Advances in the Synthesis of Ring-Fused Benzimidazoles and Imidazobenzimidazoles

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Abstract: This review article provides a perspective on the synthesis of alicyclic and heterocyclic ring-fused benzimidazoles, imidazo[4,5-f]benzimidazoles, and imidazo[5,4-f]benzimidazoles. These heterocycles have a plethora of biological activities with the iminoquinone and quinone derivatives displaying potent bioreductive antitumor activity. Synthesis is categorized according to the cyclization reaction and mechanisms are detailed. Nitrobenzene reduction, cyclization of aryl amidines, lactams and isothiocyanates are described. Protocols include condensation, cross-dehydrogenative coupling with transition metal catalysis, annulation onto benzimidazole, often using CuI-catalysis, and radical cyclization with homolytic aromatic substitution. Many oxidative transformations are under metal-free conditions, including using thermal, photochemical, and electrochemical methods. Syntheses of diazole analogues of mitomycin C derivatives are described. Traditional oxidations of o-(cycloamino)anilines using peroxides in acid via the t-amino effect remain popular.

Keywords: green chemistry; halogen; heterocycle; hydrogen peroxide; imidazole; iodine; nitrosobenzene; oxone; palladium; quinone

1. Introduction

1.1. Significance and Biological Activity

Benzimidazole is an important heterocyclic pharmacophore. Therapeutic interest was aroused in the 1950s, when 5,6-dimethylbenzimidazole was discovered as a degradation product of vitamin-B₁₂ (Figure 1) [1,2]. Benzimidazoles possess a wide range of biological activities [3–6], significantly utilized as APIs in medicines [5], and as pesticides [6]. Moreover among the ring-fused benzimidazoles, pyrimido[1,2-a]benzimidazoles were discovered as effective corticotropic releasing factor-1 (CRF-1) receptor antagonists for potential treatment of mental health disorders [7]. A pyrrolo[1,2-a]benzimidazole was found to be the most effective cyclin-dependent kinase 4/6 (CDK4/6) inhibitor evaluated within a series of highly functionalized five to seven-membered alicyclic ring-fused benzimidazoles [8]. Other ring-fused benzimidazoles, the imidazobenzodiazepines were investigated as poly(ADP-ribose) polymerase (PARP-1) inhibitors, allowing reduction of hyperglycemia with neuroprotective effects in animal models [9].

The most widely studied of the applications of ring-fused benzimidazoles and imidazobenzimidazoles is as bioreductive antitumor agents, when the fused benzene part is an iminoquinone or quinone functionality (Figures 2–5). Indole-based natural products often inspire analogue synthesis, including imidazoquinoxaline analogues of the marine natural product, wakayin [10], a topoisomerase I inhibitor (Figure 2).
agents do not contain conventional DNA damaging functionality, and cytotoxicity may
as DT-diaphorase), which is a popular target for anti-cancer studies [20]. Many anti-cancer
ligatory two-electron reductase NAD(P)H:quinone oxidoreductase 1 (NQO1, also known
electron reduction reversed by oxygen [19]. Many solid tumors also over-express the ob-
has published extensively on aziridinylpyrrolo[1,2-
alkylating aziridinyl moiety leads to cytotoxicity via the FANC DNA-repair pathway [12,13],
ductive antitumor antibiotic, which cross-links to DNA [11]. Incorporation of the DNA-

Figure 1. Benzimidazole in vitamin B12 and biologically-active ring-fused systems [1–9].

Figure 2. Imidazole-based wakayin analogues [10].

Mitomycin C (MMC) has, however, attracted most interest, as the archetypal biore-
ductive activation. NADPH-
alkylation aziridinyl moiety leads to cytotoxicity via the FANC DNA-repair pathway [12,13],
and the benzimidazole analogue of the bioactivated form of MMC (aziridinomitosene) was
prepared and evaluated against breast cancer cell lines [14] (Figure 3). Skibo has published
extensively on aziridinylpyrrolo[1,2-α]benzimidazolequinones (PBIs) as DNA-cleaving
agents with analogues showing melanoma-specific cytotoxicity [15–18].

Figure 3. Aziridinyl-functionalized antitumor agents [11–18].

One or two-electron reductases are responsible for bioreductive activation. NADPH-
cytochrome c (P450) reductase is predominant under hypoxic conditions with the one-
electron reduction reversed by oxygen [19]. Many solid tumors also over-express the
obligatory two-electron reductase NAD(P)H:quinone oxidoreductase 1 (NQO1, also known
as DT-diaphorase), which is a popular target for anti-cancer studies [20]. Many anti-cancer
agents do not contain conventional DNA damaging functionality, and cytotoxicity may be due to the formation of reactive oxygen species. Pyrido[1,2-a]benzimidazolequinone 1 (Figure 4) is more than 300 times more cytotoxic under hypoxic conditions than the clinical drug, MMC (Figure 3), with cytotoxicity for alicyclic ring-fused benzimidazoles correlated to reductive potentials [21,22]. Highly conjugated naphthyl fused benzimidazolequinone 2 leads to increased stability of reduced intermediates leading to specificity towards human cancer cell lines over-expressing NQO1 [23].

