Synthetic Studies Towards Spirocyclic Imine Marine Toxins Using N-Acyl Iminium Ions as Dienophiles in Diels–Alder Reactions

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1 Introduction

Oceanic microalgae can proliferate to concentrations of up to millions of cells per liter under favorable conditions, forming ‘algal blooms’ which are often visible as patches of red or brown on the ocean surface. Often, the species of microalgae involved produce toxins which accumulate in the tissues of feeding shellfish and can then pass on to humans when they are ingested. Harmful algal blooms are responsible for the death of wildlife and human illness and are therefore considered an increasing global health concern.

The toxins produced by marine dinoflagellates often display unique biological activities, rendering them potential pharmacological candidates. In particular, cyclic imine toxins have garnered particular interest from the synthetic community due to their synthetic complexity and potent activity. Named after their common structural motif (shaded in blue, Figure 1), these toxins can be further classified by the size of the cyclic imine ring, with either 7,6-spirocyclic systems (as in the spirolides, e.g. spiriolide A (1), pinnatoxins, and portieratoxins) or 6,6-spirocyclic systems (as in gymnodimine A (2) and recently isolated kabiromine (3)) being the most common. Portimine A (4), isolated in 2013, is an example of a cyclic imine toxin bearing a 5,6-spirocyclic imine system. Reviews regarding the classification and biological evaluation of these toxins are numerous. In particular, the toxicity displayed by the pinnatoxins, portieratoxins, and spiriolides stem from their ability to activate Ca2+ channels, while gymnodimine and members of the spirolide family have demonstrated affinity for nicotinic acetylcholine (nACh) receptors. Meanwhile, portimine A (4) displays high in vitro cytotoxicity and promotes apoptosis via caspase-3 activation in P388 leukemia cells.
The spirocyclic imine motif is postulated to be an important component of the active pharmacophore of these toxins, since spirolides E and F (derivatives of spirolide A (1) possessing a hydrolyzed keto amine motif)\textsuperscript{14} and a spirocyclic amine analogue of gymnodimine\textsuperscript{15} all lacked biological activity. Work by Romo and co-workers suggests that the cyclic imine component acts as a masked enamine, and the mechanism of action may involve reaction of this latent nucleophile.\textsuperscript{16} Therefore, there is immense value in these spirocyclic imine fragments as pharmacological probes, and convenient synthetic access to these systems is highly sought after.

**Biographical sketches**

**Jared L. Freeman** completed his PhD in 2019 from the University of Auckland under the supervision of Distinguished Professor Dame Margaret Brimble and Dr. Daniel Furkert. His PhD focused on the synthesis of an antibiotic macrocyclic natural product. He is currently undertaking a postdoctoral research position at Bayer AG in Frankfurt am Main, Germany, where his research interests currently include the synthesis of novel herbicides to address the issue of highly resistant weeds.

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**Dame Margaret Brimble** is Director of Medicinal Chemistry and a Distinguished Professor at the University of Auckland where her research focuses on the synthesis of bioactive natural products and peptide chemistry. She has published over 500 papers and named as an inventor on more than 50 patents. She was inducted into the American Chemical Society Medicinal Chemistry Division Hall of Fame in 2019 and was elected a Fellow of the Royal Society (London) and awarded the RSC Sosnovsky Award in Cancer Therapy in 2018. She won the 2012 RSNZ Rutherford, MacDiarmid, and Hector Medals. She is past president of IUPAC Organic and Biomolecular Division III and an associate editor for Organic Letters. She discovered the first drug ‘trofimentide’ to treat Rett syndrome that is in phase III clinical trials with Neuren Pharmaceuticals (http://www.neurenpharma.com) and co-founded the company SapVax to develop self-adjuvanting cancer vaccines (https://sapvaxllc.com).
since 2003, our group has remained interested in the prevalence of Diels–Alder strategies in reported syntheses of spirocyclic imine fragments by groups including Romo, Murai, Nakamura, and White. These methods often involved the use of α-exo-methylene lactam/lactone dienophiles, which ultimately afforded spirolactam/lactone ring systems and required further elaboration to the desired spirocyclic imine motif.

