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REVIEW

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Development of novel transition metal-catalyzed synthetic approaches for the synthesis of a dihydrobenzofuran nucleus: a review

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The synthesis of dihydrobenzofuran scaffolds bears pivotal significance in the field of medicinal chemistry and organic synthesis. These heterocyclic scaffolds hold immense prospects owing to their significant pharmaceutical applications as they are extensively employed as essential precursors for constructing complex organic frameworks. Their versatility and importance make them an interesting subject of study for researchers in the scientific community. While exploring their synthesis, researchers have unveiled various novel and efficient pathways for assembling the dihydrobenzofuran core. In the wake of extensive data being continuously reported each year, we have outlined the recent updates (post 2020) on novel methodological accomplishments employing the efficient catalytic role of several transition metals to forge dihydrobenzofuran functionalities.

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1 Introduction

Dihydrobenzofuran scaffolds have been recognized as key structural motifs as they are an integral part of a vast range of biologically active compounds and have attracted significant consideration in the medicinal field.¹ The dihydrobenzofuran core is composed of two rings, *i.e.*, the aryl and dihydrofuran ring (Fig. 1). This unique structural feature makes them ideal candidates for the development of novel pharmaceutical agents.²

Owing to their high medicinal profile, the synthesis of various naturally occurring products containing a dihydrobenzofuran core has gained significant attention in organic and pharmaceutical chemistry.³ Several bioactive natural products are composed of the



Fig. 1 General structure of dihydrobenzofuran 1 and dihydroisobenzofuran 2.

2,3-dihydrobenzofuran framework, such as (+)-decursivine 3,4 (+)-lithospermic acid 4,⁵ pterocarpan 5,⁶ (+)-conocarpan 6,⁷ bisabosqual A 7,8 and caraphenol A 8,9 which have been explored by several synthetic practitioners. They exhibit many biological activities such as anti-malarial, anti-HIV, hepatoprotective, antiinflammatory and antifungal activities. Similarly, furaquinocins 9 (ref. 10) are also dihydrobenzofurans constituting natural products, which act as antihypertensive agents and inhibit platelet coagulation and aggregation. Furthermore, rubioncolin A 10 and rubioncolin B 12 are used to treat cough, uterine hemorrhage, bladder and kidney stones.11 Cancer is one of the most prevailing and deadly diseases, and researchers are undertaking untiring efforts to discover and develop efficient anti-cancer agents.12-16 (-)-Nocardione 11 (ref. 17) is a Cdc25B tyrosine phosphatase inhibitor that has been observed to display cytotoxic activities (Fig. 2).

Moreover, there are many synthetic dihydrobenzofuran derivatives demonstrating a variety of intriguing medicinal properties, including the imidazolium compound (cytotoxic) **13**,¹⁸ diesters (anti-leishmaniasis) **14**,¹⁹ GPR4 agonist **15**,²⁰ triazole (antitubercular) **16**,²¹ and the drugs prucalopride **17** (used to treat constipation)²² and efaroxan **18** (α_2 -adrenoceptor antagonist)²³ (Fig. 3).

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There have been many processes reported for the synthesis of dihydrobenzofurans in the past decades that involve the use of different catalytic systems, *i.e.*, acid or base catalyzed synthesis,^{24–26} organocatalysis,²⁷ salt catalyzed synthesis²⁸ as well as electrochemical and photochemical synthesis.^{29,30} Transition metal (TM)-catalyzed reactions involving the

Review

synthesis of dihydrobenzofuran skeletons are one of the most reliable and beneficial methods.³¹ In the past years, several synthetic researchers have designed and reported facile methodologies for the efficient synthesis of dihydrobenzofurans. In this regard, Blum³² et al. in 2014 reported the Ru-catalyzed photochemical synthesis of dihydrobenzofuran derivatives 26 through the reaction of phenols 19 and alkenes 20 via oxidative [3 + 2] cycloadditions. Similarly, Fe-catalyzed facile generation of dihydrobenzofurans via the Claisen rearrangement of allyl aryl ethers 21 was reported by the research group of Sakate³³ in 2018. Furthermore, Henry and coworkers³⁴ in 2018 described an Fe and Cu dual catalysis protocol for the synthesis of 2,3-dihydrobenzofurans using substituted phenylethan-2'-ol 22. Ircatalyzed synthesis of 2,3-dihydrobenzofuran was reported by Ohmura and coworkers³⁵ in 2019, which proceeded via the intramolecular cycloaddition of the C-H bond of o-methyl ether 23 to the C-C double bond. In 2020, the research group of Fang³⁶ disclosed the efficient synthesis of chiral dihydrobenzofuran-3-ols via the Cu-catalyzed intramolecular reaction of aryl pinacol boronic esters 24. In the same year, Li³⁷ et al. envisioned the Ni-catalyzed synthesis of chiral 2,3-dihydrobenzofurans using ortho-substituted aryl halides 25 (Fig. 4). In addition, multiple other TM-catalyzed approaches have also been reported in the literature.³⁸⁻⁴⁰

The TM-catalyzed synthesis of dihydrobenzofuran derivatives has gained tremendous interest in the field of organic chemistry as it exhibits high efficiency, *i.e.*, producing desired products in high yields under ambient reaction conditions.⁴¹ Several reviews regarding the synthesis of dihydrobenzofurans have been published to date. In 2019, Laurita *et al.*⁴² published a review on the synthesis of 2,3-dihydrobenzofurans covering the data from 2012 to 2019. Similarly, Dapkekar⁴³ *et al.* in 2022 also reported a review focusing on the methodological developments regarding the synthesis of dihydrobenzofurans and



Fig. 4 Different transition metal-induced synthetic routes for the synthesis of dihydrobenzofurans.

dihydroisobenzofurans. Besides all these published reviews, no particular review has been published concerning solely the TM-catalyzed synthesis of 2,3-dihydrobenzofurans. Herein, the recent data on the synthetic methodologies involving the TM-catalyzed synthesis of dihydrobenzofurans is summarized (reported within 2021–2024).

2 Review of literature

Transition metal-catalyzed transformations represent an advancing domain over the past few years. TM-mediated synthetic pathways have several advantages over conventional protocols.⁴⁴

2.1. Rh-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Rhodium is one of the widely recognized metals in catalysis, which is extensively employed in organic synthesis and industrial processes. Among the range of synthetic methodologies represented for the synthesis of dihydrobenzofuran frameworks, Rh-catalyzed synthesis is a part of the most reliable approaches since they proceed under mild reaction conditions with high yield ranges. TM-catalyzed direct C-H bond functionalization has been proven as a powerful approach for the synthesis of target molecules in an unambiguous and stepeconomic way.45 In particular, the combination of C-H bond functionalization with C-C bond activation to facilitate the rapid synthesis of functionally diverse frameworks from comparatively simple precursors has garnered considerable focus.46 Considering the significance of utilization of TMmediated C-H functionalization towards the synthesis of several heterocycles, Zhang47 et al. in 2021 described the Rh(III)catalyzed construction of cyclic 3-ethylidene-2,3dihydrobenzofuran skeleton 29 with remarkable regioselectivity and chemoselectivity. In their novel approach, N-phenoxyacetamides 27 were reacted with cyclopropylidenemethyl alkenes 28 via subsequent C-H functionalization and [3 + 2] annulation in the presence of [Cp*RhCl₂]₂ and NaOAc (as a base) to synthesize desired products 29 in moderate to high yields (37-80%) (Scheme 1).

In 2022, Song⁴⁸ *et al.* developed a novel protocol leading towards the synthesis of dihydrobenzofuran derivatives employing coupling partners associated *via* asymmetric C–H functionalization. In their methodology, substituted



 $R^2 = 4-OMeC_6H_4, 4-FC_6H_4$

Scheme 1 Synthesis of cyclic 3-ethylidene-2,3-dihydrobenzofuran 29.

phenoxyacetamides **31** were made to react with diazooxindoles **30** in the presence of the rhodium-based catalyst (RhCp*Cl), exploiting cesium carbonate as a base in dioxane solvent. As a result, spirooxindoyl-substituted dihydrobenzofuran derivatives **32** were obtained in 49–76% yields (Scheme 2). The methodology was also employed for the synthesis of various heptacyclic molecules.

Sun⁴⁹ et al. in 2022 envisioned another rhodium-catalyzed difunctionalization of p-substituted olefinic arenes 35 to synthesize 2,3-dihydrobenzofuran derivatives 39, 40 and 41. Olefinic arenes 35 were made to react with unsaturated reactants such as isocyanates 38, dioxazolones 37 and internal alkynes 36 via a tandem reaction. In their synthetic methodology, twofold C-H activation of 35 took place at the ortho and *meta* positions. In the first step, the already present directing group (DG) on 35 resulted in the installation of the alkene coupling partner on the ortho position. This further acted as a relayed directing group for the second activation of C-H, which resulted in the cyclization of olefins at the para position to form the desired products (benzofuran derivative) 39, 40 and 41. Moderate to high yields (40-86%) were obtained when the reaction was carried out in the presence of t-AmOH (as solvent), CsOPiv (as an additive), and AgOAc (as an oxidant) at 120 °C for 12 h (Scheme 3). The mechanism of the reaction was assumed to proceed via the interaction of olefinic arenes 35 with Rh catalyst. Further, alkyne insertion took place to generate intermediate 42, followed by the dissociation of the bonds and C-H bond activation of intermediate 42. In the next step, alkene insertion of 42 resulted in the formation of another intermediate 43, which underwent reductive elimination to furnish the desired products 39 (Scheme 4). The synthetic utility of the above mentioned methodology was also examined.

Wei⁵⁰ *et al.* in 2022 envisioned a rhodium-catalyzed [3 + 2] annulation of 2-alkenylphenols **44** and *N*-phenoxyacetamides **45**

in the presence of base additive $(Zn(OAc)_2)$ along with methanol (as a solvent) to access 2,3-dihydrobenzofurans **48** in excellent yields (90%). The proposed mechanism deduced that intermediate **46** was formed *via* reversible C–H/N–H bond cleavage of **45** assisted by Rh active catalyst, followed by the insertion of **44**, to furnish intermediate **47**. After subsequent oxidative addition, intramolecular hydrogen transfer and finally nucleophilic 1,4addition gave the desired dihydrobenzofuran skeletons **48** (Scheme 5).

In the same year, another rhodium-catalyzed approach for the asymmetric synthesis of dihydrobenzofuran derivatives 51 was put forward by Yu⁵¹ et al. In their methodology, aryl-joined alkenes 49 and substituted dioxazolone 37 were subjected to chiral rhodium-promoted carboamidation, using copper acetate as an additive in the presence of AgSbF₆ and dichloroethane to attain the enantioselective synthesis of dihydrobenzofurans 51 in 44-83% yield with up to 98.5:1.5 enantiomeric ratio. The reaction mechanism was proposed to undergo C-H activation and migratory insertion, followed by oxidative addition to generate intermediate 50. The resulting intermediate 50 was then believed to execute reductive elimination and protonation to achieve enantioenriched 2,3-dihydro-3-benzofuranmethanamides 51 (Scheme 6). To examine the synthetic utility of this novel approach, derivatization of the synthesized compounds was also carried out.