Figure 4. Benzimidazolequinones targeting over-expressed reductases in solid tumors [21–23].

Ring-fused imidazo[4,5-f]benzimidazolequinones 3a and 3b are NQO1 substrates [24,25], with 3a, at the National Cancer Institute (NCI), showing specificity towards the killing of melanoma cell lines (Figure 5) [24]. Our group was the first to provide viable synthetic protocols for accessing ring-fused imidazo[5,4-f]benzimidazoles, enabling evaluation of quinone and iminoquinone derivatives for toxicity against cancer cell lines [26–29]. Compared to alicyclic ring-fused analogues 4a and 4b, the oxygen atom of the 1,4-oxazino ring was found to increase toxicity of 4c [27]. Iminoquinone 5a isolated from the Fremy oxidation to prepare 4b, was unexpectedly the most potent imidazobenzimidazole, with more than 12 times greater cytotoxicity towards a prostate cancer cell line (DU145) than a normal fibroblast cell line (GM00637) [26]. More intensive cytotoxicity assays, computational docking, and NCI COMPARE analysis on 5a, revealed good correlation with NQO1 [28]. In contrast, isomeric imidazo[4,5-f]benzimidazole 5b was inactive against the NCI 60 cell line panel [29].

Figure 5. Imidazobenzimidazolequinones and iminoquinones [24–29].

1.2. Available Synthetic Methods

The categories of syntheses of ring-fused benzimidazoles 6 are according to the cyclization reaction (Scheme 1). Oxidative cyclizations from aniline or anilide derivatives is the most studied route (Route A) and is presented in context with the plethora of other syntheses that build the benzimidazole moiety (Routes B–D). The section on Route A is sub-divided into syntheses of benzimidazole and imidazobenzimidazole scaffolds. Lastly, there is a section on syntheses, which begin with the benzimidazole moiety (Route E), sub-divided according to reaction (type) conditions. This is not an exhaustive review, and the reader should consult reviews on polycyclic benzimidazoles for comprehensive lists of syntheses [30–33]. Since the late 1990s, Aldabbagh et al. have worked on the discovery of new ring-fused benzimidazoles and synthetic methods, and the collated articles related to their research are reviewed herein. A reviewer recommended a Scifinder® search of “benzimidazole-fused”, which was completed, and significant references are incorporated. For brevity, full papers are cited and not the preceding communication article. A historical perceptive is taken and analysis of the most significant contributions to the field is carried...
out. In particular, methodology that forms a variety of ring-fused benzimidazoles is of interest, rather than procedures that give mainly the benzimidazole core.

Scheme 1. Categorizing available synthetic methods according to the cyclization reaction A-E.

2. Syntheses of Ring-Fused Benzimidazoles and Imidazobenzimidazoles

2.1. Oxidations of o-Cycloaminoanilines and Anilide Derivatives (Route A)

There are distinct differences in the reaction mechanisms and conditions for ring-fused benzimidazole and imidazobenzimidazole formation warranting sub-division. Benzimidazoles form by oxidative cyclization of anilines via nitrosobenzene intermediates; in contrast, cyclization to give the ring-fused imidazobenzimidazole must begin from anilides and proceed via amine-N-oxide intermediates under acidic conditions.

2.1.1. Forming Ring-Fused Benzimidazoles

In 1908, Spiegel and Kaufmann reported that Caro’s acid (peroxymonosulfuric acid, \(\text{H}_2\text{SO}_5\)) oxidized 5-nitro-2-(piperidin-1-yl)aniline to 7-nitro-1,2,3,4-tetrahydropyrido[1,2-a]benzimidazole [34]. In the absence of the nitro-substituent, no oxidative cyclization occurred. Caro’s acid was already known to oxidize anilines to nitrosobenzenes [35], so supporting the idea of a nitroso intermediate. The prominent 20th century chemist and creator of Adam’s catalyst, Roger Adams with Nair refined this methodology, and accessed a range of five to seven-membered ring-fused benzimidazoles in good to high yields using peroxytrifluoroacetic acid generated in situ from \(\text{H}_2\text{O}_2\) and trifluoroacetic acid (TFA) (Scheme 2) [36]. Six-membered cyclization yields were higher, when the anilines contained a nitro-substituent. Meth-Cohn and Suschitzky [37] soon refuted the observation made by Nair and Adams that acyl derivatives do not undergo cyclization to give benzimidazoles. These workers showed a range of anilide derivatives (formyl, acetyl and benzoyl) underwent oxidative cyclizations using peroxytrifluoroacetic acid or performic acid (\(\text{H}_2\text{O}_2\) and \(\text{HCO}_2\text{H}\)). Meth-Cohn preferred the use of \(o\)-cyclic amine substituted anilides as substrates for making ring-fused benzimidazoles [37,38]. Meth-Cohn commented that Nair and Adams [36] had possibly formed the anilide in situ, due to the initial addition of TFA followed by \(\text{H}_2\text{O}_2\) [38]. Mechanisms were proposed for benzimidazole formation from anilide, via an amine-N-oxide rather than the nitroso intermediate (see Section 2.1.2). Later
the use of anilide derivatives as substrates for preparation of ring-fused benzimidazoles would become commonplace (Scheme 3) [13,15,27,37–40].

