Keywords:
bicyclo[1.1.1]pentanes - bioisosteres - propellanes - radical reactions - small-ring systems
“Escaping from flatland”, by increasing the saturation level and three-dimensionality of drug-like compounds, can enhance their potency, selectivity and pharmacokinetic profile. One approach that has attracted considerable recent attention is the bioisosteric replacement of aromatic rings, internal alkynes and tert-butyl groups with bicyclo[1.1.1]pentane (BCP) units. While functionalisation of the tertiary bridgehead positions of BCP derivatives is well-documented, functionalisation of the three concyclic secondary bridge positions remains an emerging field. The unique properties of the BCP core present considerable synthetic challenges to the development of such transformations. However, the bridge positions provide novel vectors for drug discovery and applications in materials science, providing entry to novel chemical and intellectual property space. This Minireview aims to consolidate the major advances in the field, serving as a useful reference to guide further work that is expected in the coming years.

1. Introduction

Over the past three decades, the application of bicyclo[1.1.1]pentane (BCP) derivatives in materials science as molecular rods,[1] molecular rotors,[2] supramolecular linker units,[3] liquid crystals,[4] FRET sensors[5] and metal–organic frameworks[6] has been extensively investigated and is the subject of a recent review.[7] Since the seminal reports of Barbachyn (1993)[8] and Pellicciari (1996),[9] the BCP motif has also emerged within drug discovery as a valuable bioisostere for internal alkynes, tert-butyl groups and mono-substituted/1,4-disubstituted arenes.[10] The appeal of the abiotic BCP fragment, among related systems such as cubanes and higher bicycloalkanes, originates from its ability to add three-dimensional character and saturation to compounds. Increasing the fraction of sp3-hybridised carbon atoms in a drug molecule, Fsp3, has been found to make a lead oral drug compound “more developable”[11] and correlates positively with clinical success.[12] To this end, multiple research groups have documented the increased or equal solubility[13] of lead compounds that can be achieved through such bioisosteric replacements. Increasing the solubility and potency of a medicine can reduce the therapeutic dose required, potentially avoiding drug–drug interactions and drug-induced liver injury through metabolic activation. Bioisosteric replacement has also been identified as a strategy to circumvent Markush structure patent claims on drug candidates.[14]

Multiple reviews have addressed the 12 + unique synthetic approaches that have been reported for construction of the BCP framework.[7,10,17] In particular, carbene insertion into the central bond of bicyclo[1.1.0]butanes, and nucleophilic/radical addition across the central bond of [1.1.1]propellanes, have emerged as the two most practical and scalable methods (Scheme 1). Methodology now exists to install most fragments of interest in drug discovery at the bridgehead (1,3) positions. In contrast, methodology for substitution of the bridge (2,4,5) positions remains underdeveloped.[19] Most reported methods forge the key substituent bond prior to construction of the BCP core itself. For reasons outlined in Section 2.2, examples of the direct functionalisation of the bridge positions in a controlled manner are rare.

Bridge-substituted BCP derivatives represent novel chemical space and provide novel vectors for ligand–target interactions in drug discovery. As successful drug discovery becomes more challenging year on year,[19] accessing new chemical space becomes increasingly important. Introduction of the abiotic BCP motif may also provide a new strategy for overcoming genetic resistance to existing therapeutic compounds. Brown has recently commented on this issue,[20] highlighting that abiotic scaffolds may prove important in the discovery of anticancer, antibiotic, antiviral and antiparasitic drugs, where the causative agents can be highly proficient at developing resistance to established treatments.

Aspects of the bridge functionalisation of BCP derivatives have been previously addressed by Michl and co-workers.[17a] Herein, we seek to update this work and provide a summary, from a synthetic perspective, of the known methods for the preparation of bridge-substituted BCP derivatives reported up to mid-2021. The discussion is organized by the nature of the substituent installed. Further distinction is drawn between strategies that install the substituent before synthesis of the BCP core, and those proceeding via direct functionalisation of the hydrocarbon skeleton.
2. Structural Chemistry of the Bicyclo[1.1.1]pentane Core

2.1. General Considerations

The BCP core is highly strained yet remarkably stable, with a strain energy of 65.17a–68.21 kcal mol⁻¹ for the parent hydrocarbon (cf. 27.5 kcal mol⁻¹ for cyclopropane21) and thermal stability up to ca. 300°C17a,22. This strain primarily arises from significant destabilizing overlap of the rear lobes of bridgehead orbitals directed at substituent positions.23 In many cases, this transannular communication attenuates and directs the reactivity of the cage.

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John Murphy was born in Dublin and educated at the University of Dublin (TCD) and the University of Cambridge. After postdoctoral fellowships at Alberta and Oxford, he was appointed as Lecturer, then Reader, at the University of Nottingham. Since 1995, he has held the Merck-Pauson Professorship at the University of Strathclyde. His interests are in reaction mechanisms and synthesis.