Sinha⁵² *et al.* in 2023 demonstrated a rhodium-promoted synthesis of dihydrobenzofurans **55**. Imidazole (directing group)-substituted allyloxy aryls **52** were subjected to intramolecular regioselective hydroarylation in the presence of $[RhCp*Cl_2]_2$ catalyst, employing cesium acetate as a base in methanol:water (2:1) solvent. As a consequence, imidazoleconstituting dihydrobenzofurans **55** were attained in moderate to excellent yield (52–91%). The reaction mechanism involved coordination with the rhodium catalyst, followed by



Scheme 2 Synthesis of bispirooxindoyl dihydrobenzofurans derivatives 32



DG = 5-MePy, 5-OMePy, 5-FPy, 5-CIPy, 5-CF3, 4-MePy, 4-OMePy, 4-FPy, 4-CIPh, fused PhPy, pyridine R¹ = 4-MePh, 4-OMePh, 4-FPh, 4-CIPh, 4-BrPh, 4-⁷BuPh, 4-CF₃, 4-CO₂EtPh, 4-CNPh, 3-MePh, 3-OMePh, 4-FPh, 3-CIPh, 3-BrPh, 3-CO₂EtPh, Ph, (Me)₂Ph, 2-Naph, 2-furan, 2- thiophene, C₂H₄OTIPS, Me, Et, *n*-Pr, C₂H₄OMe, CH₂OMe R² = 4-MePh, 4-OMePh, 4-FPh, 4-CIPh, 4-BrPh, 4-⁷BuPh, 4-CF₃, 4-CO₂EtPh, 4-CNPh, 3-MePh, 3-OMePh, 4-FPh, 3-CIPh, 3-BrPh, 3-CO₂EtPh, Ph, (Me)₂Ph, 2-Naph, 2-furan, 2- thiophene, R³ = PhCH₃, PhOCH₃, Ph, Ph⁶Bu, Naph R⁴ = PhOCH₃, PhCH₃, PhF, PhCI

Scheme 3 Synthesis of diverse dihydrobenzofuran derivatives 39-41.

the activation of the C–H bond to generate rhodacycle intermediates 53 & 53*. The rhodacycle intermediate 53 was then supposed to undergo migratory insertion to generate the seven-



Scheme 4 Proposed mechanism for the synthesis of dihydrobenzofuran derivatives **39**.

membered intermediate 54, which underwent contraction to furnish the target molecules 55 (Scheme 7).

Oxa-Michael addition reaction is one of the significant reactions of Michael addition for the construction of C-O skeletons and is used for several methodological approaches towards dihydrobenzofurans synthesis.53 In this regard, Zhu54 et al. in 2021 accomplished the synthesis of diverse stereoisomers of asymmetric 2,3-dihydrobenzofurans 58, 59, 60 and 61 via one pot, stereodivergent dual catalysis. In their novel methodology, arylvinyldiazoacetates 56 were reacted with substituted aminophenols 57 in a relay catalytic system to synthesize asymmetric products (58, 59, 60 and 61) in excellent yields (up to 99%) with exceptional enantiomeric and diastereomeric excess (99% ee, 99:1 dr respectively). In this regard, an initial C-H functionalization was attained employing a rhodium catalyst consisting of ligand 64, followed by the utilization of other organo-catalysts 62 and 63 that induced oxa-Michael addition to furnish the desired isomers of chiral 2,3dihydrobenzofurans (58, 59, 60 and 61) (Scheme 8). In addition to gram scale synthesis, the derivatization of the generated products was also carried out as these compounds were transformed into pharmaceutically important and enantiopure amino alcohols.



$$\begin{split} \mathsf{R}^1 = 5\text{-}\mathsf{Me}, 5\text{-}{}^t\!\mathsf{Bu}, 5\text{-}\mathsf{F}, 5\text{-}\mathsf{CI}, 5\text{-}\mathsf{Br}, 5\text{-}\mathsf{CO}_2\mathsf{Me}, 5\text{-}\mathsf{NO}_2, 6\text{-}\mathsf{F}, 6\text{-}\mathsf{CI}, 7\text{-}\mathsf{Me}, 7\text{-}\mathsf{Br}, 5\text{-}\mathsf{CF}_3, 5\text{,}6\text{-}\mathsf{fused}\,\mathsf{C}_4\mathsf{H}_4, \\ 5\text{-}\mathsf{C}_2\mathsf{H}_4\mathsf{N}\mathsf{H}\mathsf{Boc}, \mathsf{H}, 5\text{,}6\text{-}\mathsf{fused}\,\mathsf{dodecahydro-}3H\text{-}\mathsf{cyclopenta}[a]\mathsf{naphthalen-}3\text{-}\mathsf{one}\\ \mathsf{R}^2 = 3\text{-}\mathsf{CO}_2\mathsf{Me}, 3\text{-}\mathsf{NO}_2, 3\text{-}\mathsf{F}, 3\text{-}{}^t\!\mathsf{Bu}, 3\text{-}\mathsf{Me}, 3\text{-}\mathsf{Br}, 4\text{-}\mathsf{F}, 4\text{-}\mathsf{CI}, 3\text{-}\mathsf{tBu}, 3\text{-}\mathsf{CI}, 5\text{-}\mathsf{C}_2\mathsf{H}_4\mathsf{N}\mathsf{H}\mathsf{Boc}, \mathsf{H}, \\ 5\text{,}6\text{-}\mathsf{fused}\,\mathsf{dodecahydro-}3H\text{-}\mathsf{cyclopenta}[a]\mathsf{naphthalen-}3\text{-}\mathsf{one} \end{split}$$

Scheme 5 Synthesis of 2,3-dihydrobenzofuran derivatives 48.



Scheme 6 Asymmetric synthesis of 2,3-dihydro-3-benzofuranmethanamides 51

Metal carbenes are diverse intermediates that enable particular C-C bond forming transformations including C-H insertion,55 ylide formation,56 and aromatic addition reactions.⁵⁷ The asymmetric synthesis of 2,3-dihydrobenzofurans involving the insertion of carbene precursors into C(sp³)-H bonds potentially provides access to various crucial compounds like dihydrobenzofurans.58 In this regard, in 2021, Bi59 et al. designed a new method for the efficient assembly of asymmetric fluoroalkyl 2,3-disubstituted dihydrobenzofurans 66 in good to excellent yields (53-99%) with remarkable diastereomeric and enantiomeric ratio (>20:1 dr, 68:32-98:2 er) via a rhodiumcatalyzed protocol. In their synthetic approach, fluoroalkyl Ntriftosylhydrazones 65 were used as carbene precursors, which were transformed into the desired products 66 via the intramolecular insertion of carbene into the α -C(sp₃)-H bond of ether (Scheme 9). This efficient protocol was also used for the

gram-scale synthesis of diverse 2,3-dihydrobenzofuran derivatives. Bi and coworkers also synthesized various enantioselective natural and bioactive molecules **67**, **68**, **69** and **70** employing the above methodology (Fig. 5).

Buckley⁶⁰ *et al.* in 2021 reported the synthesis of novel diastereoselective dirhodium carboxylate catalysts and employed them in the construction of 2,3-dihydrobenzofuran skeletons **72**, **73**. In their novel methodology, the stereoselective C–H insertion reaction of aryldiazoacetates **71** took place in the presence of catalytic amount of dirhodium catalyst **74** to furnish 2,3-dihydrobenzofuran units **72** & **73** with excellent *trans* enantiopurity (84% ee) and *trans* diastereoselectivity (>91:9 dr) in toluene (Scheme 10).

In 2021, Hong⁶¹ *et al.* accomplished the Rh and asymmetric phosphoric acid **79** catalyzed synthesis of 2,3-dihy-drobenzofurans **78** *via* stereoselective Mannich type



R¹ = H, 4-F, 4-Cl, 4-Br, 4-NO₂, 5-OMe, 6-F, 6-Br, 6-I, 6-NO₂, 6-CF₃, 6-OMe R² = Me, Et, ⁱPr, Ph, cyclohexyl





$$\label{eq:R1} \begin{split} &\mathsf{R}^1=\text{2-F, }3\text{-}OMe, \text{3-Me, }3\text{-}Br, \text{ }4\text{-}OMe, \text{ }4\text{-}Me, \text{ }4\text{-}CI, \text{ }4\text{-}CF_3, \text{ }3\text{,}4\text{-}OMe, \text{ }3\text{,}5\text{-}diOMe, \text{ }3\text{,}4\text{-}triOMe, \\ & 3\text{,}4\text{-}fused \text{ }C_4\text{H}_4, \text{ }H, \text{ }furanyl, \text{ }thiophenyl \end{split}$$

R² = R³ = Me, Et, Bn, piperidine, pyrrolidine, morpholine, 1-benzylpiperazine, 1-benzylpyrrolidine, 1-(benzyloxy)piperidine, azepane





Scheme 9 Synthesis of asymmetric fluoroalkyl dihydrobenzofurans 66.



interception of phenolic oxonium ylides. In their novel methodology, diazo-containing phenolic derivatives **75** were made to react with imines **76** to synthesize the desired dihydrobenzofuran motifs **78** in low to excellent yield range (35– 90%) with exclusive diastereoselectivity (>20:1 dr) and enantioselectivity (>99% ee) values. The formation of oxonium ylide intermediate **77** took place by the addition of phenolic OH, followed by the interception of 77 with imines **76** to generate Mannich-type products **78** in the presence of chiral phosphoric acid **77** (Scheme 11). Further transformations of the synthesized compounds were also carried out to demonstrate the synthetic efficacy of the mentioned approach.

In 2022, Hu⁶² et al. presented a strategy for the efficient synthesis of 3-hydroxyoxindole incorporating 2,3-dihydrobenzofuran derivatives 83 via a Rh-catalyzed protocol. In their novel approach, diazo-containing phenolic compounds 80 were reacted with isatin 81 through an aldol type addition reaction to afford desired scaffolds 83 in moderate to excellent yields (58-98%) with exclusive diastereoselectivity (81: 19-95:5 dr). Carbene species generated from diazo compound, underwent intramolecular cyclization with hydroxyl group to form oxonium ylide intermediate 82. The isatin 81 added on the Si-face of 82 via aldol-type reaction to furnish desired 2,3-dihydrobenzofuran scaffolds 83 (Scheme 12). Two synthesized compounds 84 and 85 were found to exhibit anticancer activities against human colon cancer cells with 15.99 μ M and 14.48 μ M IC₅₀ values, respectively (Fig. 6). In order to demonstrate the practicality of the synthetic strategy, several spiro heterocyclic and amide products were



Scheme 10 Synthesis of the 2,3-dihydrobenzofuran skeleton 72 and 73



Scheme 11 Synthesis of diastereoselective 2,2-disubstituted dihydrobenzofuran 78





Fig. 6 Structures of compounds $84\ \ensuremath{\vartheta}$ 85 exhibiting anticancer potential.

also obtained from the synthesized 2,2-disubstituted dihydrobenzofurans 83.

