Advances in mercury(II)-salt-mediated cyclization reactions of unsaturated bonds

Sumana Mandal, Raju D. Chaudhari and Goutam Biswas*

Abstract
The synthesis of complex cyclic compounds is extremely challenging for organic chemists. Many transition-metal-salt-mediated cyclizations are reported in literature. Hg(II) salts have been successfully employed in cyclizations to form complex heterocyclic and carbocyclic structures that are impossible to synthesize with other transition metal salts. In this review, we have summarized cyclization reactions that are performed with Hg(II) salts. These salts are also successfully applied in stoichiometric or catalytic amounts to form complex cyclic structures and natural products.

Introduction
The use of transition metal reagents has found considerable applications in organic synthesis [1–4] and has radically changed the realm of chemical science. It also provides a powerful tool for the construction of complex molecular frameworks [5-7]. A plethora of reviews involving transition metals such as Pd(II) [8-10], Ru(II) [11-13], Rh(III) [14-16], Mn(II) [17-19], Au(II/I) [20-22], Ag(I) [23-25] etc. in both cascade and sequential reactions have been published. Electrophilic Hg(II) salts are important reagents in organic synthesis and there is published literature establishing this fact [26-29]. However, the main drawback of Hg(II) salts, as compared to other transition metal salts, is their increased toxicity [30,31]. Hg(II) salts on the other hand, are very cheap in comparison to other transition metal salts (Table 1) and one of the soft Lewis acids of the periodic table [32]. Hg(II) salts have already manifested some unique reactivity and therefore attracted increasing interest from chemists [33]. Many examples involving Hg(II) salts with unsaturated bonds in presence of various nucleophiles giving rise to various types of products are abound in the literature. Utilization of Hg(II) salts in the intramolecular cationic cyclization of olefinic, acetylenic, and allenic substrates having aromatic rings, nucleophiles, and heteroatoms in the neighborhood were well documented. Hg(II) reagents were also often employed in the important cyclization step during total syntheses of many...
natural products [34]. Despite its wide application in ring formation reactions, only a few review articles on Hg(II)-salt-mediated cyclization reactions are available in the literature [35]. This review describes the intramolecular cyclization of unsaturated compounds in the presence of stoichiometric/catalytic amounts of Hg(II) salts. The classification of this review is based on the following topics.

- Cyclization reactions involving the stoichiometric amount of Hg(II) salts.
- Cyclization reactions involving the catalytic amount of Hg(II) salts.
- Total synthesis involving Hg(II)-salt-mediated cyclization reactions.

### Table 1: Comparison of price of reagent grade Hg(II) salts with other transition metal salts.

| Name of salts           | CAS No.     | Price $/g\textsuperscript{a}   |
|-------------------------|-------------|--------------------------------|
| mercury(II) chloride HgCl\textsubscript{2} | 7487-94-7  | 0.67\textsuperscript{b}        |
| mercury(II) acetate Hg(OAc)\textsubscript{2} | 1600-27-7  | 1.35\textsuperscript{b}        |
| mercuric triflate Hg(OTf)\textsubscript{2} | 49540-00-3 | 12.7\textsuperscript{c}        |
| mercury(II) perchlorate Hg(ClO\textsubscript{4})\textsubscript{2} | 304656-34-6 | 2.83\textsuperscript{c}      |
| aurochloride AuCl\textsubscript{3}   | 13453-07-1 | 112\textsuperscript{d}         |
| silver perchlorate AgClO\textsubscript{4} | 7783-93-9  | 8.24\textsuperscript{c}        |
| platinum chloride PtCl\textsubscript{2} | 10025-65-7 | 115\textsuperscript{d}         |

\textsuperscript{a}Prices are from http://www.sigmaaldrich.com (as on 05-08-2021).
\textsuperscript{b}100 g, \textsuperscript{c}25 g, \textsuperscript{d}5 g pack size.

The generalized mechanism for cyclization reactions are, alkenes/alkynes 1 initially react with Hg(II) salts (HgX\textsubscript{2}) leading to the formation of a mercurial carbonium ion 2 followed by the attack of an intramolecular nucleophile giving rise to a cyclized mercury–halide complex compound 3 (Scheme 1).

**Review**

**Cyclization reactions involving stoichiometric amounts of Hg(II) salts**

Cyclization involving alkenes (>C=C<)

In 1900 and 1901, the Sand and Biilmann group had first reported the Hg(II)-mediated cyclization of allyl alcohol using Hg(II) nitrate (Hg(NO\textsubscript{3})\textsubscript{2}) in two separate publications [36,37]. The cyclized product thus formed was further treated with iodine to get trans-2,5-bis(iodomethyl)-p-dioxane (4a). The formation of trans-isomer 4a as the major product and cis-isomer 4b as a minor product was later confirmed by Summerbell et al. by repeating the same experiments (Scheme 2) [38].

