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Chemical synthesis and application of aryl dihydronaphthalene derivatives

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Aryldihydronaphthalenes (ADHNs) and their derivatives are widely found in many types of natural products, bioactive compounds, and functional materials, and are also important synthetic intermediates in organic chemistry, attracting widespread attention from both organic and pharmaceutical chemists. In the past two decades, the chemical synthesis and biological activity of ADHNs and their derivatives have become two hot spots. This review summarizes the synthetic protocols of ADHN derivatives, introduces some representative examples of the reaction mechanism, and focuses on the research progress of ADHNs in natural product chemistry and chemical biology since 2000.

1. Introduction

Lignans, naturally occurring bioactive chemicals, are always formed by the oxidative dimerization of two phenylpropanoid units.¹ Although the molecular skeleton of lignans consists of only two phenylpropane (C6–C3) units, lignans exhibit remarkable structural diversity, primarily attributed to variations in the oxidation levels of their side chains and the substitution patterns on their aromatic rings.² Aryldihydronaphthalene (ADHN) is an important subtype of the lignan

family, which represents a potential drug lead due to the potent activity of its analogues against many diseases. According to the distribution of the double bonds and pendent phenyl rings in the dihydronaphthalene structure, ADHN lignans can be classified into: 1-phenyl-1,2-dihydronaphthalene (1), 1-phenyl-1,4-dihydronaphthalene (2), 2-phenyl-1,2-dihydronaphthalene (3), 2-phenyl-1,4-dihydronaphthalene (4), 3-phenyl-1,2-dihydronaphthalene (5) and 4-phenyl-1,2-dihydronaphthalene (6) (Fig. 1).

ADHNs are not only widely found in natural products, functional materials and bioactive substances,³ but also very important intermediates in organic synthesis, which can be used for the rapid preparation of highly functional and structurally diverse aryltetralins/aryltetrahydronaphthalenes and aryl naphthalenes.⁴ In view of the significant applications of ADHN derivatives, their synthetic methods have garnered

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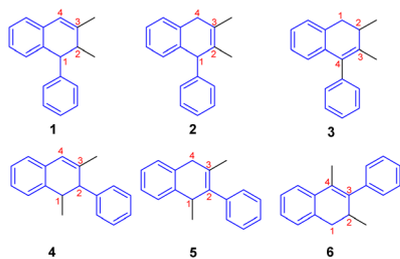


Fig. 1 Six isomers of ADHNS.

significant attention within the realm of chemistry, and this subclass of lignans stands as one of the most thoroughly studied substructural types of lignans.

Traditionally, the preparation methods of ADHNS include: (1) oxidation by different oxidants: for example, (a) FeCl_3/O_2 oxidizes two molecules of cinnamic acids or cinnamates to obtain dilactones, which rearrange to form ADHNS under acidic conditions;⁵ (b) horseradish peroxidase $\text{HRP-H}_2\text{O}_2$ or $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$ can serve as oxidants to deliver ADHNS;⁶ (2)

intramolecular didehydro-Diels–Alder (DDDA) reactions of styrene-ynes by means of refluxing acetic anhydride;⁷ (3) aryl tetrahydrofuran reacts with perchloric acid and acetic acid;⁸ (4) the oxidative coupling of ethyl ferulates generates the 1,4-diphenyl-1,3-butadiene intermediates, which then undergoes intramolecular cyclization catalyzed by aluminum trichloride.⁹ Although these methods documented in previous literature have provided viable avenues for the synthesis of ADHNS, their further utilization is constrained by stringent reaction conditions, poor functional group tolerance, and low yields. In addition, traditional approaches tend to be narrowly focused on attaining the target compounds, while neglecting the exploration of structural diversity. The aforementioned constraints impede the synthesis of derivatives based on ADHNS scaffolds, thereby preventing pharmaceutical chemists from further investigating the structure–activity relationships. As modern synthetic techniques pivot around modular and diverse synthesis, it is imperative to develop efficient and convenient approaches for building the useful ADHNS frameworks.

In recent decades, significant research progress has been made in the synthesis of ADHNS. However, comprehensive overviews encompassing the synthetic methodologies and applications of ADHNS skeletons remain scarce. This review classifies the approaches for synthesizing ADHNS derivatives into two primary categories: intramolecular and intermolecular reactions. Specifically, each reaction mode is further delineated based on its specific type, for instance, intramolecular reactions will encompass intramolecular cyclization, Heck and elimination reactions, *etc.*, whereas intermolecular reactions involve coupling reactions and addition reactions, and so forth. This review will introduce the intramolecular and intermolecular reactions involved in the synthesis of ADHNS, focusing on their applications in the total synthesis of natural products and the construction of bioactive molecules, and summarize their profound implications in the field of biology.



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Shiqiang Xu received his master's degree in medicinal Chemistry from Tongji Medical College of HUST in 2001. After 6 years working in the R&D department of CR DOUBLE-CRANE (WUHAN) pharmaceutical CO., Ltd, he joined the School of Medicine, Wuhan University of Science and Technology. His main research interests include the design, synthesis and activity study of small molecule compounds against cancer.



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Chemistry and Resource Evaluation in Hubei Province. His research interest is the discovery of bioactive metabolites from fungi and plants as well as the innovative drugs.



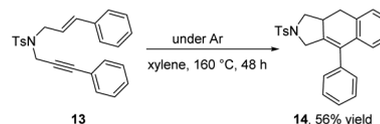
2. Chemical synthesis of ADHN derivatives

2.1 Intramolecular reactions

2.1.1 Cyclization reactions. Cyclization reaction is one of the most powerful synthetic transformations in organic synthesis, which can rapidly synthesize complex scaffolds and construct multiple C–C and C–hetero bonds in one pot.¹⁰ Therefore, cyclization reaction is regarded as a critical step in organic chemistry research. According to the external conditions used in the process, it can be divided into the following types, including heat, microwave, acid, photochemical, bases and transition metals.

Heating, particularly when combined with microwave irradiation, serves as an efficacious means to facilitate diverse reactions,¹¹ notably the renowned intramolecular dehydrogenative Diels–Alder transformation. In 2010, Ruijter *et al.* reported microwave-assisted intramolecular Diels–Alder reactions of tetrahydroisoquinolines **7** and tetrahydro- β -carbolines **10** to provide the polycyclic alkaloid-type compounds **8–9** and **11–12**, respectively (Scheme 1A and B).¹² Unfortunately, the reaction is often accompanied by inseparable byproducts **9** or **12**. 10 examples were given in this conversion (**7a–e** and **10a–e**), while the formation of a mixture of regioisomers made accurate assessment of yields and product ratios impossible in this case (**7c** and **10c**). The substrate with dimethylamino group (**10d**) did not afford any cyclization product **11d** or **12d**. Although the authors attempted to selectively obtain rearrangement (**8** and **11**) and oxidation products (**9** and **12**) in the presence of inert atmosphere or dehydrogenators (*e.g.*, S, Pd/C), respectively, no complete selectivity was observed.

Shortly after, in 2011, Matsubara, Kurahashi and coworkers proposed an example of ADHN **14** by promoting the reaction through heating at 160 °C (Scheme 2).¹³ The reaction process is simple and can proceed without the addition of Lewis acids, transition metals, or oxidants. In their research, they primarily

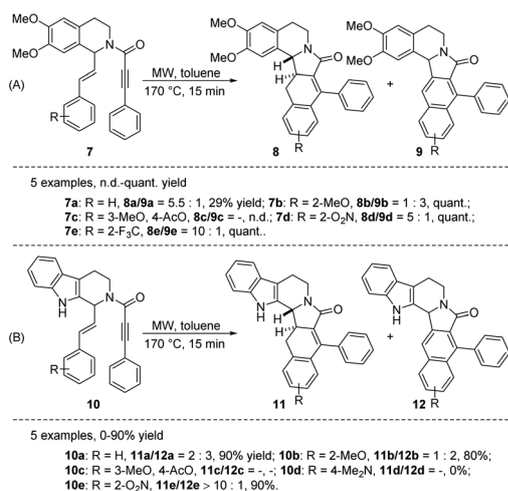


Scheme 2 Synthesis of ADHNs *via* heat-promoted dehydrogenative cycloaddition.

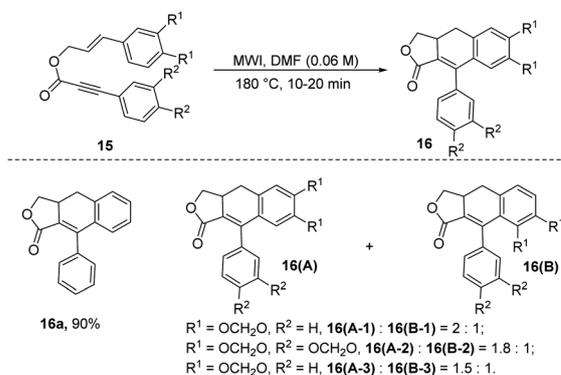
introduced the dehydrogenative cycloaddition reaction of silyl-substituted dieneynes and demonstrated that the presence of a silyl group attached to the alkyne moiety drives the occurrence of the dehydrogenation reaction.

In 2014, Brummond and coworkers proposed a solvent-regulated intramolecular dehydro-Diels–Alder (DDA) reaction of styrene precursors **15** to selectively produce ADHN lactones **16** (Scheme 3).^{4b} The styrenyl precursor **15** was subjected to microwave irradiation (MWI) at 180 °C for 20 minutes. This is the first report of entirely selective formation of ADHN lactones through the DDA reaction of styrene-ynes. Interestingly, the solvent exerts a pivotal influence on the product selectivity in the intramolecular DDA reaction. Employing DMF as the reaction solvent results exclusively in the formation of ADHN lactones **16**, while PhNO₂ selectively yields aryl-naphthalene lactones (not shown here). The authors speculated that the hydrogen atom donor properties of DMF may be a factor contributing to the selectivity of the DDA reaction in DMF, while the oxidation capacity of PhNO₂ may explain the exclusivity of aryl-naphthalene lactones products.

In 2022, our group developed an intramolecular didehydro-Diels–Alder (DDDA) reactions of styrene-ynes **17** (Scheme 4).¹⁴ Although the preparation of ADHN skeletons using styrene as a substrate is generally considered a challenging task, this reaction achieved moderate to good yields under relatively mild reaction conditions (80 °C) without any transition metal. This promising result may be explained by the critical propiolamide moiety, which was previously reported by our research group.¹⁵ The study showed that the argon atmosphere played an important role in the selective generation of ADHNs, with 2,6-di-*tert*-butyl-4-methylphenol (BHT) serving as an antioxidant. The synthetic value of this protocol was validated through



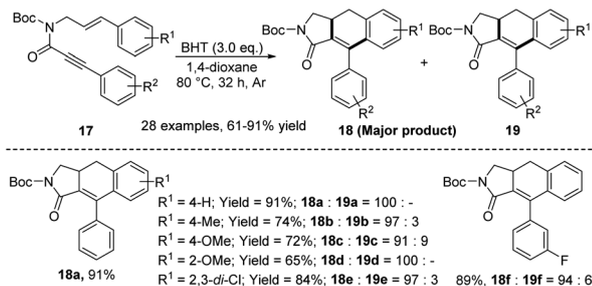
Scheme 1 Synthesis of ADHNs *via* microwave-assisted intramolecular Diels–Alder reactions of styrene-ynes.



Scheme 3 Synthesis of ADHNs through DMF-regulated intramolecular dehydro-Diels–Alder reactions of styrene-ynes.



Review

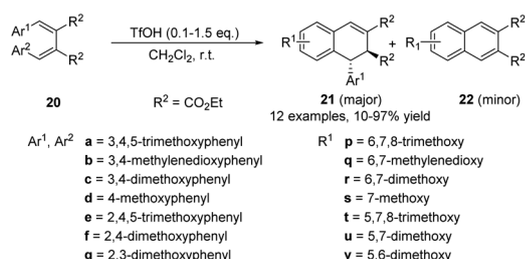


Scheme 4 Synthesis of ADHNs via intramolecular DDDA reaction of styrene-ynes.

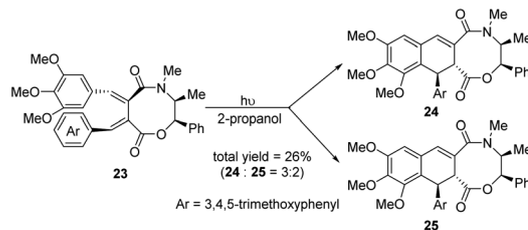
a gram-scale reaction (1.0 g, 82% yield), showcasing the successful deacylation of the *N*-bound acyl group and the oxidation of ADHN with oxygen (not shown here).

