

Cite this: *Chem. Sci.*, 2025, 16, 2961

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 26th November 2024

Accepted 24th January 2025

DOI: 10.1039/d4sc08017c

rsc.li/chemical-science

Unifying principles for the design and evaluation of natural product-inspired compound collections†

Frederik Simonsen Bro  and Luca Laraia *

Natural products play a major role in the discovery of novel bioactive compounds. In this regard, the synthesis of natural product-inspired and -derived analogues is an active field that is further developing. Several strategies and principles for the design of such compounds have been developed to streamline their access and synthesis. This perspective describes how individual strategies or their elements can be combined depending on the project goal. Illustrative examples are shown that demonstrate the blurred lines between approaches and how they can work in concert to discover new biologically active molecules. Lastly, a general set of guidelines for choosing an appropriate strategy combination for the specific purpose is presented.

Introduction

Natural products (NPs) are an important source of bioactive small molecules. They have co-evolved with their biosynthetic proteins, thus exploring biologically relevant chemical space and encoding inherent biological relevance, as a result of their ability to bind biomolecules and cross cell membranes. In many, though not all, cases they have also evolved to be stable,

at least for the duration of their intended bioactivity. Consequently, NPs were the first examples of therapeutics. NPs, their derivatives, and compounds inspired by them are and have been the foundation of organic and medicinal chemistry and play a major role in drug discovery.^{1–8} Importantly, one third of approved drugs since 1981 fall into one of these categories, highlighting the historical and continuing impact of NPs in this area.⁹

Despite the obvious benefits of NPs, there are limitations in terms of drug discovery. Accessing natural products by isolation or total synthesis (TS) can sometimes be laborious and involve inefficient processes, while often not delivering enough material for biological evaluation and structure–activity relationship

Department of Chemistry, Technical University of Denmark, 2800, Kongens Lyngby, Denmark. E-mail: luclar@kemi.dtu.dk

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4sc08017c>



Frederik Simonsen Bro

Frederik Simonsen Bro achieved his BSc in chemistry at the Technical University of Denmark (DTU) in 2019 which included a bachelor's project with Prof. Robert Madsen working on iron-catalysed dehydrogenation of alcohols. In 2021 he completed his MSc in chemistry at DTU carrying out his master's project with Assoc. Prof. Luca Laraia in the synthesis of an alkaloid-inspired library. Subsequently, he carried out his PhD studies in

the same lab focussing on the development of inhibitors of sterol transport proteins through synthesis of sterol-inspired libraries, which he completed in 2024, and continued as a postdoctoral researcher in the same group.



Luca Laraia

Luca Laraia received his MSci at Imperial College London (2009) before completing a PhD in chemical biology at the University of Cambridge (2014). He subsequently moved to the Max Planck Institute of Molecular Physiology as an Alexander von Humboldt fellow and subsequently project leader, before embarking on his independent career at the Technical University of Denmark as an assistant professor. Since 2021 he has

been an associate professor for chemical biology and medicinal chemistry. His group's work lies at the interface of chemistry and biology, including the synthesis of natural product-inspired compounds and the study and modulation of sterol-mediated processes.



(SAR) analysis. The reasons for this are low concentration and extraction from complex mixtures when isolating NPs from natural sources, and (often) multi-step and low yielding total syntheses of NPs. Lastly, the selection criteria for nature are different from the selection criteria in drug discovery. As such, NPs have evolved for their producing organisms, not human therapeutic applications.^{5,7} Though NPs cover biologically relevant chemical space, their restricted natural selection means that they only cover a limited fraction of NP-like chemical space.¹⁰ Thus, the vast majority of biologically relevant, NP-like chemical space remains to be explored. In fact, investigation of the surrounding chemical space of an NP can be more beneficial than investigating the NP alone in terms of drug discovery.² Consequently, strategies to synthesise NP-derived and -inspired compounds in a practical and efficient way are in demand to navigate new NP-like chemical space and obtain highly bioactive compounds that can serve as drug candidates or tool compounds.

To meet the demand, several strategies to synthesise compounds derived from or inspired by NPs have emerged. Diversity-oriented synthesis (DOS)^{11,12} focusses on characteristics typical of NPs, including a high fraction of sp³-hybridised carbons (Fsp³) and several stereogenic centres, but is not necessarily based on an NP or NP scaffold. The similar privileged-substructure-based DOS (pDOS)^{13,14} is based on a privileged scaffold with proven biological relevance that is not necessarily derived from an NP. For DOS and pDOS, molecular

scaffold diversity is a key point, which is also the case for activity-directed synthesis (ADS).^{15,16} The compounds resulting from the strategies including pseudo-natural product (PNP) synthesis,^{17,18} biology-oriented synthesis (BIOS),^{19,20} function-oriented synthesis (FOS),²¹ and pharmacophore-directed retrosynthesis (PDR)²² are all based on NP fragments, scaffolds, or pharmacophores. The total synthesis (TS) of NPs is guided by target molecules (TMs). However, the focus on a single TM limits the exploration of chemical space. This has been a driving force for the establishment of synthetic approaches that investigate the chemical space surrounding a guiding NP, which include complexity-to-diversity (CtD),²³ dynamic retrosynthetic analysis (DRA),^{2,24} diverted total synthesis (DTS),^{25,26} two-phase synthesis (TPS),²⁷ and analogue-oriented synthesis (AOS).²⁸ The recently described diverse PNP (dPNP) strategy²⁹ combines PNP and DOS/CtD, and thus originates from NP fragments (see Fig. S1–S13† for graphical illustrations, explanations and examples of the individual strategies).

To navigate the plethora of approaches outlined above, we have found it helpful to separate them based on the qualitative similarity of the core frameworks generated to those found in NPs,³⁰ as highlighted by some representative examples (Fig. 1).^{22,31–46} It is important to note that several quantitative computational approaches for assessing NP-likeness have been developed. For example, the NP-score uses the prevalence of specific atom-centred fragments to compare NPs to fully synthetic compounds.⁴⁷ The NP character of compound