Cyclization of biallyl ether 5 in presence of mercuric acetate (Hg(OAc)\textsubscript{2}) in an aqueous medium was reported to synthesize a diastereomeric mixture of 2,6-disubstituted-\textit{p}-dioxane 6. The outcome of the reaction was much generalized with no detailed discussion about the ratio of diastereomeric products [39]. Later, Summerbell and co-workers modified the reaction conditions to synthesize 2,6-disubstituted dioxane derivatives 6 (cis/trans 16:1). Unlike 2,5-disubstituted dioxane derivatives, here the cis-isomer was the major product. A higher ratio of the cis-dioxane 6 was achieved by increasing the reaction time and acidity of the reaction medium, while elevated temperature showed no effect (Scheme 3) [40].
The mercuricyclization was also employed in the field of carbohydrate chemistry for the synthesis of α-α-C-glycopyranosyl derivatives. The reaction between carbohydrate alkene precursor \(7\) and \(\text{Hg(OAc)}_2\) proceeds with high stereoselectivity to give the \(\alpha\)-α-\(C\)-glycopyranosyl derivative (1,5-trans-isomer) \(8\) as a single isomer [41]. Treatment of compound \(8\) with sodium borohydride (\(\text{NaBH}_4\)) under phase transfer conditions (PTC) yields compound \(9\) as the only product. The selectivity of \(\alpha\)-stereochernistry was primarily due to the strong directing effect of the neighboring benzyl ether group with the \(\text{Hg(OAc)}_2\). When cyclic mercuric halide \(8\) was treated with \(\text{NaBH}_4\) and oxygen (\(\text{O}_2\)) in DMF oxidative demercuration takes place to give alcohol \(10\) in quantitative yield (Scheme 4).

The methodology utilized in Scheme 5 had been successfully employed for the preparation of \(\alpha\)-glycopyranosyl derivatives of common and uncommon sugars like α-gluco, α-manno, β-altro, β-idio, α- and β-gulo, β-talo, α-galacto, and α-allo \(\alpha\)-glycopyranosides [42]. Vinylated derivatives of aldopentoses \(11\) were treated with \(\text{Hg(OAc)}_2\) affording the corresponding cyclized pyanosymethylmercuric chloride derivatives \(12\) (Scheme 5). This methodology can be used to synthesize rare \(\alpha\)-glycosyl carbohydrates from easily available sugars.

In a similar manner, stereoselective cyclization of \(\alpha\)-glycosyl amino acid derivative \(13\) using mercuric trifluoroacetate \(\text{Hg(TFA)}_2\) at room temperature was performed. The reaction proceeds primarily through stereoselective cyclization to give \(\alpha\)-α-\(C\)-glycopyranosyl amino acid derivative \(14\) as the major product [43]. Nevertheless, \(\alpha\)-mannopyranosyl derivatives cannot be achieved in a similar manner as reductive elimination forms during the mercury removal process (Scheme 6).

Mercury(II) salts had been effectively used to synthesize five-membered furanose derivatives with high stereoselectivity. Nicotra et al. developed \(\text{Hg(OAc)}_2\)-mediated cyclization of hydroxy-alkene derivative \(15\) to form α-α-ribose derivative \(16\) at room temperature (Scheme 7). They had confirmed the formation of the α-isomer of α-C-ribofuranosyl \(16\) predominantly over the β-isomer (\(α/β 95:5\)) [44].

Scheme 4: Cyclization of carbohydrate alkene precursor.

Scheme 5: Hg(II)-mediated synthesis of C-glycopyranosyl derivatives.
In contrast, when Reitz and co-workers carried out a ring formation of benzylated C-arabinofuranoside derivative 17 in presence of 1.4 equiv of Hg(OAc)$_2$ and sodium bromide (NaBr) at room temperature, then β-α-arabinose derivative 18 was formed as the major product (Scheme 8) [45]. The high stereoselectivity of β-derivative 18 at the anomeric position was predominantly due to the presence of the benzyl groups at the C-2 and C-3 positions of the starting material.

It had been cited in many publications that the stereoselectivity of products formed due to Hg(II)-salt-mediated cyclization reactions of alkene-alcohol derivatives depends on several factors: the nature of the Hg(II) salts [46], the starting materials [47], and the effect of H$_2$O (protic solvent) in the reaction media [48].

Pulido et al. had developed a conversion of allylsilyl alcohols 19 to diastereomeric mixtures of tetrahydrofuran derivatives 20A and 20B (Scheme 9) [47]. It was reported that more substituted alkyl groups present in allylsil alcohol 20b direct the selective formation of diastereomeric products. They also observed that changes in Hg(II) salts result in different ratios of the cis- and trans-isomer in the cyclized products [47]. The differences in cis and trans ratios were primarily due to the basicity of anions associated with Hg(II) salts [46].

A similar type of reaction methodology was employed for the formation of a bicyclic nucleoside analog. 4'-C-vinylribofuranoside derivative 21 on treatment with Hg(TFA)$_2$ followed by reduction with NaBH$_4$ leads to the formation of bicyclic nucleoside derivative 22 (Scheme 10) [49].
Pyrrrolidine and piperidine derivatives were also synthesized by Hg(II)-salt-induced cyclization. *N*-Isopropyl-1-aminohex-4-ene (23) on treatment with 1 equiv HgCl₂ followed by reduction with NaBH₄ yielded pyrrolidine 24 and piperidine derivative 25 in the ratio of 7:3 [50]. *N*-Methylaniline derivative 26 undergoes cyclization with HgCl₂ and gave cis-mercuro chlorides 27 and 28 as isolated products (Scheme 11) [51].