In 2001, Charlton *et al.* reported an acid-catalyzed cyclization of 2,3-dibenzylidenesuccinates **20** (Scheme 5),¹⁶ which has been exploited to prepare the naturally occurring lignans (\pm)-cagayanin and (\pm)-galbulin, with overall yields of 18% and 23%, respectively. It was found that the reactivity of the diesters was highly dependent on the position of the alkoxy substituents on the aryl rings. By prolonging the heating time or using excessive triflic acid, the yield of the products **21** was reduced and the dearylated compounds **22** was induced. Since (*E,E*)-dibenzylidenesuccinates can be easily prepared by the Stobbe condensation, it becomes an attractive substrate for the formation of lignans. Subsequently, in 2004, the same group prepared mandelate ester from the same raw material, 2,3-dibenzylidenesuccinate, and then transferred the (–)-ephedrine cyclic amide ester **23** to obtain optically active ADHNs **24** and **25** through an asymmetric photocyclization reaction (Scheme 6).¹⁷

In 2010, Nishii *et al.* reported the ring-expansion of phenyl hydroxy methyl cyclopropane carboxylates **26** and its diaryl analogs **28** mediated by Lewis acids, resulting in the corresponding 1,2-dihydronaphthalene-3-carboxylic acid esters **27** and **29**, respectively (Scheme 7A and B).¹⁸ Various Lewis acids such as TiCl_4 , SnCl_4 , TBDMSOTf , $\text{BF}_3 \cdot \text{Et}_2\text{O}$, $\text{Yb}(\text{OTf})_3$ and $\text{Sc}(\text{OTf})_3$ were screened in the conversion. The results showed that $\text{Yb}(\text{OTf})_3$ and $\text{Sc}(\text{OTf})_3$ were the most optimal choices among the lanthanide triflates with the highest yields (90% and 95%). EDC (1,2-dichloroethane) was identified as the best solvent. The authors cited several examples (**27a–e** and **29a–d**) to demonstrate the substrate scope, with yields ranging from 60% to 99%.



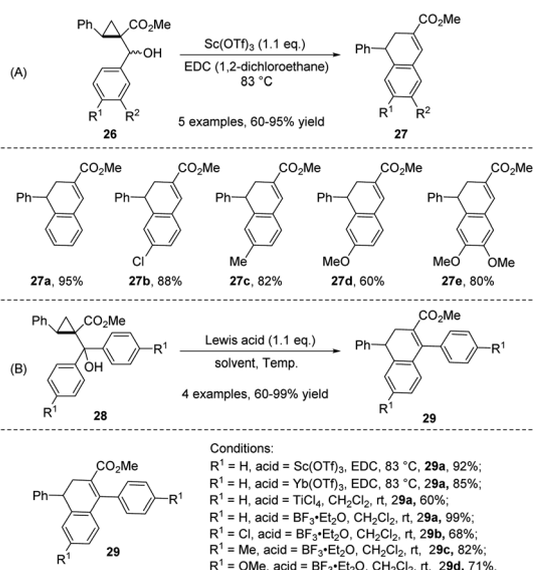
Scheme 5 Synthesis of ADHNs via acid-catalyzed cyclization of diethyl *E,E*-dibenzylidenesuccinates.



Scheme 6 Synthesis of ADHNs via photochemical closure of styrene-ynes.

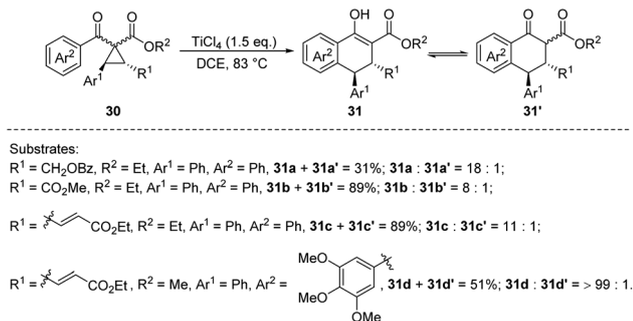
Subsequently, in 2017, the same group reported the asymmetric homo-Nazarov-type cyclization of racemic donor-acceptor (D–A) cyclopropanes **30** mediated by TiCl_4 to synthesize poly-substituted ADHNs **31** and **31'** (Scheme 8). The OH groups of the products **31** are in equilibrium with their corresponding keto-isomers **31'** and is subsequently converted to the corresponding triflates. Notably, the stoichiometric amount of TiCl_4 significantly improved the yield, and the optimal result (92%) was achieved using 1.5 equivalents of TiCl_4 at 83 °C. The practicability of the reaction was verified through further transformations of the above mentioned triflates, such as Suzuki–Miyaura coupling and Sonogashira coupling.

In 2014, Li *et al.* showed the construction of ADHN skeletons via the acid-catalyzed cyclization of **32**, providing the target ADHNs **33** (similar to the structure of **21**, Scheme 5) accompanied with a small amount of dearylated products **34** (Scheme 9).¹⁹ Differed from Charlton's work (Scheme 5), this work used compound **32** as the substrate, and it was treated with either perchloric acid in acetic acid at room temperature or *p*-toluenesulfonic acid in toluene under reflux. As a possible mechanism, they suggest that the tetrahydrofuran ring in **32** is opened under acidic conditions to form a quinone methide intermediate, which then undergoes an intramolecular nucleophilic

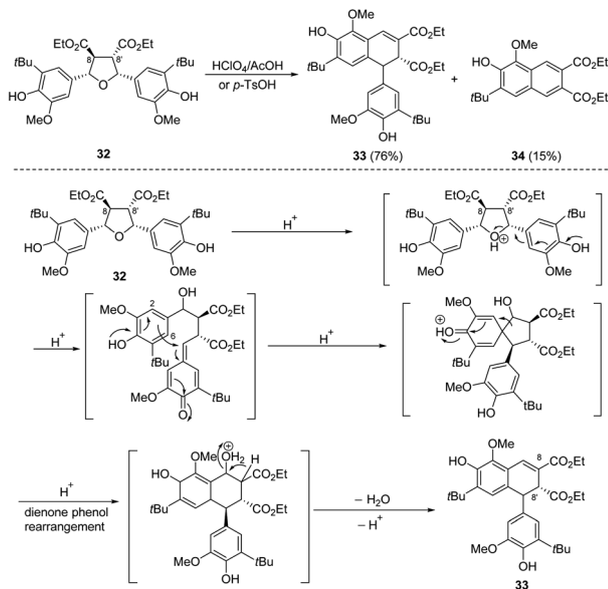


Scheme 7 Synthesis of ADHNs via Lewis acid-mediated ring-expansion reaction of cyclopropane carboxylates.





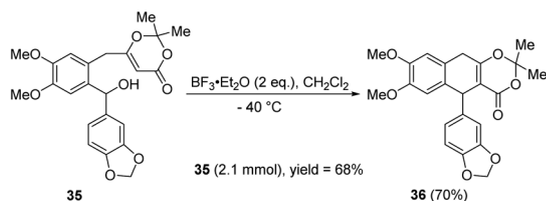
Scheme 8 Synthesis of ADHNs via TiCl_4 -mediated ring-opening cyclization of D–A cyclopropanes.



Scheme 9 Synthesis of ADHNs via acid-catalyzed cyclization of tetrahydrofuran.

attack to provide the five-membered spiro structure. During the subsequent dienone-phenol rearrangement of the spiro intermediate, the oxygen-substituted side-chain migrates and H_2O is eliminated, leading to the formation of ADHN 33.

In 2022, the group of Zhang and Chen disclosed a method for the cyclization of 35 promoted by boron trifluoride diethyl etherate (Scheme 10).²⁰ In this work, the authors initially optimized the conditions for preparing 36 from benzhydrol 35. Both Brønsted acids and Lewis acids were evaluated, and boron

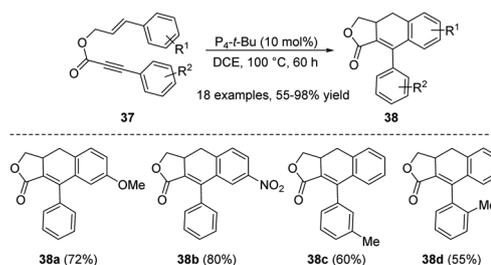


Scheme 10 Synthesis of ADHNs via Brønsted acid-catalyzed cyclization of benzhydrol.

trifluoride diethyl etherate was proved to be the best one. The relatively low temperature was conducive to production, $-40\text{ }^\circ\text{C}$ being better than $0\text{ }^\circ\text{C}$, $-30\text{ }^\circ\text{C}$ and $-50\text{ }^\circ\text{C}$. The reaction exhibited good functional group tolerance and good to excellent yields. The applicability of the reaction was further validated by a scaled-up reaction (2.1 mmol).

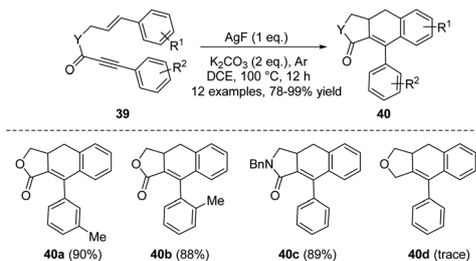
The cascade cyclization reaction strategy has attracted considerable attention in view of the convenience in the synthesis of polysubstituted polycyclic compounds.²¹ The traditional protocols mainly focused on acetic anhydride or base-mediated cyclization, while the high loadings of acetic anhydride^{7a,d} or base^{7d} result in toxic byproducts, limiting their applications. Recently, some progress has been made in base-catalyzed cyclization reactions. In 2010, Li *et al.* demonstrated the intramolecular cascade reactions of aryl-substituted enynes 37 catalyzed by $\text{P}_4\text{-}t\text{-Bu}$, resulting in the synthesis of 9-aryl-3a,4-dihydronaphtho[2,3-*c*]furan-1(3*H*)-ones 38 (Scheme 11). In this reaction, a number of bases such as Cs_2CO_3 , K_3PO_4 , NaOH , NaOEt , DABCO , DBU and K_2CO_3 were examined, while all of them were less effective than 10 mol% $\text{P}_4\text{-}t\text{-Bu}$. Solvents such as toluene, tetrahydrofuran, and *N,N*-dimethylformamide were also tested, and none of them succeed to provide a higher yield than DCE. The optimal reaction temperature for this transformation was $100\text{ }^\circ\text{C}$. Noteworthy, the electronic properties of aryl allyl groups had little influence on the conversion. The authors suggested that an intramolecular Diels–Alder cyclization mechanism might be involved in this process.

Subsequently, in 2011, the same group reported a silver fluoride-promoted intramolecular cascade reaction of aryl-substituted 1,6-enynes 39 to afford ADHN lignan lactone 40 (Scheme 12).²² The reaction was carried out at $100\text{ }^\circ\text{C}$ under an argon atmosphere, using a mixture of $\text{AgF}/\text{K}_2\text{CO}_3$ as the base and $\text{CH}_2\text{ClCH}_2\text{Cl}$ as the optimal solvent. When the alkynyl allyl ester moiety is simultaneously linked with electron-donating groups and electron-withdrawing groups, the corresponding high-yield products can be obtained. Unfortunately, however, enyne ethers 39d failed to complete this conversion (40d, 0% yield). The author suggested that the reason why silver ions can promote the reaction is probably due to their ability to coordinate with triple bonds, instantly activating them, and thus facilitating the reaction. Among silver salts, AgF likely exhibits the best effect due to its strong alkaline nature, while other silver salts are neutral or weakly acidic. Transition metal catalysts are undoubtedly one of the most fundamental and useful



Scheme 11 Synthesis of ADHNs via $\text{P}_4\text{-}t\text{-Bu}$ -mediated intramolecular Diels–Alder reactions of aryl-substituted enynes.



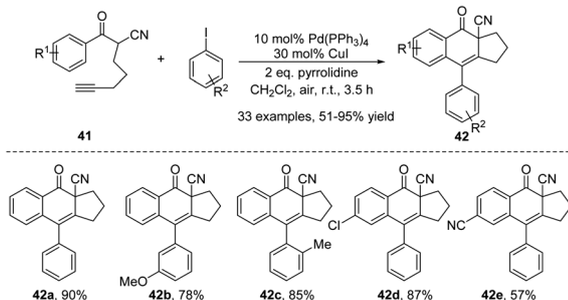


Scheme 12 Synthesis of ADHNs via silver fluoride-promoted intramolecular cascade reactions of aryl-substituted 1,6-enynes.

tools for organic chemists, playing a pivotal role in enriching reaction pathways and significantly broadening the range of substrates. They are widely used to form carbocyclic and heterocyclic frameworks as precursors for building a variety of organic molecules.²³ In 2013, Shia *et al.* developed a general approach to construct ADHNs **42** via the palladium(0)/copper(i)-catalyzed radical cyclization of **41** (Scheme 13).²⁴ Although aryl 1-cyanoalk-5-ynyl ketones **41** and aryl iodides were used as the starting materials, the Pd(0) catalyzed Sonogashira coupling reaction generated the crucial intermediate 1-cyanoalk-5-ynyl ketone, which underwent a [4 + 2] intramolecular cyclization in the reaction. Air was vital for the radical cyclization of the intermediate, facilitating the production of the target products **42**. Under the optimal conditions, both electron-withdrawing and electron-donating groups were tolerable, yielding nearly equivalent product amounts.