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2.2. Orbital Hybridisation and C–H Bond Strengths in the BCP Core

The hybridisations of bridge and bridgehead carbon orbitals used for substituent bonding on the parent hydrocarbon have been established as sp².5 and sp².0 respectively through JCH coupling constant analysis,24 in good agreement with values obtained from ab initio25 and semiempirical26 calculations. The high s character of these orbitals has two effects: 1) bicyclopent-1-yl and bicyclopent-2-yl fragments are net electron-withdrawing,17a and 2) BCP C–H bonds have a large homolytic bond dissociation energy (BDE), greater than that of typical saturated hydrocarbons. Indeed, Maillard and Walton have demonstrated that hydrogen atom abstraction by tert-butoxyl radicals proceeds more rapidly from cyclopropane than from the parent BCP.27
Furthermore, although simple consideration of hybridisation suggests that the bridge C–H bonds should be weaker than bridgehead C–H bonds, only the bridgehead radical is observed by EPR on exposure of BCP to tert-butoxyl radicals. This regioselectivity is remarkable considering that the bridge and bridgehead radicals have approximately equal calculated enthalpies of formation,[28] and that the bridge hydrogens have a threefold statistical advantage over the bridgehead hydrogens. Aside from the bridgehead C–H bonds being more sterically accessible,[29] the selectivity arises from the aforementioned transannular interactions within the BCP core. The transition state for hydrogen atom abstraction by an electronegative radical is polarised, with development of partial positive charge on the adjacent carbon. During hydrogen abstraction from a bridgehead site, this positive charge can be extensively stabilised by the transannular bridgehead orbital interaction.[37a] In contrast, the bridge carbon does not have access to such stabilisation and so hydrogen atom transfer (HAT) from this position is kinetically inhibited. Moreover, if one bridgehead position is functionalised with an electron-withdrawing group, this intracage stabilisation is compromised and so the expected selectivity of C–H abstraction from a bridge position is returned (see Section 4.1).[30]

In summary, the high BDE of BCP C–H bonds means that a direct HAT approach for decoration of bridge positions would likely be confined to a very limited range of substrates; they could possess no other alkyl C–H bonds, and blocking of unfunctionalised bridgehead positions may be necessary. More recently, attempts at directed C–H activation chemistry on the bridge positions have also proven unsuccessful.[33] Consequently, prefunctionalisation at the bridge positions is expected to be necessary for the divergent introduction of substituents.

2.3. Reactive Intermediates on BCP Bridge Positions

The generation and properties of reactive intermediates (carbocations, carbanions and radicals) on the bridgehead and bridge positions of BCP derivatives have been extensively discussed by Michl.[37] Key points relevant to the synthesis of bridge-functionalised derivatives are:

1. Bridge-centered carbenium ions are unstable, undergoing rapid and irreversible skeletal rearrangement to ring-opened products (see Sections 3.1, 4.1 and 5.2). As such, they are not viable intermediates for the preparation of bridge-substituted BCP derivatives.

2. Bridge-centered carbanions are kinetically stable against ring opening, and are putative intermediates in the Haller–Bauer cleavage and Birch reduction of 2-substituted BCP derivatives (see Section 5.2).

3. Bridge carbon-centered radicals are kinetically stable against β-scission to afford cyclobutyl products, even though this process is calculated to be strongly exothermic.[32]

3. Oxygen Substituents

3.1. Bicyclo[1.1.1]pentan-2-ol and Derivatives

Methods for the preparation of BCP bridge alcohols are rare. Attempts at direct C–H oxidation using persulfate salts have been unsuccessful.[30] The parent bicyclo[1.1.1]pentan-2-ol (14b) has been most efficiently accessed through Baeyer–Villiger oxidation of mixed ketones 12 (see Section 5.1), hydrolysis of the resulting esters and separation of the isomeric alcohols (Scheme 2A).[30] Compound 14b has also been accessed in low yield through reduction of the corresponding ketone (see Section 3.2). The 2-substituted phenyl analogue (16a) is more accessible,[33] representing the major product of the photolysis of cyclobutyl phenyl ketone via (thermally reversible)[33a,b] Norrish–Yang (NY)[44] cyclisation (Scheme 2B). The NY cyclisation strategy is compatible with electron-poor and electron-neutral substrates (16b–f), but is less tolerant of electron-rich systems such as vinyl and 2-furyl cyclobutyl ketones.[35]

While the lithium salt of the parent bicyclo[1.1.1]pentan-2-ol is stable in aprotic solvents (Scheme 2A),[49] increased strain at the bridge position means that 2-substituted bicyclo[1.1.1]pentan-2-ol alcohols are unstable. Exposure of alcohols 16a and 17 to catalytic NaOMe in methanol results in rapid cycloreversion to the corresponding cyclobutyl phenyl ketones,[33b,36] analogously to the ring-opening of cyclopropyl alcohols.[37] Deuterium-labelling studies have identified two distinct pathways for ring opening (Scheme 3); however, both are driven by release of ring strain in the transition state.[31c] Ring opening occurs rapidly at cryogenic temperatures; attempts at O-alkylation of the alkoxide of 16a with methyl iodide or dimethyl sulfate have been unsuccessful.[31a] Alkylation is presumably retarded by the decreased nucleophilicity of the alkoxide from the electron-withdrawing effect and steric demand of the BCP moiety.