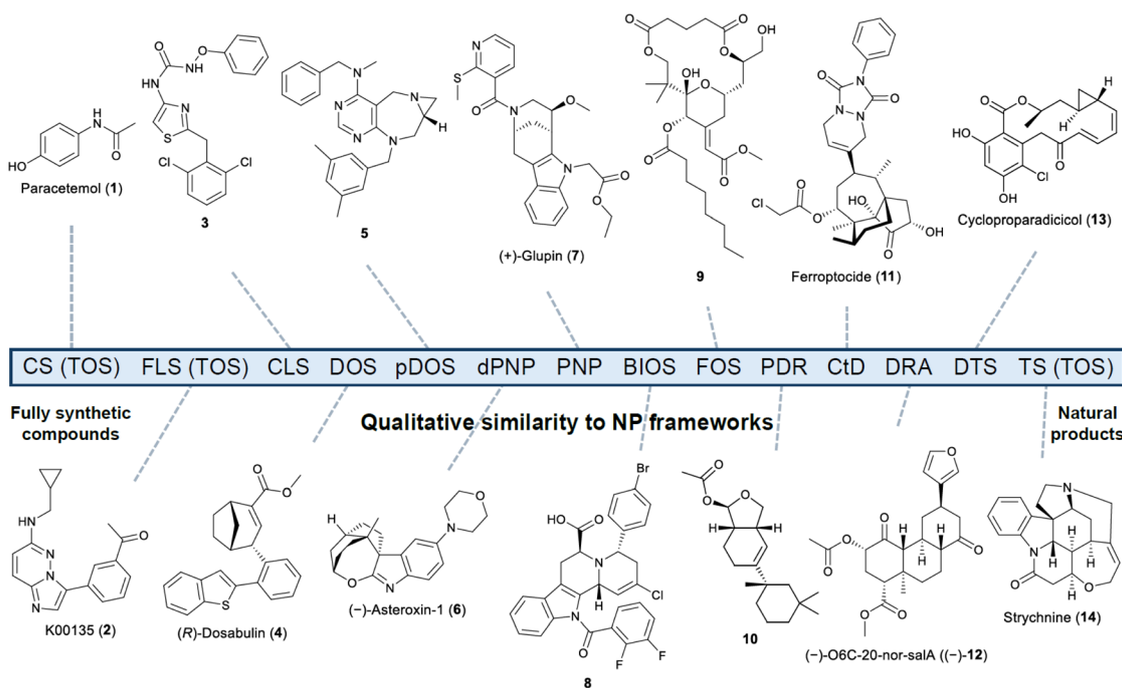


Fig. 1 Continuum of qualitative similarity to NP frameworks of compounds designed *via* the different strategies with representative examples: conventional synthesis (CS): paracetamol (1); focussed library synthesis (FLS): K00135 (2);³¹ combinatorial library synthesis (CLS): 3;³² diversity-oriented synthesis (DOS): (*R*)-dosabulin (4);³³ privileged-substructure-based diversity-oriented synthesis (pDOS): 5;³⁴ diverse pseudo-natural product (dPNP): (-)-asteroxin-1 (6);³⁵ pseudo-natural product (PNP): (+)-glupin (7);³⁶ biology-oriented synthesis (BIOS): 8;^{37,38} function-oriented synthesis (FOS): 9;³⁹ pharmacophore-directed retrosynthesis (PDR): 10;²² complexity-to-diversity (CtD): ferroptocide (11);⁴⁰ dynamic retrosynthetic analysis (DRA): (-)-O6C-20-nor-salA ((-)-12);^{45,46} diverted total synthesis (DTS): cycloproparadicol (13);^{41,42} total synthesis (TS): strychnine (14).^{43,44} TOS = target-oriented synthesis.



collections is a continuum between fully synthetic compounds (no guiding NP) and unmodified NPs. Guided by retrosynthetic analysis, fully synthetic drugs and libraries of drug candidates can be synthesised by conventional synthesis (CS) and focussed or targeted library synthesis (FLS), respectively, and NPs are synthesised by TS.^{48,49} These are examples of target-oriented synthesis (TOS) which by definition lacks diversity due to the single target approach. Nonetheless, diversity might arise for FLS and TS when several targets are synthesised in the same study. This is particularly evident for divergent approaches in TS where a common intermediate is used in the synthesis of several members of an NP compound class.⁵⁰ When a single compound of interest is not known, combinatorial library synthesis (CLS) provides quick access to many compounds. However, although combinatorial libraries can afford complexity giving the compounds slightly higher NP-character, structural diversity is often still limited.⁵¹ Here it should be noted that complexity alone does not guarantee bioactivity,⁵² nor is it always a clear predictor of properties that would be of interest to medicinal chemists, including solubility and oral bioavailability.⁵³ Increased complexity has been correlated with increased selectivity,⁵⁴ however, increased molecular weight and lipophilicity both correlate with increased promiscuity.^{55,56} As such, a careful balance of parameters should be targeted when employing such metrics in library design strategies. Overall, many ways to calculate complexity have been reported and we refer the reader to recent publications for a more detailed discussion and their application.^{52,53,57,58}

The increased focus on diversity and privileged scaffolds in the DOS and pDOS strategies brings the resulting compounds closer to NPs than combinatorial approaches. However, there is no strict requirement for the compounds to be NP-like, and design considerations are often governed by the synthetic accessibility of the resulting products. The privileged scaffolds incorporated can also be fully synthetic in origin, providing key differences to strategies delivering compounds with a higher qualitative similarity to NPs. The PNP strategy is based on the recombination of NP fragments, and as such the resulting scaffolds are not typically found in NPs, even though their constituent fragments are. This gives them a lower NP score, according to the definition of Ertl *et al.*⁴⁷ BIOS is for the most part based on actual NP scaffolds, thus bringing the resulting analogues closer to NPs compared to DOS and PNP. Since dPNPs are the result of different combinations of PNP and diversification strategies, they are not necessarily NP scaffolds and may thus be less likely to have frameworks found in NPs than both PNP and BIOS-derived compounds. The FOS, PDR, CtD, DRA, DTS, TPS, and AOS strategies can give compounds that are very close to, or some distance from, actual NPs. For example, a key difference of CtD to other strategies is the frequent use of ring distortion reactions, which can sometimes steer compounds far away from NPs in chemical space. While being beneficial in terms of targeting unexplored chemical space, it carries an inherently greater degree of uncertainty as to the utility of the resulting compounds in biological screens. It is difficult to say which strategy provides compounds with greater NP character, as the resemblance to the parent/guiding NP

varies from case to case, thus placing them somewhere in between BIOS and TS.

It should be noted that the definitions of most strategies are open to a degree of interpretation and often overlap, making the distinction between standalone strategies and umbrella terms difficult. For example, CtD could be considered a subset of DOS or its own strategy. Additionally, while TPS is rooted in TS, in principle it can be applied to diversify natural product skeletons/scaffolds. Lastly, DOS and FOS can be grouped together with TOS as general umbrella terms.⁵⁹ In principle, any strategy aiming for structural diversity in an efficient manner could be described as DOS,^{11,12} while any strategy that delivers compounds which recapitulate or even enhance the activity of a natural product through a simplified scaffold can be described as FOS.^{21,60,61} Despite this, principles for the design of DOS and FOS libraries applied as standalone strategies have been described, and representative examples are included in Fig. 1. We view this flexibility as an advantage in enabling chemists to make bolder choices in their library syntheses.

Several excellent reviews and perspectives have been published on the design of NP-inspired compound collections using specific strategies and approaches.^{2,4,5,7,18,24,62-67} These showcase recent findings, examples, and thoughts in the field relating to individual approaches. In this perspective, our goal is to identify unifying principles across a range of library synthesis approaches. We will highlight such principles with appropriate case studies and make the argument that the existing strategies for NP-inspired compound collections are not necessarily mutually exclusive, but rather complementary, with significant benefits existing from a more open approach by combining strategies or elements from them according to the project goal compared to the use of individual strategies in isolation. The choice of strategies for prospective projects will vary based on whether one seeks to develop new chemistry to increase the chemical and biological diversity of a screening collection, identify entirely new chemical matter for a target/phenotype, or improve potency or other properties for a ligand of a known target or phenotype. Therefore, we will also develop guidelines for assessing which combination of design strategies is most beneficial based on the project goal.

Current approaches: different or complementary?