Shortly after, in 2014, the same group reported the oxidative cyclization of aryl 1-cyanoalk-5-ynyl ketone **43** catalyzed by manganese(III) (Scheme 14).²⁵ Unfortunately, this protocol primarily focused on the acyl tandem cyclization to aryl 1-cyanoalk-5-ynyl ketones, and the ADHN product was presented (**44**). Compared with the traditional styrenyl DDA reaction, this reaction is characterized by mild reaction conditions. Furthermore, operational simplicity, high product yields and the potential synthetic elaboration shed light on its further synthetic value.

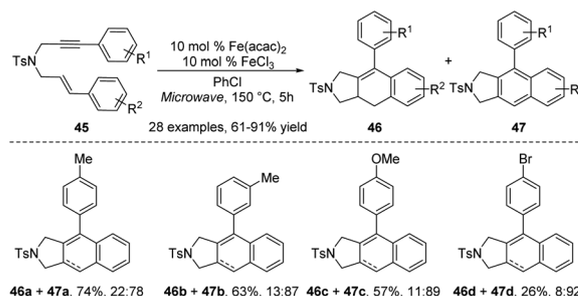
In 2018, Kang, Ahn and coworkers established an Fe(II)/Fe(III)-catalyzed microwave assisted intramolecular DDDA reaction of styrene-yne **45** (Scheme 15).²⁶ Despite their primary focus being on the synthesis of aryl naphthalenes **47**, during



Scheme 13 Synthesis of ADHNs via Pd(0)/Cu(I) catalyzed tandem annulation of aryl 1-cyanoalk-5-ynyl ketones.



Scheme 14 Synthesis of ADHNs via manganese(III)-catalyzed oxidative cyclization of aryl 1-cyanoalk-5-ynyl ketone.



Scheme 15 Synthesis of ADHNs via Fe(II)/Fe(III)-catalyzed and microwave assisted intramolecular DDDA reactions of styrene-yne.

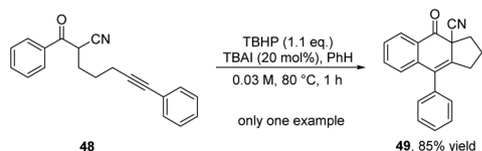
their initial experiments to identify the optimal conditions, they discovered that varying types of iron, solvents, and iron catalyst loadings significantly influenced the ratio of the ADHNs **46** and aryl naphthalenes **47**. Mechanistic investigations revealed that an Fe(III) or Fe(III) catalyst favored the dehydrogenation process, leading to the production of aryl naphthalene products.

In 2012, the Shia group described the intramolecular radical cascade reaction of α -cyano-aryl-capped alkynyl aryl alkyl ketones **48** based on radical catalytic process to offer a variety of highly functionalized ADHNs **49** (Scheme 16).²⁷ The reaction employed *tert*-butyl hydroperoxide (TBHP) as the oxidant and tetrabutylammonium iodide (TBAI) as a substitute for transition metal catalysts, rendering it a metal-free and highly efficient process. Unfortunately, this protocol mainly studied the α -cyano-TMS capped alkynyl aryl alkyl ketones substrates, paid little attention to α -cyano-aryl-capped alkynyl aryl alkyl ketones.

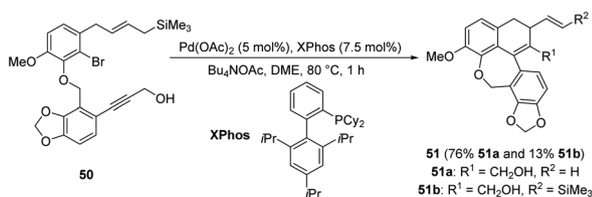
2.1.2 Coupling reactions. Over the past decades, the transition metal-catalyzed Heck reaction has emerged as an essential tool for organic synthesis.²⁸ The Heck reaction holds pivotal significance in the cyclization reaction of chain-like substrates. In 2013, Tietze *et al.* reported a domino carbopalladation/Heck reaction of an allylsilane **50** to form the ADHNs **51** (Scheme 17),²⁹ which was then converted into the natural product lignan linoxepin. This protocol is the first total synthesis of (+) and (–) linoxepin in only ten steps and an overall yield of 30% without the use of any protecting group.

Given the potential biological activity of the natural product linoxepin and the related synthetic challenges, this field has garnered significant attention from chemists. In the same year, Lautens *et al.* reported a palladium-catalyzed and base-facilitated intramolecular Mizoroki–Heck reaction of the dihydronaphthalene **52**, affording the target (+) linoxepin **53** in 76% yield (Scheme 18).³⁰ This is the first asymmetric synthesis of

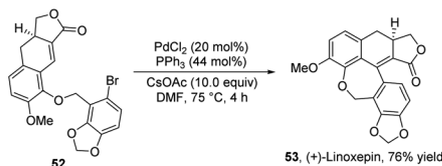




Scheme 16 Synthesis of ADHNs *via* intramolecular radical cascade of aryl alkyl ketones.



Scheme 17 Synthesis of ADHNs *via* domino carbopalladation/Heck reaction of allylsilanes.



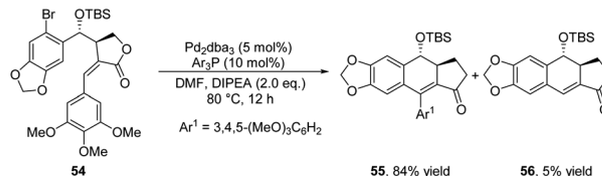
Scheme 18 Synthesis of ADHNs *via* intramolecular Mizoroki-Heck reaction of dihydronaphthalene.

(+)-linoxetine in eight steps using the authors' modified version of the Catellani reaction. The entire process is characterized by an overall yield of 30%, protecting-group-free and good enantioselectivity.

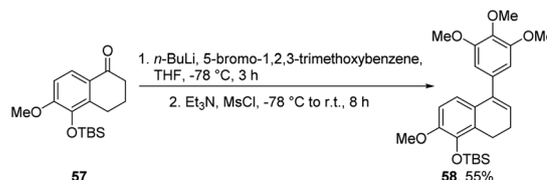
Recently, in 2022, Peng and co-workers took the same final intramolecular Heck reaction as reported by Lautens's group (Scheme 18) to prepare linoxetine.³¹ Differed from Lautens's work, Peng's group adopted a new nickel-catalyzed reductive cyclization as a key step in building a tricyclic core embedded in linoxetine (not shown here). This strategy facilitated the production of several linoxetine analogues.

In 2017, Hajra and co-workers proposed an intramolecular Heck coupling reaction of **54** to afford ADHNs **55** and dihydronaphthalene **56** (Scheme 19).³² Among the screened Pd catalysts, Pd₂dba₃ was found to be the most effective one. The ligand had strong effect on the yields of the desired products, the use of suitable ligand and base produced **55** as the major product. The suppression of Pd catalyst loading gave better yields by inhibiting the side products **56** formation. Compound **55** was a crucial synthetic intermediate. Subsequently, the authors' team furnished the total synthesis of (–)-podophyllotoxin, (–)-picropodophyllin, (+)-isopicropodophyllin, and (+)-isopicropodophyllone from the molecule **55**.

In 2018, Pinney and collaborators introduced two alternatives for the synthesis of ADHNs, one of them was the coupling reaction of 3,4-dihydronaphthalen-1(2*H*)-one **57** and 5-bromo-1,2,3-trimethoxybenzene, providing the ADHN product **58** with



Scheme 19 Synthesis of ADHNs *via* intramolecular Heck reaction.



Scheme 20 Synthesis of ADHNs *via* coupling reaction of ketone and bromo-benzene.

55% yield (Scheme 20).³³ This reaction necessitates the utilization of *n*-butyllithium (*n*-BuLi), a chemical whose handling demands rigorous safety precautions.

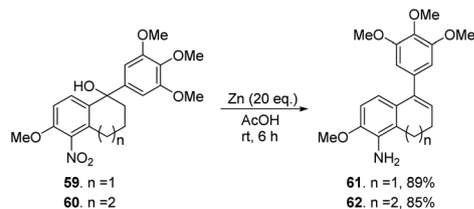
2.1.3 Elimination reactions. Pinney, Trawick *et al.* reported in 2016 that the synthesis of ADHNs **61** and **62** *via* acetic acid promoted the intramolecular elimination reaction of alcohol **59** and **60** (Scheme 21).³⁴ Notably, the addition of zinc as a nitro reductant in this reaction streamlined the process from a two-step reaction to a one-pot reaction, simplifying the steps and enhancing the productivity of the desired product. In 2018, Pinney and collaborators exhibited another pathway for the synthesis of ADHNs, which was the elimination reaction of alcohol **63** under acidic conditions, providing the target product **64** in quantitative yield (Scheme 22).³³ Distinct from their group's prior studies, this transformation was conducted under reflux conditions without the use of a zinc reducing agent.

Recently, in 2024, Peng, Xiao and co-workers reported the synthesis of dextrorotatory and levorotatory linoxetine (**66** and **66'**) through the elimination of the phenyl sulfoxides from compounds **65** and **65'**, with total yields of 50% and 55%, respectively (Scheme 23).³⁵ In this protocol, the authors employed a previously reported nickel-catalyzed intramolecular reductive cyclization cascade reaction of sophisticated β-bromo acetals with allyl tethers as the key step. In contrast to the previously reported methods for linoxetine synthesis, this work initially utilized nickel catalysis to construct the tricyclic core, which is subsequently elaborated into linoxetine *via* an intramolecular elimination reaction. Conversely, the prior method first synthesized the three-membered ring *via* nickel catalysis, and ultimately synthesized linoxetine through an intramolecular Heck reaction.

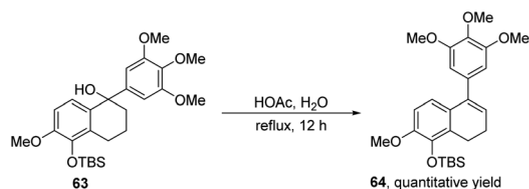
2.2 Intermolecular reactions

2.2.1 Coupling reactions. Transition metal-catalyzed cross-coupling reactions are powerful synthetic tools for the one step construction of diverse chemical bonds, offering broad and promising application prospects.³⁶ In 2009, Tringali *et al.*

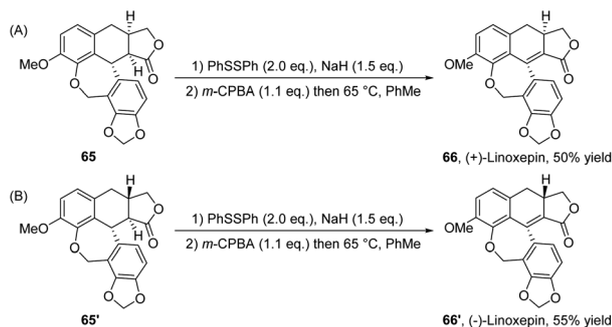




Scheme 21 Synthesis of ADHNs via AcOH mediated elimination of azabenzonornbornadienes with *N*-sulfonyl ketimines.



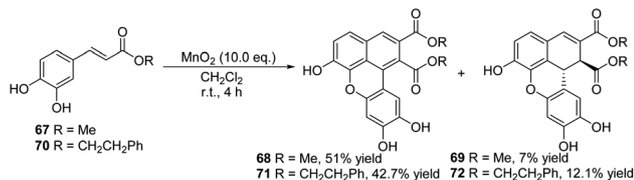
Scheme 22 Synthesis of ADHNs via elimination reaction of alcohol.



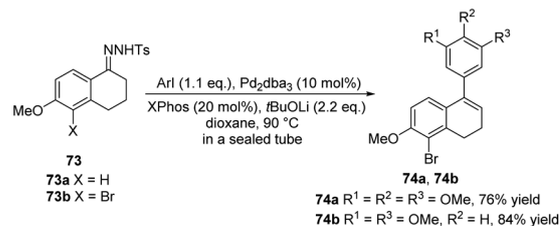
Scheme 23 Synthesis of ADHNs via intramolecular elimination of phenyl sulfoxides.

reported the biomimetic coupling reaction of caffeic acid phenethyl ester (CAPE) **67** or **70** (Scheme 24).³⁷ MnO₂ in dichloromethane functioned as an oxidant. Beforehand, CAPE had never been employed in phenolic oxidative coupling reactions. By extending the reaction time to 4 hours, a higher yield of the product was observed. Noteworthy, ascorbic acid was carefully added to quench the reaction, thereby preventing further oxidation of the products.