Moreover, the p-bromophenylcarbamate derivative of alcohol 16a has been prepared through direct reaction with the isocyanate without prior deprotonation.[33] Similarly, nitrobenzoyl esters 20[38] and 21[39a,10] have been prepared by treatment of alcohols 14b and 16a, respectively, with the
appropriate benzylic chloride. Solvolysis of these esters leads only to ring-opened products, illustrating the instability of BCP-centered carbocations. Similarly, treatment of free alcohol 16a with acetic acid produces 22 in 40% yield and the corresponding acetate in 60% yield. The mechanism of rearrangement has been studied kinetically and spectroscopically. It is proposed that departure of the leaving group is assisted by participation of an adjacent C–C σ-bond (Scheme 4). Rearrangement, presumably via housane cation II, provides cyclopentenyl cation III which may be trapped by nucleophiles. While cation I has been observed by NMR spectroscopy under cryogenic, superacidic conditions, cation II was not detected. In the absence of nucleophiles, III is observed as allylic cation IV arising from formal Wagner–Meerwein rearrangement.

In summary, access to the parent bicyclo[1.1.1]pentan-2-ol (14b) remains extremely inefficient (Schemes 2 and 5). Additionally, the instability of 14b and derivatives makes them challenging synthetic intermediates to handle, unless they can be protected as solvolytically stable functionalities under strictly neutral conditions.

3.2. Bicyclo[1.1.1]pentan-2-one and Derivatives

Ketone 24 has been prepared by Dougherty through ozonolysis of alcohol 16a. NaBH₄ reduction then provided alcohol 14b in 23% yield (Scheme 5). The poor yield of the ozonolysis reaction precludes 24 as a useful synthetic intermediate for the preparation of 2-substituted BCP derivatives. Compound 24 is thermally unstable, undergoing cycloreversion at elevated temperatures to afford allylketene and derived oligomers/nucleophilic trapping products. Cycloreversion is much more facile than for typical cyclobutanones. Ketalisation of 24 has only been achieved through irradiation in CD₃OD in the absence of base.

The related 4,5-dimethylenebicyclo[1.1.1]pentan-2-one (30) was later prepared by Dowd from tricyclopentane 25 where generation of the highly strained BCP scaffold is compensated by extrusion of dinitrogen from unstable intermediate 29 (Scheme 6). The instability of 30 similarly precludes its use as a synthetic intermediate to access bridge-substituted BCP derivatives.

4. Halogen Substituents

Only bridge-fluorinated, -chlorinated and -brominated BCP derivatives are known. While fluorine and chlorine substituents have been introduced through both dihalocar-
bene insertion into the central bond of bicyclo[1.1.0]butane precursors and through direct reaction of BCP derivatives with the elemental halogens, bromine has only been introduced through dibromocarbene insertion.[30,43]

4.1. Bridge-Chlorinated derivatives

The dichlorocarbene insertion method of Cherkofsky[43a] (Scheme 1) is arguably the most convenient approach to bridge chlorine-substituted BCP derivatives. Both chloroform[43a] and sodium trichloroacetate[13d] are viable carbene precursors. Monodechlorination is readily achieved using tin[30] or silicon[13d] hydride radical chemistry, facilitated by stabilisation of the transition state for chlorine abstraction by the remaining halogen atom. Removal of the second halogen atom, to afford a bridge-unfunctionalised BCP derivative, is possible but requires forcing reaction conditions and extended reaction times.

The direct chlorination of BCP derivatives has been extensively investigated and represents the first reported direct functionalisation of the BCP core. Photochemical chlorination has been demonstrated using both BuOCl[44] and elemental Cl2[29] by Wiberg, with 22.5:1 and 7:1 selectivity for bridgehead:bridge substitution, respectively (Scheme 7). Notably, the preference for bridgehead substitution contrasts with the direct chlorination of other bicyclic hydrocarbons (e.g. norbornane) which typically give zero or little bridgehead substitution.[29]

Repeating the chlorination with Cl2 at higher chlorine concentration increased the total yield to 50%, with a broadly similar product distribution except with zero production of 39.[29] This result was initially taken to affirm the kinetic stability of the BCP bridgehead radical: only when the supply of Cl2 is limited is the radical sufficiently long-lived to rearrange to give the ring-opened cyclobutane product. However, this was later disputed by Della and co-workers,[45] who argued that 39 instead arises from heterolysis of the bridgehead C–Cl bond of 1-chlorobicyclo[1.1.1]pentane and rearrangement of the resulting cation.

In 1989, Michl demonstrated the bridge 2,2-dichlorination of 1,3-disubstituted BCP derivatives with gaseous Cl2 (Scheme 8A).[30] Bicyclo[1.1.1]pentane-1-carboxylic acid, bearing no substituent at one bridgehead site, also underwent selective bridge dichlorination. As discussed in Section 2.2, this arises from the electron-withdrawing carboxyl group decreasing the stabilisation conferred to the bridgehead radical through the 1,3-transannular interaction. No significant quantities of monochlorinated products were observed in any case: this was rationalised through stabilisation of the

Scheme 7. Direct chlorination of the parent BCP hydrocarbon.