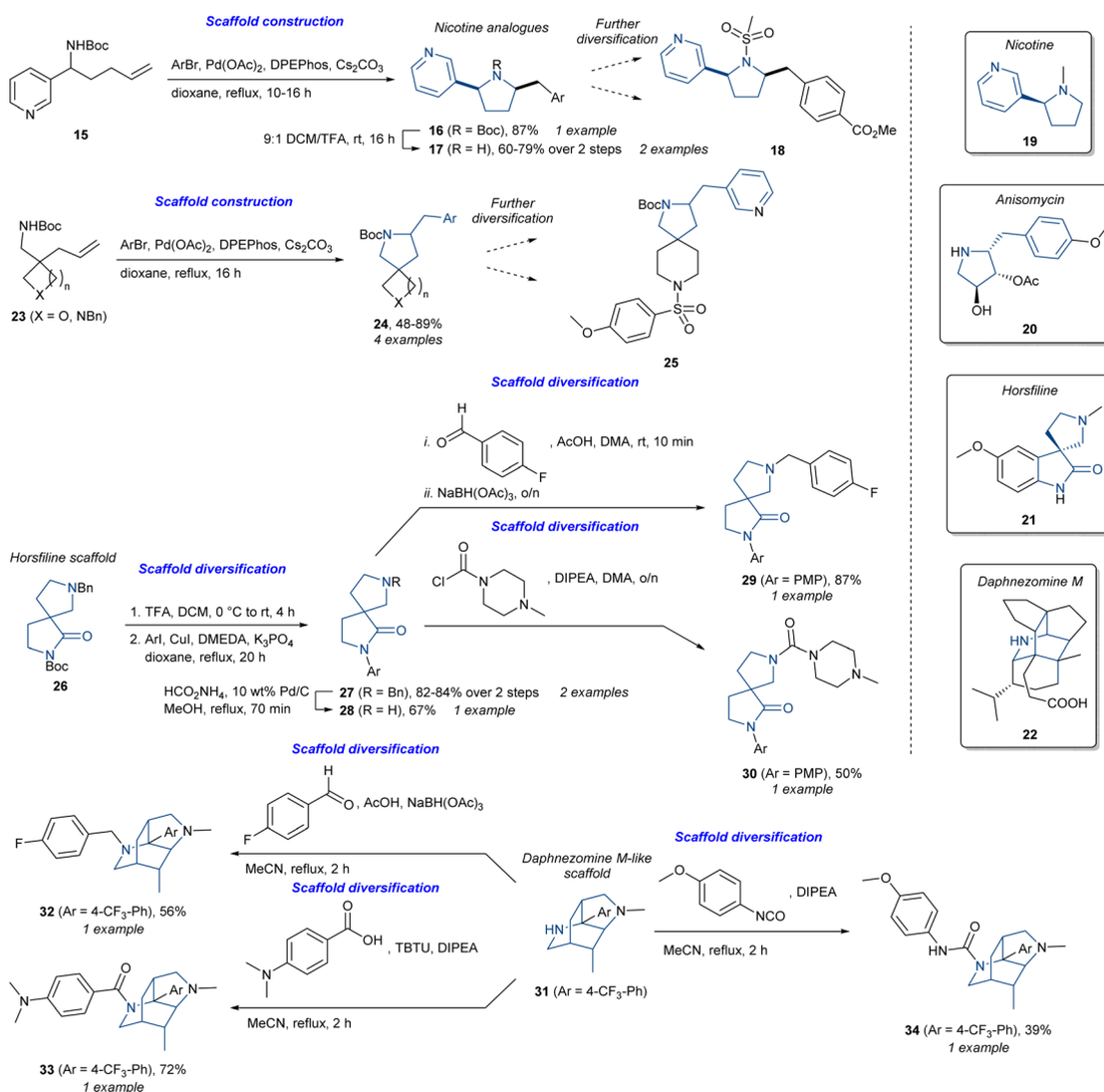
The different approaches and strategies outlined so far have developed as a consequence of different project goals and information available at the project outset. Key considerations include the availability of target or phenotype information, as well as the availability of known ligands, and particularly NPs, as starting points for design. However, the common denominator for all the approaches and strategies is the use of NPs themselves or their characteristics to develop and identify bioactive molecules, whether this is in a targeted or completely unbiased approach. The need for well-defined strategies and the differentiation between them provides theoretical frameworks that simplify and structure a project, ideally allowing fast



access to desired compounds. Differentiation between strategies should thus make it easier to find the right approach for a specific project goal. However, we have found that in many cases the approach ultimately used by research groups implicitly combines components from several different strategies under a larger “umbrella” approach, even though the initial strategy was presented as a single defined approach. More recently, research groups including our own have explicitly targeted the combination of strategies for specific applications.

To highlight how various approaches work well together and have more similarities than differences, we have chosen illustrative examples based on different combinations of strategies and the outcomes they present. We have chosen to structure the initial discussion based on chemical strategies, rather than biological outcome, with the latter being addressed in a subsequent section (*vide infra*). In this regard, the synthesis of six alkaloid-inspired libraries (Scheme 1)⁶⁸ highlights the use of

multiple strategies. These include the synthesis of nicotine (19) analogues (16 and 17) and spirocyclic analogues (24) containing a benzylic-substituted pyrrolidine as found in anisomycin (20) using a complexity-generating Pd-catalysed aminoarylation reaction starting from 15 or 23. Following *tert*-butyloxycarbonyl (Boc) deprotection of 16, compounds 17 could undergo different diversification reactions to access additional analogues including sulfonamide 18. In a similar fashion, 24 could also be diversified into additional analogues such as sulfonamide 25. In terms of synthetic design, this can be classified as substrate-based DOS. However, the simplified nicotine analogues could be considered as BIOS analogues and the pyrrolidine analogues as PNP (anisomycin fragment) or pDOS (pyrrolidine as a privileged scaffold).⁶⁹ The analogues could be further diversified using traditional diversification methodologies. Similarly, the authors also diversified the horsfiline (21) scaffold 26 using Ullmann–Goldberg cross-coupling to give 27.



Scheme 1 Synthesis of alkaloid-inspired compounds.⁶⁸ DIPEA = *N,N*-diisopropylethylamine, DMEDA = *N,N'*-dimethylethylenediamine, DPE-Phos = bis[(2-diphenylphosphino)phenyl]ether, PMP = *p*-methoxyphenyl, TBTU = *N,N,N',N'*-tetramethyl-*O*-(benzotriazol-1-yl)uronium tetrafluoroborate.

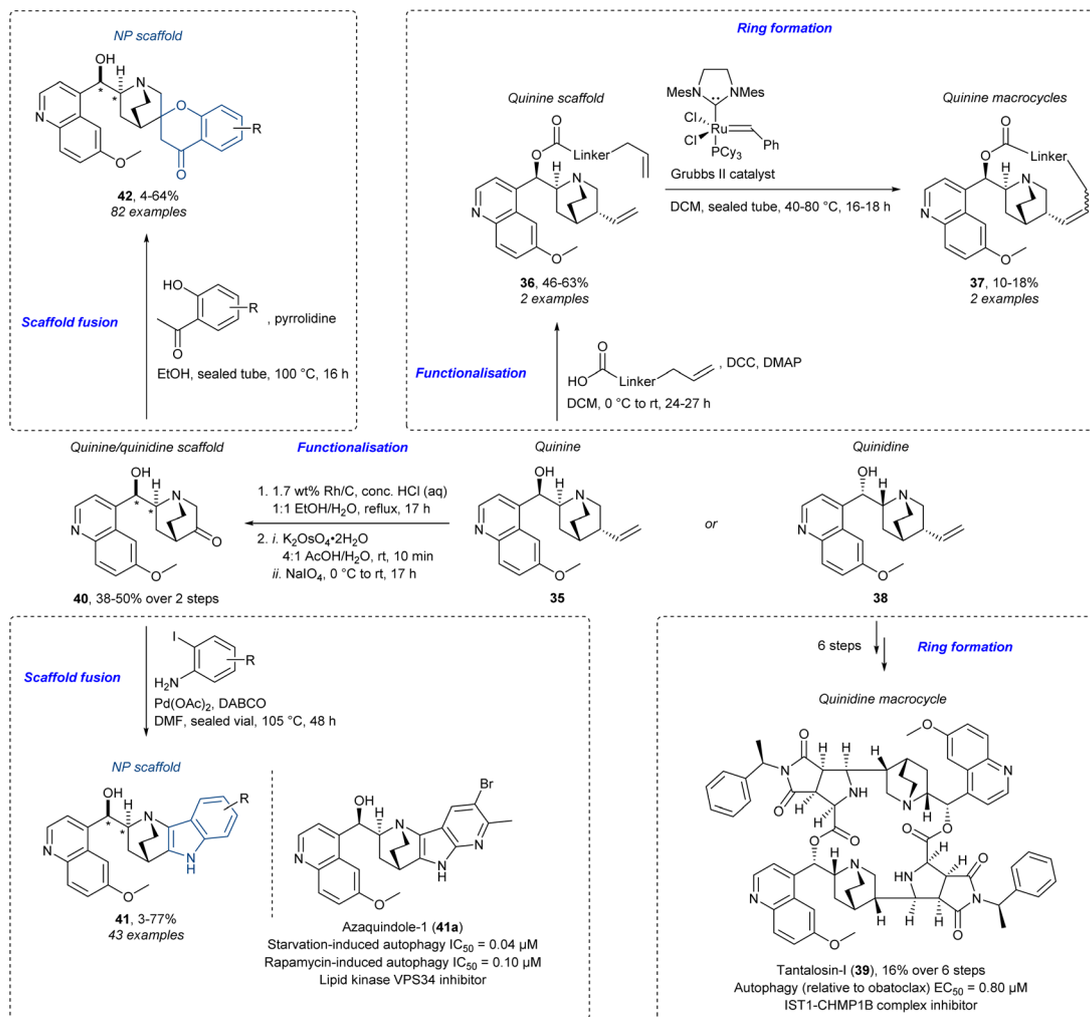


Following benzyl deprotection, **28** could then undergo either reductive amination affording **29** or urea formation to give carbamide (**30**). This shares ideas from PNP and BIOS by using the horseshine scaffold as a starting point for analogue synthesis. Furthermore, applying different reagents from **28** to generate different analogues could be classed as reagent-based DOS. The last example using the daphnezomine **M** (**22**)-like scaffold **31** also used the reagent-based DOS line of thought. Reductive amination afforded **32**, amide coupling afforded **33**, and the reaction with isocyanate afforded carbamide **34**. In total, six different libraries with different scaffolds were synthesised. Overall, this elegantly demonstrates substrate- and reagent-based DOS with the incorporation of ideas from BIOS and PNP using alkaloid scaffolds to enhance biological relevance.