Alternatively, direct access to a spirocyclic imine could be achieved via a Diels–Alder reaction of an α,β-unsaturated iminium ion dienophile. From 2005 to 2007, intramolecular Diels–Alder approaches to gymnodimine A (2) and symbioimine involving α,β-unsaturated iminium ions were reported by Kishi, Snider, and Thomson. In particular, both Snider and Thompson groups utilized a highly reactive N-acyliminium dienophile which was generated in situ from Lewis acid activation of a stable N-acetyl enecarbamate precursor. Intermolecular variants of the Diels–Alder reaction, however, were relatively unexplored prior to our initial investigations of N-acyliminium ion chemistry in 2008. Reaction of an exocyclic α,β-unsaturated N-acyliminium ion 5 with an appropriate diene 6 could provide convenient access to spirocyclic imine fragment 7 similar to those found in the cyclic imine toxins (Scheme 1). It was rationalized that in situ iminium ion generation could occur through α- or γ-elimination of a nucleofuge from cyclic carbamates 8 and 9, respectively, and a scalable synthesis of stable iminium ion precursors was the focus of our initial investigations.

Scheme 1 Possible pathways for in situ generation of α,β-unsaturated N-acyliminium ion 5 and its use in Diels–Alder reactions

2.2 Early Studies Using in situ-Generated Iminium Ion Dienophiles

Our first-generation synthetic approach to iminium ion precursors 10a and 10b (Scheme 2, A), reported in 2008 involved base treatment of the corresponding six- or seven-membered N-CBz lactam 11a and 11b, followed by alkylation of the amide enolates with Eschenmoser’s salt and Hofmann elimination. Reduction of the α-methylene lactams 12a and 12b with DIBAL-H/TMSCl and quenching with MeOH at low temperature afforded N,O-acetals 10a and 10b as iminium ion precursors, likely proceeding via

2 Strategies towards the Spirocyclic Imine Fragment of Cyclic Imine Toxins

2.1 Diels–Alder Cycloadditions of α,β-Unsaturated N-Acyl Iminium Dienophiles

The spirocyclic imine fragments of cyclic imine toxins are largely purported to arise from intramolecular Diels–Alder cycloaddition reactions. Indeed, this reflects the most convergent method for synthesizing the spirocyclic imine motif, given the ability to forge the precise stereochemical configuration of both the quaternary carbon center and the adjacent tertiary center in a single step. This factor justifies the prevalence of Diels–Alder strategies in reported syntheses of spirocyclic imine fragments by groups including Romo, Murai, Nakamura, and White. These methods often involved the use of α-exo-methylene lactam/lactone dienophiles, which ultimately afforded spirolactam/lactone ring systems and required further elaboration to the desired spirocyclic imine motif.

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iminium intermediates 13a and 13b. The modest overall yield of this sequence warranted investigation of an alternative second-generation route to the six-membered iminium precursor 10a. Later in 2009 it was discovered that an electrochemical oxidation of N-protected piperidine 14 could be employed to access hemiaminal 15 in greater yield (92% brsm) than conventional chemical oxidation methods (Scheme 2, B).41 Pihko organocatalytic methylation afforded ring-opened α,β-unsaturated aldehyde 16, and subsequent treatment with catalytic amount of Sc(OTf)3 in MeOH afforded enecarbamate 17 as an iminium ion precursor. Despite likely proceeding through identical iminium intermediate 13a, it is noteworthy that 17 was formed under thermodynamic conditions, while kinetic conditions (i.e., DIBAL-H, –78 °C) afforded its constitutional isomer 10a. This reaction sequence was then further modified in a third-generation approach to iminium ion precursor 18 while bypassing unstable aldehyde intermediates of the previous generation synthesis (Scheme 2, C).41 Iminium ion precursor 18 was synthesized in a much improved yield through anodic α-oxidation of N-protected piperidine 19, followed by NH4Cl-mediated elimination and microwave-promoted organocatalytic alkylation.