Scheme 8. A, B): Direct chlorination of BCP derivatives. C): Protodechlorination of chlorinated BCP derivatives with TTMSS (tris(trimethylsilyl)disilane). [a] Reaction mixture quenched with MeOH; yield refers to that of the obtained diester. [b] Hydrolytic workup then (COCl)2/MeOH esterification; yield refers to that of the obtained diester. [c] Isolated as the methyl ester through (COCl)2/MeOH esterification; yield refers to that of the obtained ester. AIBN = azobisisobutyronitrile; TFA = trifluoroacetic acid.
transition state for the second HAT event by the installed chlorine atom. While these studies increase the synthetic utility of the previous work of Wiber, they are still limited by the requirement for Cl₂ gas and the use of CCl₄, CFCl₃ and TFA solvents.

More recently, the same group has also accessed tri- and tetrachlorinated BCP derivatives through direct chlorination (Scheme 8B). Elevated temperatures and gaseous Cl₂ were required for effective trichlorination, while elevated temperatures and pressurised liquid Cl₂ were required for tetrachlorination. The regio- and stereoselectivity of polychlorination was also rationalised. TTMSS/AIBN reduction of 2,2,5-trichlorinated BCP 43 gave access to the 2,5-dichlorinated species, while attempted TTMSS/AIBN reduction of 2,2,5,5-tetrachlorinated BCP 44 unexpectedly led only to decomposition products (Scheme 8C).

Finally, it should be noted that bridge-chlorinated BCP derivatives likely cannot be accessed through functionalisation of 2,2-dichloro[1.1.1]propellane (47). Compound 47, prepared in 1992 by Michl and co-workers was found to be highly susceptible to oligomerisation and loss of a bridge chlorine atom, producing unstable cation 49 (Scheme 9). This loss of chlorine is consistent with that reported previously for a higher gem-dichloropropellane. By analogy with computational studies on gem-difluorinated BCP derivatives the instability of 47 could be rationalised through angular distortion of the propellane cage caused by geminal bridge substitution, which would favour an increased C-C-C valence angle at the bridge. This strain can be partially relieved either through oligomerisation to 48, or through loss of chloride and subsequent cage rearrangement. It is therefore postulated that other propellanes with bridge substituents that are competent leaving groups will also be unstable.

4.2. Bridge-Fluorinated Derivatives

Bridge-fluorinated BCP derivatives have garnered significant interest in the past two years. Unsurprisingly, direct fluorination of 1,3-disubstituted BCPs with limited gaseous F₂ is extremely unselective, producing 14 out of the 15 possible bridge-fluorinated derivatives. Exhaustive fluorination gives access to penta- and hexafluorinated derivatives in synthetically useful yields. Recent attempts at controlled fluorination on the BCP core through the electrochemical method of Baran and the amine-directed, Pd-catalysed approaches of Xu and Hrdina have all proven unsuccessful. Only via difluorocarbene insertion have bridge-fluorinated BCP derivatives been prepared in a controlled fashion.

The concurrent works of Mykhailiuk and Ma disclosed the first examples of this chemistry in 2019, using the Ruppert–Prakash reagent (TTMS-CF₂) and the Dolbier reagent (TFDA, F₅SO₂CF₂CO₂TMS) as difluorocarbene precursors, respectively (Scheme 10). Anderson has also recently demonstrated difluorocarbene insertion using a zwitterionic phosphonium carboxylate precursor. Preliminary results towards the generation of 2-chloro-2-fluorobicyclo[1.1.0]pentane derivatives, using CHFCl₂ as a chlorofluorocarbene source, were also presented by Mykhailiuk. Across both reports, the reaction scope was limited to aryl- or vinyl-based reagents (a) or (b). TFA derivatives bearing both electron-neutral and electron-poor aromatic substituents are accessible (Scheme 10).

The stability of the difluorinated BCP motif to acidic, basic, reducing and forcing palladium cross-coupling conditions was also demonstrated between the two reports. Although vinyl or aryl substituents were essential for the difluorocyclopropanation, both can subsequently be oxidised to carboxylic acids and further derivatised. This was later exploited in a further publication by Ma in 2020 demonstrating various transformations of gem-difluorinated BCPs at the bridgehead positions.

A further general limitation of dihalocarbene insertion reactions arises from the nature of the synthesis of the starting
5. Carbon Substituents

BCP derivatives bearing bridge carbon substituents have been accessed in four fundamental ways: through chlorocarbonylation of a BCP precursor; through NY cyclisation of cyclobutyl aryl ketones; from substituted [1.1.1]propellanes; and from intramolecular nucleophile displacement on a substituted cyclobutane. The synthesis of tricyclo[2.1.0.0\(^{2,5}\)]pentanes (where two bridge positions on the BCP core are directly bonded), and miscellaneous synthetic routes that afford bridge carbon-substituted BCP derivatives as minor products, are beyond the scope of this review. Both topics are addressed by Michl.[17a]

