopy.

The o-QM intermediate **211** has been photolysed with various alkenes. Irradiation of compound **210** (10^{-2} M) with compound **213** (0.1 M) in 30% aqueous CH₃CN produced the cycloadduct **214**. The cycloadduct **214** was produced in 79-87% yields at 100% conversions of the reactants (Scheme 45). Other alkene trapping reagents such as dihydropyran, ethyl vinyl ether, methyl trimethylsilyl dimethylketene acetal, and diethyl fumarate, was inadequately reactive in trapping the o-QM, which was deactivated by multiple electron-releasing substituents.

Scheme 45 Photolysis of o-QM 211 with alkene.

The immediate precursor in the photocyclization to benzoxazoline 210 is zwitterionic in character in ground state as with structure 216. From photochemistry of α -keto amides it has been shown that intermediates analogous to 216 can cyclize as well as expel phenolate or carboxylate leaving groups (Scheme 46). Uncertainty is the mechanism for elimination of carboxylate leaving groups to form o-QM 211, since the elimination is possible either from 216 as well as 210. No evidence could be obtained to support the direct elimination of carboxylate leaving groups from 216 using nanosecond laser flash photolysis methods.

Scheme 46 Mechanistic analysis for the generation of *o*-QM **211**.

Barrero and co-workers⁴⁶ reported the synthesis of bioactive pupehedione derivatives involving a hetero Diels–Alder cycloaddition of *o*-QM, which was generated by the fluoride-induced desilylation of silyl derivatives of *o*-hydroxybenzyl iodides. In this protocol two variations were investigated. At first the compound **218** was reacted with dienophiles **219** and **221** where it has been found that the reaction proceeded via the formation of **220** and **222**. It has also been revealed that *o*-QMs undergo [4+2] cycloadditions with ethers **219** and **221** acting as dienophiles during the course of the reaction (Scheme 47).

Scheme 47 [4+2] Cycloadditions of *o*-QM with ethers.

In the second attempt, variations were introduced to explore the scope of this procedure by using different oxadienic moiety. Thus, *ortho*-silyloxybenzyl iodide, **226** was obtained from commercial 2-hydroxybenzaldehyde **224**. When this compound was treated with ethyl vinyl ether, the reaction proceeded in a similar fashion to give chroman **227** exclusively (Scheme 48).

Scheme 48 Synthesis of chroman by the reaction of *o*-QM and ethyl vinyl ether.

Further unstable 3-methylenepyridin-4-one intermediate 233 has been developed by Baldwin and co-workers⁴⁷ by heating of the di-Boc-pyridin-4-one derivative 232. The intermediate 233 undergoes rearomatization followed by Diels-Alder cycloaddition with activated alkenes to give substituted pyrano[3,2-c]pyridines 234. The precursor 231 for the synthesis of 234 was obtained from commercially available 4-methoxypyridine 228. Formylation of 228 gives 229, which upon reduction furnished compound 230 (Scheme 49).

Scheme 49 Diels–Alder cycloaddition of *o*-QM with activated alkenes.

Unusual hetero Diels-Alder reactions between various electron-rich oxazoles and o-QMs have been achieved by Lindsey and Pettus.⁴⁸ In this case o-QM intermediate has been generated by deprotonation of either of the ortho-OBoc hydroxymethyl aromatic compounds (240 or 241, 0.1 M in Et₂O or THF) with tert-butyl magnesium bromide, which led to the migration of -OBoc and subsequent β -elimination (Scheme 50). Even though these o-QM species are extremely reactive, 4-methyl-oxazole 235 proved futile as a nucleophile. Instead, o-QM intermediate undergoes self-destruction via known manifolds such as Diels-Alder

dimerization and trimerization. Generation of *o*-QM **243** from compound **241** in the presence of oxazole **237** led to two adducts in a 62% combined yield. In this case, the 1,4-addition product prevails in the mixture and the [4+2] cycloadduct **245** arises in small amounts (1:10, **245**:246) as shown in Scheme 50.

Scheme 50 Hetero Diels-Alder reactions between various electron-rich oxazoles and o-QMs.

Surprisingly, silylated 4-hydroxymethyl oxazole **247** undergoes reaction with *o*-QM **242** to afford in 66% combined yield with a 3:1 mixture favouring the [4+2] adduct **249**, over the 1,4-conjugate addition product **250**. In addition, the carbonate protected 4-hydroxymethyl oxazole **248** undergoes cycloaddition with *o*-QM **243** affording benzopyran **251** in 46% yield as the only identifiable product (Scheme 51).

Scheme 51 Reaction of substituted oxazoles with *o*-QM.

The 2-amino oxazoles displaying an electron-rich substituent at the 4-position (R = H, Bn) underwent highly asynchronous reactions leading to 1,4-conjugate addition adducts, while the oxazoles displaying an electron-withdrawing substituent at the 4-position (R = -OTBS, – OCO_2Et) underwent a synchronous reaction affording the [4+2]cycloadduct (Scheme 52).

Scheme 52 [4+2] Cycloadduct of oxazole derivatives.

Meanwhile, the o-QM derived from tocopherol has been trapped with DNMM (4-methyl-3,4-dihydro-2H-[1,4]oxazine) **256** via hetero-Diels-Alder reaction with inverse electron demand and confirmed that compound **256** occurs as degradation product of N-methylmorpholine-N-oxide. As dehydro-NMM is an electron-rich olefin and o-QM is electron deficient, so hetero-Diels-Alder reaction of o-QM with ethyl vinyl ether proceeds regioselectively in a particular way that the α -carbon of the vinyl reacts with the oxygen and the vinylic β -carbon with the methylene carbon of the QM. But in the case of DNMM **256**, the situation was less unequivocal, as the compound contains both a vinyl ether and an enamine so two directions of addition with o-QMs **254** and **255** took place leading to tetracycles **257-260**. Computations on the DFT level was performed on the shortened model compounds **255**, **258**, and **260** possessing a methyl group instead of the isoprenoid side chain that predicted **260** to be 12.6 kJ/mol more stable than **258** (Scheme 53).

Scheme 53 Reaction of *o*-QM (derived from tocopherol) with oxazine.

A mechanistically interesting enantioselective [4+2] cycloaddition of o-QM with silvl ketene acetals by catalytic pathway is well described, 50 and was initiated by a chiral cinchona alkaloid-derived ammonium fluoride "precatalyst" complex to afford a variety of alkyl- and aryl-substituted 3,4-dihydrocoumarin products excellent yields with in good enantioselectivity. The electron-rich o-QM 261 was first generated by Jurd in 1977 by the reaction of sesamol and 4-methoxybenzyl alcohol under mild Friedel-Crafts conditions followed by Ag₂O oxidation. Then the newly generated o-QM **261** was treated with butyryl chloride, thermodynamic Hunig's base, and kinetic base/chiral catalyst benzoylquinidine (BQd) 262, respectively. It has been found that the high electron density on o-QM was not compatible with the mild nucleophilic zwitterionic ketene enolate 262 therefore, silyl ketene acetal 263 was used as the ketene enolate precursor to synthesize the compound 267 (Schemes 54 & 55).