For the synthesis of diastereomeric pyrrolidine derivatives, a Hg(II) salt had been used. When HgCl₂ was added to secondary methylamine derivatives 29 and 31 followed by reduction with NaBH₄ a mixture of diastereomers 30a,b and 32a,b was obtained, respectively. *trans*-Isomers were formed as major products over *cis*-isomers (Scheme 12) [52].

Five- and six-membered *N*-containing heterocyclic phosphonates were synthesized by intramolecular cyclization of alkenyl α-aminophosphonates in a similar way with the treatment of Hg(OAc)₂ [53]. The cyclized products 34A,B formed from starting material 33 were regiospecific and followed Markovnikoff’s type addition in the reaction [54-57]. It was also reported that the formation of α-phosphorylated pyrroldines mostly takes place in regio- and stereoselective ways depending on the reaction conditions (Scheme 13) [53,58].
Cyclization of 4-cycloocten-1-ol (35) with Hg(OAc)$_2$ resulted in formation of two types of fused bicyclic products, 9-oxabicyclo[3.3.1]nonane (36) and 9-oxabicyclo[4.2.1]nonane (37) (Scheme 14) [59,60]. It was observed that the regioselective synthesis of 9-oxabicyclo[4.2.1]nonane (37) was favored when NaBH$_4$ and sodium acetate NaOAc were present, while in absence of NaOAc, 9-oxabicyclo[3.3.1]nonane (36) was formed as an exclusive product.

In a similar manner, an aminomercuration reaction had been successfully employed in the cyclization of trans-amino alcohol 38 leading to the formation of (1R,2R,6R)-9-benzyl-9-azabicyclo[4.2.1]nonan-2-ol (39). The bicyclic derivative 39 through the number of consequent reactions formed a highly potent (+)/(-)-pyrido[3,4-b]homotropane (40), a bridged nicotinoid (Scheme 15) [61].

Similarly, precursor 41 at room temperature in the presence of Hg(OAc)$_2$ (1 equiv) in CH$_2$Cl$_2$ cyclized to form 2-aza- or 2-oxabicyclic mercuri derivatives. Further, treatment of intermediates with NaBH$_4$ (4 equiv) in presence of 5% aq NaOH solution for 1 hour at room temperature produced bicyclic derivatives 42 (Scheme 16) [62]. The diastereoselectivity of the products formed depends on the heteroatoms involved; azamercuration yields more selectivity than oxymercuration.

Nixon et al. reported a Hg(II)-salt-induced alkenyl hydroperoxide 43 cyclization to synthesize cyclic peroxides 44 as the major product, which on further treatment with NaBH$_4$ gives compound 45 along with the unexpected product 46 (Scheme 17) [63].

Later they had reported the reaction between alkenyl hydroperoxide 47 and aldehydes 48 to form hemiperoxyacetal 49 which subsequently reacted with Hg(OAc)$_2$ and a catalytic amount of perchloric acid (HClO$_4$) forming cyclized product 1,2,4-trioxanemercuri bromide 50. The product thus formed was reduced with NaBH$_4$ to form compound 51 (Scheme 18) [64].

Kurbanov et al. first reported the cyclization of different isoprenoid derivatives to hexahydrochromene derivatives using Hg(II) salts [65]. They had shown that trans-geranylacetone (52) produces trans-2,5,5,9-tetramethylhexahydrochromene (53) while the cis-cyclized product was formed from cis-
isoprenoid derivatives. Later, Hoye et al. used similar reaction conditions to cyclize dienes 54 with 1.1 equiv of Hg(TFA)$_2$ to yield endocyclic enol ether 55 in almost quantitative yield [66]. They had also performed experiments with different isoprenoid derivatives (carboxylic acid, ketones, alcohols, and keto esters) to form different bicyclic products (Scheme 19).

A Hg(II)-salt-promoted cyclization of amidal 56 was performed to synthesize optically active cyclic alanine derivatives 57 as 1:1 diastereomeric mixture [67]. Diastereomers were further separated by column chromatography and afforded enantiomerically pure isomers of $N$-substituted imidazolidine-4-ones 57 (Scheme 20).

In a similar manner, $N$-Cbz-protected amine 58 undergoes cyclization using Hg(TFA)$_2$ (1 equiv) and yielded a dinitrogen-containing mixture of diastereomeric alicyclic derivatives 59. The tetrahydropyrimidin-4(1H)-one-mercuri trifluoroacetate derivative 59 on successive treatment with NaBr and LiBH$_4$ gives a mixture of tetrahydropyrimidin-4(1H)-one derivatives 60 in

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Scheme 17: Hg(II)-salt-induced cyclic peroxide formation.

Scheme 18: Hg(OAc)$_2$-mediated formation of 1,2,4-trioxanes.

Scheme 19: Endocyclic enol ether derivative formation through Hg(II) salts.

Scheme 20: Synthesis of optically active cyclic alanine derivatives.
the diastereomeric ratio of 63:37 after separation by column chromatography [68] (Scheme 21).