The reactivity of six-membered iminium precursors 17 and 18 was evaluated in a series of nucleophile trapping experiments, whereby reactive iminium ions 13a and 20 were generated in the presence of a catalytic amount of Sc(OTf)3, then quenched with various nucleophiles (Scheme 3). The corresponding alkylated products 21 and 22 were afforded in high yields for various nucleophiles such as indole and silyl enol ethers.

\[
\text{Scheme 3} \quad \text{Nucleophile trapping with six-membered \(N\)-acyl iminium ions 13a and 20}^{41}
\]

Use of iminium precursors 10a, 10b, and 17 in Diels-Alder reactions to access spirocyclic systems was also reported in our 2008 study.40 Both six- and seven-membered \(N\)-carboalkoxy-\(N,O\)-acetics 10a and 10b were reacted with diene 23 following iminium ion generation using BF3·OEt2...
to deliver tricyclic N,O-acetals 24a and 24b in good yields with high endo selectivity, although simple dienes such as isoprene failed to react with the generated six- or seven-membered iminium ions (Scheme 4). Meanwhile, enecarbamate 17 reacted similarly using diene 25 to afford tricycle 26, suggesting that the reaction proceeds through in situ generation of an identical iminium ion species. These tricyclic adducts likely formed through Diels–Alder cycloaddition, in situ PMB ether deprotection, and intramolecular N,O-acetal formation through trapping of the resulting iminium intermediate. The precise order of these events, however, was unclear, as was the impact of intramolecular O-alkylation on the stereochemical outcome of the reaction. Typically, cyclic α-exo-methylene dienophiles provide exo adducts, due to a combination of steric and electronic factors associated with the s-cis dienophile configuration.42,43 The exclusive formation of endo adducts in our study was confirmed on the basis of 2D NOESY NMR analysis, which was rationalized to be due to favorable secondary orbital overlap between the diene and the s-trans-configured iminium dienophile.

These investigations demonstrated that in situ generation of iminium ions in the presence of functionalized dienes could be a promising method for accessing spirocyclic systems resembling those of cyclic imine toxins. In particular, preventing N,O-acetal formation would be required in order to further elaborate the generated fragments towards the natural products. In addition, the moderate reaction yields could be attributed to instability of the iminium ion prior to cycloaddition.

2.3 Use of More Stable Iminium Ion Dienophiles for Diels–Alder Reactions

Following these studies, in 2011, advances in intermolecular Diels–Alder reactions of iminium ions were reported by Evans and co-workers (Scheme 5).44 Remarkably, several endo- and six-membered NH-α,β-unsaturated iminium salts 27 and 28 were found to be isolable and stable to flash column chromatography. Iminium ions 27 and 28 were used in Diels–Alder reactions with simple dienes 29, with spirocyclic adducts 30 and 31 afforded in excellent yields and high endo selectivities (>90:10) due to LUMO-lowering activation of the iminium dienophiles. The dienophile counterion (X−) was found to have a significant effect on the rate and diastereoselectivity of the reaction, with the noncoordinating hexafluoroantimonate (SbF6−) counterion resulting in substantial reaction-rate acceleration. Importantly, imine α-functionalization in the form of a methyl and n-butyl substituents appeared important for the hydrolytic stability of the iminium ions.