Scheme 54 Synthesis of compound 267 utilizing silyl ketene acetal 263 and o-QM 261.

Next, reaction of *o*-QM **261** with **263** in the presence of catalytic tetrabutylammonium fluoride (TBAF) in THF afforded cycloaddition product **267** with 80% ee. Thus, it shows that the precatalyst's effect on the enantioselectivity of the reaction was not dictated by steric bulk rather by electronic effects (Scheme 55).

Scheme 55 Enantioselective synthesis of compound **267**.

A variety of chroman spiroketals have been synthesized by Pettus and co-workers⁵¹ *via* inverse-demand [4+2] cycloaddition of enol ethers with o-QMs. At low temperature *in situ* generation of o-QM allows the kinetic diastereoselective construction of these motifs providing entry to a number of unusual chroman spiroketals. When benzyl alcohol **269** (0.1 M in toluene, -78 °C) and 2-4 equivalents of the 1,3-dihydroisobenzofuran enol ether **268** was treated with t-BuMgCl, an endo-selective cycloaddition takes place as the reaction mixture slowly warms to room temperature (Scheme 56). The reaction cleanly produced the chroman spiroketal **270** in 60% isolated yield (> 8:1 dr).

Scheme 56 Synthesis of chroman spiroketal.

The simple *exo*-enol ethers readily equilibrate with their *endo*-isomers under acidic and sometimes also thermal conditions, the major product was the expected benzopyran **273** and the mono-benzannulated spiroketal **274** (as a mixture of diastereomers) was observed in 25% yield during the reaction of compound **271** with *endo*-enol ether **272**. Apparently, this was the result of cycloaddition of the intermediate *o*-QM with 4,5-dihydro-2,4-dimethylfuran (Scheme 57). ^{12,13}

Scheme 57 Synthesis of benzopyran 273 and mono-benzannulated spiroketal 274.

The intermediacy of *o*-QM has been employed to develop a concise strategy for the synthesis of isoflavans via Diels–Alder reaction between *o*-QM and aryl-substituted enol ethers **278** followed by reductive cleavage of the acetal group.⁵² The method is extended toward the total syntheses of equol, 3'-hydroxyequol, and vestitol. A reductive removal of the methoxy group from acetal **279** was achieved by using BF₃.OEt₂ and triethylsilane to give isoflavan **280** (Scheme 58).

Scheme 58 Synthesis of isoflavan **280** *via* intermediacy of *o*-QM.

It has been envisioned that the Lewis acid should be able to catalyze both the steps namely [4+2] cycloaddition and the reductive removal of the methoxy group of the acetal moiety in a one-pot transformation. In this course, the reaction of phenol **277** and enol ether **278a** with BF₃.OEt₂ in CH₂Cl₂ at 0 °C in presence of triethylsilane in the same pot gave isoflavan **280a** in 34% yield (Scheme 59).

Scheme 59 [4+2] Cycloaddition reaction of substituted phenol with enol ether 278a.

3.1.2. [4+4] Cycloaddition

A one-pot [4+4] cyclization pathway for the generation of eight-membered sultams via *in situ* generated *o*-QM was reported by Hanson and co-workers.⁵³ Here, the pairing of ambiphilic synthons in a complementary fashion was examined whereby *o*-fluorobenzenesulfonamides **281** were merged with *in situ* generated *o*-QM **283** (from compound **282**) in a formal [4+4] cyclization pathway to afford dibenzooxathiazocine-dioxides **284** under microwave (MW) conditions (Scheme 60). This is the first report representing the use of an *o*-QM in a formal hetero [4+4] cyclization.

Scheme 60 Synthesis of 5,2,1-dibenzooxathiazocine-2,2-dioxides by hetero [4+4] cyclization.

3.1.3. [5+2] Cycloaddition (phenoxonium analogous to o-OM)

Green and Pettus⁵⁴ have synthesised α -Cedrene, α -Pipitzol, and *sec*-Cedrenol by the oxidative dearomatization-induced [5+2] cycloaddition. Phenols with various *ortho*-(pent-4-enyl) substitutions were directly prepared from salicylaldehydes by low temperature *o*-QM generation and consumption. The key conversion attributed by the oxidative dearomatization of an *ortho*-(pent-4-enyl)-phenol **285** followed by an intramolecular [5+2] cycloaddition of the respective phenoxonium intermediates **287** across the attached olefin to afford the respective derivatives of skeleton **289** (Scheme 61).

Scheme 61 Intramolecular [5+2] cycloaddition of phenoxonium intermediates.

3.2. Addition of nucleophiles

Bromination of α-tocopherol **290** for the formation of **294** is not possible by simple bromination because an ethano-dimer of α-tocopherol **290** after bromination afforded pyranospirodimer of α-tocopherol **291** in >95% yields. Compound **290** and **291** constitute a largely reversible redox pair. ⁵⁵ Bromine oxidizes one α-tocopherol **290** to an *ortho*-quinone methide (*o*-QM) intermediate **292**, which is immediately attacked by the phenolic OH group of the tocopherol. It is evidently favoured over the 1,4-addition of evolving HBr by a preorganizational effect, as the ethano-bridge keeps the two reacting moieties in close distance. This acid facilitates the opening of **291** to regenerate the *o*-QM **292**. ⁵⁶ Thus, OH group was protected by TMS that continuously traps a small percentage of the *o*-QM intermediate present, which now cannot react back to give compound **291**. In this way, **291** is eventually completely converted into **293** (Scheme 62). Further reaction of **293** with elemental bromine followed by desilylation provided the target bisbromide **294**.

Scheme 62 Bis-bromination of ethano-dimer of α -tocopherol.

The pseudo three-component condensation reaction of 2-naphthol with aldehydes in the presence of various catalysts to form xanthenes has been studied widely. The reaction proceeds through the *in situ* formation of *o*-QMs with 2-naphthol acting as a nucleophile. However, the three-component condensation reactions of 2-naphthol and aldehydes with other nucleophiles are rarely reported in literature. Singh and co-workers⁵⁷ generated substituted xanthenes **297** by one-pot condensation of 2-naphthol with aliphatic/aromatic aldehydes **295** in the presence of P₂O₅ or InCl₃ as catalysts under solvent-free conditions.