In 2013, Alami, Provot and co-workers presented the coupling reaction of *N*-tosylhydrazone **73** with aryl iodides under palladium catalysis, providing the key intermediates **76a**, **b** (Scheme 25).³⁸ Then, a series of coupling reactions were



Scheme 24 Synthesis of ADHNs via oxidative coupling of caffeic acid phenethyl ester.

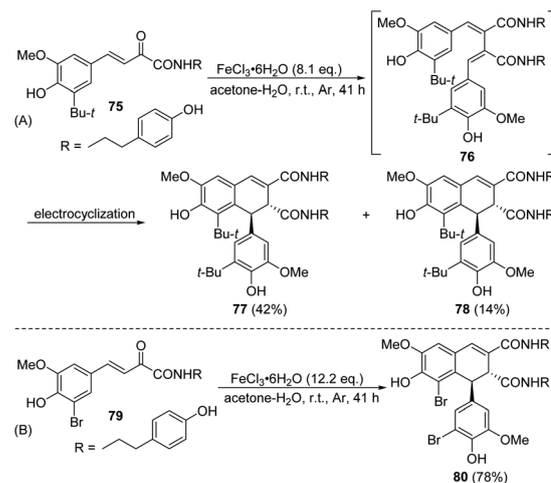


Scheme 25 Synthesis of ADHNs via coupling reaction of *N*-tosylhydrazone with aryl iodides.

performed, such as the Sonogashira–Linstrumelle reaction of **74a** with propargylic and homopropargylic alcohols, the Heck coupling reaction of **74a** with methyl acrylate, and other reactions involving **74a**, to obtain ADHN analogues with different substitutions (not shown here). These diverse ADHN analogues provide a material basis for studying their structure–activity relationships.

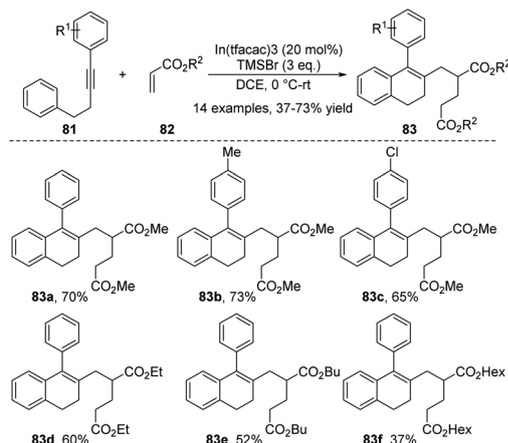
In 2015, the group of Li discovered a FeCl₃·6H₂O catalyzed oxidative coupling reaction of **75** and **79** (Scheme 26).³⁹ In contrast to the oxidation dimerization of **75** and **79** facilitated by conventional oxidants such as HPR-H₂O₂ or Cu(NO₃)₂·3H₂O, this approach boasts superior yields and selectivity, thereby broadening its applicability in the synthesis of natural products. The reaction mechanism can be best explained by the presence of the unstable intermediate **76**, which was easily generated at the start and underwent further 6π electrocyclic cyclization to form dihydronaphthalene structures under the acidic oxidative condition. The author also described the removal of *tert*-butyl protecting groups from the two coupling intermediates **77** or **78** and other related transformations. Further treatment of compound **80** with excess LiAlH₄ in THF for global debromination ultimately accomplished the concise synthesis of racemic natural ligandamide.

A combination Lewis acid system composed of two or more Lewis acids occasionally exhibits enhanced catalytic activity in organic transformations that cannot be achieved by any single



Scheme 26 Synthesis of ADHNs via regioselective oxidative coupling reaction.

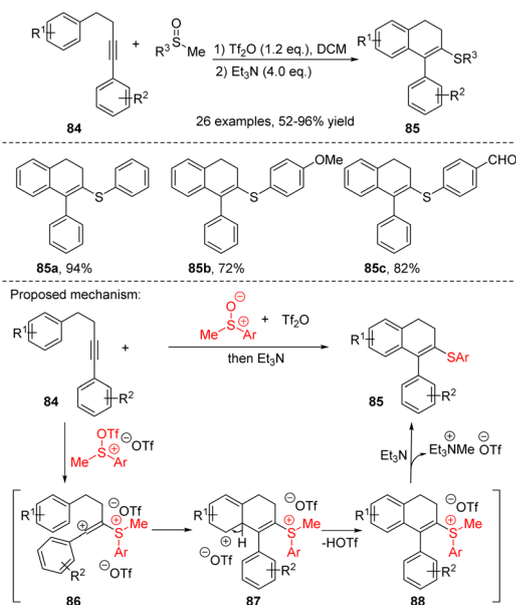




Scheme 27 Synthesis of ADHNs *via* Lewis acid catalyzed cascade cyclization reaction.

component alone. In 2018, Loh and co-workers disclosed the construction of ADHNs **83** *via* a cascade cyclization reaction of diarylalkynes **81** with acrylates **82** under the catalysis of a combined Lewis acid derived from $\text{In}(\text{III})$ salt and TMSBr (Scheme 27).⁴⁰ Both indium(III) and TMSBr are indispensable for the efficient progress of the reaction. This reaction sequence completed the formation of multiple new C–C bonds in minimal steps, aligns well with the principles of green chemistry.

During the synthesis of 3,4-dihydronaphthalene derivatives from 4-aryl alkynes, diverse electrophilic reagents can induce the cyclization process, thereby allowing the incorporation of various functional groups into the products. In 2019, Li, Xu, Du *et al.* reported a sulfur-mediated electrophilic cyclization reaction of aryl-tethered internal alkynes **84**, leading to the synthesis of 3-sulfenyl-1,2-dihydronaphthalenes **85** (Scheme 28).⁴¹ In this



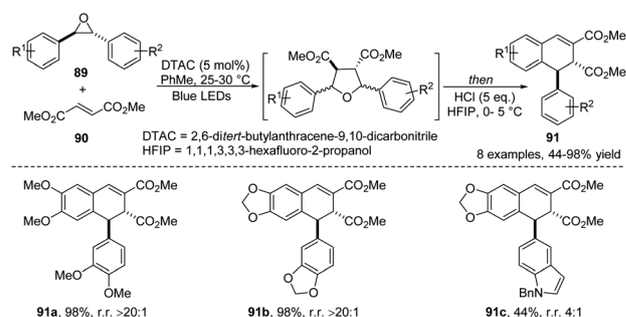
Scheme 28 Synthesis of ADHNs *via* sulfur-mediated electrophilic cyclization reaction of aryl-tethered internal alkynes.

process, the sulfoxide activated by anhydride serves as a sulfur electrophile to trigger the intramolecular electrophilic cyclization of the reactive aryl-tethered alkynes. This reaction exhibits broad functional group tolerance and excellent yields, enabling the synthesis of sulfenyl-substituted phenanthrenes, dihydroquinolines, 2*H*-chromenes, and coumarins with yields ranging from high to moderate. The proposed mechanism is illustrated in Scheme 32, the strongly electrophilic reagent sulfoxide activated by anhydride reacted with a carbon–carbon triple bond to form a vinyl carbocation intermediate **86**. The intermediate **86** can be captured by the tethered aryl ring, followed by deprotonation to generate the intermediate **87** and sulfonium salt **88**. Subsequently, demethylation was achieved using Et_3N ($\text{S}_{\text{N}}2$ process) to obtain the desired product.

In 2020, Beeler and co-workers presented a study on one-pot synthesis of ADHNs **91** *via* a concerted [3 + 2] dipolar cycloaddition reaction between epoxide **89** and dimethyl fumarate **90** (Scheme 29).⁴² In this transformation, dimethyl fumarate served as a dipolarophile. It is also noteworthy that temperature control is crucial for minimizing the undesired side products and obtaining high selectivity. In some cases, the lower yield can be rationally explained by the dearylation decomposition pathway of the resulting dihydronaphthalene into naphthalene. The obtained product **91** can be mapped as a naturally occurring lignan, which provides a basis for further study of the biological activities of ADHN derivatives.

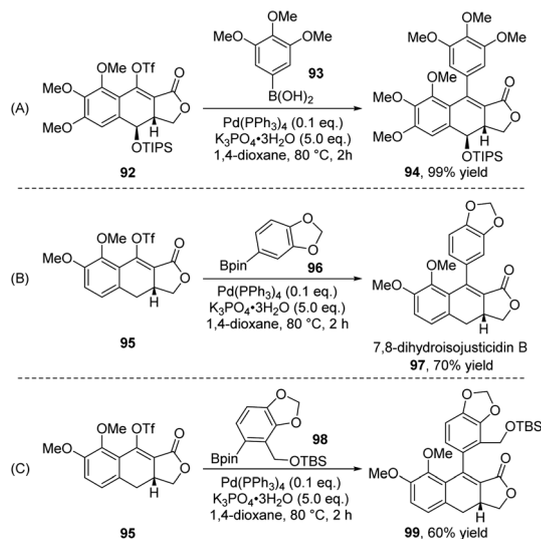
In 2021, Gao and co-workers revealed a palladium-catalyzed Suzuki–Miyaura coupling of **92** with aromatic boronic acid **93** resulting in **94** (Scheme 30A). Subsequently, a Pd^0 catalyzed Suzuki–Miyaura coupling of **95** with aryl boronic acid pinacol ester **96** or **98** afforded (–)-7,8-dihydroisojusticidin B (**97**) and the desired coupled product **99** in 70% and 60% yield, respectively (Scheme 30B and C).⁴³ The potentiality of this strategy has been well demonstrated by the preparation of ADHN-type lignans, such as linoxepin and 7,8-dihydroisojusticidin B, through the cross-coupling reactions. The ADHNs generated by this process can be further converted into aryltetralin cyclic ether lignans aglacins A, B and E *via* asymmetric photo-enolization/Diels–Alder (APEDA) reactions.

2.2.2 Addition reactions. The utilization of transition metal-catalyzed additions involving diverse nucleophiles to activate olefins has emerged as an invaluable approach in



Scheme 29 Synthesis of ADHNs *via* a concerted [3 + 2] dipolar cycloaddition of epoxide and dimethyl fumarate.



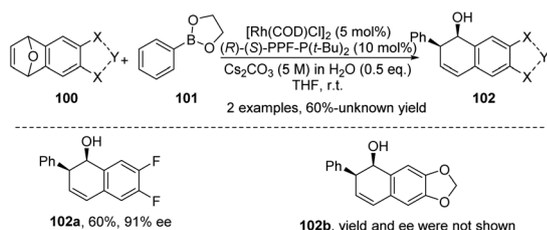


Scheme 30 Synthesis of ADHNs *via* coupling reaction of *N*-tosylhydrazones with aryl iodides.

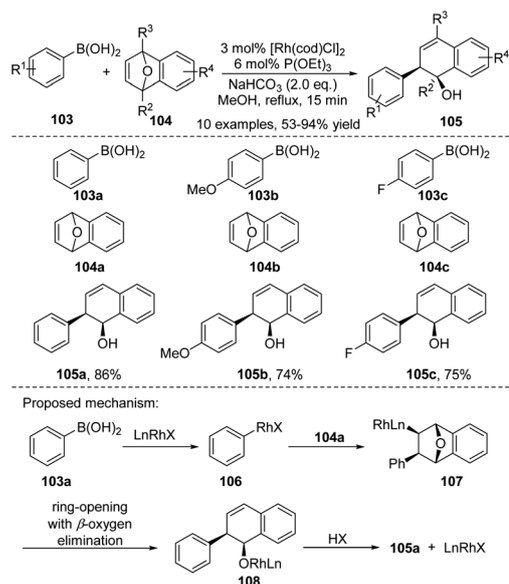
synthetic chemistry.⁴⁴ The addition of boronic acid to olefins has been recognized as a powerful tool in the field of highly enantioselective synthesis due to its convenience and simplicity, and has achieved significant development in recent years.

In 2002, Lautens and co-workers reported the first rhodium(i)-catalyzed asymmetric addition of organoboronic acids or boronic ester **101** to oxabicyclic alkenes **100** (Scheme 31).⁴⁵ The reaction features very mild reaction conditions, generating multiple stereocenters with high yields and excellent diastereoselectivity and enantioselectivity. In this scheme, arylboronic species served as nucleophilic partners. The optimization process replied that a minimal amount of water was required for the catalytic turnover of the reaction. The substrate scope study indicated that substituents on the phenyl ring *ortho* to boron were intolerant under the optimized reaction conditions. Unfortunately, only two examples of ADHN derivatives were shown in this protocol, and the reaction with **100b** led to a complex mixture of products.