Mercury(II) salts had also been successfully utilized in the cyclization of ether derivatives 61 to form stereoselectively trans-4,5-disubstituted oxazolidine derivatives 62 (Scheme 22) [69]. Later it was reported, when homologous allyloxy carbamate derivative 63 was cyclized with Hg(TFA)$_2$ then five- and six-membered rings 64 and 65 were formed with a 1:4 ratio (Scheme 22) [70]. The greater yield of the six-membered product was primarily due to the electron-withdrawing effect of ethereal oxygen which in turn destabilizes the carbocation at the β-carbon and hence the nucleophilic attack at the γ-carbon took place.

Cyclization of amide derivative 66 induced by Hg(OAc)$_2$ followed by reduction with LiBH$_4$ afforded a mixture of compounds 67A/67B [71]. The formation of endo,trans-product as a major product over the exo,cis-isomer was primarily due to a stereoinduction effect (Scheme 23). The small amount of

**Scheme 21:** Hg(II)-salt-mediated formation of tetrahydropyrimidin-4(1H)-one derivatives.

**Scheme 22:** Cyclization of ether derivatives to form stereoselective oxazolidine derivatives.

**Scheme 23:** Cyclization of amide derivatives induced by Hg(OAc)$_2$. 

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exo,cis-isomers 67Ba/67Bc was only detected by HPLC analysis of the crude reaction mixture.

Takacs et al. observed the rapid cyclization of salicylamide-derived amidal auxiliary derivatives 68 and 70 in presence of 1.5 equiv of a 1:1 mixture of Hg(OAc)$_2$/Hg(TFA)$_2$ resulting in a diastereomeric pair of cyclized products 69 and 71, respectively. It was observed that the cis-isomer was predominant in the case of the five-membered ring while the trans-isomer was predominant in the case of the six-membered ring formation (Scheme 24) [72].

Cyclization involving alkynes (-C≡C-)
Hg(II)-salt-mediated cyclizations were also observed in the case of alkynes. Arylalkynes 72 were cyclized via Hg(II)-salt-induced reactions to form benzopyran derivatives 73 [73-75]. Interestingly in the case of 1-aryloxy-4-arylthio-2-butyne derivatives 74, selective ring closure on the oxygen side afforded 6-membered chromene derivatives 75 but no thiochromene derivatives were observed [76]. Compound 75 on refluxing with trifluoroacetic acid isomerized to form exocyclic trans-compound 76. It was reported that 1,4-di(arylthiol)-2-butyne derivatives only afforded usual Hg(II)-salt-mediated hydrated products instead of cyclized thiochromene derivatives. Balasubramanian et al. had performed a mercuric-oxide-mediated cyclization of 1,6-di(aryloxy)-2,4-hexadiyne derivatives 77 to get bichromene derivatives 78 (Scheme 25) [77].

A HgCl$_2$-induced cyclization also takes place for acetylenic silyl enol ether derivative 79 forming carbocyclic compounds...
81 with good yields via intermediate 80 [78]. The cyclization of compounds 79 undergo regioselective addition with the triple bond in exocyclic alkene position leading to the formation of α-mercury ketone 80, which were later functionalized by electrophilic addition (Scheme 26).

Schwartz et al. developed a new route for the synthesis of exocyclic enol ether 83 and endocyclic enol ether 85 involving a Hg(II)-induced cyclization of acetylenic alcohols 82 and 84, respectively [79]. This work revealed that the regioselectivity of the Hg(II) addition to the alkyne may be influenced by diastereomers; in the case of cis-isomer 82 exocyclic enol ether 83 was formed while for trans-isomer 84 the reaction takes place with a much slower rate yielding endocyclic enol ether 85 as the only product (Scheme 27) [79].

trans-Acetylenic alcohol 86 on treatment with HgCl₂ (0.5 equiv) in presence of N-bromosuccinimide (NBS) undergoes cyclization yielding stable bromo alkenes 87 (Scheme 28) [80,81].

Atta et al. reported the specific cyclization of ethynyl phenols 88 in presence of HgCl₂ at ambient temperature yielding benzo-furan derivatives 89. They had reported that the electron-withdrawing electronic effect at the aromatic ring promotes the cyclization reaction, whereas there is no cyclization in the absence of the withdrawing group (Scheme 29) [82].
Scheme 29: Synthesis of benzofuran derivatives in presence of HgCl₂.

Larock et al. had performed the mercuration of 4-hydroxy-2-alkyn-1-ones 90 with HgCl₂ to get furylmercurials 91 via syn-addition of acetylene, which on carbonylation yielded the furan-containing carbonyl compound 92 (Scheme 30) [83]. It was proposed that initially mercuration of acetylene bonds via mercurinium like ions or π-complex takes place. Then the structure was stabilized through hydrogen bonding between alcohol and carbonyl groups (path a) or by hemiketal formation (path b).

Thus, the chloride ion attacks from the frontside, producing syn-addition molecules that, upon dehydration, formed furan rings.