They also reported a one-pot synthesis of tetrahydrobenzoxanthenone **299** and diazabenzoanthracenedione **300** in the presence of Indium(III) chloride catalyst by three-component cyclocondensation of aldehydes **295**, β -naphthol **296** and cyclic 1,3-dicarbonyl compounds **298** under solvent-free condition in high yields. This transformation has also been carried out effectively by P_2O_5 as a catalyst. Use of aldehyde **295**, β -naphthol **296**, and 3-oxo-3-phenyldithiopropionic acid methyl ester **301** in equimolar amounts in presence of InCl₃ gave benzo[f]chromene **302** in good yields (Scheme 63).

Scheme 63 Cyclocondensation reaction via *o*-QMs intermediates.

The nucleophilic addition of β -naphthol **296** to aldehyde **295** in the presence of InCl₃ gives naphthoquinone methide intermediate **303**, which then reacts with nucleophile like 1,3-dicarbonyl compound **298** followed by addition of the phenolic hydroxyl moiety to the carbonyl of ketone provides cyclic hemiketal, which on dehydration afforded **299** (Scheme 64).

Scheme 64 Nucleophilic addition of β -naphthol to aldehyde via o-QM intermediacy.

The *in-situ* formation of o-QM intermediate in the presence of solid silica chloride under sonication followed by nucleophilic attack of amide leading to the formation of amidoalkyl-2-naphthol has been reported.⁵⁸ Due to the collapse of the cavitation bubbles near the surface of the catalyst, the oxygen of the carbonyl group might have easily replaced the chlorine atom present on the catalyst silica chloride to give the activated aldehyde, which could be attacked by β -naphthol to give o-QM 307. Thus, the *in situ* generated o-QM 307 upon reaction with amide 308 furnished the envisioned amidoalkyl naphthol 309 (Scheme 65). This reaction was previously reported under the influence of P_2O_5 , which acted as catalyst as well as water scavenger also.⁵⁹

Scheme 65 Synthesis of amidoalkyl naphthols.

Nagawade and Shinde⁶⁰ reported the rapid synthesis of biologically significant amidoalkyl naphthols using a catalytic amount of iodine. Chlorinated solvents (dichloroethane > dichloromethane > chloroform) gave the highest yields. The yields of the end products depend on the polarity of the solvent used and after a through screening 1,2-dichloroethane was found to be the best solvent for the protocol. o-QM has also been generated by the treatment of β -naphthol, aryl aldehyde and ureas or amides in presence of catalytic amount of p-TSA at 125 °C.⁶¹ In addition to that hafnium(IV) bis(perfluorooctanesulfonyl)imide complex⁶² (Hf(NPf₂)4), NaHSO₄.H₂O⁶³ and H₃PW₁₂O₄₀ have also been shown to be effective catalysts for the same reaction with respect to reaction times and yields in the molten salt media.⁶⁴

2'-Aminobenzothiazolomethyl naphthols or 5-(2'-aminobenzothiazolomethyl)-6-hydroxyquinolines **313** was prepared by the reaction of o-QM **312** and 2-aminobenzothiazole. o-QM was generated *in situ* from the reaction of β -naphthol or 6-hydroxyquinoline and aldehyde **311** (Scheme 66). ⁶⁵

Scheme 66 Synthesis of 2'-aminobenzothiazolomethyl naphthols.

1,2-dihydronaphtho[2,1-*b*] furans and 2,3-dihydrobenzofurans **316** substituted at C-2 by an acyl or aryl group were prepared in a diastereoselective manner. The reaction goes through the formation of the *o*-quinone methide intermediate followed by Michael-type addition of the ylide **315** to the *o*-QM with a consequent intramolecular nucleophilic substitution. The reaction is applicable to a range of Mannich bases and pyridinium salts with a variety of versatile functional groups (Scheme 67).⁶⁶

Scheme 67 Diastereoselective synthesis of naphthofurans and benzofurans.

Salvadoria and co-workers⁶⁷ reported the oxidation of α -tocopherol **317** leading to o-QM species **318**, which has been reacted with bromide as a nucleophile from hydrogen bromide to afford the benzylic brominated product **319** (Scheme 68).

HO
$$R + Br_2$$

$$R = H_2C$$

$$R + Br_2$$

$$R = H_2C$$

Scheme 68 Bromination of α -tocopherol.

The bromo- α -tocopherols **317** and **320** upon treatment with nucleophilic triphenyl phosphene followed by acylation gives the furotocooherols **325** & **326** (Scheme 69).⁶⁸

Scheme 69 Synthesis of urotocooherols from bromo- α -tocopherols.

Chen and co-workers⁶⁹ reported the stereoselective synthesis of *trans*-dihydrobenzofurans **332** by the reaction of *o*-QM **328** and sulfur ylides **329**. This gave intermediate **330**, which undergoes trans-elimination–cyclization to afford the **332**. *o*-QM intermediates have been generated from 2-tosylalkylphenols **327** under the mild basic conditions (Scheme 70).

Scheme 70 Stereoselective synthesis of *trans*-dihydrobenzofurans.

Chiral biphenols were found to catalyze the enantioselective asymmetric addition of arylor alkenylboronates to *o*-QMs **333**. Thus, substituted 2-styrylphenols **336** were obtained in high yields (up to 95%) with high enantiomeric ratios (up to 98:2) in the presence of 10 mol % of 3,3'-BI₂-BINOL **335** (Scheme 71).⁷⁰

Scheme 71 Enantioselective addition of aryl- or alkenylboronates to *o*-QMs.

Kumar and co-workers^{2,71} have developed an efficient and catalyst-free approach for C–H hydroarylation of *in situ* generated *o*-QM **339** with electron-rich arenes (*tert*-aryl amines **340**) in glycerol medium. A number of 3-substituted 4-hydroxycoumarin **341** and 4-hydroxypyrone derivatives were obtained in good to excellent yields. The reaction takes place in water without any catalyst, and is highly regioselective (Scheme72). They have used coumarins, indole, and 4-hydroxypyrone derivatives to generate corresponding *o*-QMs. It is noteworthy that in all the cases *para* substituted product was obtained exclusively. The

reason of regioselectivity is the steric hindrance of *N*-alkyl group and *ortho* hydrogen of *N*,*N*-dialkylanilines, which provides the *para* selectivity.

Scheme 72 C–H hydroarylation of *in situ* generated *o*-QM.