Rh-catalyzed synthesis of heterocycles *via* [3 + 2] annulation portray an appealing strategy as it offers complete regio- and stereo-control for introducing the functional groups. Related to that, Zhong⁶³ *et al.* in 2021 demonstrated the synthesis of α -

carbon containing 2,3-dihydrobenzofuran quaternary analogues 89 via Rh-catalyzed C-H activation/[3 + 2] annulation. In their novel methodology, N-phenoxy amides 86 were reacted with propargylic monofluoroalkynes 87 in the presence of an Rh catalyst [Cp*RhCl₂]₂ and NaOAc (used as base) to afford cyclic products 89 in low to high yields (35-78%). Rhactivated C-H activation of 87 took place, followed by the regioselective insertion of alkyne to afford the five-membered species 88. Next, 88 subsequently underwent oxidative addition, reductive elimination and finally bimolecular nucleophilic substitution reaction to furnish 3-alkylidene-2,3dihydrobenzofuran moiety 89 (Scheme 13). Various spiro compounds were also synthesized from these alkylidene-2,3dihydrobenzofurans 89 to illustrate the synthetic utility of the reaction.



 $R^5 = Me, CH_2Bn, C_9H_{17}, cyclobutyl, cyclopropyl$

Scheme 13 Synthesis of 3-alkylidene-2,3-dihydrobenzofurans 89

Singh⁶⁴ et al. in 2021 also reported the Rh-catalyzed, chemodivergent synthesis of 2,3-dihydrobenzofuran derivatives 92 through the coupling of N-phenoxyacetamides 45 with alkylidenecyclopropanes 90 via C-H and C-C bond activation. This transformation was in accordance with the fact that the cyclopropanes are highly reactive in nature and underwent ring opening reactions to give five-membered heterocyclic compounds.65 Polar solvent, i.e., hexafluoroisopropanol (HFIP), induced [3 + 2] annulation to achieve the desired products 92 in moderate to high yield (52-82%). Rh-assisted coupling of 45 and 90 took place, followed by the formation and scission of the



- R² = 4-Me, 4-Cl, 4-Br, 4-Ph, 3,4-diOMe, 4-OH-3-OMe, 2-Me, 2,3-fused C₄H₄
- Scheme 14 Synthesis of 3-ethylidenedihydrobenzofuran derivatives 92

7-membered ring and subsequently β -hydride elimination to afford the intermediate 91. Next, this intermediate 91 underwent the oxidative insertion of the N-O bond, nucleophilic addition and finally deprotonation to achieve 2,3-disubstituted dihydrobenzofuran derivatives 92 (Scheme 14). The gram scale synthesis was also performed in order to demonstrate the efficacy of the synthetic strategy.

In 2021, Paymode and Sharma⁶⁶ designed a facile approach for the preparation of polycyclic 2,3-dihydrobenzofuran derivatives 95 via a rhodium-catalyzed system. In their synthetic approach, diazo compounds 93 were condensed with 2,3dihydrofurans/cyclopentenes/cyclohexenes 94 via the [3 + 2] annulation process in the presence of Rh₂(TPA)₄ to furnish desired dihydrobenzofurans 95 in moderate to high yields (34-85%) (Scheme 15). In the wake of wide-spreading bacterial diseases, synthetic chemists have employed several synthetic pathways to develop efficacious anti-bacterial agents.67 Employing the mentioned methodology, Paymode and Sharma also performed the total synthesis of naturally occurring aflatoxin B₂ 96a (Fig. 7), which exhibits potent antimitotic and antimicrobial activities.

Various tailored molecules are synthesized as a result of ring-opening reactions of cyclic compounds.68 Jiang69 et al. in 2022 carried out the rhodium-mediated one pot synthesis of benzofuran-3(2H)-ones 101 by reacting aliphatic alcohols (MeOH) with salicylaldehyde 97 and cyclopropanols 98. After interpreting the screening results, a total of three series of target heterocycles 101 were synthesized in moderate to high yield (32-70%) by independently adding the substitution on one of the three components one by one under optimized



Scheme 15 Synthesis of fused rings containing 2,3-dihydrobenzofuran derivative 95.

conditions ([CpRhCl₂]₂ catalyst, Cu(OAc)₂ oxidant & CsOAc (additive) (Scheme 15). The reaction mechanism was proposed to involve the C–H activation of substituted salicyclaldehydes **97**, followed by ligand exchange with aryl cyclopropanols **98**, β -carbon elimination and reductive elimination (cyclopropane ring opening) to result in C–H alkylation product **99**. The compound **99** was then supposed to be enolized by copper acetate, followed by β -hydride elimination and oxy-Michael addition to give benzofuranones **100**. Substituted



96a, Antimicrobial

Fig. 7 Structure of aflatoxin B₂ 96a.

benzofuranones **100** were then assumed to be reoxidized, followed by methanol involving Michael addition to afford target molecules **101** (Scheme 16).

2.2. Pd-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Palladium is a valuable grav-white metal, especially used as a catalyst due to its distinct properties such as thermal stability, resistance to poisoning, selectivity and its high surface area. Pdcatalyzed synthesis of five-membered heterocycles has been presented as the most beneficial and reliable synthetic methods as it has diverse functional group tolerance and carried out in mild reaction conditions.⁷⁰ The classical Heck coupling progresses through different steps including oxidative addition, proceeded by migratory insertion and reductive elimination. Heck coupling has provided novel methodologies towards the sophisticated aromatic substrates from organohalides and alkenes due to the stepwise development of synthetic logic.71 Over the past few decennia, Pd-catalyzed Heck coupling has been widely used as one of the crucial synthetic tools facilitating the straightforward synthesis of molecular heterogeneity and complexity.72 In 2021, Wu73 et al. disclosed an enantioselective,



Scheme 16 Synthesis of benzofuran-3(2H)-ones 101.

Pd-catalyzed method for the synthesis of 2,3-dihydrobenzofuran derivatives **105**. In their novel methodology, aryl iodide-joined alkenes **102** were reacted with *o*-alkynylanilines **103** in the presence of $Pd_2(dba)_3 \cdot CHCl_3$ along with ligand *N*-Me–Xu₃ *via* Heck/Cacchi reactions to achieve dihydrobenzofurans in excellent yields (84–97%). Asymmetric σ -alkylpalladium intermediate **104** was formed *via* intramolecular Heck coupling reaction of Pd activated **102**, followed by its reaction with **103** to generate the desired polycyclic products **105** with exceptional enantiomeric excess (84–97% ee) (Scheme 17). The gram scale synthesis

was also demonstrated along with the derivatization of the synthesized compounds.

In 2021, Sreenivasulu and Satyanarayana⁷⁴ envisioned a Pdcatalyzed, regio- and stereo-selective construction of Z/Eisomers of 1,3-dihydroisobenzofurans **108** & **109** *via* the aryl Heck coupling of *o*-substituted tertiary alcohols **106** and substituted aryl bromides **107**. It is a temperature and timedependent strategy, in which *Z*-isomer **108** was formed at 80 ° C after 4–6 hours of reaction, whereas stable *E*-isomer **109** was attained at 100 °C after 8–12 hours of reaction (Scheme 18).



- C_3H_6 -4,4-dimetriyipiperidire-2,6-dione, C_3H_6 -0xazolidire-2-
- R³ = 5-OMe, 5-Me, 5-CN, 5-CF₃, 5-F, 5-Cl, 6-CO₂Me, H
- $$\label{eq:R4} \begin{split} & {\sf R}^4 = {\sf Ph}, \, 4\text{-}{\sf OMeC}_6{\sf H}_4, \, 4\text{-}{\sf PhC}_6{\sf H}_4, \, 4\text{-}{\sf CO}_2{\sf MeC}_6{\sf H}_4, \, 4\text{-}{\sf CF}_3{\sf C}_6{\sf H}_4, \, 4\text{-}{\sf FC}_6{\sf H}_4, \, 3\text{-}{\sf OMeC}_6{\sf H}_4, \\ & {\sf PMP}, \, {\sf C}_3{\sf H}_6{\sf CI}, \, {}^n\!{\sf Bu}, \, {\sf cyclohexyl}, \, {\sf thiophene} \end{split}$$

Scheme 17 Synthesis of polycyclic dihydrobenzofurans 105.



Scheme 18 Synthesis of (Z)/(E)-3-(1-arylalkylidene)-1,3-dihydroisobenzofurans 108 and 109.

Review

Initially, Pd-assisted intermolecular Heck coupling reaction between **106** and **107** took place to generate an intermediate **110**, followed by the elimination reaction to form a palladacycle **111**. Further, this palladacycle underwent intramolecular *oxo*cyclization to furnish the desired heterocyclic products **108** & **109** in moderate to high yields, *i.e.*, 49–84% for *Z*-isomer and 68–88% for *E*-isomer, respectively (Scheme 19).

Pd/XuPhos catalyst is used as an efficient catalytic system for the stereodefined cascade Heck/intermolecular C–H alkylation reactions.⁷⁵ Taking into consideration the wide synthetic utility of borylated compounds, Wu⁷⁶ *et al.* in 2022, also developed a novel palladium promoted Heck/borylation strategy utilizing alkenes fused with aryl iodides **102** as precursors. Their synthetic protocol involved the treatment of aromatic rings substituted alkenes **102** and B₂pin₂ **112** in the presence of Pd₂(dba)₃·CHCl₃ catalyst exploiting *N*-Me–Xu₃ as a ligand (having 3,5-di-*tert*-butyl-4-methoxyphenyl group), using cesium carbonate as a base in diethyl ether and water. This protocol resulted in the efficacious and enantioselective synthesis of dihydrobenzofuran based boronic esters **113** (52–98%) with excellent enantiomeric excess (53–97% ee). This highly efficient synthetic strategy also paved routes towards the accomplishment of chromane, indoline and indane boronic esters (Scheme 20).

In 2022, Guo⁷⁷ et al. successfully designed a new Pd-catalyzed method for the synthesis of substituted 3,3-disubstituted-2,3dihydrobenzofurans via the reaction of olefin-tethered aryl iodides 102 with α,β -unsaturated ketones 114 and substituted styrenes 115. In their one-pot synthetic methodology, two sequential Heck couplings took place, leading to the construction of dihydrobenzofuran skeletons 116 and 117. Good to high yields, i.e., 87% (116) and 78% (117), with exclusive E/Z selectivity (20:1) values of target molecules were achieved when Pd(PhCN)₂Cl₂ was subjected as a catalyst along with Cy₂NMe (base) in MeCN (solvent) (Scheme 21). The reaction mechanism was proposed to proceed via the reaction of reduced Pd species [Pd(0)] and aryl iodides 102 to form an intermediate 118, which underwent subsequent intramolecular insertion reaction and olefin insertion reaction to form another intermediate 119. Next, reductive elimination reaction finally generated the desired dihydrofuran products 117 (Scheme 22).

Spirocyclic organic compounds have acquired a significant place in medicinal chemistry due to their vast biological potential.⁷⁸ Marchese⁷⁹ *et al.* in 2022 devised a modular method for the preparation of 2,3-dihydrobenzofuran-based spirocyclic



Scheme 19 Proposed mechanism for the synthesis of (Z)/(E)-3-(1-arylalkylidene)-1,3-dihydroisobenzofurans 108 and 109.