Later they had reported similar mercuration of arylacetylenes to synthesize a broad spectrum of heterocycles, namely benzofurans, benzothiophenes, isocoumarins, chromones, benzopyrans, 1,2-dihydropyranaphthalenes, coumarins, and coumestan including some physiologically active heterocyclic natural products like flavones [84]. In the presence of Hg(OAc)₂ in acetic acid, simple cyclization of ortho-substituted arylacetylenes 93, 95, and 97 yielded benzofurans 94, benzothiophenes 96, and indoles 98, respectively. When the carbonyl group was introduced between an aryl and a methoxy group (99) then six-membered isocoumarin ring 100 was formed, and when a carbonyl group was introduced in between an aryl and an alkyne group (101), chromone derivatives 102 were formed. On the introduction of the methylene moiety in between the aryl and methoxy group, 103 yielded isofuran derivatives 104 due to the change in regioselectivity during the cyclization. Dihydropyranaphthalenes 106 were synthesized by cyclization of the corresponding methoxy-arylalkyne derivative 105 (Scheme 31).

Scheme 30: a) Hg(II)-salt-mediated cyclization of 4-hydroxy-2-alkyn-1-ones to furan derivatives and b) its mechanistic pathway.
In 2005, Ghorai et al. reported the synthesis of a mixture of 6-membered as well as 5-membered heterocyclic compounds through a Hg(II)-salt-promoted unique cyclization–rearrangement reaction [85]. They had performed the HgCl₂-promoted cyclization of O-propargyl glycolaldehyde dithioacetals 107 via their dithioketals and dithioacetals to synthesize six-membered heterocycles 108 and 109 (Scheme 32). Five-membered dihydrofuryl aldehydes were also isolated as minor products in some examples.

Following a similar protocol, Biswas et al. later published the HgCl₂-mediated cyclization reaction of tethered alkynedithioacetals 110 to provide six- and five-membered carbocyclic and heterocyclic derivatives 111 and 112, respectively. They had observed that the formation of five-membered rings (112a–c)
was preferred when substituents were present at the alkyne terminus, whereas six-membered rings (111a,b) formed predominantly in case of unsubstituted alkyne dithioacetals (Scheme 33) [86]. They had reported the plausible mechanism for the formation of a six-membered pyranose ring follows ‘path a,’ while for the formation of five-membered pyrrolidine derivatives ‘path b’ was followed.

**Cyclization involving allenes (>C=C=C<)**

Apart from alkenes/alkynes, there are also examples where cyclization takes place involving allene functionalities. Some of the examples are discussed below.

Balasubramanian et al. reported the cyclization of aryl allenic ethers 113 on treatment with Hg(OTf)_2. Compound 113 reacted with 1.1 equiv of Hg(OTf)_2 at room temperature followed by reduction with alkaline NaBH_4 leading to the formation of benzopyran derivatives 114 and 115. The ratio of the formation of products depends on the temperature and time of the reaction (Scheme 34) [87].

Devan et al. had developed similar types of Hg(TFA)_2-mediated cyclizations of allene 116 at low temperature followed by reduction with alkaline NaBH_4 to form cyclized product 117 in moderate yield [88]. The reaction was believed to proceed through Hg(II) ion-promoted electrophilic cyclization (Scheme 35).

**Cyclization involving catalytic Hg(II) salts**

Cyclization involving alkenes (>C=C<)

Apart from the stoichiometric amount used in cyclization, there is abundant literature where a catalytic amount of Hg(II) salt was employed for cyclization reactions between nucleophiles and unsaturated bonds.

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**Scheme 33:** a) HgCl_2-mediated cyclization reaction of tethered alkyne dithioacetals; and b) proposed mechanism.
The synthesis of fused polycyclic ethers by the treatment of a catalytic amount of Hg(TFA)$_2$ with suitable starting material was demonstrated by Tan et al. [89]. They had reported a Hg(II)-catalyzed intramolecular trans-etherification reaction of 2-hydroxy-1-(γ-methoxyallyl)tetrahydropyran derivatives $118$ and $120$ to the corresponding bicyclic dihydropyran derivatives $119$ and $121$, respectively (Scheme 36).

Later, Namba et al. reported the synthesis of racemic vinylindoline derivatives $123$ from $N$-tosylanilinoallylic alcohol derivative $122$ by using 1–2 mol % of Hg(OTf)$_2$ in CH$_2$Cl$_2$ at room temperature [90]. An asymmetric synthesis of vinylindoline derivatives $125$ was achieved by utilizing chiral ligands like chiral binaphane (Scheme 37) [91]. They had observed that the formation of six- and seven-membered rings required elevated temperatures. Subsequently, the same group studied the cyclization of arenyne $126$ to furnish naphthalene derivative $127$. The plausible mechanism for the formation of compound $127$ proceeded consecutively with π-complex formation, Friedel–Crafts type addition, deprotonation, and finally protonation of alcohol for the elimination of water to get the final product [92].