Similar published work also shares the idea of a complexity-generating reaction followed by diversity focused reactions in a reagent-based DOS fashion.^{54,55} Focus on diversity is also a key point in activity-directed synthesis, where chemical space is explored by using reactions with multiple and diverse possible outcomes.^{15,16} The overall idea of generating diversity from

a single substrate is embedded in CtD as well, where NPs with (preferably many) diversification vectors are used to create NP-derived analogues using ring distortion, ring formation, and diversification reactions. CtD limits itself to NPs that are not the end point, but rather are complex starting points that can be diversified.²³ Thus, CtD could be seen as an example of reagent-based DOS on NPs, where simply by modifying the reagents one can access significant scaffold diversity. The idea of creating diversity from a complex starting material is thus found in both CtD and in work not related to an actual NP that could consequently be viewed as “CtD on non-NPs”. For example, pDOS has been combined with CtD by using ring distortion and ring formation to form diverse medium/macro- and bridged heterocyclic compounds containing the privileged scaffold pyrimidine.⁷⁰

Employing an NP as a starting point for complexity works well conceptually with other strategies. For example, fragment-sized NPs like the cinchona alkaloids quinine and quinidine can work well in DOS, BIOS, or PNP campaigns.⁷¹ Significant effort has been made to make quinine-derived analogues



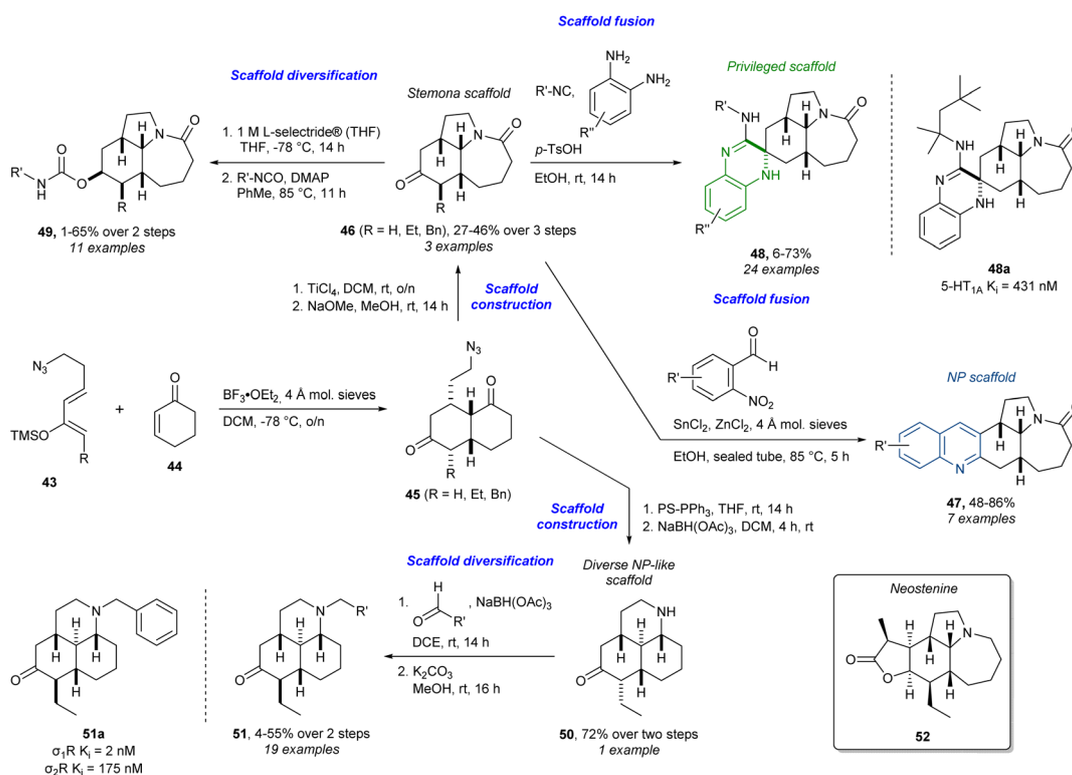
Scheme 2 Synthesis of quinine- and quinidine-inspired compounds^{73,80} and identification of the autophagy inducer, tantalosin-1 (**39**),^{77,78} and autophagy inhibitor, azaquindole-1 (**41a**).⁷⁹ DABCO = 1,4-diazabicyclo[2.2.2]octane. DCC = *N,N'*-dicyclohexylcarbodiimide, DMAP = 4-dimethylaminopyridine, Mes = mesityl.



(“quinalogs”⁷²) using different strategies (Scheme 2). Diverse and complex macrocycles can be accessed directly from quinine using a combination of CtD and DOS strategies,⁷³ which are both useful strategies in the synthesis of diverse macrocycles.⁷⁴ Initially, quinine (35) is functionalised with terminal alkene-containing linkers *via* Steglich esterification to give esters 36 which were then cyclised to form the quinine macrocycles 37 using ring-closing metathesis (RCM). Four other macrocycles were also accessed from other quinine-derived building blocks. This approach has a strong resemblance to the build/couple/pair (B/C/P) strategy: “building” quinine building blocks, intermolecular “coupling” with other functionalised building blocks, and intramolecular “pairing” of the functional groups to form the macrocycles. Macrocycles remain a desired moiety since they have proven to be a privileged class of molecules for modulating challenging targets such as protein–protein interactions in drug discovery.^{75,76} In this context, a complex quinine (38)-inspired 20-membered macrocycle, tantalosin-I (39), that induced autophagy by induction of microtubule-associated protein 1A/1B light chain 3 (LC3) lipidation through disruption of a particular part of the endosomal sorting complexes required for transport (ESCRT) called the IST1-CHMP1B complex was recently reported.^{77,78} Additional work showed how you could use the relatively small NPs quinine and quinine in a PNP setting.^{79,80} The NPs were initially transformed to the corresponding ketones 40 in two steps by Rh-catalysed isomerisation of the terminal alkene to the internal alkene followed by a Malaprade–Lemieux–Johnson oxidation. Ketones

are a strategic functional group in the synthesis of NP-inspired compounds since they serve as a suitable coupling partner for fusion with other scaffolds.⁸¹ They reported the synthesis of edge-fused indoles and azaindoles 41 from quinine and quinine *via* a one-pot imine condensation and a Hegedus–Mori–Heck reaction. This led to the identification of an autophagy and lipid kinase VPS34 inhibitor, azaquindole-1 (41a). Moreover, the spiro-fused chromanones 42 were accessed through a Kabbe reaction. Thus, different PNPs could be accessed simply by changing the reagents, the principle of reagent-based DOS. In this work there was no specific biological target in mind and the compounds were screened phenotypically. Importantly, tantalosin-I (39) and azaquindole-1 (41a) show different bioactivity to each other and to the parent NP, highlighting the value of diversifying a relatively large building block, the cinchona alkaloid framework. All the analogues in Scheme 2 come from a synthetically easily accessible quinine- or quinine-scaffold, thus having a resemblance to BIOS-derived compounds. This shows how the different strategies can overlap in a beneficial way. Another great example of this is the work on indotropans.⁸² It can be seen as a BIOS library^{20,83} but also shares ideas from the PNP strategy.⁸⁴ Whether you define it as one or the other, the final outcome was the identification of a novel class of hedgehog-signalling inhibitors and the myosin light chain kinase 1 (MLCK1) inhibitor, myokinasib.