![Diagram](image-url)

**Scheme 2.** Nair and Adams oxidative cyclizations of anilines [36].

Significant amounts of seven- and eight-membered ring-fused [1,2-α]benzimidazoles 9a and 9b were formed from nitrobenzenes 7a and 7b during the one-pot catalytic hydrogenation and acetylation to give acetamides 8a and 8b (Scheme 3a) [39]. It seemed that the conformation of these large alicyclic rings favored advantageous cyclization of nitroso intermediates formed during the hydrogenation process. Acetamides 8a and 8b were cyclized using performic acid to 9a and 9b in good yields, with the former transformed to 3-aziridinylazepino[1,2-α]benzimidazolequinone [13,39]. A readily available and safer alternative to Caro’s acid is Oxone (2KHSO₅·KHSO₄·K₂SO₄) [41]. Due to the absence of organic waste products, Oxone in the presence of formic acid gives ring-fused benzimidazole [27,40] and imidazobenzimidazole [27] adducts without the requirement for chromatography. 2-Oxa-7-azaspiro[3.5]nonane acetamide gave the spirocyclic oxetane ring-fused benzimidazole 10 in good yield by simple organic extraction from the basified aqueous mixture (Scheme 3b) [40].

![Diagram](image-url)

**Scheme 3.** Acetanilide cyclizations onto (a) azepino- and azocino-[39], and (b) spirocyclic oxetane-[40] derivatives.

Preparations of ring-fused benzimidazoles using 9- cyclic amine substituted anilines with performic acid compare favorably with the derivative aniline reaction, with Smalley et al. reporting moderate to good yields for five- to seven-membered adducts (Scheme 4a) [42]. In a more recent study, recyclable ethyl acetate (EtOAc) replaced formic acid, with aqueous effluent, organic-aqueous extraction and chromatography avoided for the preparation of pyrrolo[1,2-α]benzimidazoles from commercial 9-(pyrrolidin-1-yl)anilines (Scheme 4b) [43].
Although, the presence of strong electron-withdrawing substituents (NO₂, CN) and the six-membered cyclization required methanesulfonic acid (MsOH) to reach high yields. However, MsOH is a green acid undergoing biodegradation by forming CO₂ and sulfate [44].

![Scheme 4](image)

**Scheme 4.** Aniline cyclizations using (a) performic acid [42] and (b) acid-free conditions [43].

Alternatives to peroxide-based oxidizing systems, include MnO₂ in cold chloroform, but yields of ring-fused benzimidazoles from o-cycloaminoanilines were 15–20% due to presumed formation of azo-compounds [45]. Möhrle and Gerloff reported the use of a Hg(II) EDTA complex to deliver ring-fused benzimidazoles, in quantitative yield, apart from the morpholino compound, synthesized in 47% yield (Scheme 5) [46].

![Scheme 5](image)

**Scheme 5.** Hg(II)-mediated oxidative synthesis [46].

The cross dehydrogenative coupling (CDC) involves forming the C-N bond directly from C-H and N-H bonds under oxidative conditions with a formal loss of H₂, in a process often catalyzed by transition metals. CDC is used to describe pentamethylcyclopentadienyl Ir(III)dichloride ([Cp*IrCl₂]₂) catalyzed oxidative cyclization of o-tetrahydroisoquinoline substituted aniline derivatives (Scheme 6a) [47]. The bulk around the primary amine dictated regioselectivity. The o-cyclic amine substituted aniline gave the benzimidazo[2,1-α]isoquinoline 12, while the more hindered acetamide derivative gave the alternative kinetic product 13. The formamide has less steric bulk than the acetamide forming a mixture of the thermodynamic and kinetic products. The reaction was extended to the synthesis of pyrrolo-, pyrido-, and azepino[1,2-a]benzimidazoles, without the requirement for a ligand (Scheme 6b), but was less successful for making morpholino- and piperazino-ring-fused analogues [48].
The isoindoline and tetrahydroisoquinoline substrates are the easiest to oxidize at high temperatures, including in the presence of TEMPO in air (Scheme 7a) [49], and catalytic iron(III) [50]. The latter gave the highest yields for the benzimidazo[2,1-a]isoquinoline systems (Scheme 7b). 

Scheme 6. Ir(III)-mediated cyclizations (a) with ligand [47] and (b) without ligand [48].

Scheme 7. Thermal oxidative cyclizations mediated by (a) TEMPO/air [49] and (b) Fe(III) [50].

Aniodic oxidation gave the required iminium ion 14 for cyclization (Scheme 8) [51]. The electrolyte was n-Bu4NPF6 (20 mol%), and the anode is reticulated vitreous carbon (RVC), and Pt is the cathode in an undivided cell at a constant current of 10 mA. A Russian team reported the electrochemical oxidative cyclization with reduction of nitrobenzene for cyclization onto an o-piperidinyl-substituent to give pyrido[1,2-a]benzimidazoles [52].
intermediate of the reaction between H₂O₂ and HCl.