The installation of carbon substituents on the BCP core through intermolecular carbene insertion into a bicyclo[1.1.0]butane (Scheme 1) has not yet been demonstrated, although there are a small number of reports of a related intramolecular reaction providing [1.1.1]propellanes (see Section 5.3). Attempts at intermolecular delivery of a CH₂ fragment via photolysis with diazomethane[56] and under Simmons–Smith conditions have been unsuccessful.[13d,57] Attempted reaction of bicyclo[1.1.0]butanes with bis(carbomethoxy)carbene was also unsuccessful.[58] Experimental and computational studies of the mechanism of carbene insertion into bicyclo[1.1.0]butanes have been previously discussed.[17a,58]

5.1. Carbon Substitution through Direct Chlorocarbonylation

Acyl chloride 11b represents the first example of a BCP derivative bearing a bridge carbon substituent. The product distribution of chlorocarbonylation of 31 with oxalyl chloride corresponds to a selectivity of 17:1 for bridgehead:bridge substitution. As described previously, manipulation of mixed acyl chlorides 11 also gave first access to bicyclo[1.1.1]pentan-2-ol (14b) (Scheme 11).

\[ \text{31} \xrightarrow{(\text{COCl})_2, \text{hv}} \text{COCI} \]

\[ \text{Scheme 11. Preparation of acyl chlorides 11a,b through photochemical chlorocarbonylation of the parent BCP hydrocarbon.} \]

5.2. Carbon Substitution through Norrish–Yang (NY) Cyclisation

The synthetic utility of the NY cyclisation of cyclobutyl phenyl ketone (Section 3.1) was explored by Alexander in the 1970s[38] and later in 1993[39] by Wiberg (Scheme 12). In two steps from alcohol 16a, Alexander prepared mandelate ester 52 which underwent photochemical extrusion of CO₂ to afford 53. Jones oxidation then provided ketone 54. Under irradiation,[50] a second NY cyclisation afforded tricyclo[2.2.0.0^{2,5}]hexanol 55. Alternatively, treatment with NaN₃[50] effected formal Haller–Bauer cleavage to afford 2-phenylbicyclo[1.1.1]pentane (57). Presumably the generation of an intermediate benzyl anion promotes formation of 57 rather than the bicycloalkyl carbamate, the typical product of the Haller–Bauer reaction. Compound 57 was more conveniently accessed by Wiberg[50b] through acylation of 16a and subsequent reduction with metallic sodium.

Direct deoxygenation of 16a with Li/NH₃ was unsuccessful due to rapid cycloreversion to cyclobutyl phenyl ketone (15a) under basic conditions (see Section 3.1). Reductive deoxygenation of 16a using Et₂SiH/BF₃·Et₂O and halogenative deoxygenation under Appel conditions were also unsuccessful.[58] Both resulted in fragmentation of the BCP core, further corroborating the instability of the putative bridge carbocation. Both Alexander and Wiberg demonstrated oxidative deamination of the aromatic ring of 57 to the carboxylic acid, which provided access to 2-benzoylbicyclo[1.1.1]pentane (61) and 2-nitrobicyclo[1.1.1]pentane (60) via bicyclo[1.1.1]pentyl-2-amine.

Across two studies in 2005[59] and 2012,[61] Xia, Yang et al. reported findings from a study of enantioselective photochemical solid-state NY cyclisations of ketones 63. Asymmetry originates from the homochiral environment of the crystalline lattice of the solid salts. Ketones 63 were prepared from alcohols 62, presumably originating from carboxylic acid 59 or a derived ester. Ultraviolet irradiation of the solid ketones gave the corresponding tricyclic alcohols 64 with modest enantioselectivity (maximum 90%[60] and 60%[61] ee). Extended reaction times and/or higher reaction temperatures were necessary to achieve satisfactory conversion in the solid state; however, increased conversion was generally to the detriment of asymmetric induction.

5.3. Carbon Substitution through the Use of Substituted [1.1.1]Propellanes

[1.1.1]Propellanes are attractive intermediates for the synthesis of BCP derivatives, as their unique electronic profile[62] allows opening of the central bond with a broad variety of carbon- and heteroatom-centered radicals and nucleophiles (treatment with electrophiles[63] or transition metal catalysts[64] affords exclusively ring-opened products). As such, [1.1.1]propellanes could represent key divergent intermediates for the library synthesis of BCP derivatives in drug discovery. A variety of bridge-substituted [1.1.1]propellanes have been prepared, including 2,2′-bonded dimers and those featuring spirocyclic attachment of another ring, either through a polar cyclisation strategy[65] or through intramolec-
ular carbene insertion.\[63\] However, many of these substituents are simple hydrocarbons or contain few functional handles for further derivatisation.