Evans’ work therefore prompted a redesign of our iminium ion dienophile scaffold to incorporate a methyl substituent at the imine center (Scheme 6, A), which would serve...
to improve the hydrolytic stability of the resultant iminium ion while also providing a synthetic handle for natural product elaboration through methodology previously reported by our group.\(^4^5\) Such a substituent at the α-position would also render the iminium center less susceptible to nucleophilic attack, prohibiting the previously observed intramolecular N,O-acetal formation. As an exo-Diels–Alder reaction was required to establish the natural C6–C7 relative stereochemical configuration, we postulated that increasing steric bulk of the dienophile through bulky protecting groups on both the diene and iminium dienophile components could direct Diels–Alder cycloaddition through an exo transition state (Scheme 6, B). An electron-withdrawing N-protecting group would serve to enhance the reactivity of the resultant iminium ion, while also proving easy to remove and reveal the desired spirocyclic imine motif. Additionally, the isolation of portimine A (4, Figure 1) in 2013,\(^1^2\) which possesses a 5,6-spirocyclic imine system, presented itself as an additional application for this chemistry through accessing an analogous five-membered iminium ion, which was previously unexplored.

As a result, we reported our progress regarding the synthesis and reactivity of more stable six- and five-membered α,β-unsaturated N-acyl iminium ions in 2016\(^4^6\) and 2019,\(^4^7\) respectively. Robust syntheses of bench-stable six- and five-membered iminium ion precursors 32 and 33 were established in these studies (Scheme 7). Alkylation of iodides 34 and 35 using ethyl acetoacetate proceeded smoothly to afford β-keto esters 36 and 37. Direct cyclization of β-keto ester 36 to form six-membered cyclic enecarbamate 38 proved problematic; however, it was found that N-Boc deprotection, cyclization, and subsequent amine reprotction were required to give stable six-membered cyclic enecarbamate ester 38 in high overall yield. Comparatively, direct cyclization of the analogous two-carbon-chain compound 37 to form cyclic five-membered enecarbamate 39

\[\text{Scheme 6} \quad \text{Our revised iminium ion scaffold and proposed outcome of their use in Diels–Alder reactions}\]

\[\text{Scheme 7} \quad \text{Synthesis of six- and five-membered N-Boc cyclic enecarbamates 32 and 33 as stable iminium ion precursors}^{46,47}\]
was successful, with facile cyclization of 37 occurring during the acidic workup following alkylation. DIBAL-H-mediated reduction of the ester motif of both 38 and 39 provided allylic alcohols 40 and 41 which were initially investigated as potential iminium precursors. However, instability and low reactivity led us to favoring methylation of alcohols 40 and 41, delivering both methyl ethers 32 and 33 as convenient precursors of N-acyl iminium ions, which also proved stable long-term. Our newly developed route proved robust on gram-scale and overcame synthetic difficulties of previously reported methods, proceeding through stable intermediates (e.g., 38 and 39) suitable for long-term storage if required.

Use of precursors 32 and 33 to prepare the corresponding α,β-unsaturated N-acyl iminium ion dienophile 42 in the presence of the Lewis acid, which can then react with a range of silyloxy E/Z-dienes in the Diels–Alder reaction, were therefore investigated (Scheme 8). It was rationalized that a Z-diene configuration could also provide adducts 43 bearing the desired stereocchemical orientation through a concerted cycloaddition via an endo reactivity mode.

![Scheme 8 Proposed outcome of Diels–Alder reactions of iminium ion 42 with E- or Z-silyloxy dienes](image)