Salt of hypochlorous acid (HOCl) is the active ingredient in domestic bleach and is the

In the early 1970s, o-cyclic amine substituted anilines reacted in neat sulfuryl chloride

More recently, we heralded the use of H₂O₂ with hydrohalic acid (HX), as a convenient

Domestic bleach gave cyclization and dichlorination of aniline 15 in 56% yield (Scheme 11a),

Scheme 8. Electrochemical oxidative cyclizations [51].

Scheme 9. Synthesis of polychlorinated ring-fused benzimidazoles using SO₂Cl₂ [53].

Scheme 10. Molecular halogen (X₂) generated from H₂O₂ and HX [54–56].

Domestic bleach gave cyclization and dichlorination of aniline 15 in 56% yield (Scheme 11a),

with the lower yield attributed to the requirement for chromatography to separate the addi-
ditives in the bleach [54]. Moreover, using H₂O₂/HX a library of selectively dichlorinated and dibrominated ring-fused benzimidazoles was prepared in high yields from com-
cmercial o-cyclic amine substituted anilines, with most cases not requiring chromatography

(Scheme 11b,c). 5-Fluoro-2-piperidinylaniline was an exception, giving significant amounts

cyclization with monochlorination or monobromination. Bromination tended to be slower than chlorination, and tribrominated product 16 was isolated for the o-(pyrroloidin-
1-yl)aniline, due to difficulties in cleanly isolating 5,7-dibromopyrrolo[1,2-a]benzimidazole (Scheme 11d).

Scheme 11. One-pot ring closure with selective halogenation using (a) domestic bleach or H\textsubscript{2}O\textsubscript{2}/HCl, (b) H\textsubscript{2}O\textsubscript{2}/HCl, and (c) H\textsubscript{2}O\textsubscript{2}/HBr, and (d) five to eight-membered ring-fused adducts [54].

3,6-Dimethoxy-2-(cycloamino)anilines underwent 6-electron oxidations to afford a variety of ring-fused halogenated benzimidazolequinones, when using higher amounts of HCl or HBr relative to H\textsubscript{2}O\textsubscript{2} (Scheme 12a) [55]. The active species is the elemental halogen (X\textsubscript{2}) with water required for quinone formation (Scheme 12b). When less in situ halogen was generated, using [H\textsubscript{2}O\textsubscript{2}] > [HX], the 4-electron oxidation occurred, to give ring-fused halogenated benzimidazoles (Scheme 12c).

The use of hydroiodic acid (HI) is preferred when oxidative cyclization is required without halogenation [56], due to the relatively smaller electrophilicity of iodine [57]. Five- and seven-membered cyclizations of 3,6-dimethoxy-2-(cycloamino)anilines with H\textsubscript{2}O\textsubscript{2} and a catalytic amount of HI in EtOAc proceeded in high yield (Scheme 13a), but 1,4,6,9-tetramethoxyphenazine 17, was unexpectedly formed, as an orange precipitate from six-membered cyclizations (Scheme 13a,b) [56]. The absence of phenazine 17 from the five- and seven-membered cyclizations was consistent with previous observations that six-membered oxidative cyclizations are more difficult [36,43]. The formation of 17 was optimized by reducing the amount of EtOAc (the reaction solvent) by four-fold and by decreasing the reaction temperature to room temperature (Scheme 14). Moreover, the isolation of 17 was indicative of a nitrosobenzene intermediate in the conversion of o-(cycloamino)anilines to
ring-fused benzimidazoles via the so-called t-amino effect [58]. Syntheses of phenazines involve nitroso intermediates [56,59,60]. Recent evidence in the synthesis of ring-fused benzimidazoles, included GC-MS of the reaction mixture (Scheme 14), after 1 h, which revealed EI-MS fragmentation pattern consistent with intermediate 18 [56].

Scheme 12. One-pot transformations of anilines to halogenated ring-fused benzimidazolequinones using (a) H2O2/HX, (b) Cl2 and Br2 with water, and (c) H2O2/HX without quinone formation [55].

Scheme 13. H2O2/HI-mediated (a) five- to seven-membered cyclizations and (b) six-membered cyclizations with phenazine formation [56].
3,4-Dihydro-1H-[1,4]oxazino[4,3-a]benzimidazole can however be prepared in good yield in the absence of acid from 2-(morpholin-4-yl)aniline using Oxone at room temperature (Scheme 15) [61]. Sampling the reaction at short reaction times (within 2 min), gave 4-(2-nitrophenyl)morpholine (20), which formed through advantageous air oxidation of the nitroso intermediate 19. The proposed mechanism postulates that KHSO₅ (the active ingredient of Oxone) is catalytic, consistent with the use of catalytic amounts of HI in the H₂O₂/HI-mediated oxidative cyclization of 3,6-dimethoxy-2-(cycloamino)anilines (Schemes 13 and 14) [56].