Antonio and Freccero⁷² have reported the synthesis and the unexpected photochemical reactivity of a new class of naphthalimide derivatives (NIs) **342** bearing quinone methide precursor (QMPs) at the imide moiety by irradiation of the NI core (Scheme 73). Photogeneration of the transient QM **343** through a PET mechanism between the phenolic precursors and the NI core was rationalized. Such reactivity represents the first case of non-direct phototriggering of QMs *via* photoinduced electron transfer (PET) involving a tethered peripheral moiety.

Scheme 73 Phototriggering of QMs *via* photoinduced electron transfer (PET).

Freccero and co-workers²⁶ have reported the photo induced generation of new binol quinone methides 349, which undergo mono- and bisalkylation of free nucleophiles by product distribution analysis and laser flash photolysis in water solution using binol quaternary ammonium derivatives 346 and 352 as photoactivated precursors (Schemes 74 & 75). N and S nucleophiles are powerfully competitive in alkylation with the hydration reaction.

Scheme 74 Addition of nucleophiles on Binol quinone methides.

The DNA cross-linking potency of the water-soluble binol quaternary ammonium salt 346 was investigated as a pH function and compared to that of other quaternary ammonium salts proficient for photogenerated benzo-QM by gel electrophoresis. DFT calculations in the gas phase and in water bulk on the binol and benzo quaternary ammonium salts 346 and 352 evidence structural and electrostatic features of the binol derivative, which might offer a validation of its promising high photo-cross-linking efficiency.

Scheme 75 Bis-alkylation of photoinduced *o*-QM **354** and **355**.

o-QM **358** (NDI-QM) has been generated by activation of suitable quinone methide precursors (QMPs) **357** under biocompatible conditions like mild thermal digestion (40 0 C), UV-Vis irradiation and mild mono-electronic reduction (Scheme 76). The conjugated NDI-QMPs might have been acting as hybrid ligand-alkylating structures selectively targeting G-4 folding oligonucleotides.

Scheme 76 Addition of nucleophile on NDI-QM 358.

3.3. Alkylation on o-QM: DNA alkylation

The formation of QMs depends on the strength of the leaving group attached to the benzylic position of *o*-QM **360** in the presence of biological nucleophiles (dN).^{74,75} The formation and reaction of QMs are also highly responsive to the presence of electron-withdrawing and electron-donating groups. The observed trends of reactivity are consistent with the electron-deficient nature of the QM intermediate. Electron-donating groups greatly facilitate initial QM generation and its subsequent regeneration from adducts formed by the nucleophiles of deoxynucleosides (dN) that also function as good leaving groups. Conversely, electron-withdrawing groups greatly suppress initial formation of QM and its regeneration from the reversible deoxynucleoside adducts **362** (Scheme 77). The stability of *o*-QM will be influenced mainly by the electronic perturbation over it and the kinetics of the product formation also alters by reaction with deoxynucleosides.⁷⁶ Electron-rich QMs react much more slowly but selectively with nucleophiles than do the electron-poor QMs. These fundamental characteristics provide the basis for explaining the product profiles observed after DNA has been exposed to various QMs.⁷⁷ These characteristics are used to fine-tune the stability and reactivity of QM conjugates for target-promoted and gene-specific alkylation.⁷⁸

Scheme 77 Generation of deoxynucleoside and *o*-QM adducts.

Rokita and co-workers⁷⁹ reported the formation of the reversible quenching adducts **366** under physiological conditions between *o*-QM **364** and deoxycytidine **363** by selective oxidation using bis[(trifluoroacetoxy)iodo]benzene (BTI), which induces a surprising rearrangement driven by over oxidation of a derivative lacking an alkyl substituent at the 4-position of the QM (Scheme 78).

Scheme 78 Reaction of *o*-QM with deoxycytidine **363**.

The rates of alkylation of 2'-deoxyguanosine **369** and guanosine **370** with o-quinone α -(p-anisyl)methide **368** (Scheme 79), were studied by flash photolysis in a series of aqueous sodium hydroxide solutions and bicarbonate ion, t-butylhydrogenphosphonate ion, as well as in biphosphate ion buffers. Rate constants showed that the reaction of o-quinone α -(p-anisyl)-methide **368** with guanosine and deoxyguanosine to be quite fast processes. Even it was considerably faster than the biologically wasteful reaction of the quinone methide with water which is the ubiquitous medium in biological systems and hence makes the quinone methide a potent guanosine and deoxyguanosine alkylator.

Scheme 79 Alkylation of 2'-deoxyguanosine and guanosine on *o*-quinone α -(*p*-anisyl)methide.

The selenide derivatives **371** can generate covalent interstrand cross-linking with its complementary strand through the formation of *o*-QM intermediate **373** induced by periodate oxidation.⁸¹ It has been evidenced by hydroxyl radical foot printing experiment that the quinone appendage specifically alkylated the cytosine base (nuc) extending the duplex formed between the conjugate and the target strand (Scheme 80).

Scheme 80 Alkylation of *o*-QM by using cytosine base (nuc).

It is well known fact that cation on porphyrin can increase both selective binding affinity to nucleic acid or protein and solubility in aqueous conditions. Hence, with photoinducing, *o*-QM intermediate and singlet oxygen could be formed and they may play an important role for both alkylation and oxidative damage to nucleic acids or proteins. In all probability, porphyrin 377 was photochemically activated and induced to form both singlet oxygen and *o*-QM intermediate 378 which led to both oxidative damage and alkylation of the organism in the bacterial system (Scheme 81).⁸²

Scheme 81 Binding of nucleic acid or protein on o-QM having porphyrin unit.

Song and co-workers⁸³ reported three water-soluble DNA cross-linking phenol quaternary ammonium derivatives, which could inhibit the transcription *in vitro* by photo-activation. The cross-linking agent **379** could significantly be a reason for the late apoptosis of tumour cells. The twisted form and two quaternary ammonium groups could anchor DNA strands and facilitate binding of DNA strands that cross-linking between strands then proceeded through a bis(quinone methide) intermediate **380**. The *in situ* formed *o*-QM also has been trapped by ^t-BuNH₂ (Scheme 82).

Scheme 82 Trapping of bis(quinone methide) intermediate by ^t-BuNH₂.

Quinolinyl QM **387** was designed through the fluoride-induced QM generation by desilylation and elimination of acetate group from compound **386** (Scheme 83). The incorporation of the quinolinyl substitute was designed to enhance the potential interactions with DNA through possible partial intercalation and charges of the quinoline moiety and also for the investigation of the steric effect on the reactivity of QMs. Authors have reported that the N1-dG adduct **388** was selectively formed with quinolinyl QM as a favored dG adduct in 30% aqueous DMF and 10 mM phosphate buffer (pH 7.0). 84

Scheme 83 Synthesis of N1-dG adduct using quinolinyl QM and deoxyguanosine.