To probe the reactivity of six-membered N-acyl iminium ions as dienophiles in Diels–Alder reactions, an initial reaction involved the generation of the iminium dienophile from precursor 32 using BF$_3$·OEt$_2$ prior to reaction with silyloxydiene (E)-44 to provide adduct 45 as a single diastereomer in 34% yield (Scheme 9, A). Use of an excess amount of triethylamine was required to convert iminium adduct intermediate 46 into the observed enamine adduct 45. Interestingly, unpublished results from our group involved an analogous reaction using Z-diene (Z)-44 with six-membered iminium precursor 32 which afforded the same adduct diastereomer 45 in 27% yield. This appeared to corroborate a similar observation reported by Romo and coworkers in 2000, whereby Diels–Alder reaction of a six-membered α-exo-methyl lactam resulted in a single adduct diastereomer obtained, irrespective of the diene configuration. In their case, they reasoned that a formal Diels–Alder reaction, proceeding stepwise via sequential Michael additions was in fact occurring, and it was conceivable that a similar process was occurring in our reaction. Further investigation of diene protecting groups revealed that replacement of benzyl with a benzyl group on the primary alcohol of the diene [(E)-47 and (E)-48] improved the yield of the cycloaddition products 49 and 50 significantly (Scheme 9, B). Stereochemical elucidation was enabled through hydrolytic ring opening of the N-heterocycle 50, affording keto amine 51 which displayed a plausible through-space correlation between the C-6 acetyl substituent and H-1 in a 2D ROESY spectrum. This correlation corresponded to the desired natural configuration, which would have arisen from a formal exo cycloaddition. More recently, unpublished results from these ongoing studies within our group involved iminium Diels–Alder reactions using N-Ts enecarbamate precursor 52 which could be readily prepared through an analogous reaction sequence to previously reported N-Boc enecarbamate precursor 32 (Scheme 9, C). The desired iminium Diels–Alder reaction proceeded smoothly, however, stereochemical elucidation of adduct 53 through 2D NMR experiments proved ambiguous. Silyl group removal using TBAF provided crystalline ketone 54, which possessed formal endo configuration as determined by X-ray crystallography. At this stage, whether or not there was a stereochemical discrepancy between the Diels–Alder reaction products of N-Boc and N-Ts iminium ions is yet to be confirmed unambiguously, and further work will be required to determine the factors controlling the stereochemical outcome of this reaction.

The discrepancy between the reactivity of five- and six-membered iminium ions was noted when investigating the Diels–Alder reaction with diene (E/Z)-48 in our 2019 study, with the N-protecting group also playing a major role (Scheme 10). The Diels–Alder reaction of five-membered N-Boc enecarbamate 33 with diene (E/Z)-48 was initially attempted using conditions previously optimized for six-membered 32 (Scheme 10, A). Extensive decomposition was observed, and subsequent investigation of an extensive range of reaction conditions failed to afford more than a trace amount of bicyclic adduct 55. It was soon realized that the five-membered N-Boc acyl iminium ion was unstable, and modification of the N-acyl substituent appeared as a promising method to modulate the reactivity of the resulting iminium ion. N-CBz and N-Ts enecarbamates 56 and 57 were therefore synthesized using similar sequences to those used to prepare 33. In the case of N-CBz enecarbamate 56, the desired bicyclic adduct 58 was afforded in a 20% yield, and a 2D NOESY experiment revealed a through-space correlation between H-3 and H-6, suggesting the for-
formation of an endo-cycloaddition adduct. Meanwhile, an analogous reaction using N-Ts enecarbamate 57 provided the expected bicyclic adduct 59 in 5% yield, alongside a major quantity (64%) of enone 60, which likely forms through Mukaiyama–Michael addition of diene 48 to the five-membered N-Ts iminium ion 61 (Scheme 10, B). This marked the first instance that such an intermediate had been detected and isolated throughout our extensive investigation of reactive iminium ions. It was expected that enone 60 could undergo intramolecular cyclization to afford bicyclic adduct 62, and this was indeed the case, with exposure of 60 to Sc(OTf)3 facilitating cyclization to afford adduct 62 as a single diastereomer (Scheme 10, C). The configuration of 62 was determined to be the formal endo-cycloaddition product through X-ray crystallography, and 62 could be further elaborated to natural product like fragment 63, albeit with non-natural C5–C6 configuration. The isolation of enone 60 and its ability to cyclize to bicyclic compound 62 appeared to suggest that the iminium Diels–Alder reactions were in fact proceeding through a stepwise mechanism involving initial Mukaiyama–Michael addition of diene (E/Z)-48 to iminium ion 61, followed by Lewis acid mediated intramolecular cyclization. In support of this conclusion, the same adduct diastereomer was obtained irrespective of the diene geometry employed in the reaction, with use of a mixture of dienes (E)-48 and (Z)-48 affording the same, single diastereomer 62.