Subsequently, in the same year, using the similar reagents, Murakami and co-workers achieved a rhodium(i)-catalysed addition reaction of arylboronic acids **103** with oxabenzonornbornadienes **104** to afford *cis*-2-aryl-1,2-dihydro-1-naphthol stereoselectively, a series of products were obtained in good



Scheme 31 Synthesis of ADHNs *via* rhodium(i)-catalysed asymmetric addition of organoboronic ester with oxabicyclic alkenes.

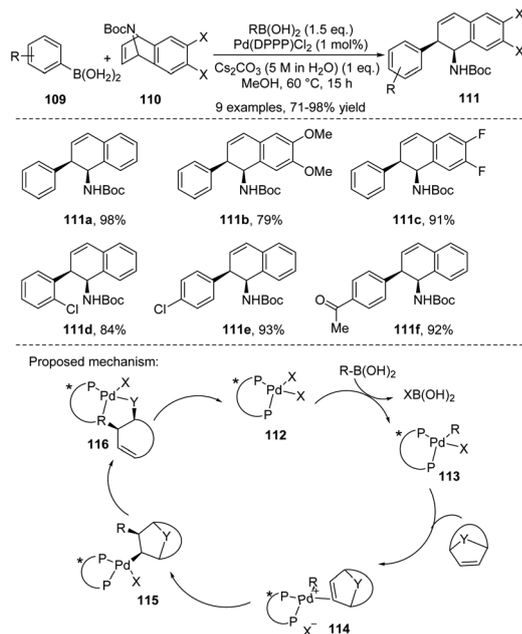


Scheme 32 Synthesis of ADHNs *via* rhodium(i)-catalysed addition reaction of arylboronic acids with oxabenzonornbornadienes.

yields without concomitant deboronation of the boronic acid (Scheme 32).⁴⁶ In this reaction, the rhodium complex was prepared *in situ* by refluxing [Rh(cod)Cl]₂ and P(OEt)₃ (Rh : P = 1 : 2) in MeOH under a nitrogen atmosphere. Interestingly, the hydrolysis deboronation of phenylboronic acid was minimized under the current conditions, as only 1.1 equivalents of boronic acid **103a** were sufficient to obtain the product **105a** in 86% yield. In previously reported rhodium-catalyzed addition reactions of this type, competitive hydrolysis degradation reactions demanded the use of excess arylboronic acids, ranging from 1.4 to 10 equivalents. The authors proposed a plausible mechanism, which initially involves a metal transformation of rhodium(i) with phenylboronic acid **103a** to generate phenylrhodium(i) **106**. This process might be promoted by a base. Subsequently, the addition of the phenyl-rhodium linkage across a carbon-carbon double bond of **104a** from the *exo*-side to give **107**. β-Oxygen elimination then proceeds to open the furyl ring. The resulting rhodium(i) alkoxide **108** underwent protonolysis with methanol or phenylboronic acid **103a** to produce the alcohol **105a** and regenerate rhodium(i) for a new catalytic cycle.

Due to the challenges encountered in Lautens' previously reported reaction (Scheme 26), particularly pertaining to the addition of heteroaryl boronic acids, which frequently leads to the formation of unreacted addition products and oligomers. Lautens and his colleagues have continued their efforts to further enhance the yield of these reactions. They broadened their studies by investigating the boronic acid addition reaction with different metals. In 2003, the authors' group reported a palladium(ii) catalyzed ring-opening addition of arylboronic acids **109** to heterobicyclic alkenes **110** (Scheme 33).⁴⁷ Under these optimized condition, a wide variety of arylboronic acids were tolerated, leading to the target products with excellent yields (71–98%). The reaction optimization process revealed that metal

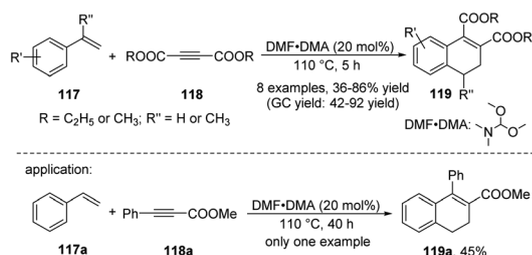




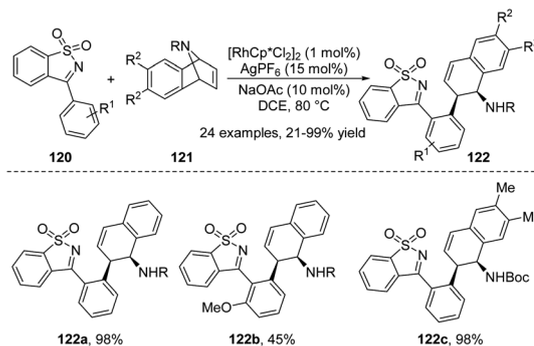
Scheme 33 Synthesis of ADHNs via palladium(II) catalyzed ring-opening addition of arylboronic acids with heterobicyclic alkenes.

catalysts such as nickel and platinum were unreactive, inorganic bases and water were necessary for the good reactivity, and methanol had a promoting effect on the reaction. Among the ligands screened, DPPP (1,3-bis(diphenylphosphino)propane) showed the best effect. The authors' group proposed a plausible mechanism, firstly, in methanol solvent, the reactive Pd(ligand)(OMe)₂ **112** was generated. Then, the aryl boron underwent a transmetalation process to give intermediate **113**. After the dissociation of the X ligand, the heterobicyclic alkene substrate (Y = oxygen or protected nitrogen) complexes to form the cationic complex **114**. Finally, β-heteroatom elimination after carbopalladation results in the ring-opening product.

In 2007, Hua *et al.* demonstrated the DMF·DMA (*N,N*-dimethylformamide dimethyl acetal) catalyzed cycloaddition reaction of vinylarenes **117** with electron-deficient alkynes **118** for the highly efficient synthesis of 1,2-disubstituted-3,4-dihydronaphthalenes **119** (Scheme 34).⁴⁸ The main advantages of this protocol are the availability of substrates, the low cost and stability of the DMF·DMA, and the ability to conduct the catalytic reaction under air. The screening of reaction



Scheme 34 Synthesis of ADHNs via DMF·DMA-catalyzed cyclic addition of vinylarenes with acetylenedicarboxylate.



Scheme 35 Synthesis of ADHNs via rhodium(III)-catalyzed ring-opening addition reaction of azabenzonorbornadienes with *N*-sulfonyl ketimines.

conditions showed that the catalysts used and the ratio of **117** to **118** had significant effects on the product yield. Notably, the reaction exhibited complete regioselectivity, with only **119a** being detected by GC and GC-MS.

In 2019, Fan, Chen *et al.* reported the formation of 2-aryl dihydronaphthalene derivatives **122** through a rhodium(III)-catalyzed ring-opening addition reaction between azabenzonorbornadienes **120** and cyclic *N*-sulfonyl ketimines **121** (Scheme 35).⁴⁹ The reaction exhibited good efficiency and excellent regioselectivity under the redox-neutral conditions. It has proved that the use of the additive is essential, solvents other than DCE, including tetrahydrofuran, acetonitrile, 1,4-dioxane, and toluene, are detrimental to the conversion process. The assessment of substrate scope showed that under optimized conditions, various electron-donating substituents such as methyl, ethyl, tertiary butyl, methoxyl, or phenoxy on the phenyl ring of the *N*-sulfonyl ketimines were well tolerated, yielding the products in good to excellent yields. Nevertheless, the conversion of oxabenzonorbornadienes proved unsuccessful, resulting in the formation of naphthalen-1-ol as the major byproduct.

3. Applications of ADHNs

3.1 Synthetic applications

The ADHN, aryl naphthalene and aryl tetrahydronaphthalene skeletons are widely existed in natural products and bioactive compounds. Among which the ADHN skeletons are especially useful because it can be transferred into naturally occurring aryl naphthalene and aryl tetrahydronaphthalene molecules *via* a one-step reaction. Therefore, the synthesis and transformation of ADHNs for the establishment of aryl naphthalene and aryl tetrahydronaphthalene have attracted much attention among organic chemists, and many elegant synthetic methods for constructing ADHN-type natural products have been reported.

3.1.1 Synthesis of ADHNs-type natural products and bioactive compounds. Previously, natural product chemists have reported a large number of natural products or bioactive compounds containing ADHN skeletons (Fig. 2). Subsequently,



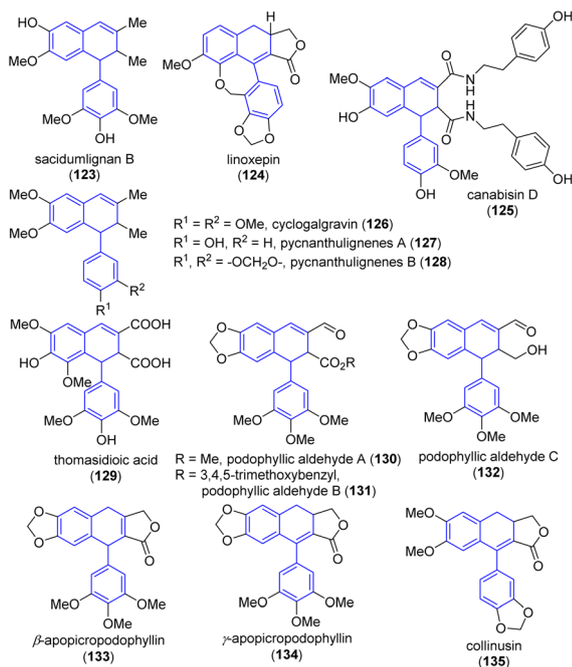


Fig. 2 Natural products or bioactive compounds containing ADHN skeletons.

organic chemists shifted their attention towards its synthesis, reporting numerous facile and efficient synthetic methods.

Sacidumlignan B (123) (Fig. 2) is a natural product belong to typical 2,7'-cycloignans, which have already attracted broad attention from the synthetic community. In 2005, Yue and co-workers isolated sacidumlignan B from the EtOH extract of the plant *Sarcostemma acidum* (Roxb.) in Hainan Island of China and confirmed its relative configuration.⁵⁰ In 2012, Ramana and co-workers first reported the asymmetric total synthesis of the naturally occurring sacidumlignan B and determined its absolute configuration.⁵¹ In 2020, the Peng group reported a base-promoted addition of *N,N*-dimethylacetamide (DMA) to 1,1-diarylethylenes, and successfully applied this method to the total synthesis of (–)-sacidumlignan B with an overall yield of 3%.⁵² Although this protocol avoids the use of transition metal catalysts, providing simple operation and broad substrate scope, the route comprises 16 steps. Therefore, they continued their research to explore more efficient strategies. Encouragingly, in 2022, the same group first reported the racemic synthesis of this molecule, featuring seven steps (46% overall yield) from a known compound (38% overall yield, nine steps from commercially available syringol/syringaldehyde).⁵³

Linoxetine (124) is a natural ADHN lignan, which was first isolated from the dichloromethane extract of aerial parts of *Linum perenne* L. (Linaceae) and first named by Schmidt's group in 2007. Its absolute configuration was determined using CD spectroscopy combined with DFT calculations.⁵⁴ In 2013, Tietze *et al.* reported the first total synthesis of (+)- and (–)-linoxetine in only ten steps without the use of any protecting groups, achieving an overall yield of 30%.²⁹ In this transformation, the Sonogashira reaction and the domino carbopalladation/Heck

reaction of allylsilane played significant roles in delivering the target compound. Later, in the same year, Lautens and co-workers reported for the first time an eight-step asymmetric synthesis method of (+)-linoxetine, which was achieved through the modified version of Catellani reaction for the synthesis of this natural product with high enantioselectivity and without protective groups.³⁰ Shortly thereafter, in 2014, the Tietze group further reported the enantioselective total synthesis of the natural lignan (+)-linoxetine, starting from commercially available raw materials through a total of 11 steps with an overall yield of 27%.⁵⁵ The key steps of which include the domino carbopalladation/Mizoroki–Heck reaction for the formation of the pentacyclic system, asymmetric hydroboration, and oxidative lactonization. In 2015, Nagasawa, Yamanaka and co-workers reported an organocatalytic oxidative kinetic resolution approach for the enantioselective synthesis of (+)-linoxetine from tetralone-derived β -ketoesters in only six steps.⁵⁶ In 2021, Gao *et al.* reported an asymmetric photoenolization/Diels–Alder (PEDA) reaction between electron-rich 2-methylbenzaldehydes and unsaturated γ -lactones, which allowed the construction of (+)-linoxetine in six steps.⁴³ Recently, in 2024, Peng, Xiao and co-workers reported a nickel-catalyzed reductive tandem cyclization of the elaborated β -bromo acetal with a dibenzoxepin scaffold to realize enantioselective total synthesis of linoxetine.³⁵

Canabisisin D (125) is a naturally occurring ligandamide which can be found in fruits or seeds of several plants. In 2015, Li *et al.* reported a regioselective oxidative coupling reaction for the synthesis of natural (\pm)-canabisisin D, in which, *tert*-butyl or bromine atom was introduced into the C-5 of *N-trans*-feruloyltramine precursor to hamper the 8–5 coupling product.³⁹ This biomimetic synthetic strategy can be extensively investigated in the synthesis of a variety of naturally occurring ligandamides for the further study of their structure–activity relationships.