Furthermore, relatively few substituted propellanes have then been opened to afford monomeric bicyclo[1.1.1]pentane derivatives.\[64a,d–h,65a,66\]

Some have been polymerised to afford [\(n\)]staffane chains; this has been recently reviewed by Bräse.\[67\] Co-polymerisation of substituted propellanes with acrylates has also been studied, as well as the preparation of bridge-substituted-BCP-containing polyamide polymers.\[67\]

Simple examples of the functionalisation of bridge-substituted propellanes were provided by Szeimies,\[64a,65a\] with more recent advances from the groups of Kaszynski,\[64f\] Mazal,\[64g,h\] Baran\[31b\] and Ma.\[68\]

Kaszynski\[64f\] and Mazal\[64g,h\] demonstrated the preparation of 1,2,3,4-tetrasubstituted BCP derivatives via functionalisation of substituted [1.1.1]propellanes accessed from benzvalene (65). In one approach (Scheme 13A),\[64g\] diimide reduction of benzvalene afforded alkane 66. The acidity of the bicyclo[1.1.0]butane scaffold then enabled construction of the propellane motif in four steps according to the original method of Szeimies.\[64a\] Photochemical radical opening with biacetyl furnished intermediate 68, whose acyl groups were oxidised via haloform reaction to provide diacid 69. Subsequent treatment with liquid Cl\(_2\) resulted in exclusive halogenation of the two-carbon bridge over the BCP core, consistent with the BDE trends described previously (Section 2.2). Treatment of the obtained mixed chlorides with sodium amide furnished olefin 70, which underwent ozonolysis in high yield to afford cis-endo-tetrasubstituted BCP derivative 71. Compound 71 then served as a common precursor for the synthesis of four other tetrasubstituted analogues (72–75) through conventional carbonyl manipulations. Acid 73 spontaneously formed anhydride 76 during isolation, which went to completion on heating above 80°C for several hours.

In the second approach (Scheme 13B),\[64f,h\] osmium-mediated dihydroxylation of 65 followed by ketalisation delivered dioxolane 77. Construction of the propellane motif, then photochemical radical opening with biacetyl furnished bridged BCP 79. After oxidation to 80,\[64h\] the dioxolane moiety was deprotected prior to Swern oxidation to furnish unstable 1,2-diketone 81. Condensation of 81 with \(o\)-phenylenediamine afforded stable BCP-fused quinoxaline 82. Alternatively, the ketone moieties of 79 were transformed into vinyl phosphates or nonaflates, whereupon elimination with LDA furnished bis-alkyne 83.

In 1993 and 1995 reports, Schlüter and co-workers reported the synthesis of five propellanes containing a protected and free hydroxymethyl functionality (Scheme 14).\[64d,e\] These were accessed through a modified Szeimies procedure employing functionalised olefin 86. Aside from the synthesis of [\(n\)]staffane polymers, the synthetic chemistry of these propellanes would surprisingly not be explored until late 2020 in the works of Ma\[68\] and Baran.\[31b\]

The work of Ma delineated the synthesis of analogues of propellane 88 through a similar method. The generated propellanes (95) were then subjected to the strain-release amination conditions of Baran\[99\] furnishing 1-amino-2-functionalised BCP products (96) (Scheme 15). MOM, TBS and

![Scheme 12. NY cyclisation of cyclobutyl phenyl ketone and manipulation of resulting alcohol 16a. The synthesis of compounds 14b/24 and 60 (described in Sections 3.1 and 6, respectively) are also included for clarity. “quant” = quantitative; R* = enantiopure chiral fragment.](https://www.angewandte.org)
Scheme 13. Synthesis of bridge-substituted BCP derivatives via substituted propellanes prepared from benzvalene. LDA = lithium diisopropylamide; TMS = trimethylsilyl; pTsOH = p-toluenesulfonic acid; nonaflate = nonafluorobutanesulfonate.
benzyl O-protecting groups were all compatible with the synthetic sequence. A dioxolanylmethyl substituent at C2 was also tolerated. A small range of secondary amines were proficient substrates, including allylic/benzylic amines and piperidine derivatives; however, sterically hindered amines, cyclopropylamines and pyrrolidine derivatives were unsuccessful. After methoxymethyl cleavage, the alcohol functionality of bicyclopentane 97 was derivatised to the corresponding fluoride, azide, thioacetate and N-(4-fluorophenyl)carbamate under standard conditions.

These studies represent the first peer-reviewed investigation into a convenient and general strategy for the synthesis of bridge-substituted BCP derivatives. However, the chemistry is currently limited by its poor performance with more complex amines and the difficulty of achieving simultaneous 1,3-disubstitution across the propellane; preliminary attempts at aminoalkylation under the conditions of Gleason,[70] and aminoalkenylation under the Zweifel conditions of Aggarwal,[71] were unsuccessful and extremely low-yielding, respectively. For the bioisosteric replacement of an ortho- or meta-substituted benzenoid fragment this may not be necessary, but having equal access to both 1,2-disubstituted and 1,2,3-trisubstituted BCP derivatives would greatly increase the synthetic utility of the chemistry and broaden the chemical space it can access. Restricting studies to the synthesis of 1,2-disubstituted BCP derivatives can arguably never fully capitalise on the three-dimensionality of the BCP core.

In the report of Baran, propellane 88 was prepared analogously to the Schlüter approach; treatment with chloroiodomethane under the conditions of Anderson[72] then delivered compound 101 in 33% yield (Scheme 16). Radical de-iodination provided 102, whose chloromethyl handle was then derivatised to the corresponding acetate (103), alcohol (104), carboxylic acid (105) and Cbz-protected amine (106).