N-Acyli iminium ions are well established as versatile intermediates for the synthesis of nitrogen-containing compounds, and their chemistry remains a promising area of research, particularly towards complex alkaloid synthesis, with extensive reviews regarding this field have been reported.49,50 Taken together, these findings appeared to strongly suggest that the originally hypothesized concerted Diels–Alder cycloadditions of α,β-unsaturated N-acyl iminium ions may in fact proceed through a stepwise mechanism. Despite providing adducts bearing non-natural relative stereochemical configuration, we demonstrated easy
access to highly reactive iminium ions from stable precursors and their use in a highly convergent approach to access spirocyclic systems. Additionally, the ability to access intermediate enone 60 in the recent study (Scheme 10) offers the possibility for asymmetric control of the subsequent cyclization step, which is an ongoing area of investigation by our group.

2.4 Other Notable Strategies towards Spirocyclic Imines

Although Diels–Alder cycloadditions were utilized as the most efficient strategy to construct the spirocyclic imine fragments, other synthetic methods, developed by our group and others, have been utilized to construct the spirocyclic ring systems of members of the spirolides,26 gymnodimine,51 and, more recently, portimines.21 As an alternative, indirect approach to forge spirocyclic imine fragments, these methods often involve construction of the ‘lower’ cyclohexene ring bearing the required stereochemistry at the quaternary center, with tethered side chains installed to later form the nitrogen-containing ‘upper’ ring.

In 2005, Zakarian and co-workers reported a synthetic approach to the spirocyclic imine ring system of the pinnatoxins, whereby dihydropyran 64 underwent a cascade sigmatropic rearrangement to afford chiral ketone 65, thus constructing the chiral quaternary center present in the pinnatoxins (Scheme 11, A). Subsequent elaboration of ketone 65 to azide 66 was followed by an aza-Wittig reaction, which facilitated cyclization of the seven-membered imine ring of 67.52 In 2010, our research group reported the synthesis of the spirocyclic imine fragment of members of the spirolides, whereby the cyclic imine formation was also achieved using a late-stage aza-Wittig cyclization (Scheme 11, B). The synthesis involved a microwave-assisted Diels–Alder cycloaddition between Danishefsky’s diene (68) and α,β-unsaturated ester 69, affording cyclohexenone 70 in 65% yield as a mixture of three diastereomers (5:2:1). Following successful formation of the ‘lower’ ring, subsequent transformations of the desired (S,S)-adduct 70 to keto-azide...
was followed by an aza-Wittig cyclization reaction, affording spirocyclic imine 72 possessing the same relative stereochemistry as members of the spirolide family.26

The Guillou group has reported several synthetic approaches to the 6,6-spirocyclic imine motif of the gymnodimine family since 2011.51,53 Their most recent work in 2014 hinged on a palladium-catalyzed decarboxylative allylation reaction of 73 to forge the required quaternary stereocenter in high yield and enantioselectivity (Scheme 12). This was followed by a microwave-assisted alkene isomerization to afford azidoalkene 74 which subsequently underwent a [3+2] cycloaddition to afford spirocyclic imine 75 in moderate yield. Importantly, 6,6-spirocyclic imine 75 incorporates the requisite functional handles to be further elaborated to a viable spirocyclic imine intermediate for the synthesis of the gymnodimines.

Also in 2011, our research group reported a new approach to model spirocyclic imines which relied on quaternary center construction via an alkylation reaction, followed by a gold-catalyzed intramolecular hydroamination reaction to construct the cyclic imine system (Scheme 13).54 This mild chemical transformation appeared ideally suited for late-stage construction of spirocyclic imine motifs. Alkylation of methyl cyclohexanecarboxylate (76) using alkyl iodides 77a–c afforded the corresponding alkylated adducts 78a–c which were subsequently functionalized to afford amino alkynes 79a–c in good overall yield. With amino alkynes 79a–c in hand, a large range of cyclization conditions were trialed, culminating in successful hydroamination of amines 79a and 79b through reaction with Au(PPh3)SbF6 and triethylamine at high temperature, generating spirocyclic imines 80a and 80b in good yield. However, hydroamination proved unsuccessful for amine 79c, with complex mixtures generated instead, thus this approach was not feasible for accessing 7,6-spirocyclic imine fragments (e.g., 80c).