Cyclogalgravin (126) is a natural product which was extracted from *Araucaria angustifolia* knots.⁵⁷ The air-dried and ground roots of *P. angolensis* were extracted with a mixture of CH_2Cl_2 – CH_3OH (1 : 1) at room temperature to provide pycnanthulignenes A (127) and B (128).⁵⁸ In 2011, the Barker and Rye group reported the first asymmetric synthesis of three types of natural ADHNs such as cyclogalgravin (125), pycnanthulignene A (127) and pycnanthulignene B (128).⁵⁹ A common chiral precursor, aza-Claisen-derived amide, was the starting material for each of these structurally ADHNs. In more detail, the synthesis of pycnanthulignenes A and B began from aldehyde, then it led to the corresponding alcohol, which was then be treated with mesyl chloride to give the ADHN intermediate, the further deprotection of the methoxymethyl (MOM) ether or treatment of alcohol with mesyl chloride gave the desired pycnanthulignene A and pycnanthulignene B in 86% and 95% yield, respectively. In 2020, Beeler and co-workers reported an *in situ* Finkelstein reaction with NaI and single electron transfer (SET) reduction with Zn of mesylated alcohol to offer pycnanthulignene B in 73% yield.⁴²

In 2002, Niwa and co-workers reported the synthesis of thomasidiolic acid 129 from dehydrosinapinic acid dilactone *via* an α,β -unsaturated γ -lactone-type dimer in phosphate buffer



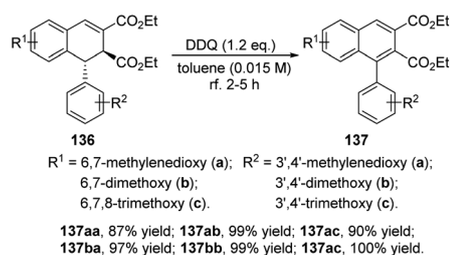
(pH 7.4).⁶⁰ In this work, sinapinic acid was used as the starting material. Differed from the previous reported synthetic methods (under acid conditions),^{5b} this approach finished the target conversion under neutral conditions. Later, in 2008, using sinapinic acid as the same starting material, Orlandi and co-workers achieved enantioselective biomimetic synthesis of thomasidioic acid through asymmetric biomimetic oxidation of the intermediate phenol, yielding a racemic mixture of *trans*-thomasidioic acid dimethylesters with a 70% yield.⁶¹

In 2014, Nishii and co-workers first reported the total synthesis of three (+)-podophyllin aldehydes **130–132**, with corresponding yields and enantiomeric excess (ee) of 85%, 72% and 97%, 95% ee, 95% ee, and 95% ee, respectively.⁶² The overall yields of (+)-podophyllin aldehydes A, B, and C were 30% (16 steps), 26% (16 steps), and 43% (8 steps), respectively. The key steps of which were the organocatalyzed highly enantioselective cyclopropanation and Lewis acid-mediated chiral transfer ring expansion of the cyclopropane. Notably, the transformation was accomplished with remarkable enantiomeric control.

In 2018, Peng *et al.* reported the regiodivergent oxidation of (+)-deoxypodophyllin in the synthesis of bioactive ADHNs (+)- β - and (+)- γ -apopodophyllins (**133–134**).⁶³ The *in situ* *syn*-elimination of the phenylselenoxide generated by the oxidation of α -phenylselenide provided (+)- β -apopodophyllins **133** in 88% yield. In contrast, the subjection of β -phenylselenide to *m*-chloroperoxybenzoic acid (*m*-CPBA) led to (+)- γ -apopodophyllins **134** in 88% yield.

The same year, Giri and co-workers reported a Ni-catalyzed tandem cyclization/cross-coupling regioselective dicarbon functionalization of unactivated olefins, and applied this method to the concise synthesis of (\pm)-collinusin **135** on a gram-scale.⁶⁴ In this protocol, the authors utilized the highly functionalized (2-aryoyl)zinc iodide as the coupling partner, the carbonylbutyrolactone core was synthesized in one pot and two step procedure, affording a yield of 70% (2.5 mmol scale, 0.672 g). Subsequently, the carbonylbutyrolactone was treated with lithium diisopropylamide (LDA) and SOCl₂ to furnish (\pm)-collinusin with a 65% yield. Noteworthy, the concise synthesis of (\pm)-collinusin *via* this method is feasible due to the ready access to arylzinc reagents containing carbonyl groups.

3.1.2 For the synthesis of aryl-naphthalenes. Arylnaphthalene lactones are natural products which can be isolated from a wide range of plants and have the significant biological activities including cytotoxicity, antimicrobial, diuretic, and ion channel blocking. ADHN skeleton serves as a versatile and important precursor for the formation of aryl-naphthalene-type natural products and bioactive compounds. DDQ (2,3-dichloro-5,6-dicyano-1,4-benzoquinone) is a useful oxidant that has found widespread application in numerous oxidation processes. In 2000, Cow and co-workers reported a DDQ facilitated oxidation of aryl-1,2-dihydronaphthalenes **136** for the preparation of aryl-naphthalenes **137** (Scheme 36), which then underwent hydrolysis, reduction and lactonization reaction to afford a series of natural lactones containing aryl-naphthalene skeletons, such as taiwanin C, chinensin, justicidin B, dehydroanhydrodiphyllotoxin, and 7-deoxydiphyllin.^{4a} In 2014,

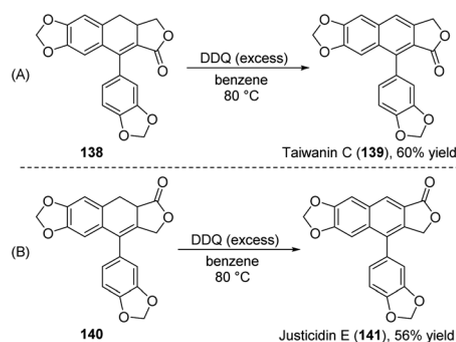


Scheme 36 Synthesis of aryl-naphthalenes *via* DDQ-mediated oxidation of ADHNs.

Shin, Seo and co-workers reported a facile and regioselective intramolecular Diels–Alder reaction of 3-arylprop-2-ynyl 3-aryl-propiolates **138** and **140**, followed by a subsequent DDQ oxidation process, yielding aryl-naphthalene lactones, taiwanin C **139** and justicidin E **141**, with yields of 60% and 56%, respectively (Scheme 37A and B).⁶⁵ Notably, the current method showcases exceptional regioselectivity when compared to previously reported approaches.

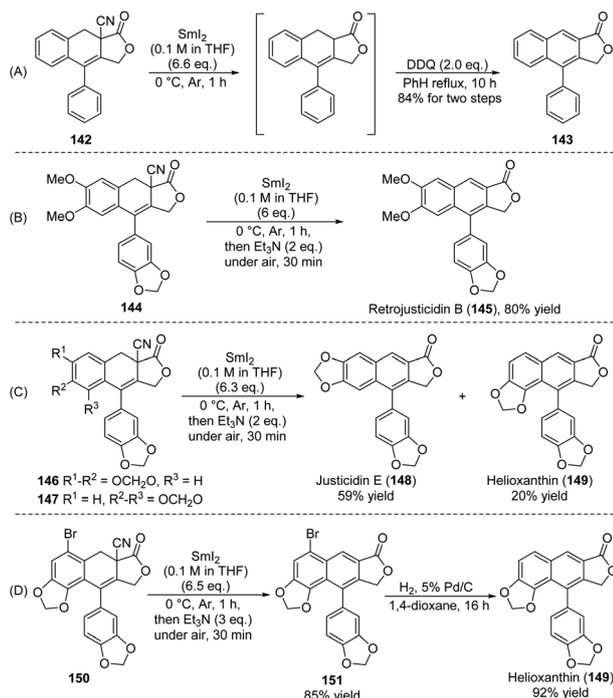
In 2015, Shia, Lin and co-workers reported a SmI₂ mediated decyanation reaction of α -cyano lactone **142**, followed by DDQ oxidation, to furnish the aryl-naphthalene lactones **143** (Scheme 38A).⁶⁶ Then, a similar approach was represented by this group, and it was used to construct the natural aryl-naphthalene lignans, retrojusticidin B **145**, justicidin E **148**, and helioxanthin **149** (Scheme 38B and D).

In 2022, Zhang, Chen and co-workers reported a versatile and flexible approach toward the synthesis of natural aryl-naphthalene lactone lignans, including justicidin B **154**, taiwanin C **157**, and justicidin E **148**, from ADHNs precursors (Scheme 39A and B).²⁰ In this channel, an aryl-alkyl Suzuki cross-coupling was introduced for the formation of the dioxinone unit, a cation-induced cyclization occurred to construct the aryl dihydronaphthalene, and base-mediated oxidative aromatization to furnish the aryl-naphthalene core. For example, the treatment of **152** or **155** with sodium methoxide in MeOH under air followed by the addition of Tf₂O and diisopropylethylamine (DIPEA) in CH₂Cl₂ produced the first common intermediate **153** or **156**, both with yields of 45%. In this work, an oxidative (by air) aromatization occurred under

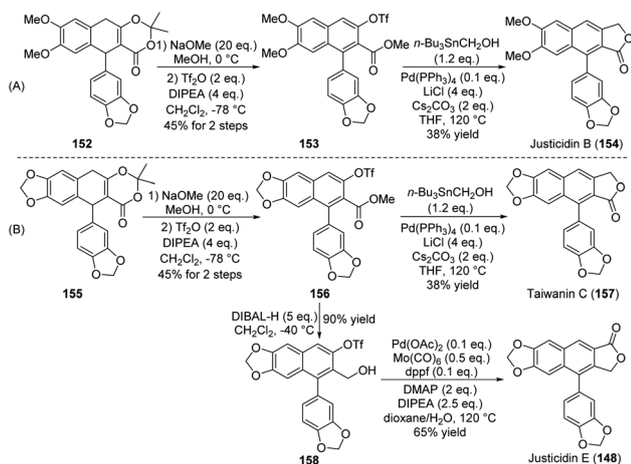


Scheme 37 Synthesis of aryl-naphthalene lactones, taiwanin C and justicidin E *via* DDQ-mediated oxidation of ADHNs.





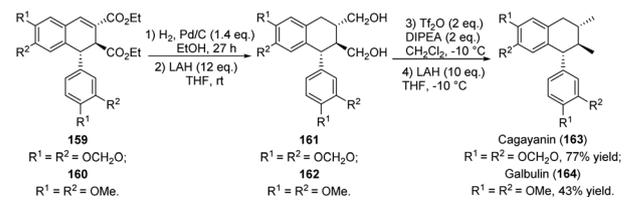
Scheme 38 Synthesis of arynaphthalene lactones, retrojusticidin B, justicidin E, and helioxanthin via Sml_2 -mediated reaction of ADHNs.



Scheme 39 Synthesis of arynaphthalene lactones, justicidin B, taiwanin C and justicidin E.

strong basic conditions. Then, in the presence of $Pd(PPh_3)_4$, Cs_2CO_3 , and LiCl, a Pd-catalyzed Stille cross-coupling reaction was performed between triflates **153** or **156** and tributylstannyl methanol. This was followed by spontaneous lactonization reaction, providing the naturally occurring justicidin B (**154**) or taiwanin C (**157**). Subsequently, reduction of **156** using diisobutylaluminum hydride (DIBAL-H) provided the alcohol **158** in 90% yield. The natural justicidin E (**148**) was furnished in 38% isolated yield via an improved Pd-catalyzed carbonylative lactonization of triflate **158** with $Mo(CO)_6$.