Similarly, in addition to the abovementioned cinchona alkaloids, stemona-inspired compounds have also been targeted using diverse strategies (Scheme 3).^{85–87} This work is



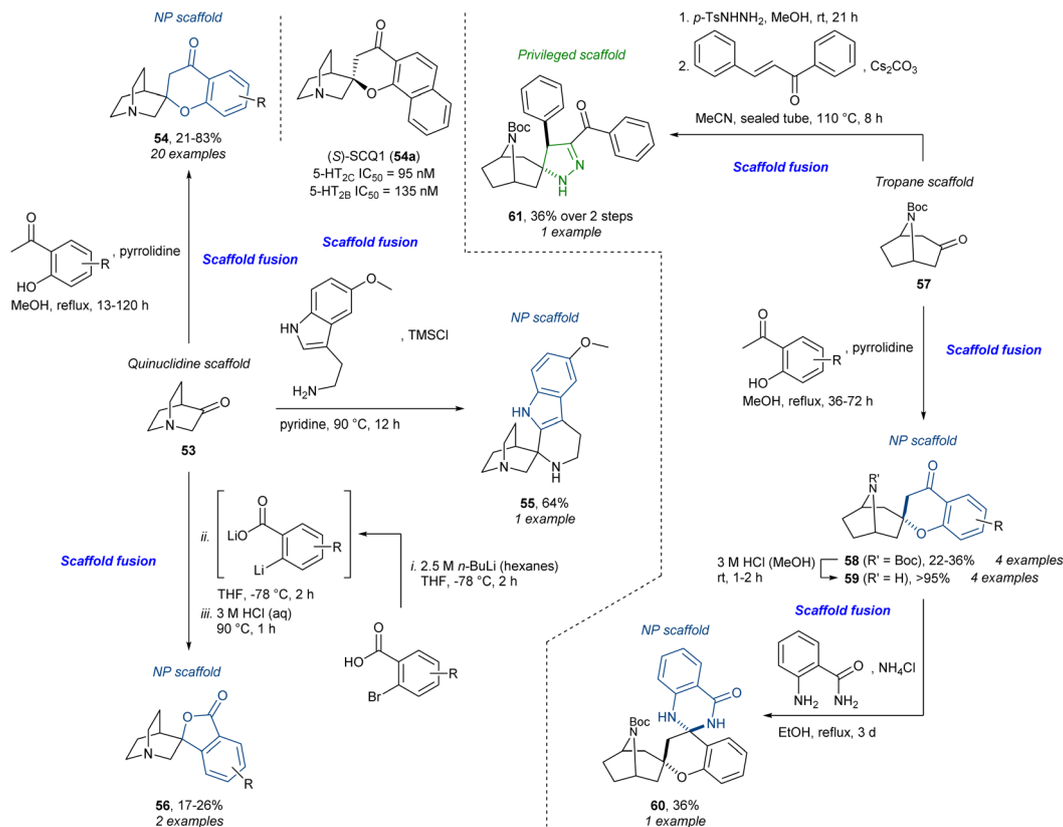
Scheme 3 Synthesis of stemona-inspired compounds and identification of 2-HT_{1A} ligand 48a and σ_1 R and σ_2 R ligand 51a.^{85–87} L-selectride® = lithium tri-*sec*-butylborohydride, PS-PPh₃ = polystyrene bound triphenyl phosphine.



particularly interesting as it can be considered an early example of the PNP strategy before it was formally defined, with elements from BIOS and (p)DOS. Here, the core scaffolds containing a strategic ketone are synthesised from scratch and not derived from an NP as above. Firstly, a Lewis acid-catalysed Diels–Alder reaction between diene **43** and dienophile **44** afforded the bicyclic azido diketones **45** which could undergo a Schmidt reaction by further treatment with a Lewis acid. Careful choice of Lewis acid and its equivalents allowed for the tandem Diels–Alder/Schmidt reaction to go all the way to the tricycle. Epimerisation with base of the alkyl group of the diastereoisomeric mixture of α -alkylated tricyclic ketones afforded the thermodynamically favoured β -alkyl stemona scaffold **46**. This could then be subjected to a reductive Friedländer quinoline synthesis to afford edge-fused quinoline-analogues **47** in a PNP-fashion. By definition the PNP approach only allows the fusion of NP fragments. However, fusion with privileged scaffolds not found in NPs, as implied in the pDOS approach, to access a large number of scaffold combinations, can enable the identification of compounds that modulate diverse targets and processes in a selective way. Even though this work predates the PNP concept, this line of thought can be seen in the access of the spiro-fused 3,4-dihydroquinoxalines **48**, a privileged scaffold, *via* a multicomponent reaction (MCR) with *o*-phenylenediamines, the ketone **46**, and isocyanides.⁸⁸ The 3,4-dihydroquinoxaline **48a** was found to have a binding affinity (inhibitory constant (K_i)) of 431 nM to the 5-hydroxytryptamine

(serotonin) 1A receptor (5-HT_{1A}). Furthermore, subjecting PNPs to general synthetic diversification strategies as in DOS can give access to even more diverse and biologically relevant compounds. This idea is visible in the synthesis of the carbamates **49**, which is an important structural motif in medicinal chemistry.⁸⁹ The ketone **46** was diastereoselectively reduced with L-selectride® to give an alcohol which was then reacted with isocyanates to give the carbamates **49**. Other analogues were accessed from another tricyclic ketone **50**, synthesised from **46** *via* an aza-Wittig reaction. The tricyclic ketone scaffold **50** is not found in NPs, but it is NP-like in terms of complexity. From this ketone, reductive amination yielded the tertiary amines **51** where analogue **51a** showed a K_i of 2 nM to the sigma-1 receptor (σ_1R) and 175 nM to the sigma-2 receptor (σ_2R). Several other analogues with different functionalities were also accessed and in total the library consisted of 104 stemona analogues. Using the tricyclic core of stemona alkaloids in the synthesis shares a lot of ideas from BIOS. In addition, some similarity to reagent-based DOS is obvious since the analogues are derived from two core ketone scaffolds by changing reagents and conditions. One may even classify this example as an exhaustive DTS identifying **46** as the advanced intermediate. In fact, **46** has been used in a TS *en route* to the stemona alkaloid neostenine (**52**).⁹⁰

Our own work on a tropane- and quinuclidine alkaloid-inspired compound collection⁹¹ is conceptually similar to the above. Using the commercially available quinuclidine scaffold



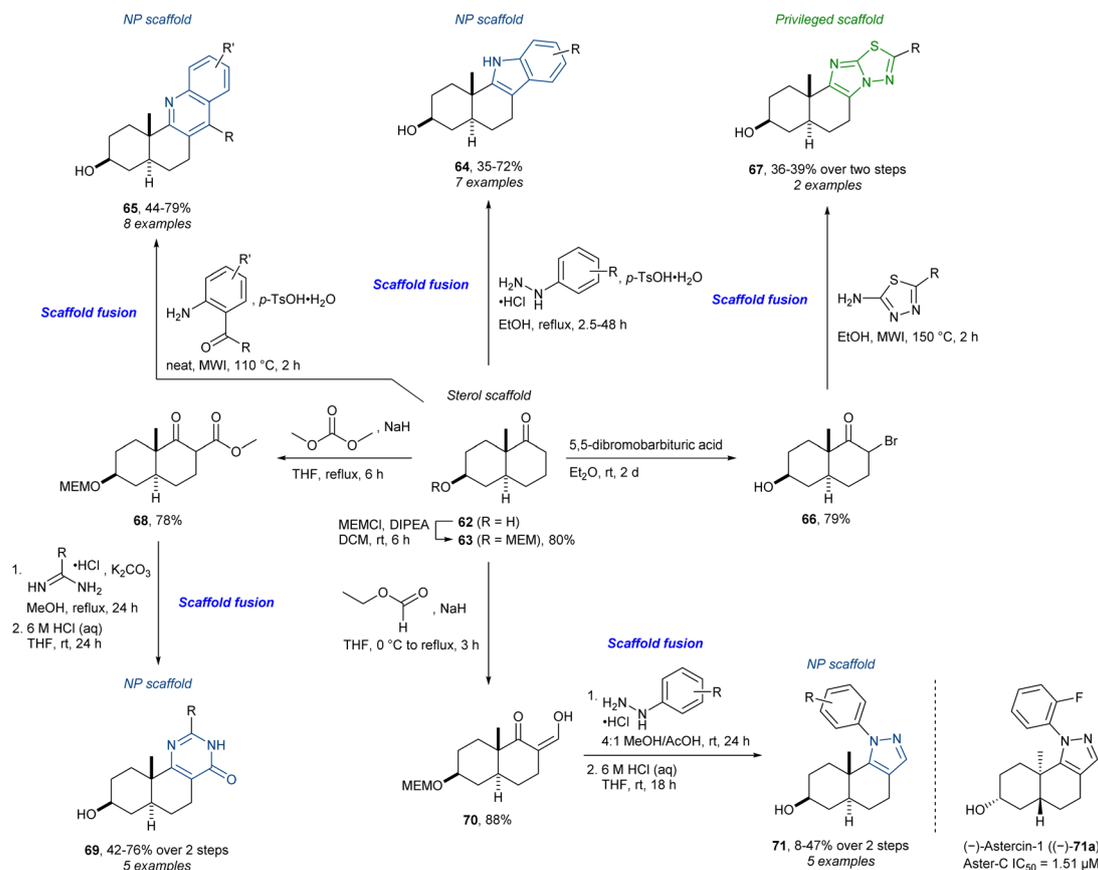
Scheme 4 Synthesis of tropane- and quinuclidine alkaloid-inspired compounds and identification of dual 2-HT_{2B/C} antagonist (S)-SCQ1 (**54a**).⁹¹ TMS = trimethylsilyl.