A novel G-quadruplex (G-4), ligand/alkylating hybrid structures **391** tethering the naphthalene diimide moieties to quaternary ammonium salts of Mannich bases, as quinonemethide precursors **389** was activated by mild thermal digestion (40 °C). The bissubstituted naphthalene diimides were capably synthesized and their reactivity as activatable bis-alkylating agents was explored in the presence of thiols and amines in aqueous buffered solutions (Scheme 84). During alkylation it has been found that the reversible process is a precondition to DNA alkylation, which in turn strengthen the G-quadruplex structural reorganization.

389
$$Y = NR_2$$
 390 391 $Y = NR_3$ R''

Scheme 84 DNA alkylation *o*-QM precursors to generate G-quadruplex.

Freccero and co-workers⁸⁶ also reported a photo activation of new BINOL-amino acid and -amino ester conjugates acquiesced alkylating and DNA cross-linking agents with high photoefficiency and superior cytotoxicity. While using nucleosides as nucleophiles, only the conjugate adducts with deoxycytidine at the N3 (dC-N3), deoxyadenosine at both N1 and NH₂ (dA-N1 and dA-NH₂), and deoxyguanosine at NH₂ (dG-NH₂) are stable in water

solution. Conversely, the alkylation adducts at N7 of both dG and dA is thermodynamically unstable, since the reversal of the alkylation becomes a competitive process.

In due course the alkylation reaction of 9-methyladenine and 9-methylguanine (as prototype substrates of deoxyadenosine and -guanosine) with the parent o-QM have been studied by using DFT at the B3LYP/6-311+G(d,p) level. The guanine oxygen atom (O^6) (ΔGq gas) 5.6 kcal mol⁻¹) followed by the adenine N1 (ΔGq gas) 10.3 kcal mol⁻¹) are more nucleophilic while other centres have comparatively lower nucleophilicity. But water as a solvent diminishes the nucleophilicity of both 9-methyladenine N1 (DGq solv) 14.5 kcal mol⁻¹) and 9-methylguanine O6 (DGq solv) 17.0 kcal mol⁻¹) as a consequence it reverts the purine base nucleophilicity. Regarding product stability, calculations predict that only two adducts of o-QM with 9-methyladenine, at NH₂ and N1 positions, are in lower energy than reactants, in the gas and water phases. However, adduct at N1 can easily be dissociated in water. In particular, the oxygen alkylation adduct becomes slightly unstable in water (ϕG_{solv}) +1.4 kcal mol⁻¹), and the N7 alkylation product remains only moderately more stable than free reactants (ϕG_{solv}) -2.8 kcal mol⁻¹).

Pettus *et al.*⁸⁸ have reported the total synthesis of all of the known chiral Cleroindicins from the known substituted benzaldehyde **392**, which has been treated with [(2-(trimethyl-silyl)ethoxy)methyl]lithium **393** resulted in Boc migration, and afforded an intermediate phenoxide. This phenoxide generated the *o*-QM by *in-situ* elimination, which then underwent succeeding *in situ* 1,4-reduction with sodium borohydride to produce the phenol **394** in 68% yield. Mitsunobu coupling of compound **394** with amide **395** afforded the inverted amide **396** in an 85% yield and >99% ee (Scheme 85).

Scheme 85 Synthesis of chiral Cleroindicins from the substituted benzaldehyde 392.

3.4. Coupling reactions

The coupling of o-QM have been done by redox reactions of vitamin E in 1,2-dichloroethane with oxidants in water/1,2-dichloroethane interface.⁸⁹ Here, the oxidation reaction of α -tocopherol (TOH) with MnO₄ in DCE has been explained by the series of reactions (Eqs. 1-3). Further the formation of α -TD in hexane and o-QM of α -TOH produced during the oxidation of α -TOH dimerizes easily to α -TD as shown in scheme 86.

$$3\alpha - TOH_{DCE} + 2MnO_{4 DCE} + 4H_{2}O_{DCE} \rightarrow 2MnO_{2 DCE} + 3\alpha - TO_{DCE} + 5OH_{DCE}$$
(1)

$$\alpha - TO^{+}_{DCE} + OH_{DCE} \rightarrow \alpha - QM_{DCE} + H_{2}O_{DCE}$$
(2)

$$2\alpha - QM_{DCE} \rightarrow \alpha - TD_{DCE}$$
(3)

Scheme 86 formation and dimerization of α -TD.

3.5. Complex formation with metals

Korthals and Wulff⁹⁰ reported that the reactions of Chromium carbene complexes of type **398** with enynyl propargyl ethers **397** in the presence of base will generate hexahydrodibenzopyrans **399** after an oxidative workup. The involvement of a [4+2] cycloaddition of a Cr complexed *o*-QM intermediate was inferred from the observed high asymmetric induction (Scheme 87).

$$(OC)_5Cr$$

$$398$$

$$(i-Pr)_2EtN, toluene$$

$$FeCl_3/DMF$$

$$399 (3R, 4R) Enantiomer$$

$$R^1= TIPS$$

$$= TBS$$

$$= TBS$$

$$= Tr$$

$$R^2 = Bn; R^3 = Ph$$

$$R^2 = Bn; R^3 = C_8H_5O$$

Scheme 87 [4+2] Cycloaddition of a Cr complexed *o*-QM with enynyl propargyl ethers.

Compound **401** was prepared through the Diels-Alder reaction of the *o*-QM intermediate **406**. Methyl group at the C-9 position of *o*-QM **406** puts into the effect for the formation of a single diastereomer from the Diels-Alder reaction possessing a chair transition state in an *exo*-cycloaddition where the CH₃ is in an equatorial position. Such a chair transition state is possible in **406**, where the cycloaddition occurs *anti* to the chromium. But in case of diastereomer **407**, the Cr is on the opposite face of the *o*-QM where the chair transition state for the *exo*-cycloaddition *anti* to the metal has an axial methyl. The (3*S*,5*S*)- enyne **403** is the incompatible case and gives a mixture of diastereomers the two most predominate of which are formed *via* an *exo* cycloaddition with a chair transition state with an axial methyl **407** and an *endo*-cycloaddition with a chair transition state with an equatorial methyl **408** (Scheme 88).

Scheme 88 Diastereoselactive synthesis of hexahydrodibenzopyrans *via* a chair transition state of a Cr complexed *o*-QM.

Wulff and co-workers⁹¹ also disclosed the synthesis of allocolchicinoids that involves the benzannulation reaction of Fischer chromium carbene complexes and alkynes with high to moderate diastereoselectivity. One of the diastereomer **409** was recemized but not the other **413**. The intermediacy of an *o*-quinone methide chromium tricarbonyl complexes **411** & **412** and the intramolecular hydrogen bonding assist the reversible dissociation of methanol in compound (aS, 7S) **409** to give (aR, 7R) **410**. Due to the intramolecular H-bonding it becomes possible only in diastereomer **409**, while the opposite diastereomer **413** would not participate and thus retains its enantiomeric purity (Scheme 89).