A Hg(OTf)$_2$-mediated cyclization was utilized for the synthesis of 1,4-dihydroquinoline $129$ possessing a quaternary carbon center from $128$ [93]. The reaction was reported via a seven-membered bicyclic hemiaminal as mentioned in the mechanism. This catalytic rearrangement protocol was successfully applied for the construction of complex carbon frameworks from various tosylanilinoallyl acetals. 4H-Chromene derivatives were also synthesized starting from phenol derivatives (Scheme 38).
Cyclization involving alkynes (–C≡C–)

Marson et al. had developed the synthesis of substituted furans 133a–c promoted by catalytic use of Hg(II) salt through cyclization of secondary and tertiary 1-alkynyl-2,3-epoxy alcohols 131a [94]. This is an example of a Hg(II)-salt-catalyzed rearrangement of 1-alkynyl-2,3-epoxy alcohols to substituted furans. The furan 133a was formed by dehydration of intermediate 132a through the corresponding oxonium cation. When R³ is an alkyl group, direct dehydration was not possible because of blocking. Later they reported a similar transformation of
thiiranes to provide a wide variety of substituted thiophenes. They had synthesized substituted thiophenes $\text{133b}$ starting from thiiranes $\text{131b}$ via cyclization utilizing the catalytic amount of HgO and dil. $\text{H}_2\text{SO}_4$ at room temperature [95]. Initially, the Hg(II)-salt-catalyzed the formation of intermediate $\text{132b}$ which further proceeded by dehydration to yield thiophenes $\text{133b}$ as the final product (Scheme 39).

Starting with 1-alkynyl-2,3-epoxy alcohols $\text{135}$, Marson et al. had reported a Hg(II)-salt-catalyzed rearrangement to produce 2,3-disubstituted-2,3-dihydropyranone derivatives $\text{136}$. The stereochemistry of substituents at 2,3-positions of 2,3-dihydropyranone $\text{136}$ was controlled by cis- and trans-configuration of the epoxide of starting materials (Scheme 40) [96].

Scheme 38: a) Synthesis of 1,4-dihydroquinoline derivatives by Hg(OTf)$_2$ and b) plausible mechanism of formation.

Scheme 39: Synthesis of Hg(II)-salt-catalyzed heteroaromatic derivatives.

Scheme 40: Hg(II)-salt-catalyzed synthesis of dihydropyranone derivatives.
Several unsaturated lactones had been synthesized from alkynoic acids via Hg(II)-salt-catalyzed cyclization reaction. For example, simple 4-pentynoic acid derivatives 137 afforded γ-methylene butyrolactones 139 in good yields via the formation of organomercurial compounds 138 using catalytic mercuric oxide, mercuric acetate, or mercuric trifluoroacetate as shown in Scheme 41 [97].

Alkyne carboxylic acids also undergo oxymercuration reactions to form furan- and pyran-like derivatives. When γ-alkyne carboxylic derivative 140 was refluxed with HgO the cyclization took place to give product 141 in almost quantitative yields [98,99]. Compound 142 under refluxing conditions gave spirocyclic compound 143 as the exclusive product (Scheme 42) [99]. Propargylic triols 144 undergo Hg(OTf)₂-catalyzed cyclization reaction, to produce monounsaturated spiroketal 145.

Finally in the case of 1,2-dihydroxy-3-alkynes 146 comparable reaction conditions leads to the formation of 2,5-disubstituted-furan 147 rather than mono-unsaturated spiroketals [100].

Interestingly when 1,4-dihydroxy-5-alkyne derivatives were subjected to a Hg(OTf)₂-catalyzed cyclization then oxacyclization takes place to form tetrahydropyran derivatives (Scheme 43) [101]. Later it was shown that alkynyl diol 150 when treated with 20 mol % Hg(OTf)₂ followed by Et₃SiH afforded bispyranoyl ketone 151, but when Hg(OTf)₂ was increased (1 equiv) then fused pyran-oxocane derivative 152 was isolated [102].

Six-membered morpholine derivatives were also synthesized by catalytic Hg(II)-salt-induced cyclization. Yamamoto and co-workers published the intermolecular cyclization of alkynyl-
carboxylic acid 153 to produce 6-membered morpholine type ring compound 154 and compound 155 [103]. The stereochemistry of the chiral amino acid was not conserved in the cyclized product hence it leads to the formation of racemic products with moderate yields (Scheme 44).

1,6-Enynes 156 underwent smooth hydroxylative carbocyclization in the presence of catalytic Hg(OTf)₂. As an example, 3-methylene five-membered cyclic derivatives 157 were synthesized by Nishizawa et al. via the Hg(OTf)₂-catalyzed hydroxylative carbocyclization of 1,6-enyne 156 (Scheme 45) [104]. It was observed that from 1,7-enynes, six-membered rings were formed in low to moderate yield while from 1,8-enynes, uncyclized hydrated products were isolated as major products along with different byproducts.

In a Hg(OTf)₂-catalyzed process, 1,6-allenynes 158 were cycloisomerized to generate allenes 159 in moderate to good yield (Scheme 46) [105]. However, depending on the substituents, allene and/or unexpected triene were produced as a main product for disubstituted 1,6-allenynes. It was hypothesized based on experimental evidence that alkynes would first form a π-complex with Hg(OTf)₂, followed by vinylmercuration, demercuration, and eventually isomerizes to allene.
In 2003, a carbocyclization with a catalytic quantity of mercury salt was used to efficiently synthesize dihydronaphthalene derivative 161 from exemplifying benzyl derivative 160 [106]. The reported methodology was an example of the Hg(OTf)$_2$·(TMU)$_3$ complex promoting a moderate and efficient procedure for arylalkyne cyclization to directly afford dihydronaphthalene derivatives (Scheme 47). Later, Friedel–Crafts type reaction of alkynylfuran 162 and 164 were...
reported in presence of the Hg(OTf)$_2$·0.1 Sc(OTf)$_3$ complex (5 mol %) to form six- (163) and seven-membered rings (165) in good yield. For the cyclization at 2-position of the furan the Hg(OTf)$_2$·(TMU)$_3$ complex was used as catalyst [107].