53 and tropane scaffold 57 several scaffold fusions could be carried out, again taking advantage of a reactive ketone (Scheme 4). Using the Kabbe reaction, the spirochromanones 54 and 58 could be accessed. The spirochromanone-quinuclidine analogue 54a was identified as a selective dual serotonin 2B (5-HT_{2B}) and 2C receptor (5-HT_{2C}) antagonist which was termed (S)-SCQ1. Chromanone-tropane 58 could be further diversified condensing with 2-aminobenzamide to give spirocyclic 2,3-dihydroquinazolinone 60. From the quinuclidine ketone 53, the Pictet-Spengler reaction afforded the spiro-fused tetrahydro-β-carboline (tryptoline) 55. Additionally, the spirophthalides 56 were accessed from 2-bromobenzoic acid in a three-step sequence going through a lithium-halogen exchange, nucleophilic attack, and intramolecular Fischer esterification. Lastly, the tropane 57 could be converted to the *N*-tosylhydrazone which could participate in a 1,3-dipolar (3 + 2) cycloaddition with chalcone to give the spiro-pyrazoline 61, a privileged scaffold. The total number of analogues was 58 including six additional other scaffolds not presented here. The overall strategy was presented as a mixture of PNP and DOS (reagent-based); however, the use of privileged scaffolds and core alkaloid skeletons makes the resemblance to pDOS and BIOS striking. Together with the aforementioned stemona alkaloid library, this is a representative example of the construction of compound libraries with specific target(s) in mind. This

contrasts with the phenotypic screening approach as described in the synthesis of, for example, the “quinalogs”.

As illustrated above, the incorporation of privileged scaffolds and diversity-generating strategies ((p)DOS) into a PNP approach can be very beneficial. Our recent work⁹² is an example of the PNP strategy where privileged scaffolds are introduced to access additional scaffolds. By the fusion of a *trans*-decalin sterol scaffold with several NP scaffolds and “unnatural” privileged scaffolds, a range of sterol-inspired analogues was synthesised (Scheme 5). The ketone 62 was synthesised as a key precursor containing the *trans*-sterol scaffold. The Fischer indole synthesis afforded the indoles 64. Furthermore, the quinoline-fused analogues 65 could be synthesised following a microwave irradiation (MWI) assisted Friedländer quinoline synthesis. *Via* the α-bromoketone 66, the imidazothiadiazoles 67 were isolated. The Pinner pyrimidone synthesis yielded the analogues 69 from the β-ketoester 68. Lastly, the β-ketoaldehyde 70 was used in a Knorr pyrazole synthesis affording 71. In addition, nine other scaffolds were accessed affording 65 sterol-inspired compounds in total. The pyrazole-fused analogues led to the identification of the potent and selective Aster-C inhibitor, (–)-astercin-1 (71a). Interestingly, the active enantiomer has the “unnatural” AB-ring stereochemistry. This indicated the importance of synthesising the library as racemic mixtures from the beginning. This is important for libraries where the goal is



Scheme 5 Synthesis of sterol-inspired compounds and identification of Aster-C inhibitor (–)-astercin-1 (71a).⁹² DIPEA = *N,N*-diisopropylethylamine, MEM = 2-methoxyethoxymethyl.

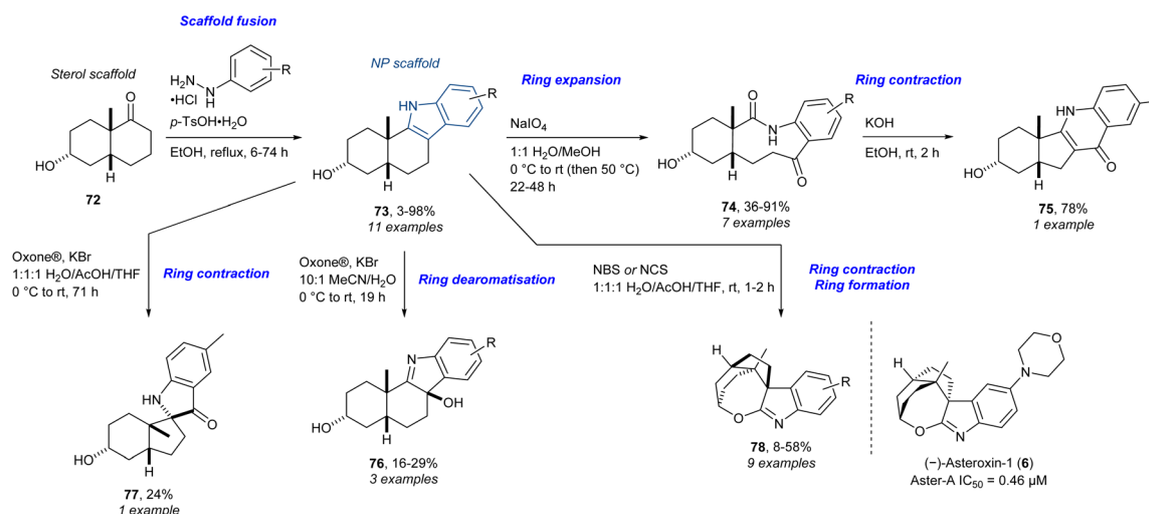


biological, as well as chemical, diversity, as this is more likely to be obtained by doubling the total number of compounds and probing enantiomeric differences. This is relevant for libraries where analogues are derived from scaffolds that are synthesised *de novo* in contrast to analogues derived from an NP source directly, since the majority of NPs are naturally produced as single enantiomers.

Following the *trans*-decalin sterol-inspired library, we decided to target a *cis*-decalin sterol-inspired compound collection,³⁵ the reasoning being that compound libraries with diverse diastereochemical attributes can result in diverse biological profiles and different biological activity.^{93,94} In addition to similar edge-fused analogues targeted in the *trans*-decalin library, additional analogues were targeted using the CtD strategy. In this context, there is no reason why the CtD strategy should be limited to NPs.^{95–97} NP fragments or NP-inspired compounds that are accessible in sufficient quantities are also excellent substrates for the ring distortion reactions used in the CtD approach. Indoles have proven to be useful scaffolds for the CtD strategy.^{98–102} Thus, we employed a ring distortion strategy on the PNPs to access dPNPs (Scheme 6). The *cis*-fused decalone **72** was synthesised and used as the primary sterol scaffold. The indoles **73** were accessed by the Fischer indole synthesis in a similar manner to the *trans*-fused library. The indoles could be ring-expanded to afford the ketolactams **74** in a Witkop oxidation. The resulting ketolactams could be ring-contracted upon treating with base yielding **75** though the Camps quinolone synthesis. Oxidative ring dearomatisation of the indoles gave the 3-hydroxyindolenines **76**. The ring contraction through an oxidative rearrangement of the indoles afforded spiro-pseudoindoxyl **77**. The spirooxepinoindoles **78** were obtained by a different oxidative ring contraction in tandem with an intramolecular ring-forming condensation. The *cis*-fused sterol-inspired library consisted of 69 compounds in total. The morpholine-substituted spirooxepinoindole (–)-asteroxin-1 (**6**) was identified as a potent and selective Aster-A inhibitor. Again,