Scheme 89 Synthesis of allocolchicinoids involving Fischer chromium carbene complex and alkynes.

QMs could be stabilized by the steric bulk or by co-ordination to electron-rich transition metal complexes. The mechanism of an enantioselective palladium-catalyzed alkene difunctionalization reaction give the evidence of rapid ligand exchange between palladium and copper as well as a correlation between ligand electronic nature and enantioselectivity. The substrate having an alkene with an adjacent *ortho*-phenol and a linked nucleophile **415** was used. *ortho*-Phenol not only coordinate to the Pd(II) intermediate **416**, preventing hydride elimination, but it also allowed for the *in situ* formation of an electrophilic quinone methide **418**, necessary for the second functionalization (Scheme 90). So this approach contrasts with others as oxidation of the substrate, not the Pd centre promotes the addition of the second nucleophile.

Scheme 90 Enantioselective palladium-catalyzed difunctionalization of alkene.

Pd(II) catalyze the aerobic reductive coupling reactions of organometallic reagents and styrenes through o-QM intermediate. ⁹³ Compound **A** initially oxidizes the alcoholic solvent to generate the Pd-hydride intermediate **B**, which coordinates with alkene **419** to give cationic complex **C**. The alkene inserts into the Pd-hydride yielding **D** and **D**'. As the consequence of formation of the benzylic addition product only, it has been emphasized that intermediate **D**' proceeds to product via formation of o-QM **E** with concomitant reduction of Pd(II) to Pd(0) and generation of **420**. Subsequently ethanol reacts with the o-QM **E** to liberate the ether product **420** and the Pd(0) **F**, which has been reoxidized by CuCl₂ or O₂ (Scheme 91).

Scheme 91 Pd (II) catalyzed reductive coupling of organometallic reagents and styrenes *via* o-QM intermediate.

Phenols when bound to pentaammine osmium(II) **424**, readily react with aldehydes **425** to generate the complexes of *o*-QMs **427**. While these complexes are remarkably stable, when exposed to air or dissolved in water. Treatment with CAN oxidizes the metal and releases the intact QM ligand, which can be trapped by a suitable dienophile **428** to give the compound **429** (Scheme 92).⁹⁴

Scheme 92Trapping of Os-complexed *o*-QM by different alkenes.

Whenever the phenol complex was treated with acetaldehyde, it gave the *o*-QM **430** with small amount of acetaldehyde complex [Os(NH₃)₅(acetaldehyde)]²⁺. Reaction of complex **430** with an excess of 3,4-dihydropyran and 1.5 equiv of CAN in acetonitrile provided tetrahydropyranochromene **431** (Scheme 93).

Scheme 93 Reaction of Os-complexed *o*-QM with 3,4-dihydropyran.

The intramolecular Diels-Alder reaction was accomplished by using the o-QM **432**, which was formed by treatment of the phenol complex **424** with (R)-citronellal and pyridine. Compound **432** after treatment with CAN leads to the demetallation and cycloaddition to form the benzo[c]chromene **433** in 52% yield as a single diastereomer (Scheme 94).

Scheme 94 Diastereoselective synthesis of benzo[*c*]chromene.

Cycloaddition reactions of activated olefins and styrenes 435 with o-quinine methides 436 in the presence of PtCl₄ and AuCl₃ gave the wide range of 2-alkyl or 2-aryl chromans 437. Good diastereoselectivity (up to >99:1) of the chromans were obtained (Scheme 95). 95

Scheme 95 Cycloaddition reactions of activated olefins with *o*-QM.

The reactivity⁹⁶ of an *o*-QM can be understood from M^{III} resonance form in which the carbon can be viewed as bearing a negative charge (Scheme 96). It has been emphasized that the stabilisation of *o*-QM could be correlated to the degree of deviation of the carbonyl/alkene segment of the *o*-QM from planarity with the coordinated diene segment, which is a consequence of metal back-bonding into the aromatic system.

Scheme 96 Resonating forms of metal *o*-QM complex.

Metal stabilised o-QM showed a complete reverse effect, the methylene carbon becomes nucleophilic upon transition metal coordination. For example, in Cp*Ir(o-QM) 442 the exocyclic carbon undergoes nucleophilic substitution reactions, e.g. with iodine or methyl propiolate, and [3+2] dipolar cycloadditions as in the case of the electron-poor dienophile N-methylmaleimide 443 (Scheme 97). The partial charge and "hinge" angle follow the order Ru < Co < Rh < Ir. The Ru and Co complexes predicted high reactivity. Mono substituted nucleophilic metal complexes were isolated, which could establish the importance to the application of this chemistry for the stereoselective synthesis.

Scheme 97 Cycloaddition reaction of Cp*Ir(o-QM) with N-methylmaleimide.

Knoevenagel reaction of 2-hydroxy-1,4-naphthoquinone, 4-hydroxycoumarin, and 4-hydroxy-6-methylcoumarin with paraformaldehyde generated respective quinone methides **447**, **448** and **449**. They applied these *in situ* generated intermediate in hetero-Diels–Alder reaction with β -vinyl-*meso* tetraphenylporphyrinatozinc(II) **446** leading to macrocycles containing 5,10-dioxobenzo[g]chromene, 5,6-dioxobenzo[h]-chromene **450**, pyrano[3,2-c]coumarin **451**, and benzopyran **452** motifs at the β -position (Scheme 98).

Scheme 98 Hetero-Diels–Alder reaction of *o*-OMs with β -vinyl-*meso* tetraphenylporphyrinatozinc(II).

3.6. Rearrangement reactions

The chemical transformation of fluorogenic BCC triggered by thiols proceeds through a tandem benzoquinone reduction and QM type rearrangement reactions, which are spontaneous and irreversible at physiological temperature in aqueous media. ⁹⁸ Generally, the intracellular reduction of the quinone moiety **453** produces the corresponding hydroquinone **454** that instinctively releases the deactivated drug and gives compound **455** via a QM-type rearrangement (Scheme 99).

Scheme 99 Generation of deactivated drug *via* QM-type rearrangement.

QM-type rearrangement reaction sequence had been employed as part of the fluorescence releasing mechanism for latent fluorophores or the drugs release mechanism of the prodrugs as shown in scheme 100.

Scheme 100 Drugs release mechanism of the prodrugs.