Compounds 103 and 104 were further used to prepare analogues of six drugs containing ortho- or meta-substituted benzene rings, including MTP inhibitor lomitapide (107), Hh signalling inhibitor sonidegib (109) and histamine H1 antagonist meclizine (111) (Figure 1). For the latter and three other

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**Scheme 14.** Synthesis of propellanes 88 and 90–93. Reagents and conditions: i) NaH, triethyl phosphonoacetate, 83%; ii) LiAlH4, AICl3, 90%; iii) dimethoxymethane, PhMe3N+Cl−, 30%; iv) Ac2O, Et3N, DMAP; v) MeLi, 57%; vi) HCl, MeOH, 90%; vii) TBSiCl, imidazole, 95%; viii) MeLi, 55%; ix) TBAF; x) Ac2O, Et3N, DMAP; xi) BnNCO, Et3N.

**Scheme 15.** Preparation of C2-functionalised bicyclo[1.1.1]pent-1-yl-amines. DB18-c-6 = dibenzo-18-crown-6.

**Scheme 16.** Preparation of compounds 101–106. DMM = dimethoxy methane; DPPA = diphenylphosphoryl azide.
examples, both constitutional isomers about the BCP framework were prepared, differing in which substituents of the benzene ring occupy the bridge and bridgehead positions. For the direct appendage of aromatic rings to the BCP core, conversion of BCP carboxylic acids to $N$-hydroxyphthalimide (NHP) esters permitted application of the group thermal decarboxylative Negishi arylation chemistry (generalised in Scheme 17). In summary, the work of Baran broadens the range of synthetic transformations known for decoration of the BCP core, and provides the first examples of bridge-substituted BCP analogues of drug compounds. Future work will help elucidate the ortho vs. meta character of 1,2-disubstituted BCPs in this context. The work confirms the viability of thermal decarboxylation of redox-active esters on the bridge positions of BCPs (previously demonstrated only on the bridgehead positions), but the full synthetic potential of the generated radical is yet to be explored. As discussed, future work focusing on 1,2,3-trisubstitution of the BCP system will further increase the value of the developed methods within drug discovery.

5.4. Carbon Substitution through Intramolecular Nucleophilic Displacement

In early 2021, Qin presented an elegant contemporary adaptation (Scheme 18B) of the original intramolecular displacement methodology employed in the first synthesis of bicyclo[1.1.1]pentane (Scheme 18A). Cyclobutanones 118 were accessed through the group’s boron-preserving cross-coupling of aldehydes 117 followed by ketal hydrolysis. Conversion of 118 to mesitylsulfonyl hydrazones, followed by addition of Cs$_2$CO$_3$, furnished intermediate diazo compounds 119. Trapping of the pendant boron atom delivered high-energy zwitterionic intermediates 120, which rearranged to afford 2-substituted bicyclo[1.1.1]pentanes 121 via 1,2-metalate rearrangement with extrusion of dinitrogen. A key benefit of the developed process lies in its robustness: in contrast to reactions employing propellanes, this reaction can be performed in air with only a small decrease in yield. Additionally, as stoichiometric water is generated during sulfonylhydrazone formation (Scheme 18B, step ii), the reaction is entirely tolerant of trace water in reagents and solvents. A range of secondary BPin esters were viable substrates, providing bridge alkyl- (122) and cycloalkyl- (123–125) substituted BCP derivatives. A primary BPin ester could also be cyclised to afford the bridge-unsubstituted product. Tertiary BPin esters also successfully underwent cyclisation, providing geminal bridge-disubstituted products (e.g. 126). Heteroaryl (125), isopropyl ester (127a), vinyl (127b), ethynyl (127c), methyl, amide and carbamate groups were tolerated as substituents on the cyclobutane ring (R). A positive correlation was identified between the steric demand of the “R” substituent and the reaction yield: this is a consequence of increased preference for the BPin-containing side chain to adopt the pseudo-axial position required for cyclisation. This requirement means that cyclobutanones with small “R” substituents are not viable substrates; compounds 128a,b, with R = BPin (A-value = -0.42 kcal mol$^{-1}$) [75] and R = H, respectively, were not accessible using this methodology. The introduction of a bridge methyl substituent was also demonstrated through α-substitution on the cyclobutaneone precursor, rather than substitution adjacent to the boron centre. Finally, asymmetric borylation under the conditions of Aggarwal provided an enantioenriched secondary BPin ester cyclisation precursor, which was cyclised with only 3% erosion of enantiopurity to afford BCP derivative 129. To our knowledge, this represents the first example of asym-

![Figure 1. Structures of lomitapide (107), sonidegib (109) and meclizine (111) and their BCP-containing analogues (108, 110, and 112a/b, respectively).](image)

![Scheme 17. General approach to arylation of the BCP core through decarboxylative Negishi arylation. DIC = diisopropylcarbodiimide; DMAP = 4-(dimethylamino)pyridine.](image)
metric functionalisation of the bridge positions of BCP derivatives.

The BPin group of compound 123 was derivatised to hydroxyl, vinyl and heteroaryl functionalities under standard conditions. Matteson reaction \[76\] provided the homologated primary BPin ester, while conversion to the trifluoroborate salt further enabled proto-deboronation and Suzuki cross-coupling reactions. The trifluoroborate salt could also be employed in another round of boron-preserving cross-coupling chemistry with a ketone, providing another means of formal homologation of the C–B bond.