3.1.3 For the synthesis of aryltetrahydronaphthalenes. The ADHN skeleton is also a versatile and important precursor for



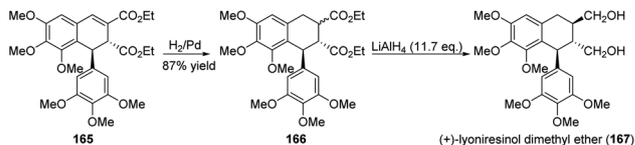
Scheme 40 Synthesis of aryltetrahydronaphthalene, cagayanin and galbulin via reduction reaction of ADHNs.

the formation of aryltetrahydronaphthalene-type natural products and bioactive compounds. In 2001, Charlton *et al.* reported an acid-catalyzed cyclization of 2,3-dibenzylidenesuccinates for the synthesis of lignans (\pm)-cagayanin **163** and (\pm)-galbulin **164** (Scheme 40),¹⁶ in which two important ADHNs-type intermediates, **159** and **160**, were used as the precursors. The present method should be applicable to the syntheses of other lignans and related compounds.

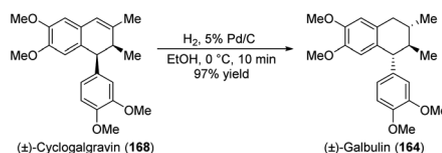
In 2004, the same group used the same starting material, 2,3-dibenzylidenesuccinate, and showed an asymmetric photocyclization to produce a chiral ADHN. Subsequent hydrolysis and re-esterification provided the precursor **165**, the reduction reaction of the double bond of **165** offered ester **166**, then providing (+)-lyoniresinol dimethyl ether **167** (Scheme 41) as a major product, along with its 2,3-*cis* diastereomer as a minor product.¹⁷

Similarly, in 2013, Peng and co-workers reported the synthesis of (\pm)-galbulin (**164**) via hydrogen and Pd/C catalyzed reduction of cyclogalgravin (**168**) (Scheme 42).⁶⁷ Notably, the cyclogalgravin emerged as a pivotal precursor facilitating the collective synthesis of stereodivergent 2,7'-cyclo-lignans, featuring either *anti-anti* or (*anti*)-*syn-anti* substituent arrangements within the tetrahydronaphthalene scaffold.

In 2020, Studer and co-workers also presented the synthesis of natural product (+)-Galbulin (**164**). The remarkable transformation of the ADHN precursor was outlined in Scheme 43,⁶⁸ in which the incorporation of pinacol into **169** yielded the corresponding boronic ester **170**. Notably, the extreme instability

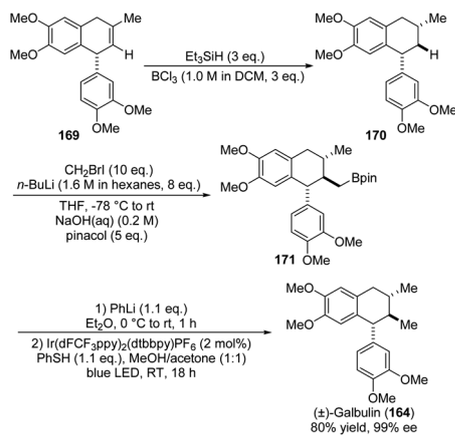


Scheme 41 Synthesis of aryltetrahydronaphthalene, lyoniresinol dimethyl ether via reduction reaction of ADHNs.



Scheme 42 Synthesis of aryltetrahydronaphthalene, galbulin via reduction reaction of cyclogalgravin.



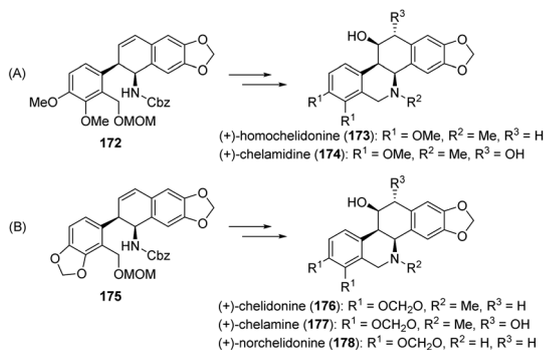


Scheme 43 Synthesis of aryltetrahydronaphthalen, galbulin via oxidation under photoredox conditions.

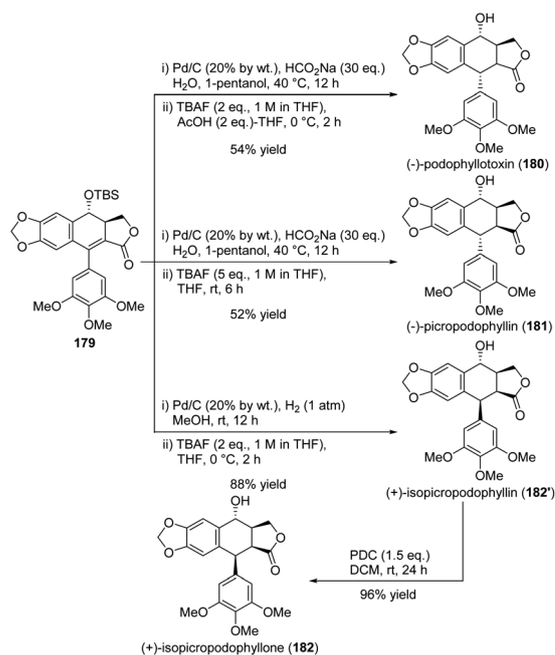
of **170** necessitates its immediate progression through the subsequent reaction sequence without any further purification. Unfortunately, due to potential steric hindrance, **170** failed to transform into the boronic ester **171**. Then, transesterification with pinacol yielded the pinacol boronic ester **171**. Ultimately, the final protodeboronation of **171** led to (\pm)-galbulin (**164**). Despite the formidable challenge posed by the final step, they successfully achieved this through the oxidation under photoredox conditions and the subsequent capture of the generated primary alkyl radical with a thiophenol.

In 2008, Lautens and co-workers completed the first enantioselective total synthesis of aryltetrahydronaphthalen-type natural products, (+)-homochelidonine **173**, (+)-chelamidine **174**, (+)-chelidonine **176**, (+)-chelamine **177** and (+)-norchelidonine **178** from ADHNs precursors **172** and **175** by several steps (Scheme 44A and B, the details were not shown here).⁶⁹ The synthetic method reported here provided possibility to prepare structural analogues of the hexahydrobenzo[*c*]phenanthridine alkaloids with potentially improved pharmacological properties.

Podophyllotoxin and its natural analogues, isolated from *Podophyllum* species, have been extensively studied over the past two centuries due to their significant biological activities and structural diversity. However, the catalytic and



Scheme 44 Enantioselective total synthesis of aryltetrahydronaphthalen, (+)-homochelidonine, (+)-chelamidine, (+)-chelidonine, (+)-chelamine and (+)-norchelidonine from ADHNs.

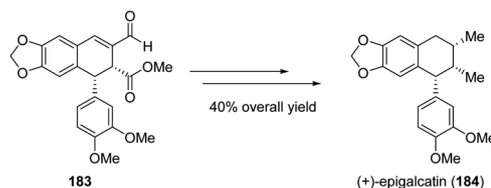


Scheme 45 Synthesis of aryltetrahydronaphthalen, (-)-podophyllotoxin, (-)-picropodophyllin, (+)-isopicropodophyllin, (+)-isopicropodophyllone from ADHN.

enantioselective concise total synthesis of **180** remains an unmet challenge. In 2017, Hajra and co-workers reported the first catalytic enantioselective synthesis of (-)-podophyllotoxin **180**, (-)-picropodophyllin **181**, (+)-isopicropodophyllin **182'**, (+)-isopicropodophyllone **182** (Scheme 45) from a common ADHN intermediate.³²

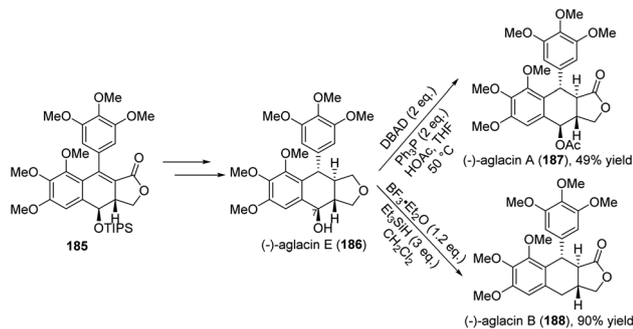
Previously, (+)-epigalcatin has not yet been synthesized or found in natural sources. In 2018, Czarnocki and Lisiecki presented the first total synthesis of (+)-epigalcatin **184** using ADHN precursor **183** as the raw material, achieving a three step synthesis yield of 40% (Scheme 46).⁷⁰ With *L*-prolinol as the chiral source, the target product **184** was synthesized in 11 steps with an overall yield of 5%. The product **184** can be obtained in a highly stereoselective manner from piperonal, 3,4-dimethylbenzaldehyde, and diethyl succinate. In this scheme, a key intermediate cyclolignan was formed through the continuous flow photocyclization of a chiral atropisomeric 1,2-bisbenzylidenesuccinate amide ester. This strategy may allow access to cyclolignan analogues that are inaccessible from natural plant sources.

In 2021, Gao *et al.* reported the first asymmetric total synthesis and absolute configuration revision of naturally occurring aryltetralin cyclic ether lignans aglacin A, B, and E



Scheme 46 The first total synthesis of (+)-epigalcatin from ADHN.





Scheme 47 The first asymmetric total synthesis of aglacin A, B, and E from the ADHN precursor.

from ADHN precursor 185 based on the asymmetric photoenolization/Diels-Alder (PEDA) reaction between electron-rich 2-methylbenzaldehydes and unsaturated γ -lactones (Scheme 47).⁴³ The Lewis acid $[Ti(O^iPr)_4]$ played a pivotal role in the PEDA reaction by activating the inert dienophiles and controlling the diastereoselectivity. Upon obtaining aglacins A (187) and B (188), aglacins A (187) and B (188) can be derived through the manipulation of the C7 hydroxyl group of aglacins A (187) and B (188).

3.2 Bioactivity applications

The structural diversity of ADHN-type lignans determines their extensive range of biological activities. Currently, pharmaceutical chemists have reported the antitumor, antimicrobial, antiviral and neuroprotective activities of ADHNs derivatives. The overview of their biological activities is shown in Fig. 3.

3.2.1 Antitumor activity. Tumors pose a formidable threat to human health,⁷¹ with high incidence and mortality rates. Their primary harm lies in their uncontrolled proliferation, which disrupts the structure and function of normal tissues and organs, triggering dire consequences such as pain, bleeding, infection, and organ failure. Traditional therapeutic interventions include surgery, chemotherapy, and radiotherapy.⁷² Chemotherapy, renowned for its capacity to achieve comprehensive systemic treatment, efficacy, and versatile modes of administration, is an indispensable clinical option in cancer

therapy.⁷³ Despite the availability of several effective anti-tumor chemotherapeutic drugs, the lack of selectivity, the intrinsic or acquired resistance of tumors to chemotherapy, as well as severe toxic and side effects, have consistently posed major obstacles to chemotherapy. Therefore, the development of novel anti-tumor drugs with high selectivity, no cross-resistance, and low toxicity remains a synthetic goal in the field of bioorganic chemistry.⁷⁴ Some representative antitumor agents containing ADHNs are listed in Fig. 4.

Cyclolignans podophyllotoxin and its derivatives etoposide and teniposide are widely used anticancer drugs, exhibiting good clinical effects in treating various neoplasms, such as testicular and small-cell lung cancer, lymphoma, leukaemia, and Kaposi's sarcoma.⁷⁵ Nevertheless, limitations such as myelosuppression, the emergence of drug resistance, and cytotoxicity towards normal cells still persist. Therefore, it is necessary to develop new cyclolignans.

Previously, (-)-podophyllic aldehyde A (189), a form of cyclolignans, was tested by Gordaliza's group at the National Cancer Institute (USA) against a panel of 60 diverse cancer types. Notably, it emerged as one of the most potent compounds, displaying highly selective cytotoxicity, particularly towards colon carcinoma lines.⁷⁶ Subsequently, to further substantiate the impact of the aldehyde group on antineoplastic selectivity and establish the structure-activity relationships of these compounds, the same research team prepared and tested other aldehydes derived from podophyllotoxin. Gordaliza and

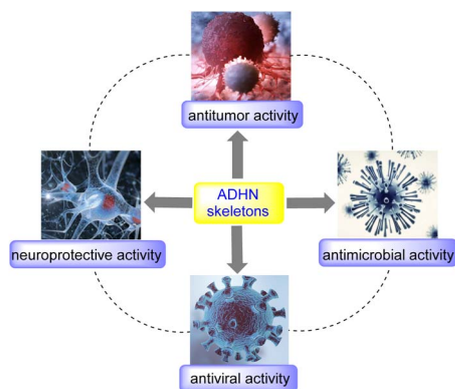


Fig. 3 The summary of biological activities of ADHNs derivatives (the images are sourced from the website: <https://cn.bing.com>).