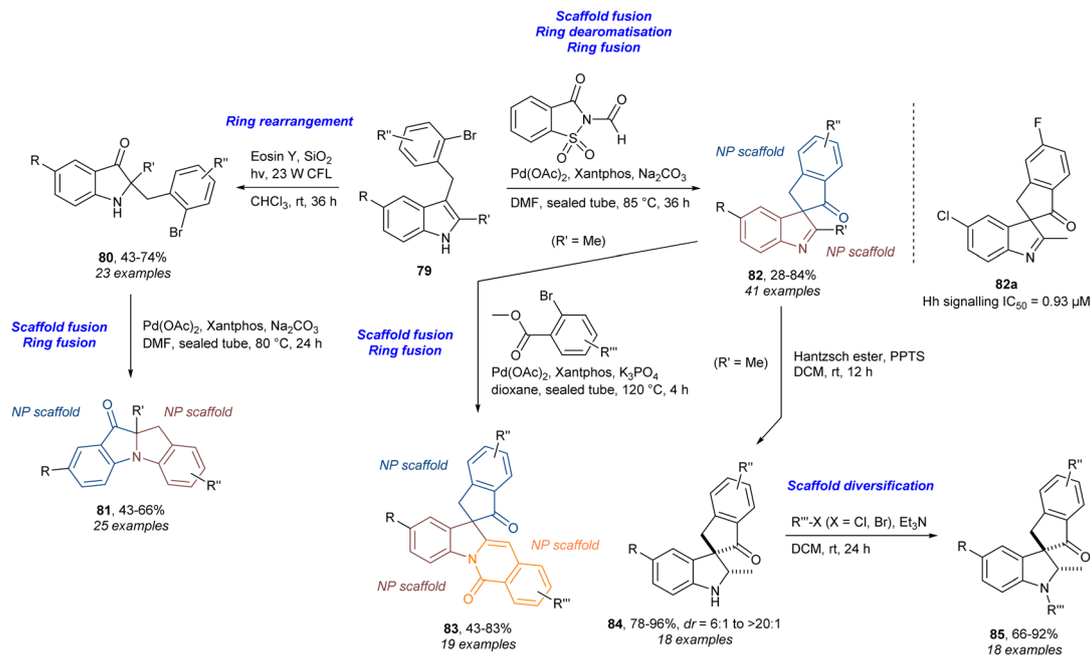
the active enantiomer featured the unnatural stereochemistry at the AB-fusion, though one can argue that the resulting scaffold scarcely resembles a steroid in structure, while retaining its bioactivity features. In this case, modifying the oxidants afforded a range of diverse scaffolds from a PNP, where one could argue that the CtD component of the library is an example of reagent-based DOS. Importantly, the work on *trans*- and *cis*-fused sterol-inspired compounds is another example of a target-based screening campaign, where biological diversity was sought within a specific class of proteins, rather than across the whole proteome.

In addition to our own work, several other research groups have also combined PNPs with diversity strategies (DOS) and ring distortion (CtD) in an explicit manner.^{29,103,104} In this context dPNPs were first defined as the combination of PNP and DOS/CtD giving compound collections that incorporate both biological relevance and scaffold diversity.²⁹ In this work (Scheme 7), the indoles **79** underwent photocatalysed ring rearrangement to give pseudoindoxyls **80**. These products could be fused to give **81** through an intramolecular Buchwald–Hartwig cross-coupling. The starting indoles **79** could also be subjected to a Pd-catalysed carbonylation/intramolecular indole dearomatisation cascade to give the ring-spiro-fused indoly-lindanones **82**. The analogue **82a** was identified as an inhibitor of Hedgehog (Hh) signalling. The analogues **82** could be further diversified by ring-edge-fusion to give indoline-indanone-isoquinolines **83** *via* a combined Pd-catalysed arylation and amidation. Additionally, reduction of the indolenine functionality in **82** afforded the spiro-indoline-indanones **84** with high diastereoselectivity. Further diversification of this class was achieved by substitution at the nitrogen using different halides to give **85**. A number of other compound classes were produced to afford a compound collection of 154 analogues in total. This work is a good example of using diversity strategies from DOS (here B/C/P in particular) to make diverse scaffolds which by design fall under the category of PNPs. Thus, the diversity is



Scheme 6 Synthesis of sterol-inspired compounds and identification of Aster-A inhibitor (–)-asteroxin-1 (**6**).³⁵ Oxone® = $\text{KHSO}_5 \cdot 0.5\text{KHSO}_4 \cdot 0.5\text{K}_2\text{SO}_4$, NBS = *N*-bromosuccinimide, NCS = *N*-chlorosuccinimide.





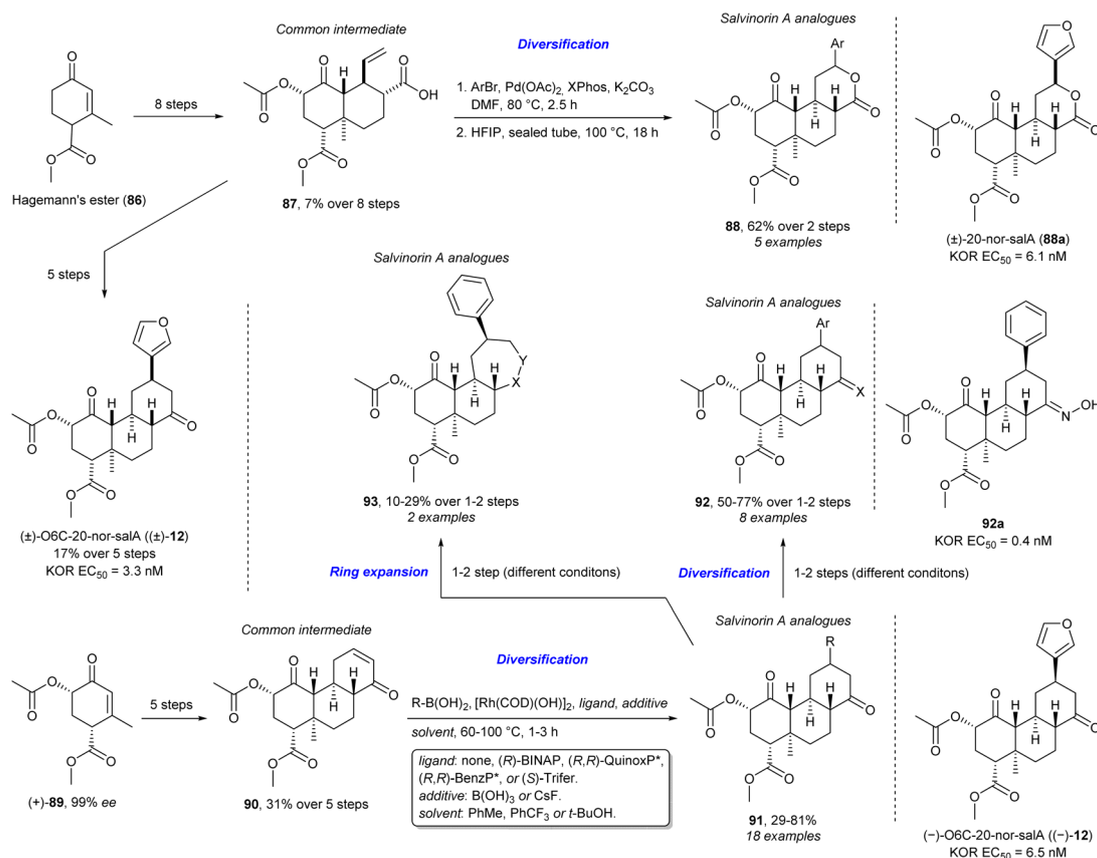
Scheme 7 Synthesis of dPNPs and identification of Hh signalling inhibitor **82a**.²⁹ CFL = compact fluorescent lamp, Eosin Y = 2-(2,4,5,7-tetrabromo-6-oxido-3-oxo-3H-xanthen-9-yl)benzoate, Hantzsch ester = diethyl 1,4-dihydro-2,6-dimethyl-3,5-pyridinedicarboxylate, PPTS = pyridinium *p*-toluenesulfonate, Xantphos = (9,9-dimethyl-9H-xanthen-4,5-diyl)bis(diphenylphosphane).

introduced in the design of the PNPs. This is slightly different from the work on (–)-asteroxin-1 where the diversity is introduced to the resulting PNPs *via* ring distortions afterwards and not directly into the design of the initial PNPs.