Scheme 20 Synthesis of 2,3-dihydrobenzofuranyl boronic esters 113.



- R^1 = H, 5-F, 5-Cl, 5-Br, 5-Me, 5-CF₃, 5-COOMe, 6-COOMe, 6-Cl, 6-Br, 4,5 fused C₆H₄, 6-1-(piperidin-1-yl)ethan-1-one R^2 = Me,COOMe, COOEt, H
- $R^{3} = 4-FC_{6}H_{4,} 4-CIC_{6}H_{4,} 4-BrC_{6}H_{4,} 4-CH3C_{6}H_{4,} 4-OCH3C_{6}H_{4,} 4-OPhC_{6}H_{4,} 4-tBu, 4-CF3C_{6}H_{4,} 4-NO_{2}, C_{6}H_{5,} 3-FC_{6}H_{4,} 2-FC_{6}H_{4,} naphthyl$

Scheme 21 Synthesis of 3,3-disubstituted-2,3-dihydrobenzofuran 116 and 117.



Scheme 22 Proposed mechanism for the synthesis of 3,3-disubstituted dihydrobenzofurans 117.

compounds **121**, **122** and **123** *via* palladium catalysis. In their synthetic approach, *o*-substituted aryl iodides **120** were transformed into corresponding bis-heterocyclic spirocycles **121**, **122** and **123** by following the Mizoroki–Heck-type reaction pathway in the presence of Cs_2CO_3 (used as base). Initially, the oxidative addition of **120** took place, followed by migratory insertion and C-H activation to form an intermediate, which on reductive elimination generated spirocyclic dihydrobenzofurans **121**, **122** and **123**. Low to high yields (31–94%) were observed using this methodological approach (Scheme 23).

Another palladium-catalyzed method for dihydrobenzofurans was given by Kang⁸⁰ *et al.* in 2022. They carried out the palladium-mediated aminocarbonylation by treating aryl iodide-linked alkenes **124** with diversely substituted nitro compounds **125** and **126**. The reaction took place smoothly utilizing $PdCl_2$ as a catalyst, $Mo(CO)_6$ for the release of CO, triphenylphosphine as the ligand, DIPEA as base in tetrahydrofuran solvent without any addition of additive and reducing agent (Scheme 24). The synthetic pathway was proposed to follow the certain steps, *i.e.*, oxidative addition, intramolecular carbopalladation, linking and integration of CO to form intermediate **131**. This was followed by the attack of amine (obtained by the reduction of substituted nitro compound) and reductive elimination to gain dihydrobenzofuran derivatives **127** and **128** in 51–88% yields (Scheme 25).

In 2023, another palladium-promoted asymmetric synthesis of dihydrobenzofurans *via* Heck/Tsuji–Trost was put forward by Zhang's group.⁸¹ In the newly developed synthetic pathway, substituted dienes **133** were treated with halo-substituted phenols **134** using $Pd_2dba_3 \cdot CHCl_3$ as catalyst and TY-Phos as ligand using sodium phenoxide in dichloromethane, which afforded alkenyl-substituted dihydrobenzofurans **135** in 35–99% yield with enantiomeric excess (73–97 ee) (Scheme 26). To demonstrate the practicality of the novel synthetic approach, the gram scale synthesis of **135** was performed. Moreover, the derivatization of the synthesized compounds towards the synthesis of natural products, *i.e.*, tremetone and fomannoxin, was also performed.

The Pd-catalyzed synthesis of 2,3-dihydrobenzofurans involving the insertion of carbene precursors into $C(sp^3)$ –H bonds potentially provides access to various crucial compounds like dihydrobenzofurans.⁸² In 2023, Ding⁸³ *et al.* reported a rarely explored 1,5-Pd/H shift to synthesize dihydrobenzofurans. They treated alkynes-substituted compounds **136** (obtained from 2-bromo-3-hydroxy benzaldehydes) with R–B(OH)₂/R– BPin, substituted alkynols **137** and *N*-sulfonyl hydrazones **138** *via* palladium catalyzed (PdCl₂ & Pd(OAc)₂ respectively) cascade reaction. The reaction proceeded *via* the 1,5-Pd/H shift approach between the vinyl and acyl moiety to furnish acyl palladium species. The acyl palladium intermediates were then



X = O, NTs, N

R¹ = 7-CO₂Me, 7-Me, 8-Cl, 8-SO₂-4-MeC₆H₄, H R² = 4-Cl, H







129, 73%

subjected to decarbonylation, followed by the reaction with nucleophiles to yield dihydrobenzofuran derivatives **139**, **140** and **141** in efficient yields *via* decarbonylative alkenylation, arylation and alkynylation (Scheme 27). The novel synthetic strategy highlights the significant synthetic efficacy, facilitating the production of disubstituted and polycyclic products.

Annulation reactions simply refer to the process of combining two or more molecular fragments, often cyclic structures, to form bridged or fused ring systems. Pd-catalyzed annulation reactions involving the synthesis of arene-fused furan heterocycles have tremendous importance in organic synthesis.⁸⁴ In this perspective, Zhou⁸⁵ *et al.* in 2021 prepared

130, 59%



Scheme 25 Proposed mechanism for the synthesis of carbamoylsubstituted 2,3-dihydrobenzofurans **128**.

a library of polycyclic dihydrobenzofurans **146** *via* Pd-catalyzed annulation reaction. In their novel methodology, alkenyl ethers **142** and alkynyl oxime ethers **143** underwent cyclization reaction in the presence of $Pd(OAc)_2$ along with $CuCl_2$ (used as an oxidant) and tetrabutyl ammonium bromide (as co-catalyst) to afford 2,3-dihydrobenzofuran derivatives **146** in moderate to excellent yields (41–86%) (Scheme 28). This synthetic approach covers a broad range of substrate scope (32 examples). Various synthetic transformations of the synthesized compounds took place to illustrate the synthetic efficiency of the procedure. Similarly, in 2022, Houghtling⁸⁶ *et al.* reported a palladiumcatalyzed protocol for the synthesis of variety of dihydrobenzofuran derivatives **148**, in which a novel urea ligand **149** was utilized to enhance the product yield (50–72%). In this context, substituted 2-bromophenols **134** were treated with 1,3dienes **147** in the presence of NaO^{*t*}Bu (as base) in a 9 : 1 ratio of PhMe and anisole (used as solvent) at 110 °C (Scheme 29).

In 2023, Sun⁸⁷ et al. reported the palladium-catalyzed chiral [4 + 1] cyclization reaction to synthesize enantioselective dihydrobenzofurans 152. In their synthetic methodology, cyclic vinyl methylene ketone 151 and leaving group substituted aryl ethers 150 were subjected to a series of allylation reaction and C-H activation reaction in the presence of palladium catalyst and ligand L, utilizing cesium fluoride as a base in 2-methyl tetrahydrofuran and acetone. As a result, the enantiomeric synthesis of a library of dihydrobenzofurans 152 was achieved in 38-89% vield up to 99:1 enantiomeric ratio. The reaction mechanism was supposed to involve the palladium mediated decarboxylation of substituted vinyl methylene carbonate, oxidative process, O-nucleophilic conversion, asymmetric concerted metalation-deprotonation, protonation and reductive elimination to yield target molecules (Scheme 30). Various derivatives of 152 were synthesized in order to elaborate the synthetic efficiency.

In recent times, synthesis of various heterocycles and carbocycles have been achieved by employing alkyne cyclization. In these cyclization reactions, palladium-based catalysts have gained great significance. In 2023, Sun⁷⁰ *et al.* reported the synthesis of dihydrobenzofurans **145** by treating tosyl



- $\begin{array}{l} {\rm R}^{1} = {\rm H, Ph, 2-MeOPh, 2-MePh, 2-CF_{3}Ph, 3-MePh, 3-CIPh, 4-OMePh, 4-OCF_{3}Ph, 4-'BuPh, \\ {\rm 4-PhC_{6}H_{4}, 4-CIPh, 4-CF_{3}Ph, 2,4,6-MePh, naphthyl, thiophenyl, cyclohexyl, C_{2}H_{4}Ph, \\ {\rm C_{2}H_{2}(CH_{2})_{10}Br, C_{2}H_{2}C_{5}H_{11}, \ C_{2}H_{2}(CH_{2})_{5}OTBS, (R)-4,8-dimethylnona-1,7-diene, OTBS, \\ {\rm fused C_{4}H_{6}, } \end{array}$
- $R^2 = H$, Me $R^3 = H$, Me
- R⁴ = 4-Me, 4-F, 5-OMe, 5-^tBu, 5-Me, 5-F, 5-CF₃, 6-OMe, 6-Me, 6-F, 6-COOMe, 7-OMe, 7-Me, 7-Ac, fused C₄H₄

Scheme 26 Synthesis of 2,3-dihydrobenzofurans 135.



Scheme 27 Synthesis of 2,3-dihydrobenzofurans 139-141.

hydrazones **143** and aryl-substituted joined alkynes **144** in the presence of bis(triphenylphosphine)palladium(π) dichloride catalyst, using Cs₂CO₃ as a base and tricyclohexylphosphine PCy₃ as a ligand in toluene, followed by the addition of dienophile, *i.e.*, dimethylbut-2-ynedioate. The reaction mechanism involved the intramolecular carbopalladation, migratory insertion, γ -hydride elimination and Diels–Alder ([4 + 2] cycloaddition) reaction to furnish the spirocyclobutane-substituted dihydrobenzofurans **145** (Scheme 31).

Wu^{ss} *et al.* in 2021 developed a Pd-catalyzed protocol for the synthesis of 2,3-dihydrobenzofuran derivatives **158** *via* $C(sp^3)$ –H and $C(sp^2)$ –H intramolecular coupling. In their novel methodology, alkyl phenyl ethers **156** were used as efficient starting materials, which underwent subsequent $C(sp^3)$ –H and $C(sp^2)$ –H bond activation and reductive elimination in the presence of 1,4-dibenzoquinone (BQ), AgOAc and LiOAc (as base) to furnish desired heterocyclic products **158** in moderate to excellent yields (33–99%) (Scheme 32). The developed protocol was also utilized further for the gram scale synthesis of target molecules.

Reddy⁸⁹ *et al.* in 2021 proposed a Pd-catalyzed highly efficient protocol for the preparation of 2,2,3-trisubstituted dihydrobenzofuran derivatives **162** *via* intramolecular condensation. In their novel methodology, 2-hydroxyphenyl-substituted enones **159** were reacted with diazo compound **160** in the presence of [Pd(cinnamyl)Cl]₂ along with MeSO₃H (used as an additive) to afford dihydrobenzofurans **162** in moderate to excellent yields (51–91%). Active Pd catalyst interacted with diazo compounds **160** to give carbene species, which on nucleophilic addition with **159** gave oxonium ylides that existed in equilibrium with the corresponding zwitterions **161**. Zwitterions **161** were then trapped *via* Michael addition to afford desired products **162** (Scheme 33). Further, a number of substituted dibenzofuran products were also synthesized in order to demonstrate the synthetic efficiency of the novel approach.