Jana et al. reported one-pot sequential amination of nitrosoarenes with alicyclic amines, followed by oxidative cyclization to give ring-fused benzimidazoles (Scheme 16a–c) [62]. Oxone oxidation of commercial anilines gave the nitrosoarene substrates. Nucleophilic aromatic hydrogen substitution (SNArH) led to the pyrrolo[1,2-c] benzimidazoles (Scheme 16a). Nucleophilic ipso-substitution of nitrosonaphthols with cyclic amines and subsequent oxidative cyclization delivered a diverse range of ring-fused naphthoimidazoles (Scheme 16b). The authors proposed a mechanism with mass spectrometry detecting the o-cyclic amine substituted phenyl hydroxylamine 21, which oxidized to the o-cyclic amine substituted nitrosoarene 22 (Scheme 16c). A formal 1,5-hydride shift gave the key iminium intermediate 23 for oxidative cyclization [58]. In contrast to our recent work [43,56,61], the nitrosoarene cyclizations, were carried out using elevated temperatures and under acidic conditions [62].
Scheme 16. Sequential aminations of nitrosoarenes to prepare (a) alicyclic ring-fused benzimidazoles, (b) naphthoimidazoles, and (c) the proposed mechanism [62].

2.1.2. Forming Ring-Fused Imidazobenzimidazoles

Oxidative cyclizations in this class date back to the 1950s and 1960s [37,63], and Skibo et al. used the performic acid mediated protocol to convert diacetamides to dipyrrolo- and dipyrido-ring-fused imidazo[4,5-f]benzimidazoles, but yields were low (Scheme 17a) [24,25]. One-pot double alkyl radical cyclizations onto the 2- and 6-positions of imidazobenzimidazole gave dipyrrolo-, dipyrido-, and diazepino-ring-fused imidazo[4,5-f]benzimidazoles and imidazo[5,4-f]benzimidazoles in 47–90% yield (see Section 2.5.3) [26]. However, the radical cyclization route cannot easily give imidazobenzimidazoles containing two different fused rings. Oxone in acid enabled the synthesis of symmetrical and unsymmetrical ring-fused imidazo[4,5-f]benzimidazoles and imidazo[5,4-f]benzimidazoles (Scheme 17b–d) [27,29,61]. Yields were higher than older oxidative methods [24,25,37,63], perhaps due to the easier work up, allowing isolation of the imidazobenzimidazole directly from the acidic reaction mixture using precipitation with solid Na₂CO₃ [27]. It is noteworthy that 49 and 55% yield for the spirocyclic oxetane ring-fused isomers, represent ≥70% yield for each ring closure [61].
Anilide reactant and acidic conditions are a prerequisite for oxidation to the imidazo[4,5-f]benzimidazole, with attempts to cyclize 4,6-di(piperidin-1-yl)-1,3-phenylenediamine (24) giving an intractable mixture (Figure 6) [61]. Meth-Cohn proposed the oxidative cyclization of acetamides to benzimidazole derivatives occurs via the amine-N-oxide intermediate 25 [38]. Isolated amine-N-oxides undergo acid-mediated benzimidazole and imidazobenzimidazole formation [27,61].

Figure 6. Key chemical structures in resolving the mechanism [27,37,38].

The diamine-N-oxide intermediate 26 for imidazo[5,4-f]benzimidazole formation was isolated by Fagan [27]. The X-ray crystal structure of 26 showed hydrogen bonding between the amide NH and the amine N-oxide residues, supporting the absence of the amide NH peaks in the $^1$H NMR spectra of solutions of amine N-oxides [27,38,61]. This is contrary to the orientation of the amine-N-oxide 25 adopted in Meth-Cohn’s Polonovski-type reaction...
Our proposed mechanism begins with oxidation of the cyclic amines of di-acetamide 27 to the Fagan amine-N-oxide orientation 26 (Scheme 18). Protonation in acidic media gives the imidols, upon loss of water. The double intramolecular nucleophilic imidol addition onto the iminum ion leads to the ring-fused imidazo[5,4-f]benzimidazole [27].

An acid-catalyzed cyclization mechanism was proposed for conversion of dimorpholine N-oxide 26 to the [1,4]oxazino[4,3-a]benzimidazole (Scheme 19a) [27]. In the absence of an external oxidant, oxidation is possible through the internal conjugated system. MsOH allowed conversion of amine N-oxide 28 to the imidazo[4,5-f]benzimidazole, where presumably another molecule of 28 acts as oxidant (Scheme 19b) [61].

2.2. Reductions of Nitrobenzene-o-Cycloamines (Route B)

The reduction under acidic conditions of the aromatic nitro-group with cyclization onto the adjacent cycloamine substituent dates to the 1960s and employ ZnCl2/Ac2O [16, 64,65], TiCl4/HCl [66], and Fe/AcOH [67]. There are cyclizations using Pd-catalysis with CO [68] or H2 [69]. Recent metal-free conditions use visible light, phenylthiourea as catalyst and PhSiH3 as reductant [70], and electrochemical cyclizations [52]. Thermal annulation using nitrobenzene substrates are possible with neat 1,2,3,4-tetrahydroisoquinoline (THIQ) (Scheme 20a) [71], and cyclizations occur using I2/HCO2H [72] (Scheme 20b). For the former reaction, the authors speculated on THIQ acting as a hydride donor after the initial SNAr and redox cyclization, while HI generated in situ, is the active catalytic species in the latter reaction, acting as a strong Brønsted acid and reductant [71,72].
the initial SnAr and redox cyclization, while HI generated in situ, is the active catalytic species in the latter reaction, acting as a strong Brønsted acid and reductant [71,72].