Silva and Bozzelli ⁹⁹ have reported that *o*-QM, a 6-methylene-2,4-cyclohexadiene-1-one is an important intermediate in lignin and alkyl benzene combustion, and so the thermal decomposition of *o*-QM is pertinent to the combustion of transportation fuels (containing toluene) and of biomass and low-rank coals (containing lignin). It has been demonstrated a potential pathway leading to *o*-QM **463** formation by the reaction of the benzyl radical **464** with O₂, which would also be important in toluene combustion (Scheme 101).

Scheme 101Combustion of toluene.

The formation of tropone *via* a hydroxyphenylcarbene intermediate **467** from *o*-QM undergoes an intramolecular hydrogen shift to produce 2-hydroxphenylycarbene **467** in a concerted process (Scheme 102).

Scheme 102 Synthesis of tropone from *o*-QM.

Eventually it has been considered the mechanism for fulvene (+ CO) formation. The ring opened product **470** cyclises to the bicyclic intermediate bicyclo[3.2.0]hepta-3,5-dien-1-one **471**, which rearranges to yield spiro[2.4]hepta-4,6-dien-1-one **(472)**. This intermediate subsequently dissociates to fulvene **473** and CO (Scheme 103).

$$CH_2$$
 CH_2 CH_2

Scheme 103 Synthesis of fulvene.

It has been assumed¹⁰⁰ that the structure obtained after proton transfer is a resonance hybrid of two possible forms, zwitterionic (*ortho*-quinoid, **474B**) and keto enes (*ortho*-quinoid, **474C**) as shown in Scheme 104. The wave function of the real structure is a linear combination of the wave functions of two resonance forms. The results demonstrate that not only resonance enhances the strength of hydrogen bonds, but also the formation of intramolecular hydrogen bonds leads to more efficient resonance. The value of this stabilization can be estimated as a difference in energy between the proton transfer state and the open one, which is about 5 kcal/mol in *N*-methyl-2- hydroxybenzylideneamine.

Scheme 104 Resonating structures of zwitterions (ortho-quinoid, and keto enes).

Reaction of guaiazulene¹⁰¹ **475** with 2-methoxybenzaldehyde **476** in methanol in the presence of hexafluorophosphoric acid at 25°C gives (3-guaiazulenyl)(2-

methoxyphenyl)methylium hexafluorophosphate 477 in 93% yield. Similarly, reaction of 475 with 3-methoxybenzaldehyde under the same reaction conditions affords (3-guaiazulenyl)(3-methoxyphenyl)methylium hexafluorophosphate 478 (91% yield) or with 4-methoxybenzaldehyde gives (3-guaiazulenyl)(4-methoxyphenyl)methylium hexafluorophosphate 479 (97% yield). The crystal structures of these monocarbenium-ion compounds, possessing interesting resonance forms are stabilized by the 3-guaiazulenyl and anisyl (2-, 3-, or 4-methoxyphenyl) groups (Scheme 105).

Scheme 105 Reaction of guaiazulene with methoxybenzaldehyde and resonating structure of products **477-479**.

The regiochemical product of Pictet-Spengler cyclization reactions focussed toward the preparation of the pentacyclic core of the ecteinascidin class of antitumor antibiotics has been explored on two different phenolic substrates. Reduction of the β-lactam unit in compound 480 go through the coordinated lithium complex 481, which also prevent the over-reduction of the corresponding aldehyde group. Following aqueous quench with NH₄Cl, condensation of the secondary amine group onto the hemi-aminal unit 481 led to the key iminium ion intermediate 482. The expected hydrogen bonding between the C-4 benzylic amine and the E-ring phenolic residue directs the Pictet–Spengler cyclization *ortho* to the phenol. Finally, spontaneous elimination of the C-4 benzylamine group occurs through the intermediacy of *o*-QM 483 to afford the C3–C4 unsaturated pentacyclic compound 484 as the major product (Scheme 106).

Scheme 106 Pictet-Spengler cyclization reaction for the synthesis of C3–C4 unsaturated pentacyclic compound **484**.

3.7. Tautomerism

After oxidation quercetin **485** exists in four tautomeric forms **486-489**, one *o*-quinone isomer **486** and three quinone methide isomers **487**, **488** and **489**. Due to the relatively high abundance of the quinone methide isomers, glutathionyl adduct formation occurs at positions C6 and C8 of the quercetin A ring to give 6-GSQ **490** and 8-GSQ **491** (Scheme 107).

Scheme 107 Glutathionyl adduct formation at the quercetin ring.

Depending on the orientation of the indole rings around the single bond, it has been predicted that the pairs of syn-periplanar (sp) and anti-periplanar (ap) rotamers for the o-quinone, quinone methide, and quinonimine tautomers of dimers 493-495, two geometrical isomers (E and Z) are possible for the extended forms. In order to resolve a reasonable entrant among these structures for the observed two-electron oxidation products of the dimers, all tautomeric quinones of dimers 493-495 were geometry-optimized in vacuum at the PBE 0/6-31+G(d,p) level of theory. The pentahydroxybiindolyl 498 was isolated from unprecedented hydroxylated indole derivative by oxidation of dimer 494 (Scheme 108).

Scheme 108 Structures of dimers of indole and hydroxybiindolyl.

Pestacin occurs naturally as a racemic mixture of (S) **499** and (R) **501** enantiomers. This racemization likely occurs post-biosynthetically as enzyme mediated reactions with stereospecificity. A racemization mechanism proceeds through a cationic intermediate **500**, which has the desirable feature of being stabilized by different resonating structures. These intermediates are further stabilized by their ability to tautomerize into the three neutral *o*-QMs **501-503** (Scheme 109).

Scheme 109 Resonating structures of Pestacin.

It has been identified that certain aryl quinones experience a tautomeric isomerization to an intermediate QM because of the low barriers of rotation about central aryl-quinone C-C bond, which bring a significant implications in considering the design of synthetic routes leading to natural aryl quinones such as hibarimicin B. ¹⁰⁶ The rotational barrier of phenol is significantly lowered due to stabilization of the transition state structure leading to the interconversion of **504** and **506** by tautomerization to intermediate QM **505** (Scheme 110).

Scheme 110 Tautomerization in quinone methide.

3.8. Photochemical reactions

The photochemistry of a number of p-biphenyls and terphenyls substituted with hydroxyl and hydroxymethyl groups in aqueous solution has been studied in detail by Wan and coworkers. They reported that the simple hydroxyl group can strongly activate appropriately substituted benzenes or biphenyls towards dehydroxylation due to its strongly electron-donating nature as well as carrier of an acidic proton. The intermediates formed are the corresponding quinone methides (QMs) or biphenyl quinone methides (BQM, **508** & **511**) as shown in Scheme 111.