2.3. Cu-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Copper-based catalysts are most active, cheap and widely used catalysts in organic transformations and have various oxidation states. The reductive elimination of Cu(m) to generate C–X (X = heteroatom) bond is well studied in the literature.⁹⁰ Cucatalyzed synthesis also plays a crucial role in the synthesis of diverse dihydrobenzofuran scaffolds.⁹¹ Synthesis *via* deconstructive insertion reaction involving the generation of new structures *via* the bond cleavage of easily accessible scaffolds has recently attained growing attention from synthetic practitioners.⁹² In this regard, Zheng⁹³ *et al.* in 2022 reported the Cucatalyzed synthesis of pyridine-fused dihydrobenzofuran derivatives **167**. In their synthetic protocol, coumarins **163** and oximes **164** were reacted in the presence of CuBr (catalyst) along with Li₂CO₃ (used as an additive) to afford 2,3-dihydrobenzofurans **167** in low to high yields (16–85%) *via* the







destructive insertion of **164** into **163**. This synthetic methodology provides broad functional group tolerance. Cu-assisted oxime-Cu-iminyl radical was converted into α -carbon radical, which underwent radical addition to give intermediate **165**, followed by intramolecular addition to carbonyl group and oxidation to yield the desired products **167** (Scheme 34). The synthesized compounds were further transformed into a number of useful products to illustrate the proficiency of the developed synthetic route.

[3+2] cycloaddition reactions are pericyclic reactions, which involve the addition of a double bond system to a three-atom ring system. These cycloaddition reactions are amongst the most convenient synthetic protocols for the synthesis of fivemembered heterocyclic ring compounds.⁹⁴ Considering this, in 2021, Jing⁹⁵ *et al.* described the synthesis of a range of enantioselective 2,3-dihydrobenzofurans **170** *via* the Cu/SPDOcatalyzed synthetic route. In their novel approach, quinone esters **168** and substituted styrenes **169** underwent [3 + 2]cycloaddition in the presence of SPDO-ligated Cu(OTf)₂ to furnish the desired dihydrobenzofuran moieties **170** in good to excellent yields (86–96%) with extraordinary enantioselectivities (86–99% ee) (Scheme 35). Natural products **171** and **172** were also synthesized by utilizing the mentioned methodological strategy (Fig. 8).







Scheme 31 Synthesis of polycyclic 2,3-dihydrobenzofurans 155.



Scheme 32 Synthesis of fused cyclic dihydrobenzofurans 158.

An enantioselective and diastereoselective pathway towards the synthesis of dihydrobenzofurans **163** was given by Zhu⁹⁶ *et al.* in 2023. They exploited copper(π)/SPDO complex (chiral spirocyclic pyrrolidine (oxazoline)) catalyst for carrying out asymmetric [3 + 2] cycloaddition reaction between 2,3-dihydrofuran **173** and quinone esters **168** utilizing copper triflate Cu(OTf)₂ and ligand L in toluene, wet tetrahydrofuran or mestylene solvent, to enable the asymmetric synthesis of benzofuran derivatives 175. The reaction mechanism was suggested to move forward with the formation of intermediate 174, on the coordination of substrates and catalyst. Intermediate 174 was then believed to undergo further transformations to furnish the synthesis of target molecules (Scheme 36). The developed synthetic route was further explored towards the synthesis of olefin variants-substituted dihydrobenzofurans (A & B) (Fig. 9) and naturally-occurring aflatoxins 96, *i.e.*, (–)aflatoxin B₂ 96a & (–)dihydroaflatoxin D₂ 96b (Fig. 10).

The five-membered O-containing compound, *i.e.*, 2benzylidene-1-benzofuran-3-one, called aurone, showed a number of pharmacological activities like antidiabetic, antiviral and anticarcinogenic.⁹⁷ In 2021, Devi *et al.*⁹⁸ employed copper bromide-promoted synthesis of these 2-benzylidene-1benzofuran-3-one derivatives **179**. For this purpose, they furnished chalcones **179** by treating aromatic aldehydes **177** with substituted acetophenone **176** utilizing montmorillonite K10 clay as a catalyst. The synthesized chalcones **179** were then subjected to copper bromide cyclization by employing dimethylformamide/water (7:3) as a solvent to attain substituted benzofuranone heterocycles **179** (Scheme 37).

In the same year, Mitsudo⁹⁹ *et al.* designed an efficient Cucatalyzed protocol for the synthesis of dihydrofuran-fused thienoacenes **182** and **183** *via* C–O cyclization. In their novel methodology, 2-(benzo[*b*]thiophen-2-yl)phenols **180** and 2-(benzo[*b*]thiophen-3-yl)phenols **181** underwent the



 $\begin{array}{l} {\sf R}^1 = {\sf 5}\text{-}{\sf Me}, \ {\sf 5}\text{-}{\sf OMe}, \ {\sf 5}\text{-}{\sf F}, \ {\sf 5}\text{-}{\sf CI}, \ {\sf 6}\text{-}{\sf Me}, \ {\sf H} \\ {\sf R}^2 = {\sf Me}, \ {\sf C}_6{\sf H}_5, \ {\sf P}\text{-}{\sf Me}{\sf C}_6{\sf H}_4, \ m\text{-}{\sf Me}{\sf OC}_6{\sf H}_4, \ {\sf Et}, \ {\sf furanyl}, \ {\sf thiophenyl} \\ {\sf R}^3 = {\sf C}_6{\sf H}_5, \ {\sf 4}\text{-}{\sf Me}{\sf C}_6{\sf H}_4, \ {\sf 4}\text{-}{\sf f}{\sf Bu}{\sf C}_6{\sf H}_4, \ {\sf 4}\text{-}{\sf Ph}{\sf C}_6{\sf H}_4, \ {\sf 4}\text{-}{\sf FC}_6{\sf H}_4, \ {\sf 3}\text{-}{\sf Cl}{\sf C}_6{\sf H}_4, \ {\sf 3}\text{-}{\sf CF}_3{\sf C}_6{\sf H}_4, \\ {\sf 3}, {\sf 4}\text{-}{\sf dicl}{\sf C}_6{\sf H}_3, \ {\sf 2}\text{-}{\sf Me}{\sf C}_6{\sf H}_5, \ {\sf 3}, {\sf 4}\text{-}{\sf fused} \ {\sf C}_4{\sf H}_4, \ {\sf COCH}_3, \end{array}$

Scheme 33 Synthesis of 2,2,3-trisubstituted dihydrobenzofuran derivatives 162.



 $\begin{aligned} \mathsf{R}^1 &= 6\text{-NEt}, \ 6\text{-OMeC}_6\mathsf{H}_4, \ 6\text{-BrC}_6\mathsf{H}_4, \ 6\text{-BrC}_6\mathsf{H}_4, \ 6\text{-BrC}_6\mathsf{H}_4, \ 5\text{-CH}_3, \ 5\text{-FC}_6\mathsf{H}_4, \ 4,5\text{-fusedC}_4\mathsf{H}_4, \\ &4,5\text{-diOMe}, \ 5,7\text{-}^{t}\mathsf{Bu}, \ 5,7\text{-diCl}, \ 5,7\text{-diBr}, \ 5,7\text{-dil}, \ 6\text{-OC}_9\mathsf{H}_{17} \\ \mathsf{R}^2 &= \mathsf{CO}_2\mathsf{Me}, \ \mathsf{COPh}, \ \mathsf{COC}(\mathsf{C}_3\mathsf{H}_9), \ \mathsf{PO}(\mathsf{EtO})_2, \ \mathsf{CO}_2\mathsf{Et} \\ \mathsf{R}^3 &= 4\text{-}^{t}\mathsf{BuC}_6\mathsf{H}_4, \ 4\text{-MeC}_6\mathsf{H}_4, \ 4\text{-OMeC}_6\mathsf{H}_4, \ 4\text{-BrC}_6\mathsf{H}_4, \ 4\text{-IC}_6\mathsf{H}_4, \ 4\text{-MeSC}_6\mathsf{H}_4, \\ \mathsf{R}^3 &= 4\text{-}^{t}\mathsf{BuC}_6\mathsf{H}_4, \ 4\text{-MeC}_6\mathsf{H}_4, \ 4\text{-OMeC}_6\mathsf{H}_4, \ 4\text{-BrC}_6\mathsf{H}_4, \ 4\text{-BrC}_6\mathsf{H}_4, \ 4\text{-MeSC}_6\mathsf{H}_4, \end{aligned}$

- 4-CO₂MeC₆H₄, 4-CNC₆H₄, 4-CF₃C₆H₄, 3-CIC₆H₄, 3-CNC₆H₄, 2-MeC₆H₄, 3,4-diMeC₆H₄,
 - $4-SO_2N(C_3H_7)_2C_6H_4$, C_2H_2Ph , C_6H_5 , benzodioxole, furanyl, thiophenyl, benzofuran,
- 1,Me-1*H*-indole, pyrridine, 1-Me-1*H*-pyrrole, 1-Me-1*H*-pyrazole,

3-(1-(3-(2-methoxyphenoxy)propyl)piperidin-4-yl)benzo[d]isoxazole

Scheme 34 Synthesis of 2,3-dihydrobenzofuran-fused pyridones 167.

degenerative cyclization process to afford thieno[3,2-*b*]furans **182** (12–86%) or thieno[2,3-*b*]furans **183** (53–95%) respectively. Similarly, substrates **181** afforded heteroacenes **184** and **185** having π -expanded system *via* double C–O cyclization. The catalytic amount of Cu(OAc)₂, NaOAc (as base) and PhCOOH (as acid) were used to achieve the efficient yields of target molecules (Schemes 38 and 39).

Oxidative cross coupling is a strategy where two phenolic moieties are coupled together *via* an oxidative process to generate new C–C bonds.¹⁰⁰ Cu-catalyzed intermolecular oxidative cross coupling reactions have attracted much attention in organic synthesis to generate new dihydrobenzofuran units. In 2021, Dong¹⁰¹ *et al.* developed a biomimetic, Cu-catalyzed synthesis of neolignane analogs comprising of 2,3-dihydrobenzofuran units **190**. In their novel methodology, substituted *para*-alkenyl phenols **188** were cross coupled with a number of electron rich phenols **20** to furnish 8-5' neolignan correspondents **190** exclusively. Cuassisted radical formation of phenols **(188, 20)** took place, which coupled to generate quinone methide intermediate





189, followed by the addition of -OH to **189** to afford the desired 2,3-dihydrobenzofuran skeletons **190** in excellent yields (24-95%) with notable diastereoselectivity values (>20:1 dr) (Scheme 40). Through the developed synthetic route, Licarin A **191** was also synthesized, which exhibits anti-inflammatory activity (Fig. 11).