The proposed structure of upenamide (81) contains a 6,6-spirocyclic imine derived fragment (Scheme 14). In 2013, Taylor and co-workers reported a 19-step linear sequence from commercially available meso-anhydride 82 for
the synthesis of 6,6-spirocyclic imine 83 as a key fragment of upenamide (81, Scheme 14).55 To this end, 82 was converted into syn-methyl ester 84 over 7 steps in 55% yield. The subsequent construction of the spirocyclic imine ring system involved a base-mediated alkylation of cyclohexene 84 with iodoazide 85. Staudinger reduction of azide 86, and lactam formation from amine ester 87 afforded bicyclic compound 88, which was eventually elaborated to spirocyclic imine 83.

In particular, the alkylation of cyclohexyl methyl ester 84 using iodoazide 85 reported by Taylor and co-workers appeared relevant for our own investigations as a means to streamline our previously reported synthesis of spirocyclic imine ring systems through the direct alkylation of ester 76 with iodoazides 85 and 89 (Scheme 15, A). Indeed, alkylation of 76 using iodide 85 provided the desired alkylated adduct 90 in high yield. Interestingly, the analogous reaction using iodoazide 89 failed to deliver the expected alkylated adduct 91, instead affording α,β-unsaturated N-vinyl amide 92. Following this observation, our study expanded the substrate scope to confirm this unexpected result and soon realized this reaction to be a remarkable and direct synthesis of industrially important α,β-unsaturated N-vinyl amides 93 from esters 94 and N-vinyl amides 95 from aldehydes 96 (Scheme 15, B). The reaction most likely proceeds through the in situ formation of an N-vinyl azide 97, which undergoes facile azide–enolate [3+2] cycloaddition, followed by rearrangement and nitrogen extrusion (Scheme 15, C). This work was published in 201756 and remains an ongoing area of research within our group, using differentially substituted iodoazides and their reaction with aldehydes and esters to construct unique N-vinyl amide building blocks.

2.5 Recent Efforts towards the 5,6-Spirocyclic Imine Marine Toxin Portimine A

Among the cyclic imine marine toxin family, portimine A (4) represents the smallest member (5,6-spirocyclic imine) identified to date with a unique biological profile. It exhibits high in vitro cytotoxic activity (P388 cells, EC_{50} = 2.7 nM), yet low in vivo toxicity, setting it apart from the rest of cyclic imine toxin families which exhibit less specific toxicity.57 In 2019, Fujiwara and co-workers reported the first synthetic study towards portimine A (4, Scheme 16), in which the synthesis of the cyclohexene fragment 98 of the natural toxin was achieved in 6.5% overall yield over 16 steps. The sequence involved the construction of the tertiary center of 99 through conjugate addition of vinylmagnesium bromide to α,β-unsaturated ketone 100, followed by diastereoselective dihydroxylation of the vinyl group and acetal formation to provide acetal 101. Installation of the diene moiety through a Grignard addition–dehydration sequence afforded diene 102, which was subsequently elaborated to provide racemic cyclohexene fragment 98.21

Meanwhile, the most recent study from our group in 2019 has demonstrated 2-bromo-1,3-butadiene systems (e.g., 103, Scheme 17) as highly effective substrates for tandem Diels–Alder–transition-metal cross-coupling reaction sequences.28 Intermolecular cycladdition of 2-bromodiene 103 with a variety of activated dienophiles was shown to proceed in generally high yield, with good to excellent endo diastereoselectivity. The resulting vinyl bromide cycloducts 104 readily underwent subsequent Stille and Suzuki cross-couplings under standard conditions to provide access to novel substituted cyclohexene products 105. Of particular importance to this review is our application of this study to successfully assemble the spirocyclic imine motif 106 of portimine A (4), whereby the Diels–Alder reaction of diene 103 and enal dienophile 107 was used to forge the
Scheme 15  Unexpected formation of N-vinyl amide 92 and related study by Brimble and Furkert et al.56