Scheme 111 Photochemical synthesis of compounds 509 and 512.

There was no pH effect on photosolvolysis quantum yield of the hydroxyterphenyls **513** in the pH 4-12 region, so the most likely mechanism for photosolvolysis is formation of excited state phenolate followed by dehydroxylation to give terphenyl quinone methides (TQM, **514**), which after reaction with methanol gave compound **515** (Scheme 112).^{107a}

Scheme 112 Reaction of methanol with terphenyl quinone methide 514.

Compound **516** with competing photosolvolytic reaction generated compound **517** by the ESIPT process with or without water. ¹⁰⁸ At higher water content, an intermediate **518** (with one or more water molecules) underwent the loss of hydroxide ion from the benzylic position with the assistance of the hydronium ion to generate quinone methide **519**, and finally over

all solvolysis (attack by CH₃OH at the terminal quinone methide position) gave compound **520** (Scheme 113).

Scheme 113 Photosolvolytic reaction of quinone methide 519.

The compound 9-(2'-hydroxyphenyl) anthracene **521** undergoes proficient photoaddition of water and alcohols at the 9- and 10-positions of the anthracene moiety to give isolable triphenylmethanol or triphenylmethyl ether type products. ²¹ The reaction is supposed to ensue via water-mediated formal excited state intramolecular proton transfer (ESIPT) from the phenolic OH to the 10-position of the anthracene ring generating an o-QM intermediate that is observable by nanosecond laser flash photolysis, and is trappable with nucleophiles. Upon irradiation in deuterated solvents led to incorporation of one deuterium atom at the methylene position in the photoaddition product, and partial deuterium exchange of the 10-position of recovered starting material, which is reliable with the proposed formal excited state proton transfer mechanism (Scheme 114). The deuterium exchange and photoaddition have got maximum quantum efficiency in 5 M water (in CH₃CN or CH₃OH), and there was no reaction being observed in the absence of a hydroxylic solvent, thus represents the sensitivity of this type of ESIPT to solvent composition. Once formed, QM 522 is transformed to 523 (hydration product) or returns to 521 via enolization. These two reaction pathways from 522 are really not astonishing apart from the isolation of the stable anthracene hydrates and alcohol adducts (Scheme 114).

Scheme 114 Photoaddition of hydroxylic solvents at 10-position of the anthracene.

In general, caged carbonyl compounds¹⁰⁹ are prepared by their acetalization with (2,5-dimethoxyphenyl)ethylene glycol followed by oxidative demethylation to produce corresponding (1,3-dioxolane-4-yl)-1,4-benzoquinones. (2,5-Dihydroxyphenyl) ethylene glycol acetals of aldehydes and ketones **524a-e** upon irradiation (at 300 nm) resulted in their efficient cleavage and regeneration of the carbonyl compound. This proceeds *via* photo induced transfer of the phenolic proton to the dioxolane oxygen in the 3-position accompanied by the C-O bond cleavage and the formation of an intermediate *o*-QM **525**. *o*-QM is very reactive and could undergo rapid hydration or tautomerization to form corresponding unstable hemiacetals **526** and/or **527**. The hydrolysis of **526** and **527** liberates the carbonyl compound **528** (Scheme 115).

Scheme 115 Regeneration of the carbonyl compound by irradiation of ethylene glycol acetals.

The dibenzo[1,4]dioxins **529** underwent a photochemical aryl-ether bond homolysis to generate reactive 2-spiro-6-cyclohexa-2,4-dienone **531** and subsequent biphenylquinone intermediates. Under steady-state irradiation, the 2,2'-biphenylquinones **532** undergo

excited state hydrogen abstraction from the organic solvent to give the corresponding 2,2-dihydroxybiphenyls **533** (Scheme 116).

Scheme 116 Photochemical aryl-ether bond homolysis of dibenzo[1,4]dioxin.

The flash photolysis of methyl 2-hydroxyphenyldiazoacetate **534** in dilute aqueous perchloric acid solution and acetic acid and biphosphate ion buffers generated a transient species that was identified as *o*-quinone-α-carbomethoxymethide **535** (Scheme 117). This structural assignment is based upon solvent isotope effects, the form of buffer catalysis, UV absorption maxima, and the identity of decay rate constants with those determined for the transient obtained by flash photolysis of other more conventional quinone methide precursors namely the benzyl alcohol methyl 2-hydroxy mandelate **536** and its acetate, 20-acetoxy-2-hydroxyphenylacetate **537**.

Scheme 117 Photolysis of methyl 2-hydroxyphenyl diazoacetate.

New (2-adamantyl)naphthol derivatives **536-540**, quinone methide precursors (QMP, Scheme 118) were synthesized and their photochemical reactivity was investigated by preparative photolyses, fluorescence spectroscopy, and laser flash photolysis (LFP). All derivatives undergo photosolvolysis via QM intermediates. Presence of adamantly moiety results in different reactivity pattern compared to the non-susbtituted structure. They also

investigated the anti-proliferative activity of the QMPs on three human cancer cell lines HCT 116 (colon), MCF-7 (breast), and H 460 (lung). Exposure of cells treated with 5 to 300 nm irradiation led to enhanced anti-proliferative effect on MCF-7 cell line, classifying 5 (or QM5) as a potential lead for further anti-proliferative studies.¹¹²

Scheme 118 Quinone methide precursors of (2-adamantyl)naphthol derivatives.

4. Concluding remarks

In this review, we present the power and diversity of applications of ortho-quinone methides (o-QMs), which have become powerful, fascinating, and highly efficient intermediates in organic synthesis. Recent advances toward the generation, reactivity and application of o-QMs in organic synthesis offer a bright future for the development of novel methodologies for the synthesis of natural and man-made compounds. The cases cited in this review have been selected to highlight the most promising applications of o-QMs in organic synthesis. It can be seen that many well-established different methodologies were successfully applied to o-QMs by worldwide laboratories to more environmentally benign approaches. This area is clearly expanding in several aspects and will reveal spectacular applications in the near future, ranging from organic synthesis, material chemistry, fine chemicals, and pharmaceuticals to the synthesis of natural products. Definitely, much more remains to be done in this field, and in the next few years, we will see many new, exciting findings in o-QMs chemistry. Moreover, it will lead to the serendipitous discovery of many more new reactions-for as much as we may know, chemistry always has new and intriguing surprises in store. Therefore, we sincerely hope that this overview will be of interest for the synthetic community as well as for the general readers. Despite the flurry of recent interest in this species, it is unlikely that we have seen a peak, and it is probable that in future years oQMs will be viewed in the same light as radicals, carbenes, benzynes or similar reactive intermediates.

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