2.4. Ni-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Nickel, which is also regarded as the "spirited horse" of TM catalysis, is a well-known metal catalyst used in organic

synthesis.102 Enantioselective Ni-catalyzed reductive Heck coupling of tethered alkenes has been utilized to furnish a range of benzene-fused heterocyclic rings bearing a quaternary stereogenic center.¹⁰³ In reference to this, Cerveri¹⁰⁴ et al. in 2021, synthesized a library of asymmetric 2.3dihydrobenzofuran-3-ylacetic acids 193 via a tandem, enantioselective Ni-catalyzed Heck cross coupling reaction. In their novel methodology, CO₂ fixation took place when o-aryliodines 102 were cyclized in the presence of $[(L)_2Ni(H_2O)Cl]Cl$ (used as pre-catalyst), Zn (as reducing agent), tetrabutylammonium iodide (TBAI) and trimethylsilyl chloride (TMSCl) (as additives) to afford asymmetric dihydrobenzofuran derivatives 193. Complex formation of compound 102 and NiL-precatalyst was carried out to form intermediate 192, followed by subsequent Zn-mediated reduction and CO₂ insertion to afford the target compounds in moderate to high yields (47-69%) with up to 99% enantiomeric excess (Scheme 41).

Pyrano-fused cyclic organic compounds are ubiquitously employed in medicinal chemistry owing to their unique



Scheme 36 Synthesis of 2,3,3a,8a-tetrahydrofuro[2,3-b]benzofurans 175.



Fig. 10 Structures of naturally-occurring dihydrobenzofuran derivatives 96a and 96b.

biologically active nature. There are mainly two types of these heterocycles based on the point of attachment of oxygen atom, *i.e.*, C3–C2 & C2–C1. In 2022, Bhardwaj¹⁰⁵ *et al.* reported a novel and efficient methodology by reacting enopyranoses **195** and iodine substituted phenols **194** under the action of nickel catalyst using cesium carbonate as a base in the presence of triphenylphosphine and dimethylformamide. This one-pot synthetic approach resulted in the straightforward synthesis of pyrano *cis*-fused dihydrobenzofurans **198** (47– 75% yield). The reaction mechanism was proposed to proceed *via* the oxidative addition and carbometallation to form an intermediate **196**, followed by β-OAc elimination and allylic rearrangement to furnish the desired dihydrobenzofurans **197** (Scheme 42).

As a key component of several dicarbon-functionalization reactions, alkene aryl-acylation or aryl-carbamoylation facilitates the synthesis of structurally significant heterocycles comprising of carbonyl compounds.¹⁰⁶ In this respect, Wang¹⁰⁷ *et al.* in 2022 described a novel approach to afford dihydrobenzofuran derivatives **200** employing nickel-catalyzed aryl carbamoylation and aryl acylation of aryl iodide-joined alkenes **124**. Aryl iodide-joined alkenes **124** were subjected to treatment with aryl-substituted isocyanates **199** *via* Nicatalyzed aryl carbamoylation to result in the efficient synthesis of substituted dihydrobenzofurans **200** (62–73%) (Scheme 43).

In 2021, Lin¹⁰⁸ *et al.* reported the Ni-catalyzed asymmetric synthesis of 2,3-dihydrobenzofuran derivatives **204** *via* the reductive aryl allylation process. In their synthetic approach, aryl iodides **102** were reacted with cyclic vinyl ethylene carbonates **203** to achieve cyclic products **204** in moderate to good yields (46–67%) with remarkable enantioselectivity values (>98% ee). Alkene-tethered aryl iodides **102** underwent oxidative addition with Ni catalyst to generate intermediate **201**, followed by olefinic migratory insertion to form cyclic intermediate **202**. Next, vinyl ethylene carbonates **203** were oxidatively added to the intermediate **202**, proceeded by reductive elimination to access the desired dihydrobenzofurans **204** (Scheme 44).

Geng¹⁰⁹ *et al.* in 2021 reported the Ni-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives **207** *via* carbonylative synthesis. In their novel methodology, *ortho*-substituted aryl iodides **124** reacted with alkyl halides **205** in the presence of Ni(acac)₂ (as a catalyst), 4,4'-di-*tert*-butyl-2,2'-bipyridyl (**L**) (as ligand), Mo(CO)₆ (as carbonylating agent) and Mn (as reductant) to furnish desired products **207** in low to excellent yield (38–87%). In the first step, activated Ni was oxidatively added to *ortho*-substituted aryl iodide **124**, followed by intramolecular addition to the carbonyl group and addition of alkyl halide **205** to generate intermediate **206**. The intermediate **206** was further subjected to reductive elimination to afford the 2,3-dihydrobenzofuran derivatives **207** (Scheme 45).

Electrification is an optimistic approach of replacing traditional energy-demanding chemical processes with greener substitutes and thus alleviating carbon emissions.¹¹⁰ Utilizing this strategy, Déjardin¹¹¹ *et al.* in 2021, proposed an electroreductive, Ni-catalyzed synthesis of 2,3- dihydrobenzofuran derivatives **211** based on domino reaction. In this regard, intramolecular carbonickelation of propargylic aryl halides **208** was carried out, followed by the subsequent cyclization and nucleophilic addition of benzaldehydes **209** to furnish



Scheme 37 Synthesis of 2-benzylidene-1-benzofuran-3-ones 179.



184

185

Scheme 38 Synthesis of furan-fused thienoacenes 182–185.



Scheme 39 Proposed mechanism for the synthesis of furan-fused thienoacenes 182.

substituted 2,3-dihydrobenzofuran **211** (11–90%). The reaction was highly stereo and regio-selective. Maximum yield (90%) was observed when the reaction proceeded at 80 °C in the presence of DMF (as a solvent) (Scheme 46).

2.5. Au-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Au-containing catalytic systems are extensively studied in organic synthesis as they have exclusive low temperature oxidation activity. Au-catalyzed synthesis of heterocycles *via* [3 +

2] annulation portrays an appealing strategy as it offers complete regio- and stereo-control for introducing the functional groups. Utilizing this strategy, Liang¹¹² *et al.* in 2021 described the synthesis of polycyclic dihydrobenzofuran derivatives **215** *via* gold and Cu-catalyzed tandem reaction. In their synthetic approach, propargyl alcohols **212** were coupled with pyridylhomopropargylic alcohols **213** *via* a number of processes, *i.e.*, 5-*endo-dig* cyclization, Meyer–Schuster rearrangement and Friedel–Crafts-type reactions to furnish 2,3dihydrobenzofurans **215** in low to high yields (14–81%) (Scheme 47).

In 2023, Morita¹¹³ *et al.* carried out the efficient synthesis of dihydrobenzofurans by involving gold-based NHC catalyst (permethylated β -cyclodextrin-tagged *N*-heterocyclic carbene-gold catalyst) in water. They utilized the NHC catalyst due to the hydrophobic nature of β -CD,¹¹⁴ which was expected to enhance the solubility of substrates with non-polar moieties in water. Moreover, NHC-based gold catalyst was assumed to activate the alcoholic functionality in benzylic alcohols **217** and carbonyl functionality in *p*-quinones **216** to facilitate the synthesis of dihydrobenzofurans **197**. Substituted *p*-quinones **216** were treated with isoeugenols *via* [3 + 2] cycloaddition reaction in the presence of β -CD–NHC–AuCl catalyst, utilizing AgNTf₂ (as an additive) in the excess of water to result in the efficient asymmetric synthesis of target molecules **218** in 31–81% yield (Scheme 48).



 R^{2} = 3-OMe, 3,5-diOMe, 2-Cl, 3-Cl, 3-Br, 2-Br-6- R^{2} = Me, C₂H₅, 3,5-diOMeC₆H₃, H

 $R^3 = NMe_2$, NHC_6H_5 , NHMe, OMe, $COCH_3$, C_3H_5 , $NHCH_2C_6H_5$, morpholine, furan-2-yl-methanamine,

(piperazin-1-yl)ethanone, cyclopentanamine

Scheme 40 Synthesis of neolignan analogs comprising of 2,3-dihydrobenzofuran 190.



Fig. 11 Structure of licarin A 191.

Wang¹¹⁵ *et al.* in 2021 accomplished the gold-catalyzed [2 + 3] cycloaddition of phenols **20** and substituted-1,3-enynes **219** to furnish dihydrobenzofuran scaffolds **221** in low to high yields (25–81%). In their synthetic methodology, the *ortho*-selectivity of phenols **20** was attained for the first time *via* electrophilic aromatic substitution employing **219** as an α -oxo vinyl gold carbenoid analogue. Catalytic amounts of ^tBuX-PhosAuCl (phosphine ligated aurum catalyst) and 2,6-

dichloropyridine *N*-oxide **222** (as an additive) were utilized in dichloroethane to achieve the desired products **221**. Gold and ligand **222** assisted 1,3-enynes **219** underwent concerted $S_N 2$ reaction and proto-deauration upon the addition of substituted phenols **20** to generate intermediate **220**. The intermediate **220** was converted to the final product **221** *via* oxa-Michael addition (Scheme 49).

Du¹¹⁶ *et al.* in 2023 disclosed a gold-catalyzed, one-pot construction of benzofuran-3(2H)-one skeletons 226 using the phenomenon of cycloisomerization of alkynyl phenols 223. In their novel methodology, *o*-alkynyl phenols 223 were reacted with substituted alcohols 224 in the presence of Ph₃PAuCl (catalyst), Selectfluor (as an oxidant) and TfOH (as an additive) to afford benzofuranones 226 in moderate to good yields (22–76%). *o*-Alkynyl phenols 223 were activated and oxidatively added to its phenolic part to form a cyclic intermediate, which was oxidized by Selectfluor. In the next step, reductive elimination took place, followed by the nucleophilic substitution



Scheme 41 Synthesis of 2,3-dihydrobenzofuran-3-ylacetic acids 193.

Review





reaction to generate intermediate **225**. This intermediate then underwent ketal hydrolysis to furnish the final products **226** (Scheme 50).

2.6. Ru-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Ruthenium, though being the rarest earth metal, is widely used as a catalyst in organic transformations. The Ru-catalyzed synthesis of dihydrobenzofuran derivatives via C-C and C-H bond functionalization has gained tremendous attention in organic synthesis. Directing groups accompanied enantioselective functionalization of C-H bonds promoted by TM catalysts has gained significant utility in several organic transformations. In 2023, Sau¹¹⁷ et al. carried out the synthesis of chiral dihydrobenzofuran derivatives 228 via rutheniumcatalyzed C-H functionalization of 3-(allyloxy)benzamides 227 in the presence of AgSbF₆, using copper acetate (as an additive) in 1,2-dichloroethane. The synthesized optically active dihydrobenzofurans were then subjected to treatment with in substituted alkynes 36 acetic acid to afford dihydrobenzofuran-fused isoquinolinones 228 in high enantiomeric excess (up to 97:3 er) along with a side product 229 (Scheme 51).

Similarly, Pannilawithana *et al.*¹¹⁸ utilized ruthenium catalyst-mediated synthesis of dihydrobenzofuran heterocycles by treating varied phenols **20** with substituted aldehydes **230**. The coupling reaction was efficiently carried out in the presence of carbon monoxide and dichloroethane as CO addition greatly enhanced the yield of benzofuran derivatives **231** (64–93%) in comparison to alkylation products **232** (side product) (Scheme 52).

Utilizing the annulation reaction strategy, another ruthenium-catalyzed approach for the facile synthesis of dihydrobenzofurans was demonstrated by Phukon¹¹⁹ *et al.* in 2023.