Scheme 16  Synthesis of the cyclohexene fragment 98 of portimine A (4) by Fujiwara et al.21
cyclohexene quaternary center. Adduct 108 then underwent Seyferth–Gilbert homologation, N-phthalimide deprotection, and gold-catalyzed hydroamination, employing conditions previously developed by our group.\textsuperscript{54} to afford bicyclic imine 109. Subsequent Stille coupling successfully installed a vinyl substituent to construct bicyclic fragment 106 resembling the 5,6-spirocyclic imine system of portimine A (4), albeit with non-natural (R,R)-relative stereochemical configuration. Future efforts will focus on understanding factors governing the diastereoselectivity of Diels–Alder reactions using 2-bromo-1,3-butadienes, and methods to reverse the inherent endo selectivity and favor adducts bearing the same stereochemical configuration seen in the cyclic imine toxims.

2.6 Construction of Another Challenging Motif of Portimine A

While the spirocyclic imine fragments of cyclic imine toxims command particular attention from synthetic chemists, each toxin of the family also bears a complex polyketide motif, representing another formidable synthetic task. At the time of Fujiwara and co-workers’ synthesis of the cyclohexene fragment of portimine A (4), the polyketide motif of the natural product remained a challenging target, with the syn-1,3-dihydroxyketone motif in particular (shaded in blue of 4, Scheme 18), identified by our group as an important synthetic issue to address before a total synthesis could be realized. The syn-dihydroxyketones are embedded in a wide range of biologically active natural products, however, development of stereoselective synthetic methods to assemble these structures has proven a challenging task. In 2019, we reported our development of a highly stereoselective synthesis of syn-dihydroxyketone motifs 110 from readily available propargylic alcohols 111 (Scheme 18).\textsuperscript{58} The reaction sequence involved regioselective cyclization of propargylic alcohols 111 with incorporation of triketone 112 to give enol dioxolanes 113 that then underwent highly diastereoselective epoxidation to give spiroepoxide intermediates 114. Hydrolysis or acetylation of the spiroepoxides 114 then cleanly afforded syn-dihydroxyketones 110 or the corresponding monoacetylated dihydroxyketones 115 as single diastereomers. This methodology also included a telescoped one-pot protocol with no loss of overall yield or diastereoselectivity comparing to the three-step sequence, with wide scope for application in the stereoselective synthesis of complex molecular architectures.

3 Conclusion and Future Perspectives

Taken together, this review details multiple synthetic investigations by our research group towards cyclic imine toxims. In particular, iminium chemistry has been investigated and has proved to be an efficient method for direct access to spirocyclic imine fragments, albeit bearing non-natural relative stereochemical configuration. In parallel, we also investigated alternative approaches to spirocyclic imine fragments, including alkylation–hydroamination sequences and Diels–Alder–hydroamination sequences. Along the way, an efficient method to synthesize N-vinyl amide building blocks was serendipitously discovered, and recently, a novel approach to access the challenging 1,3-syn-dihydroxyketone motif present in many bioactive natural products, including portimines, was established.

Recent years have seen a resurgence in interest in the spirocyclic imine fragment of cyclic imine toxims, with multiple new reports since 2017 validating the importance of these structural motifs. Diels–Alder approaches appear the most efficient means of accessing spirocyclic imine fragments, with the iminium Diels–Alder reaction in particular providing rapid access to spirocyclic imine systems. Further work is required to investigate the factors governing the stereochemical outcome of iminium Diels–Alder reactions, since the means with which to precisely control this will enable access to adducts bearing the natural stereochemical configuration.
Scheme 18  Diastereoselective synthesis of syn-1,3-dihydroxyketone motifs 110 by Furkert and Brimble et al.59

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