Scheme 20. Reductive cyclizations using (a) thermal conditions [71] and (b) I2/HCO2H [72].

2.3. Using Aromatic Amidines, Lactams, and Isothiocyanates (Route C)

Non-redox cyclizations of o-haloarylamidines give ring-fused benzimidazoles, using CuI as catalyst [73,74] (Scheme 21a) or strong base (Scheme 21b) [8,75], where nitrogen displaces the o-halogen. Pyrrolo[1,2-a]benzimidazole was reported from an o-lactam substituted aniline cyclization using di-t-butyl sulfoxide/NBS via an aza-Wittig type-reaction (Scheme 21b) [76]. 1-(2-Isothiocyanatophenyl)pyrrolidines cyclize under thermal acidic conditions to benzimidazothiazepines (Scheme 21c) [77].

Scheme 21. Synthesis from (a) o-haloarylamidine [75], (b) a lactam derivative of o-phenyldiamine [76], and (c) a rearrangement of aryl isothiocyanates [77].

Hypervalent iodine(III) reagents allow cyclization of aryl amidines onto non-functionalized benzenes (leaving group = H) [78,79]. The overall dehydrogenative process is proposed to proceed via a homolytic aromatic substitution (HAS) of nitrogen-centered radicals giving pyrrolo- and pyrido[1,2-a]benzimidazoles (Scheme 22a) [78]. Kosher’s reagent (PhI(OH)OTs)
gave a range of thiazino and oxazino ring-fused benzimidazoles (Scheme 22b) [79]. The oxazino ring-fused benzimidazole required a longer reaction time, and the use of 4Å molecular sieves.

![Scheme 22. Cyclization of aryl amidines using (a) phenyliodine(III) diacetate [78] and (b) Kosher’s reagent [79].](image)

A combination of Chan−Lam and Ullmann type couplings were said to give the fully unsaturated pyrido[1,2-α]benzimidazoles using 2-aminopyridine and 2-iodoarylboronic acids via an amidine intermediate (Scheme 23) [80].

![Scheme 23. Synthesis of pyrido[1,2-α] benzimidazoles via amidine intermediates [80].](image)

### 2.4. Condensations (Route D)

This includes the traditional bimolecular condensation of 1,2-phenylenediamines with aldehydes (Scheme 24) [81,82]. Rh-catalyzed cyclization of N-alkenyl-1,2-diaminobenzenes with CO/H₂ gas gave alicyclic ring-fused benzimidazoles via a hydroformylation intermediate [83].
Scheme 24. Acid-catalyzed condensation of 1,2-phenylenediamine with o-phthalaldehyde and glutaraldehyde [82].

2.5. Annulations onto Benzimidazoles (Route E)

A widely reported category for the synthesis of ring-fused benzimidazoles is annulations onto the 1- and 2-positions of benzimidazoles. This section reviews syntheses over the past 20 years, sub-divided according to the employed reaction (type) conditions.

2.5.1. Base-Mediated Methods

Early examples of annulation of the benzimidazole moiety cyclized 2-haloalkylbenzimidazoles under basic conditions through N-1 benzimidazole deprotonation to give pyrrolo-, pyrido- and azepino[1,2-α]benzimidazoles [84,85]. bromoethylsulfonium salt disintegrated under basic conditions with loss of diphenyl sulfide, upon reaction with benzimidazole-2-methanols to give oxazino[4,3-α]benzimidazoles (Scheme 25) [86]. Extension of the latter sulfur ylide approach, achieved thiazino- and piperazino-adducts, but in lower yields.

Scheme 25. Annulations [4 + 2] of benzimidazoles using bromoethylsulfonium salt [86].

2.5.2. Transition Metal and Lewis Acid Catalyzed Methods

InCl$_3$-catalyzed the synthesis of benzimidazole-fused 1,4-oxazepines by intramolecular addition of a pendant alcohol onto an in situ-generated imine (Scheme 26) [87].

Scheme 26. Synthesis of benzimidazole-fused 1,4-oxazepines [87].

Starting from 2-arylbenzimidazoles and aryl iodides, a tandem Pd(II)-catalyzed C-H arylation and aerobic oxidative C-H amination sequence gave a variety of benzimidazole-fused phenanthridines (Scheme 27) [88]. Arylated intermediate 29 underwent C−H activation in a seven-membered palladacycle with the nitrogen of benzimidazole to yield the aminated products.
Scheme 27. Pd(II)-catalyzed synthesis of benzimidazole-fused phenanthridines [88].