Scheme 43 Synthesis of substituted dihydrobenzofurans 200



Scheme 44 Synthesis of 2,3-dihydrobenzofuran 204.



Scheme 45 Synthesis 2,3-dihydrobenzofuran derivatives 207.



In the developed synthetic methodology, substituted naphthols 234 were made to react with substituted sulfoxonium ylides 233 *via* the cyclization reaction, which involved the employment of 1,4-dioxane (as a source of $-CH_2$) and copper acetate to synthesize dihydronaphthofurans 235 in 51–68% yield (Scheme 53).

In 2021, Yuan¹²⁰ *et al.* successfully designed a multicomponent, photoredox catalytic protocol that was applied for the construction of dihydrobenzofuran ring systems **240**. For this purpose, 2-vinyl phenols **44**, *N*-alkoxypyridinium salts **236** and sulfur ylides **237** were coupled in the presence of Ru(bpy)₃Cl₂- \cdot 6H₂O as an efficient photocatalyst along with CuI/ligand (L) in DABCO. Ru catalyst facilitated the formation of sulfonium ylide and alkoxy radical that got transformed into benzylic radical intermediate **238** by reacting with *in situ* formed 2-vinylphenolate. Benzylic intermediate **238** further interacted with sulfonium ylides **237** to form **239**, followed by nucleophilic substitution reaction to furnish the desired cyclized products **240** in low to good yields (26–74%) (Scheme 54).

2.7. Ir-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Iridium is the most corrosion resistance metal, which displays variable oxidation states and is utilized as a good catalyst in several organic transformations. Cross-dehydrogenative coupling (CDC) is an efficient approach to afford various organic frameworks by involving the formation of C-C bond via two inequal C-H bonds. This synthetic transformation is highly ecofriendly and requires a short duration.121 The synthesis of several sophisticated heterocycles can be achieved by the intramolecular cross-dehydrogenative coupling (first given by Fagnou and Liégault), which involves the fusion of alkyl groups to the aromatic ring. Kusaka¹²² et al. in 2022 presented the enantioselective synthesis of dihydrobenzofurans 242 exploiting intramolecular CDC $C(sp^2)$ -H/C(sp³)-H. They carried out this approach on silyl-based aryl ethers 241 using iridium-based (S)-DTBM-SEGPHOS as a catalyst and *tert*-butyl ethylene (TBE) as a hydrogen collector involving p-xylene as a solvent to furnish enantioenriched dihydrobenzofuran derivatives 243 in 53-86% yield range with up to 99% ee. The reaction was proposed to



R¹ = 10-OMe, 10-F, 11-OMe, 12-OMe, 10-OMe, 10-Me, 12-Me, 10-Et, H, 10,12-diMe, CF₃, 10-Ph, 10-COOMe, 10-NO2, 10-F, 10-Cl, 10-Br, thiophenyl $R^2 = Ph, 4-MeC_6H_4, 4-FC_6H_4, 4-CIC_6H_4, 2-CIC_6H_4, 4-OMeC_6H_4,$ R³ = Ph, 4-MeC₆H₄, 4-FC₆H₄, 4-CIC₆H₄, 4-Ph-C₆H₄, R⁴ = 6-Me, 6-Cl, 6-F, H $R^5 = Me$, diMe, H,











ОН



PCy₃

[Ru] Complex

 $R^2 = C_6H_5$, C_3H_7 , C_2H_5 $R^3 = CH_3$, C_2H_5

 R^2 , R^3 = cyclopentyl, cyclohexyl

Scheme 52 Synthesis of dihydrobenzofuran heterocycles 231.

proceed *via* oxidative addition of **241**, followed by the removal of hydrogen and ligand exchange, leading towards the synthesis of aryliridium intermediate **242**. The intermediate **242** then underwent cyclization and reductive elimination to give dihydrobenzofurans **243** in high enantiomeric excess (up to 99% ee) (Scheme 55).

TM-catalyzed intramolecular straightforward addition of aromatic C–H bonds to olefinic bonds, named hydroarylation, has presented facile pathways towards the cyclic compounds with efficient atom-economy.¹²³ Utilizing the intramolecular hydroarylation strategy, in 2021, Sakamoto and Nishimura¹²⁴ described the Ir-catalyzed, enantioselective synthesis of 2,3-



Scheme 53 Synthesis of dihydronaphthofurans 235.



2-FC₆H₄, 2-C₆H₄OMe, 2,4-FC₆H₃, naphthyl, thiophenyl

Scheme 54 Synthesis of dihydrobenzofuran derivatives 240.

dihydrobenzofuran derivatives **248** *via* C–H activated intramolecular hydroarylation. In their novel methodology, they used one-pot protocol to attain desired cyclic products **248** in good to excellent yields (62–99%). Pd-catalyst was used to generate *m*- cinnamyloxyphenyl ketones **247** from allyl carbonate **246**, followed by the Ir-catalyzed protocol to afford 2,3-dihydrobenzofuran compounds **248**. A bisphosphine ligand, *i.e.*, (*S*,*S*)-QuinoxP*, was used to attain higher enantioselectivities (>98% ee) (Scheme 56).



Scheme 55 Synthesis of dihydrobenzofuran derivatives 243.





Scheme 56 Synthesis of 2,3-dihydrobenzofuran derivatives 248



R = 5-Me, 5-OMe, 5-CF₃, 5-Ph, 5-SiMe₃, 5-Cl, 5-F, 6-Me, 6-OMe, 7-Me

In 2021, Ohmura¹²⁵ *et al.* reported the synthesis of enantioselective 2,3-dihydrobenzofuran scaffolds **251** *via* iridiumcatalyzed intramolecular hydroarylation. In their novel methodology, *tert*-butyldimethylsilyloxy-substituted aryl ethers **249** acted as starting materials that were transformed into dihydrobenzofurans **251** (featuring a stereogenic carbon at C3 position) in low to excellent yields (18–85%) with significant enantioselectivity values (33–99%) (Scheme 57).

2.8. Fe-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Being inexpensive and earth friendly, iron (Fe) and its salts have become highly enticing. These parameters led the researchers to increase focus towards the iron-based catalysis for temperate and green reactions.¹²⁶ Bashir¹²⁷ *et al.* in 2023 proposed a facile method for the synthesis of 2,3-dihydrobenzofurans **254** *via* [3 + 2] cycloaddition. In this regard, substituted hydroquinones **252** were made to react with *N*-arylated cyclic enamines **253** in the presence of a biomimetic catalyst, *i.e.*, hemin (an iron embedded porphyrin) with an oxidant partner, *i.e.*, ^{*t*}BuOOH. Low to high yields (27–91%) were obtained by employing different catalytic loadings, CH₂Cl₂ (solvent) at room temperature for 10 hours (Scheme 58). The synthesized compounds were evaluated for their anti-cancerous potential against MCF-7 cancer cells, among which the compound with 2-propyne *N*substitution exhibited potent IC₅₀ value = 27.73 μ M.

In 2021, Zhang¹²⁸ *et al.* reported the iron(III)-mediated, free radical-catalyzed cascade protocol for the synthesis of naph-thodihydrofurans **258**. In their novel approach,

Scheme 57 Synthesis of 2,3-dihydrobenzofuran 251.

Review



naphthoquinones 255 were reacted with allyl alcohols 256 in the presence of Fe(acac)₃ and Ph₂SiH₂ (used as reductant) in EtOH to access naphthodihydrofurans 258 in moderate to excellent yields (47-93%). Alkyl radical formed from 256 and naphthoquinones 255 underwent Michael addition and single electron transfer process to form intermediate 257, which was dehydrated and cyclized to yield desired products 258 (Scheme 59). The gram scale synthesis of 258 was performed to scale up the reaction and structurally important 3,3-dimethyl-2,3-dihydronaphtho[1,2-b]furan-4,5-diones were also generated from the synthesized naphthodihydrofurans 258.

2.9. Ag-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Ag-catalyzed [3 + 2] annulation reactions are the most convenient synthetic strategies for the synthesis of five-

membered heterocyclic ring compounds.⁹⁴ In this context, in 2023, Guo129 et al. executed visible-light promoted, additive-free synthesis of dihydrobenzofurans 261 and 262. They treated substituted phenols 20 with styrenes 259 and Nacylindoles 260 in the presence of nanoparticles-based silver phosphate (Ag₃PO₄) in dichloromethane or HFIP solvent, to accomplish the synthesis of dihydrobenzofurans and indolines constituting dihydrobenzofurans 261 and 262, respectively. This visible light-induced synthetic route proved to be high yielding and efficient due to the recyclable nature of nanoparticles-supported silver phosphate catalyst. On the catalytic surface, the stabilized radical cations of both substrates underwent [3 + 2] cross coupling reaction to yield the target molecules. Initially, the radical formation of substrates took place, which resulted in the formation of the radical cation intermediate. This intermediate radially



- $R^3 = Me, H$
- R^4 = Me, H, cyclohexyl
- $R^5 = 4-MeC_6H_4$, $4-OMeC_6H_4$, $4-FC_6H_4$, $4-CIC_6H_4$, $4-BrC_6H_4$, $3-CIC_6H_4$, cyclopentyl, cyclohexyl, cycloheptyl, cyclooctyl, Me, H

Scheme 59 Synthesis of naphthodihydrofurans 258





eliminates a proton to furnish the desired 2,3-dihydrobenzofuran derivatives 261 and 262 (Scheme 60). The proposed mechanism of the reaction was supposed to be initiated with the formation of radical species 265, which cross coupled with the radical cation of 259 to form an intermediate 266. Finally, this intermediate led to the synthesis of target molecules 261 upon deprotonation and cyclization (Scheme 61).

In 2021, Dias and coworkers¹³⁰ also optimized a silvercatalyzed method for the synthesis of 2,3-dihydrobenzofuran neolignans 269 and 270 via oxidative coupling strategy. In their synthetic approach, methyl p-coumarates 267 and methyl ferulates 268 were transformed into 269 and 270, respectively, in

the presence of Ag₂O (used as catalyst) and azobisisobutyronitrile (as radical inhibitor) in acetonitrile. These conditions furnished significant stability between conversion and selectivity (with 45% conversion and more than 71% selectivity) of dihydrobenzofurans 269 and 270 (Scheme 62).

2.10. Pt-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Platinum (Pt) catalysts depict astounding capacity as a carbon monoxide (CO) oxidation catalyst in the catalytic converter on account of their exceptional stability and tailoring adaptability. They are also widely utilized as electrodes in electrosynthesis. Electrosynthesis has gained significant weightage in organic



Scheme 61 Proposed mechanism for the synthesis of dihydrobenzofurans 261.



synthesis owing to the green applicability of electrons in proceeding redox reactions. However, its industrial usage is limited due to the bulk waste generation as a result of using concentrated supporting electrolytes.¹³¹ Regarding to this, Okamoto¹³² et al. in 2023 reported the synthesis of dihydrobenzofuran derivatives 271 by employing an electrochemical approach utilizing ultra-low electrolyte concentration (0.001-0.01 M). For the first time, they treated alkenes 21 and phenols 20 via electrochemically-induced [3 + 2] cycloaddition in the presence of flow microreactor using HFIP as the solvent. Bu₄NPF₆ (tetrabutylammonium hexafluorophosphate) was added with dichloromethane and acetic acid to be used as an electrolyte in the chemical reaction. Their synthetic route was highly productive and environment friendly owing to the complete elimination of waste production, thereby giving substituted dihydrobenzofurans 271 in 21-91% yield. Phenoxonium cations of 20 were formed anodically, which were further reacted with 21 to afford 2,3-dihydrobenzofuran derivatives 271 (Scheme 63).