Later it was reported that 1-alkyn-5-ones such as 166 also undergo an effective cyclization reaction to synthesize 2-methylfuran derivatives 167 with high yield in the presence of Hg(OTf)$_2$ as the catalyst under very mild reaction conditions (Scheme 48) [108].

Cyclization of 6-aminohex-1-yne 168 was performed by catalytic amounts of Hg(NO$_3$)$_2$ and generated 2-methyleneepiperidine 169 initially, which further isomerizes to form 2-methyl-1,2-dehydropiperidine (170, Scheme 49) [109].

Similarly, in 2007 Kurisaki et al. had also prepared indole derivatives 172 with excellent yields from 2-ethynylaniline derivatives 171 upon the treatment of catalytic amounts of Hg(OTf)$_2$ at room temperature. It was an example of cycloisomerization of 2-ethynylaniline derivatives utilizing mild reaction conditions (Scheme 50) [110].

Rong et al. had demonstrated the Hg(II)-salt-catalyzed enolate umpolung reaction for the efficient synthesis of various 3-indolinones and 3-coumaranones 174. They had proved that the reaction mechanism proceeds via activation of the alkynyl group with Hg(OTf)$_2$ salt and addition of 2-chloropyridine N-oxide. The resulting activated alkynyl complex was demercurated, followed by the SN$_2'$ reaction thus formed undergoes demercuration to yield 3-coumaranone (Scheme 51) [111].

Recently, a Hg(OTf)$_2$-catalyzed one-pot cyclization of nitroalkyne 175 and alkyne had been reported to synthesize indole derivatives 176. Based on this strategy, the one-pot method to synthesize indole derivatives had been developed [112]. Similarly, benzo[c]isoxazole was also formed in excellent yields with high selectivity using this strategy. In these transformations, two Hg-carbene intermediates were proposed to be involved (Scheme 52).

Mercury-catalyzed reactions were also well known for the formation of various complex scaffolds like tricyclic pyrazinones.
Scheme 52: a) Hg(OTf)$_2$-catalyzed one pot cyclization of nitroalkyne and b) its plausible mechanism.

from the corresponding starting materials. For example, Zhang et al. showed that refluxing anilide 177 in presence of a catalytic amount of Hg(OAc)$_2$ and 90% formic acid gave the tricyclic heterocyclic scaffold 178 [113]. It involved a two-step process with the rearrangement of the primary cyclization products (Scheme 53).

Scheme 53: Synthesis of tricyclic heterocyclic scaffolds.
In 2013, Lin et al. reported a Hg(II) chloride-mediated cyclization reaction of 2-alkynylphenyl alkyl sulfoxide 179 to synthesize benzothiophene derivatives 180 with good yields [114]. In this case, the reaction was believed to proceed via the initial formation of metal carbenoids followed by a sequential C–H insertion and then oxidation (Scheme 54). This methodology was later successfully utilized for the total synthesis of raloxifene and benzo[b]thiophene derivatives [115].

Cyclization involving allenes (>C=C=C<)
Hg(II) triflate salts had also been successfully employed for the aryllallene 181 cyclization by Yamamoto et al [116]. The catalytic pathway was proved to involve the direct H-transfer to the vinylmercury complex from the aromatic ring. It involved Hg(OTf)$_2$-catalyzed cyclization of aryl 1,1-disubstituted allenes with the formation of a quaternary carbon center followed by the formation of a cationic vinylmercury intermediate (Scheme 55).

For the synthesis of stereoselective tetrahydropyran derivatives 184, Hg(II)-catalyzed cyclization proved to be more effective than silver(I)-salt-mediated cyclization. It showed that methyl-substituted allenes undergo efficient cyclization to form polycyclic ethers under Hg(OTf)$_2$-catalyzed conditions at lower temperature (Scheme 56) [117].

Following a similar strategy, mercury chlorate (Hg(ClO$_4$)$_2$) had been employed successfully as a cheap alternative to precious metals salts in the cyclization of α-allenol derivatives 185 to differently substituted 2,5-dihydrofurans 186 in an efficient and selective manner. It was also shown that from enantiopure alleny derivatives, the desired pure cyclized product was generated by utilizing the above reaction conditions (Scheme 57) [118].

**Mercury(II)-salt-mediated cyclization in total synthesis**

Apart from previously described cyclization reactions, there were examples where a Hg(II)-salt-mediated cyclization had
Scheme 56: Stereoselective synthesis of tetrahydropyran derivatives.

Scheme 57: a) Hg(ClO₄)₂-catalyzed cyclization of α-allyl derivatives. b) Simplified mechanism.

Scheme 58: Hg(TFA)₂-promoted cyclization of a γ-hydroxy alkene derivative.