In summary, the work of Qin provides a conceptually distinct method for the synthesis of bridge-substituted BCP derivatives. The developed methodology is operationally simple and can deliver nonsymmetrical 1,3-di-, 1,2,3-tri-, and 1,2,2,3-tetrasubstituted BCP products containing versatile functional handles for downstream derivatisation. However, a limitation is arguably its reduced divergency compared to the methods of Baran and Ma, wherein a [1.1.1]propellane provides an early common intermediate to potentially hundreds of unique derivatives. With the present method, bespoke cyclisation precursors must be synthesised (asymmetrically, if required) for each fundamental series of compounds, limiting its applicability to late-stage functionalisation. Only the BCP products post-cyclisation represent points of divergence.

6. Nitrogen Substituents

Surprisingly, none of the works of Baran, Ma or Qin considers the synthesis of BCP derivatives containing bridge nitrogen substituents. This is in contrast to the considerable progress made towards installation of nitrogen atoms at the bridgehead positions, for example, for bioisosteric replacement of anilines, common structural alerts and toxicophores in drug design. To date, nitrogen has only been installed on the bridge positions of BCP derivatives through Schmidt rearrangement of carboxylic acid 59 (Schemes 12 and 19) by Wiberg \[35b\], Alexander previously reported the same transformation in 1978 \[35b\] however, this claim was not substantiated by characterisation data for the amine product. As expected, the electron-withdrawing BCP core decreases the basicity of 130 compared to other cycloalkylamines.

Peroxyacid oxidation of 130 then furnished the corresponding nitro compound (60) in 24% yield \[35b\]. Like nitrocyclopropane and in contrast to nitrocyclopentane and nitrocyclohexane, 60 was found to be insoluble in aqueous NaOH. This is attributed to its increased pK\textsubscript{a} compared to other nitrocycloalkanes, a consequence of the reluctance of the bridge carbon atom to engage in π-bonding with associated increase in strain.

7. Boron Substituents

As part of their synthetic campaign, Baran and co-workers aimed to establish a route to a boron-containing BCP as a high-value building block. After propellane 88 was opened with benzyl iodoacetate, borylated BCP 134 was accessible via the group’s copper-catalysed decarboxylative
borylation methodology in five linear steps (Scheme 20). No transformations of 134 were then explored; however, the sequence further validates decarboxylative radical transformations as a general manifold for the bridge functionalisation of BCP derivatives.

The methodology of Qin[73] also provided BCP derivative 133 containing a bridge BPin substituent (Scheme 21). Likewise, no transformations of the bridge functionality were then explored.

8. Summary and Outlook

The works of Baran, Ma and Qin highlight two general synthetic strategies with potential to address the long-standing question of the preparation of bridge-functionalised BCP derivatives. The use of a substituted [1.1.1]propellane intermediate could facilitate divergent synthesis through the exploitation of its broad and ambiphilic reactivity profile. The application of decarboxylative radical transformations, dozens of which have now been reported,[78] adds a further dimension of divergency. This is in stark contrast to the prior art in this field, which has been sporadic and largely reliant on inconvenient and low-yielding synthetic methods.

With decarboxylative radical transformations on the BCP core successfully demonstrated, in the near future we are likely to witness the pursuit of a greater range of related transformations. The application of enabling technologies such as photoredox catalysis and synthetic electrochemistry will likely be central to this development. As such transformations often proceed under very mild conditions,[79] the late-stage bridge functionalisation of BCP derivatives may become commonplace. Judicious selection of different radical precursor groups, or the identification of synthetic sequences that allow the installation of a single radical precursor group at different stages, may also prove to be an effective strategy for the functionalisation of multiple bridge positions. In addition, the deployment of dual catalysis platforms such as metallaphotoredox catalysis, organophotoredox catalysis and mediated electrochemistry, could provide access to bridge substituted BCPs in an asymmetric fashion. This would provide a complementary approach to that illustrated by Qin, and would not require the synthesis of enantioenriched precursors.

Efficient asymmetric access to densely functionalised bicyclo[1.1.1]pentanes is likely to be transformational within drug discovery. The inclusion of such moieties into lead compounds increases their 3D character and stereochemical complexity without significantly increasing molecular weight, allowing a deeper exploration of chemical space and the design of molecules which can better complement the spatial subtleties of target proteins.[12b,80] Increased potency and selectivity of lead compounds may reduce off-target effects: drug promiscuity commonly results in adverse toxicity, a major cause of drug attrition in the clinic.[12b] “Escape from flatland” promises to be a powerful solution to this widespread and intractable problem: we are apparently on the verge of a new era in the synthetic chemistry of the bicyclo[1.1.1]pentane core.

Acknowledgements

J.M.A. is grateful to GSK for a studentship via the GSK/University of Strathclyde Centre for Doctoral Training in Synthetic and Medicinal Chemistry. The authors thank the EPSRC for funding via Prosperity Partnership EP/S035990/1 and GSK for financial support.

Conflict of Interest

The authors declare no conflict of interest.

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