There are several transition metal mediated benzimidazole-2-CH activated annulations including onto benzimidazole N-alkenyl [89–91] and N-alkynyl [92] substituents, with high enantioselectivity also achieved [90,91]. These include the use of chiral Ni/JoSPOphos manifolds (Scheme 28) [91].

Scheme 28. Asymmetric pyrido[1,2-α]benzimidazole syntheses [91].

Ru(II)-catalyzed [4+2] annulations of 2-arylbenzimidazole with styrenes yielded benzimidazo[2,1-α]isoquinolines (Scheme 29) [93].

Scheme 29. Annulations [4+2] of benzimidazoles using Ru(II)-NH and CH activation [93].

DeBoef et al. employed Pd(II)/Cu(I) catalysis with the oxidant of Cu(OAc)2 to effect oxidative dehydrogenative coupling of 1-benzyl substituted benzimidazoles (Scheme 30a) [94]. The Pd(II) C-H aryl-activation of 2-phenylbenzimidazoles with C-N coupling of two benzimidazole fragments gave benzimidazoquinazolines, with Cu(OAc)2 or air being the terminal oxidants (Scheme 30b) [95].
Scheme 30. Pd-catalyzed dehydrogenative cross-couplings to give (a) 11H-isoindolo[2,1-a]benzimidazoles [94] and (b) benzimidazoquinazolines [95].

Scheme 31. CuI-catalyzed annulations onto N-1 of benzimidazole using (a) 1,1-dibromoalkenes [96] and (b) cyanamide [97].

Cul-catalyzed an aminothiolation of 1,1-dibromoalkenes (Scheme 31a) [96]. The mechanism involves initial in situ dehydrobromination of the 1,1-dibromoalkene to 1-bromoalkyne. N-alkylation of the aromatic alkene and Cu-catalyzed C-S coupling or 5-endo-dig cyclization gives the annulated benzimidazole. More recently, 2-(2-bromoaryl)- and 2-(2-bromovinyl)benzimidazole substrates were coupled with cyanamide using Cul to give 2-aminoquinazoline and 2-aminopyrimidine hybrid structures (Scheme 31b) [97]. Similar conditions allowed Cul-mediated coupling and cyclization between 2-(2-bromoaryl)indoles and 2-aminoazoles [98], including 2-aminobenzimidazoles with loss of ammonia to give indole analogues of Scheme 30b adducts [95].

Scheme 31. Cul-catalyzed annulations onto N-1 of benzimidazole using (a) 1,1-dibromoalkenes [96] and (b) cyanamide [97].
A Pd-catalyzed reductive cyclization gave a variety of benzimidazole-fused thiazocine scaffolds (Scheme 32), but also thiazonines (9-membered rings), and thiazecines (10-membered) having an exocyclic double bond [99]. Ammonium formate is the hydrogen-donor.

![Scheme 32. Pd-catalyzed macrocyclization to give benzimidazole-fused thiazocine [99].](image)

2.5.3. Radical Cyclization Methods

We have reported annihilations utilizing the benzimidazole-2-position to form nucleophilic ylide for condensation reactions [100,101] and the benzimidazol-2-yl radical for initiator-free photochemical HAS (Scheme 33a) [102]. Photochemical intramolecular HAS of the imidazol-2-yl radical are superior to analogous Bu3SnH-mediated reactions [103].

![Scheme 33. Radical initiator-free HAS using (a) UV-light [102] and (b) Barton ester intermediates [104].](image)

Bu3SnH-mediated chain reactions enabled 6-exo-trig cyclizations of aryl or pyridinyl radicals to yield a variety of aromatic-ring fused benzimidazoles [23,105] (Scheme 34a). HAS using alkyl radical cyclizations onto benzimidazole [21], imidazo[4,5-f]benzimidazole and imidazo[5,4-f]benzimidazole [26] proceed as non-chain reactions [106–108] requiring full equivalents of Bu3SnH and azo-initiator to give the aromatic products in moderate to high yields (Scheme 34b). Single ring-fused imidazo[5,4-f]benzimidazolequinones for anti-cancer evaluation studies, were also derived via this radical cyclization route (Scheme 34c) [28]. The latter cyclization required a substrate prepared by two separate alky-
lations of imidazobenzimidazole using 1-chlorobutane and 1-chloro-4-(phenylselenyl)butane. For Bu$_3$SnH-mediated reactions, slow addition of radical initiators is required, which prevents Bu$_3$SnH reduction of the cyclizing radical. Camphorsulfonic acid (CSA) or Ac$_2$O are added to more difficult cyclizations, to activate the benzimidazole-2-position towards nucleophilic radical addition [21,26]. For the HAS onto imidazobenzimidazoles (Scheme 34b,c), exposing the reaction mixture to air for part of the reaction time gave higher yields, since oxygenated radicals are thought to be involved in the oxidative re-aromatization (or hydrogen atom-abstraction) step [26,108].

Scheme 34. Bu$_3$SnH-mediated formation of ring-fused benzimidazoles using (a) chain reaction [23,105], and non-chain reaction (b) double [26] and (c) single [28] HAS.