In 1998, a highly enantioselective total synthesis of (+)-furanomycin (190) was achieved through Hg(TFA)₂-promoted cyclization of γ-hydroxyalkene derivative 187 as an intermediate stage to give a mixture of diastereomeric 2,5-disubstituted tetrahydrofurans 188a and 188b (Scheme 58) [119].

Later for the total synthesis of ventiloquinone J (194), a Hg(II)-salt-catalyzed intramolecular cyclization reaction of the ortho-allyl alcohol 192 was involved. The reaction went through the formation of a mixture of diastereoisomers 193a and 193b. After which the inseparable mixture of products underwent further oxidative demethylation and yielded the final products (Scheme 59) [120].

Kraus and co-workers had reported the synthesis of the racemic naphthohydroquinone hongconin (197) starting from the ortho-allyl alcohol derivative 195. The starting material was cyclized using a Hg(II) salts to get an inseparable mixture of products 196a and 196b in the ratio of 1:5 [121]. The synthetic route proceeded by benzylic alcohol ortho-metalation followed by a regioselective mercuri-cyclization reaction (Scheme 60).
Scheme 59: Synthesis Hg(II)-salt-mediated cyclization of allyl alcohol for the construction of ventiloquinone J.

Scheme 60: Hg(OAc)$_2$-mediated cyclization as a key step for the synthesis of hongconin.

During the total syntheses of (±)-fastigilin C and (−)-fastigilin C (201), 2 equiv of Hg(TFA)$_2$ were used to synthesize key intermediate tricyclic furan compounds 199, 200, and 203. Hg(TFA)$_2$ helped in the desired ring-formation reaction of compound 198 to afford two cyclic compounds 199 and 200 in the ratio of 6:1 with an overall 81% yield [122]. But in the case of methoxy-substituted derivative 202 only one furan derivative 203 was formed (Scheme 61).

In 2007, Nishizawa et al. successfully utilized a Hg(OTf)$_2$-induced cyclization in the key step for the synthesis of (±)-thal-lusin (207, Scheme 62). A complex mixture of Hg(OTf)$_2$ and...
$N,N$-dimethylaniline (DMA) (1.2 equiv) was initially used for olefin cyclization to produce a regio- and a stereoisomeric mixture of acetate 205 after reduction and acetylation of the crude product. The Hg(OTf)$_2$-catalyzed isomerization of the double bond in compound 205 yielded thermodynamically favorable isomer 206 as a major product [123].

In 2010, Ravindar et al. developed the total synthesis of the steroidal natural product hippuristanol (211) starting from 11-ketotigogenin 208 (Scheme 63). They had utilized a Hg(OTf)$_2$-catalyzed spiroketalization reaction in their key step to form the desired ketal intermediate 210 [124]. Subsequently, in 2011, the same group reported another synthesis of hippuristanol (211) and its analog from easily available starting materials [125]. In this work, hippuristanol and some analogs were successfully synthesized utilizing a Hg(OTf)$_2$-catalyzed cascade spiroketalization step of the 3-alkyne-1,7-diol motif. The Hg(OTf)$_2$-catalyzed cascade spiroketalization step was proved to be more convenient than Suárez cyclization.

A Hg(TFA)$_2$-mediated cyclization was efficiently utilized for the synthesis of highly strained tricyclo[5.2.1.0$^{1,6}$]decene intermediate 214 containing a cyclobutane ring (Scheme 64). Compound 213 is an important precursor for the asymmetric total synthesis of solanoeclepin A. The formation of $\beta$-hydroxyketone 213 was achieved by Hg(TFA)$_2$-mediated cyclization of compound 212 using TFA/H$_2$O (1.7:1) as the solvent. In presence of other mercury compounds like HgO and Hg(OAc)$_2$ no product or starting material was recovered [126].

Spiro-skeleton structures are found in many natural products and synthesizing stereoselective spiro-skeletons has always been difficult for organic chemists. Morimoto and co-workers were the first to disclose the Hg(OTf)$_2$-catalyzed cycloisomerization of amino ynone to produce the azaspiro skeleton. Later, this methodology was successfully used for the synthesis of several spiro skeleton structures. Natural products such as histrioenicotoxin alkaloids 218 (Scheme 65) [127,128] and lepadoformine [129,130] were being successfully synthesized using...
Conclusion
In conclusion, this review summarizes Hg(II)-salt-mediated cyclization reactions either for direct synthesis of cyclized products or as a part of the total synthesis of important natural products. Different Hg(II) salts were used stoichiometrically or catalytically depending upon the nature of functional groups and reactants. However, the reactivity of different unsaturated bonds involved in the cyclization primarily depends on the nucleophile as well as nature of functional groups attached to unsaturated bonds. When alkenes are linked to activating groups like methoxy or hydroxy, a catalytic quantity of Hg(II) ions is required for cyclization. Nothing can be predicted in case of alkynes; however, the presence of a strong nucleophile promotes the Hg(II)-salt-catalyzed cyclization of allenes in most circumstances. In cyclization reactions, Hg(OTf)$_2$ showed to be the most effective and versatile of all Hg(II) salts. Mercury(II) salts can also be used to cyclize unsaturated bonds in a regio- and diastereoselective manner. Apart from toxicity concerns, Hg(II) salts are cheap, stable, and versatile in terms of reactivity, making them a viable option to similar transition metal catalysts.

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