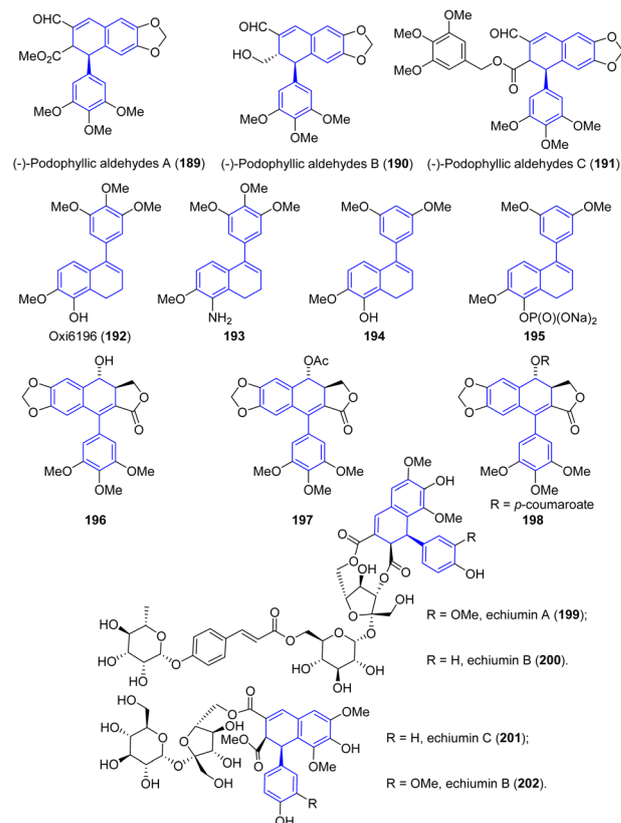


Fig. 4 Representative antitumor agents containing ADHNs.



co-workers discovered that (–)-podophyllic aldehydes B (**190**)⁷⁷ and C (**191**)⁷⁸ exhibit significant antineoplastic cytotoxicity and apoptosis-inducing activities (Fig. 4).⁶²

Vascular disrupting agents (VDAs) represent one class of vascular targeting agents (VTAs) that specifically target tumor vasculature. The anti-tumor mechanism of VDAs involves inducing microtubule depolymerization and cytoskeletal rearrangement, ultimately resulting in irreversible and selective damage to the microvasculature within the tumor microenvironment.⁷⁹ In 2004, Pinney and colleagues reported that dihydronaphthalene Oxi6196 (**192**) possesses potent cytotoxicity and can be used as an inhibitor of tubulin assembly and a VDA (Fig. 4).⁸⁰

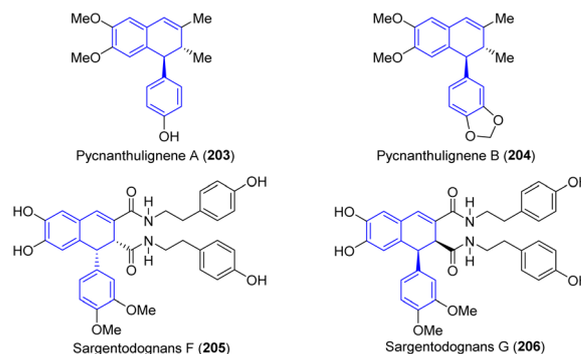
Microtubules are found in the cytoskeleton of almost all eukaryotic cell types and are directly involved in a variety of cellular functions such as cell movement, transport of organelles inside the cell, maintenance of cell shape as well as mitosis and cell replication.⁸¹ Therefore, perturbations in tubulin assembly/disassembly have emerged as a popular target for novel chemotherapeutic drugs. In 2012, Alami, Provot and co-workers designed and synthesized a series of new dihydronaphthalene derivatives. Biological evaluation of these compounds revealed that compounds **192–194** had high cell growth inhibition (at the nanomolar level) against four tumour cell lines.³⁸ The effect of the lead compound **192** on cancer cells was associated with cell cycle arrest in the G2/M phase.

Subsequently, in 2018, Pinney and co-workers found that compound **192** is a potent tubulin polymerization inhibitor (IC₅₀ = 1.0 μM) with low nM cytotoxicity against human cancer cell lines.³³ To further improve the water-solubility for *in vivo* evaluation, the phosphate salt prodrug compound **195** was synthesized and subjected to preliminary *in vitro* and *in vivo* (mouse and rat) evaluation. The results showed that the compound **195** has inhibition of tubulin polymerization (cell-free assay) and cytotoxicity *in vitro* against NCI-H460 (non-small cell lung carcinoma), DU-145 (prostate carcinoma), and SKOV-3 (ovarian adenocarcinoma) human cancer cell lines, and disruption of tumor-associated vasculature, thereby functioning as VDAs *in vivo*.

Alvarez, Salas-Vidal and co-workers demonstrated that compounds **196–198** exhibit varying degrees of cytotoxic activity against human nasopharyngeal (KB), colon (HF-6), breast (MCF-7) and prostate (PC-3) cancer cell lines, with IC₅₀ values ranging from 1.49 to 1.0 × 10⁵ μM (Fig. 4).⁸² The *in vivo* zebrafish embryos experiments indicated that these compounds may exert an antimitotic effect by disturbing tubulin. The study also showed that compounds **196–198** promote mitotic arrest and induce morphological changes in a similar manner to that of the antimitotic drugs nocodazole and podophyllotoxin.

Four previously undescribed ADHN-type lignans, echiumins A–D (**199–202**) (Fig. 4), were isolated and identified from the butanol fraction of *Echium angustifolium* Mill, displaying strong to weak antitumor activity against HepG2 and MCF7 cancer cell lines.⁸³ Among them, echiumins A and D have the strongest anticancer activities.

3.2.2 Antimicrobial activity. Bacteria and fungi are important human pathogens that cause a variety of diseases.



Scheme 48 Representative antimicrobial agents containing ADHNs.

Antimicrobial resistance poses an increasingly serious threat to public health and renders established therapies ineffective.⁸⁴ Therefore, discovering new antibacterial and anti-fungal lead compounds is imperative to address this pressing challenge.

In 2010, Nkengfack and colleagues reported two novel cyclolignene derivatives, named pycnanthulignene A (**203**) and pycnanthulignene B (**204**), which were extracted and isolated from the air-dried and ground roots of *Pycnanthus angolensis* (Scheme 48).⁵⁸ Compound **203** was evaluated *in vitro* for antimicrobial activity against a panel of drug-resistant bacteria and pathogenic fungi strains, including one Gram-positive bacteria, methicillin-resistant *Staphylococcus aureus* (MRSA, LMP805), and six Gram-negative bacteria including β-lactamase positive (L+) *Escherichia coli* (L+ EC, LMP701), (L+) *Shigella dysenteriae* (L+ SD, LMP606), ampicillin-resistant *Klebsiella pneumoniae* (ARKP, LMP803), carbenicillin-resistant *Pseudomonas aeruginosa* (CRPA, LMP804), chloramphenicol resistant *Salmonella typhi* (CRST, LMP706), and chloramphenicol resistant *Citrobacter freundii* (CRCF, LMP802). The two pathogenic fungi used were *Candida albicans* (*C. albicans*, LMP709U) and *Microsporium audouinii* (*M. audouinii*, LMP725D). Compound **203** had significant activity against all of the tested organisms, methicillin-resistant *S. aureus* being the most sensitive pathogen. The minimal inhibitory concentration (MIC) values for **203** varied from 28.7 μM (against *S. aureus*) to 230.9 μM (vs. *K. pneumoniae* and *P. aeruginosa*). In addition, the compound **203** exhibited a minimum microbicidal concentration (MMC) value against the listed bacteria and fungi that was four times lower than its corresponding MIC value. In 2016, a comprehensive review conducted by Spiteller and co-workers also concluded that the compound **203** possessed moderate antimicrobial activity against a panel of drug-resistant pathogens, particularly with an MIC of 29 μM against *Staphylococcus aureus*.^{3c}

In 2015, two novel phenolic glycosides, Sargentodognans F–G (**205–206**, Scheme 48), were extracted and isolated from a 60% ethanol extract of *Sargentodoxa cuneata*, and their structures and absolute configurations were determined by Tang's group.⁸⁵ Antimicrobial tests were performed against *S. aureus* ATCC 29213, *S. aureus* ATCC 25923, *A. baumannii* ATCC 19606, and *C. albicans* ATCC 10231, respectively. Among them, compounds **205–206** showed antibacterial activity against *S. aureus* ATCC 29213.



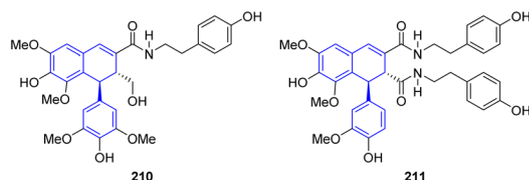


Scheme 49 Representative antiviral agents containing ADHNs.

3.2.3 Antiviral activity. Most viruses can be transmitted outside the body, where they easily attach to the surface and remain adherent, causing infection in humans. It is important to note that the emergence of drug-resistant viruses poses a major challenge to the effectiveness of treatments.⁸⁶ As a result, the design and synthesis of novel and targeted antiviral compounds is an effective strategy to reduce the risk to public health.

A review on the antiviral activities of lignans⁸⁷ prompted the Charlton group to explore the antiviral activities of several 1-arylnaphthalenes and 1-aryl-1,2-dihydronaphthalenes lignans and their analogues. In 2000, the compounds **207–209** were tested by Charlton's group using a standard plaque reduction assay to evaluate their ability to inhibit human cytomegalovirus (Scheme 49).^{4a} The cytotoxicity of the compounds were also measured using a tetrazolium salt MTT. The experimental results showed that the EC₅₀s of compounds **207–209** were in low micromolar range (EC₅₀/TC₅₀: **207** 25/>37 μM; **208** 1.4/>25 μM; **209** 22/>33 μM).

3.2.4 Neuroprotective activity. Neurodegenerative diseases pose significant challenges, especially in developed nations where diets and living standards are elevated, and are becoming more common as medical advances extend natural lifespans. Providing care for afflicted individuals incurs significant expenses, posing immense challenges for patients and their loved ones. Consequently, the development of neuroprotective therapeutics represents a pivotal area of research. In 2023, four pairs of ADHN-type lignanamide enantiomers were isolated from *Solanum lyratum* (Solanaceae) by Song, Bai and co-workers,⁸⁸ the *in vitro* neuroprotective effects of all compounds were evaluated using H₂O₂-induced human neuroblastoma SH-SY5Y cells and AchE inhibition activity. The experimental results showed that one of the enantiomers, the compound **210** (Scheme 50) had a remarkable neuroprotective effect at high concentrations of 25 and 50 μmol L⁻¹ comparable to Trolox. Another enantiomer, compound **211** (Scheme 50), showed the strongest inhibitory effect on AchE with an IC₅₀ value of 3.06 ± 2.40 μmol L⁻¹. The overall spatial structure suggested that the



Scheme 50 Representative neuroprotective agents containing ADHNs.

complex of AchE and the compound **211** was stable, indicating that the interaction between **211** and its targets may play an important role in the inhibition activity of AchE.

4. Conclusions

In summary, we have presented an overview of the research progress on ADHNs derivatives over the past two decades. Generally, the types of intramolecular and intermolecular synthesis methods for ADHNs derivatives, their reaction characteristics, applications in the total synthesis of natural products and bioactive molecules, and their biological applications are systematically introduced in this review.

Notably, the intramolecular cyclization reaction of styrenynes has gained substantial attention and made much progress in recent years owing to its atom-economy, mild reaction conditions, extensive functional group tolerance and good yields. However, some substrates for the intramolecular cyclization reactions require multiple preparation steps, which affects the convenience of the synthesis process and reduces the overall yields of the synthetic routes. Despite some alternative intermolecular strategies have been established, most of them tend to provide access to the desired products, with few comprehensive methods to achieve diversified constructions. Moreover, the chemoselectivity in obtaining ADHNs skeletons exclusively remains elusive and challenging to pinpoint, leading to increased difficulty in purification and decreased yields.

Although some biological activities such as antitumor, antimicrobial, antiviral and neuroprotective activities have been well documented by researchers, offering a reference for future related biological study and drug exploitation, the scarcity of diverse ADHNs derivatives hinders pharmaceutical chemists from exploring their structure–activity relationships and impedes the development of their biological applications and potential therapeutic values. In the next stage, more attention should be paid to the exploration of new, atom- and step-economical, chemically selective, and environmentally benign synthetic methodologies to increase the diversity of ADHNs derivatives, laying a foundation for further exploration of the organic synthesis and synthetic and biological applications of ADHNs in the future.

Data availability

All data included in this study are available upon request by contact with the corresponding author.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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