An alternative approach to HAS, is initiator-free, using in situ generated Barton ester intermediates derived from robust and readily accessible carboxylic acid substrates, giving five- to seven-membered alkyl and cyclopropyl radical cyclizations via a chain reaction mechanism (Scheme 33b) [104,109]. Most recently, benzimidazo[2,1-$a$]isoquinolin-6-ones were prepared by the addition of methyl radicals onto N-methacryloyl-2-phenylbenzimidazoles using di-tert-butyl peroxide (DTBP) initiator (Scheme 35) [110]. The tertiary adduct radical undergoes six-membered cyclization followed by oxidation of the derived cyclized radical. Earlier, tert-butyl radicals generated from pivalic acid and several other carboxylic acids with K$_2$S$_2$O$_8$/AgNO$_3$ initiated an analogous cascade [111].
2.5.4. Other Metal-Free Methods

Mal et al. reported the use of PIFA (PhI(OCOCF$_3$)$_2$) for six-membered aromatic substitutions via a proposed nitrenium ion intermediate (Scheme 36a) [112]. Cho et al. made the reaction at N-1 of benzimidazole intermolecular, with addition onto aryl isocyanate and aromatic substitution of the carbamoyl-NH to give fused quinazolinones and pyrimidinones (Scheme 36b) [113].

There are now several reports of benzimidazole syntheses using iodine in metal-free procedures [56,72,114]. Benz[4,5]imidazo[1,2-a]quinoxaline derivatives were obtained by an I$_2$-mediated oxidative condensation of 2-(benzimidazol-1-yl)aniline substrates (Scheme 37) [114].
2.5.5. Miscellaneous: Syntheses of Mitomycin Analogues

We reported the synthesis and anti-cancer evaluation of diazole analogues of MMC derivatives (Figure 3) and related indolequinones [14,21,22,115,116]. The formation of pyrrolo-, pyrido-, azepino- and azocino[1,2-
\a]benzimidazoles with a fused cyclopropane ring involves N-aziridinyl imines (Eschenmoser hydrazones) undergoing thermolysis with a loss of nitrogen and trans-stilbene (Scheme 38a) [22,115,116]. The formation of the aziridinomitosene analogue skeleton was via an intramolecular anionic aromatic ipso-substitution of an aziridinyl functionality onto the benzimidazol-2-yl position (Scheme 38b) [14]. Earlier the 2-phenylsulfanylbenzimidazole precursors gave five to seven-membered ring-fused benzimidazoles using alkyl radical cyclizations with substitution of the radical leaving group at the 2-position [117].

3. Conclusions

Over the past 20 years, significant advances in the synthesis of ring-fused benzimidazoles have occurred, notably using dehydrogenative coupling and radical cyclization. Transition metal catalysts achieve intramolecular and intermolecular aminations with benzimidazoles, with enantioselectivity for the former. Ring-fused benzimidazoles are prepared
using hypervalent iodine(III) reagents and elemental iodine under metal-free conditions. There is an increasing use of sustainable and non-metal-mediated protocols, including photochemical, electrochemical, and thermal methods. Mild oxidative conditions tend to be more effective for preparing isoindoline and tetrahydroisoquinoline-fused scaffolds. There are effective methods for incorporating heteroatoms into the fused-ring, including N, O, and S atoms, and forming alicyclic rings with additional fused cyclopropane or delicate oxetane and aziridine rings. In terms of versatility, green chemistry, and value for money, it is difficult to beat the use of H$_2$O$_2$ in traditional oxidative cyclizations of o-(cycloamino)anilines. H$_2$O$_2$ in combination with HX generates the ordinarily inconvenient elemental halogen (X$_2$) in situ, to mediate one-pot oxidative cyclization with halogenation. The latter includes one-pot approaches to potential antitumor ring-fused benzimidazole-quinones from readily accessible anilines. Oxone is a cheap alternative, with technical and environmental benefits, including the circumvention of organic waste by-products. Our present work generates X$_2$ in situ, by combining Oxone with benign NaX, to carry out one-pot halogenation with oxidative demethylation to give the dimeric quinones of ring-fused dimethoxybenzimidazole-benzimidazolequinones (DMBBQs) [118]. The DMBBQ scaffold offers unique regioselective functionalization opportunities for bis-quinone motifs, and these unique dimeric structures require investigation as bioreductive anti-tumor agents.

Mechanisms for ring-fused benzimidazole and imidazobenzimidazole formation via the $t$-amino effect are now better defined. Recent studies have shown that acid and heat are unnecessary for oxidative cyclization via a nitrosobenzene intermediate using o-(cycloamino)anilines as substrates for benzimidazole formation. This offers opportunities for further investigations into the synthesis of ring-fused benzimidazoles under nonaggressive ambient conditions using commercial anilines as starting materials. While use of anilide derivatives results in a different mechanism via an amine-N-oxide intermediate. The anilide derivative and acidic conditions are mandatory for peroxide-mediated ring-fused imidazobenzimidazole preparations. To date ring-fused imidazobenzimidazoles have only been prepared via oxidative cyclizations of anilides and radical cyclizations onto imidazobenzimidazoles, surely new synthetic methods will merge for this interesting scaffold.

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