In 2023, Guan's¹³³ group reported another catalyst free, environmentally benign electrochemical synthesis of dihydrobenzofurans 275. They treated substituted aminophenols 272 with diverse variety of olefins *via* an electrochemicallymediated approach in the presence of ${}^{n}\text{Bu}_{4}\text{NBF}_{4}$ in acetonitrile solvent to carry out the synthesis of target molecules in effective yields (33–99%). The developed route was found to be highly economical as it avoids the use of any catalyst, oxidizing agent and additive. The proposed mechanism of this reaction involved the anodic oxidation of substituted aminophenols 272 *via* single electron transfer, followed by an addition reaction with alkenes 273 to form intermediate 274. The resulting intermediate was then supposed to form the carbocation, which ultimately afforded dihydrobenzofurans 275, followed by intramolecular cyclization (Scheme 64).

2.11. V-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Vanadium (V), being a transition metal, has the capability to change its oxidation state and is used as an efficient catalyst for many reaction systems. Utilizing the V-catalyzed synthetic protocol, Wang¹³⁴ *et al.* in 2023, treated the substituted aryl acetates 277 and dihydroxy substituted naphthoic acid esters 276 *via* the [3 + 2] cascade reaction to afford dihydrobenzofuran derivatives 279 in 20–88% yield. The reaction was executed in the presence of catalytic isothiourea and vanadium oxide acetate utilizing hydrogen peroxide, *n*-Bu₄NHSO₄ and B(OMe)₃ in 1,4-dioxane. The postulated mechanism involved the formation of C1-ammonium enolate and removal of proton, followed by Michael addition and lactonization to furnish dihydrobenzofuran derivatives 279 (Scheme 65).

2.12. W-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Owing to their inexpensive and reusable characteristics, tungsten (W) catalysts are abundantly used in organic



Scheme 63 Synthesis of dihydrobenzofuran 271.







Scheme 65 Synthesis of polycyclic dihydrobenzofuran 279.

synthesis. Another visible-light promoted synthetic route to achieve dihydrobenzofurans was given in 2023 by Gowda135 et al. For this purpose, diversely substituted aromatic aldehydes 281 were made to react with alkynyl aryl ethers 280 exploiting TBADT (tetrabutylammonium decatungstate), which is a photocatalyst, responsible for carrying out 1,5hydrogen atom transfer. The reaction protocol involved the use of visible light (390 nm) in acetonitrile solvent to afford dihydrobenzofurans 283 (in 48-76% yield range) via the activation of C-H bond without any addition of additive. The plausible mechanism of this reaction was believed to include the excitation of photocatalyst via visible light, followed by the removal of proton from aromatic aldehydes 281 to give the acyl radical. The resulting radical was then assumed to execute the addition reaction with alkynyl aryl ethers 280 proceeded by 1,5-transfer of hydrogen atom to give rise to intermediate 282. This intermediate then finally gave substituted

dihydrobenzofurans **283** *via exo-dig* radical involving cyclization and subsequent reduction (Scheme 66).

2.13. Co-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives

Cobalt (Co)-based catalyst have high reducibility, high oxygen mobility and thermal stability due to which they are extensively studied and used in the field of organic synthesis. Tian¹³⁶ *et al.* in 2021 reported a facile one-pot, photosensitized and cobaltcatalyzed approach for the synthesis of 2,3-dihydrobenzofuran derivatives **289**. In their synthetic methodology, 2-propynolphenols **287** underwent semi-hydrogenation and intramolecular cyclization in the presence of a photosensitizer, *i.e.*, Ir[dF(CF₃)ppy]₂(dtbbpy)PF₆ and a cobalt catalyst (Co(OAc)₂-·4H₂O), to access 2,3-dihydrobenzofuran units **289**. This methodology covers diverse functionalities (18 examples) and gave moderate to excellent yields (54–98%). Phenolic hydroxyl



Scheme 66 Synthesis of dihydrobenzofuran derivatives 283.

played a vital role for the intramolecular cyclization with (*Z*)alkene, which was transformed from alkyne group of **287**. As per the suggested mechanism, activated Co species underwent hydrometallation with alkyne part of **287**, proceeded by protonation and oxidative addition (with phenolic hydroxyl group). The resulting intermediate was converted into intermediate **288** on migratory insertion and yielded the final products **289** *via* reductive elimination (Scheme 67).

2.14. Dual metal-catalyzed synthesis of 2,3dihydrobenzofuran derivatives

2.14.1. Pd and Cu-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives. Phthalans are structurally dihydrobenzofurans constituting organic scaffolds, which are of significant contribution in the pharmaceutical industry. In 2022, Huang¹³⁷ *et al.* reported a facile approach to procure a diverse range of substituted phthalans **294** utilizing Sonogashira coupling. Their synthetic strategy involved the reaction of substituted aryl halides **290**, nucleophiles **291** and substituted triynes **292** in the presence of catalytic amount of palladium tetrakis(triphenylphosphine) and copper iodide using diisopropyl amine as the base in tetrahydrofuran. The reaction was assumed to propagate *via* oxidative addition and palladium promoted coupling, followed by reductive elimination and base-catalyzed propargyl–allenyl isomerization to generate intermediate **293**. Intermediate **293** was then supposed to undergo PDDA cyclization, followed by subsequent regioselective nucleophilic addition to construct a wide variety of 1,3-dihydroisobenzofurans (phthalans) **294** in moderate to excellent yields (40–92%) (Scheme 68).

2.14.2. Pd and Ni-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives. Dual catalysis is one of the most



Scheme 67 Synthesis of dihydrobenzofuran derivatives 289.



Scheme 68 Synthesis of substituted phthalans 294.

dynamic strategies for the advancement of chemical reactions in organic synthesis. Lian¹³⁸ *et al.* in 2022 devised a dual catalytic protocol to synthesize 2,3-dihydrobenzofurans **298** containing *gem*-difluorovinyl framework. In their synthetic approach, the coupling of *gem*-difluorovinyl tosylates **296** and alkynyl bromoarenes **295** occurred in the presence of Pd and Ni catalyst along with ZnI₂ (as an additive) and Zn (as reductant) in DMF at 100 °C. As interpreted by the proposed mechanism, alkynyl bromoarenes **295** underwent subsequent oxidative addition and cyclization process in the presence of Pd(0) species. Next, Ni(0)-catalyzed oxidative addition of *gem*-difluorovinyl tosylates **296** took place, proceeded by the transmetallation step to generate *gem*-difluorovinyl zinc intermediate, which interacted with Pd-mediated alkynyl



R¹ = 4-Me, 3-COOMe, 4-Cl, 4-F, 4-OMe, 4-^tBu

- $R^{2} = C_{6}H_{5}, 4-PhC_{6}H_{4}, 4-C_{2}H_{4}TMSC_{6}H_{4}, 4-{}^{t}BuC_{6}H_{4}, 4-MeOC_{6}H_{4}, 4-CH_{2}CNC_{6}H_{4}, 4-COCH_{3}C_{6}H_{4}, 4-CH_{2}CNC_{6}H_{4}, 4-COCH_{3}C_{6}H_{4}, 4-MeOC_{6}H_{4}, 4-CH_{2}CNC_{6}H_{4}, 4-COCH_{3}C_{6}H_{4}, 4-MeOC_{6}H_{4}, 4-CH_{2}CNC_{6}H_{4}, 4-COCH_{3}C_{6}H_{4}, 4-CH_{2}CNC_{6}H_{4}, 4-CH_{2}CNC_{6}H_{4}, 4-COCH_{3}C_{6}H_{4}, 4-CH_{2}CNC_{6}H_{4}, 4-CH_{2}CNC_{6$
 - 4-MeCOOC₆H₄, 3-CF₃OC₆H₄, 4-FC₆H₄, 4-CIC₆H₄, 3,4 fusedC₆H₄, 4-CI-3-(4-ethoxybenzyl)benzene,
 - (5-(4-F-phenyl)thiophen-2-yl)(o-tolyl)methanone, 4-(phenylsulfonyl)morpholine, 4,4,5,5-tetramethyl-2-phenyl-
 - 1,3,2-dioxaborolane, thiophenyl, quinoline, benzo[d][1,3]dioxole
- R^3 = 3,4-fusedC₆H₃, 4-Ph, 4-Me, 3-Me, 2,3diMe, 3-Cl, 4-OCF₃

Scheme 69 Synthesis of 2,3-dihydrobenzofurans 298



bromoarenes species to form intermediate **297**. This intermediate underwent reductive elimination to furnish the desired 2,3-dihydrobenzofuran derivative **298** (Scheme 69).

2.14.3. Ti and Ni-catalyzed synthesis of 2,3-dihydrobenzofuran derivatives. Ni-catalyzed stereoselective reductive dicarbofunctionalization of alkenes is the most facile approach for the rapid generation of several asymmetric cyclic frameworks.139 Similarly, various catalysts comprising of titanium (Ti) play a crucial role in the development of dihydrobenzofuran core structure.¹⁴⁰ In 2023, Zhao¹⁴¹ et al. also utilized haloarene-substituted alkenes 300 for the asymmetric synthesis of dihydrobenzofurans 301. They reacted aromatic rings-joined alkenes with substituted benzyl alcohols 299 by employing Ni(COD)₂ and titanium chloride (TiCl₄) as catalysts in the presence of ligand, manganese, 2,6-lutidine in tetrahydrofuran achieve the enantioenriched to dihydrobenzofurans 301 in 31-51% yield with 75-87% ee (Scheme 70). The synthetic strategy was also utilized to furnish the dihydrobenzofuran derivatives 302 and 303 from clinically approved drugs, *i.e.*, bexarotene (used against T-cell lymphoma cancer) and probenecid (used in the treatment of gout), respectively.

3 Conclusion

This review article elucidates the latest developments in the transition metal-catalyzed synthesis of polysubstituted dihydrobenzofuran derivatives. Due to their exemplary output, cost effectiveness, and diversification, transition metal-mediated one-pot synthesis involving cutting-edge methodologies, *i.e.*, multiple C–C/C–O bond-forming processes in an intermolecular or intramolecular approach, annulation and insertion reactions, have been rigorously explored towards the accomplishment of these heterocycles. Various transition metals, *i.e.*, Rh, Pd, Cu, Co, Ni, Fe, Ag, Au, W, Ir, Pt, V and Ru-catalyzed robust methodologies have been scrutinized towards the synthesis of 2,3-dihydrobenzofurans derivatives along with their mechanistic details. It is aimed that this communication, by providing a comprehensive perspective of the synthetic protocols available, will assist to provoke attention in the study of dihydrobenzofurans in pharmaceutical and medicinal chemistry.

Conflicts of interest

There are no conflicts to declare.

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