As shown above, taking certain elements from NPs to generate NP-inspired compounds can certainly result in biologically relevant compounds. Nevertheless, the actual TS of NPs and their simplified derivatives has long been used to access bioactive compounds and generate SAR information about a certain pharmacologically active NP. However, while beneficial, important areas of chemical space may be missed due to synthetic limitations.⁶⁴ Alongside the strategies focused on generating a wide range of structurally different analogues, modern pragmatic takes on TS such as DTS, FOS, DRA, and PDR have also developed in order to streamline the process of investigating the chemical space surrounding an NP. These share important similarities with other strategies, especially BIOS. Recent work presents the synthesis of salvinorin A (saLA) analogues using DRA to explore the chemical space around this NP (Scheme 8).^{45,46,105} SaLA is a potent κ -opioid receptor agonist. However, its TS has been troublesome due to its complexity and instability which in turn have also made exploration of the chemical space around it and SAR studies difficult. The NP was treated as a dynamic TM to reduce synthetic complexity while retaining molecular complexity. This led to the realisation that a rational removal of the C20 methyl would ease the synthesis and stabilise the resulting compounds. The goal of easing the synthesis of the NP is also a key point in FOS and PDR. Additionally, the authors used molecular docking (another key attribute in FOS) to evaluate 20-nor-salvinorin A, which suggested that it would have similar binding to saLA. Thus, they

synthesised the common intermediate **87** in eight steps from Hagemann's ester (**86**) which could be diversified in a DTS-fashion through a Heck reaction followed by lactonisation to afford the first generation of saLA analogues **88** which were more stable than saLA. More importantly, the analogue (\pm)-20-nor-saLA (**88a**) showed similar potency and selectivity. They then synthesised (\pm)-O6C-20-nor-saLA ((\pm)-**12**) in five steps from **87** and identified that replacement of the O6 with a carbon further stabilised the compound while retaining the potency and selectivity. They then developed an asymmetric synthesis of **12** starting from (+)-**89** which could be synthesised with a 99% enantiomeric excess (ee). From (+)-**89** the common intermediate **90** was accessed in five steps. This intermediate was diversified by a Hayashi conjugate addition to give the second generation of saLA analogues **91** including the enantioenriched (–)-**12**. Further diversification of the second vector, the ketone, allowed for synthesis of oximes and alcohols (**92**) through condensation and nucleophilic addition/reduction, respectively. Consequently, a saLA analogue with improved potency (**92a**) was identified. Lastly, ring expansion of **91** in a CtD-manner through the Beckmann rearrangement and Baeyer–Villiger oxidation afforded the corresponding lactam or lactone (**93**), respectively. This second generation of saLA analogues allowed for further SAR study and exploration of the saLA chemical space through a common and diversifiable intermediate similar to DTS and AOS. In addition, some of the added functionalities are NP fragments or privileged scaffolds showing some resemblance to PNP and BIOS in this diversification step. In total, five first generation and 29 second generation saLA analogues were synthesised.





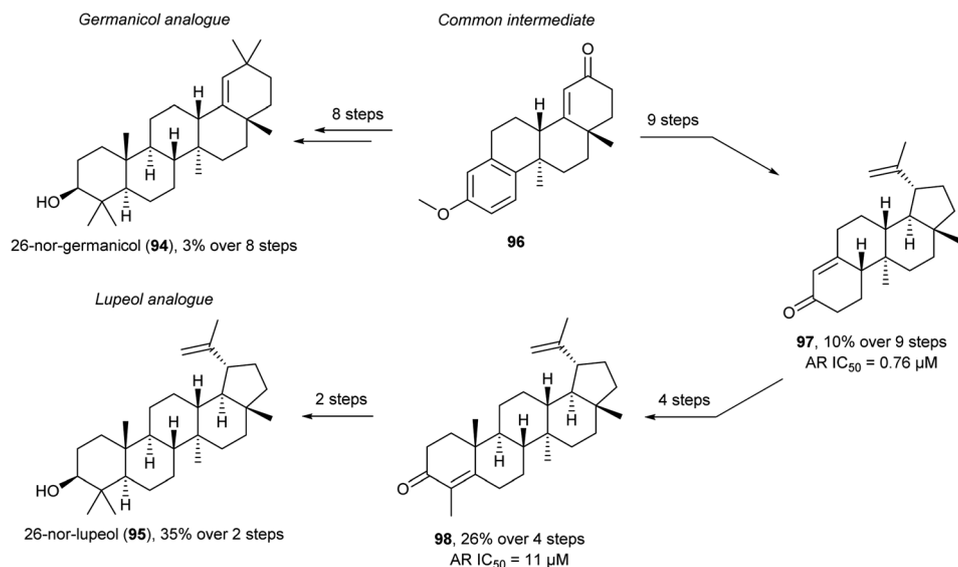
Scheme 8 Synthesis of salvinorin A analogues and identification of multiple KOR agonists.^{45,46,105} BenzP* = 1,2-bis(*tert*-butylmethylphosphino)benzene, BINAP = 2,2'-bis(diphenylphosphino)-1,1'-binaphthyl, COD = 1,5-cyclooctadiene, QuinoxP* = 2,3-bis(*tert*-butylmethylphosphino)quinoxaline, (S)-Trifer = 1,1'-bis[1-((*R*)-ferrocenyl-2-(*S*)-ethyl-1-(diethylamino)phenyl)-(*R*)-phosphino]ferrocene, XPhos = dicyclohexyl[2',4',6'-tris(propan-2-yl)[1,1'-biphenyl]-2-yl]phosphane.

In recent work methyl deletion is also used in a FOS of 26-nor-germanicol (**94**) and 26-nor-lupeol (**95**) (Scheme 9).¹⁰⁶ Here the methyl is removed to remove the synthetically difficult vicinal quaternary stereogenic centre from the synthesis. The authors speculated that the biological function of lupeol and germanicol would be retained in the nor-derivatives. They desired to reduce synthetic complexity while retaining function which is very similar to ideas of DRA and PDR. Initially, the common intermediate **96** was synthesised which allowed access to the germanicol analogue, 26-nor-germanicol (**94**), in six steps and the lupeol analogue, 26-nor-lupeol (**95**), in 12 steps overall. The synthesis of **95** went through **97** (six steps from **96**) and **98** (four steps from **97**) which was converted into **95** in two steps. Unfortunately, biological screening of the 26-nor analogues **94** and **95** was not feasible due to dimethyl sulfoxide (DMSO) solubility issues. Interestingly, screening the intermediates *en route* to the TMs, similar to the PDR, led to the identification of two substituted unnatural *ent*-estrans as androgen receptor (AR) antagonists with similar (**98**) and enhanced (**97**) potency compared to lupeol. Thus, these less structurally complex intermediates retain or improve function. The late intermediate **96** allows for the synthesis of other 26-nor analogues similar to a DTS approach to further gain SAR information.

The work on latrunculin analogues¹⁰⁷ should also be mentioned. In this work, the authors simplify and streamline the synthesis including yet another methyl deletion. They identify a simplified analogue that shares similar actin-binding properties to the most active member of the latrunculin family. The work is presented as a DTS but ideas from FOS, DRA, PDA, and even BIOS are easy to identify. The research on analogues of sinularia NPs¹⁰⁸ led to the identification of new compounds with interesting cytotoxicities and selectivities against cancer cell lines. The compounds contain the tricyclic cores as found in sinularia NPs but in general less complex, and one of the compounds could serve as a useful intermediate towards additional analogues. The work is published as an example of PDR, but again it shares a lot of similarities to DTS, AOS, FOS, PDA, and BIOS.

The majority of examples outlined so far focus on terpenes and alkaloids as guiding NP targets for library synthesis. However, polyketides and more specifically polyether ionophores (PEIs) have received considerably less attention. This is most likely in large part due to the absence of a bioactivity-defining scaffold, requiring larger synthetic efforts to approach NP complexity. Recent work has addressed this challenge by deconstructing natural PEIs to smaller fragments and reconstructing them in new ways to create structural

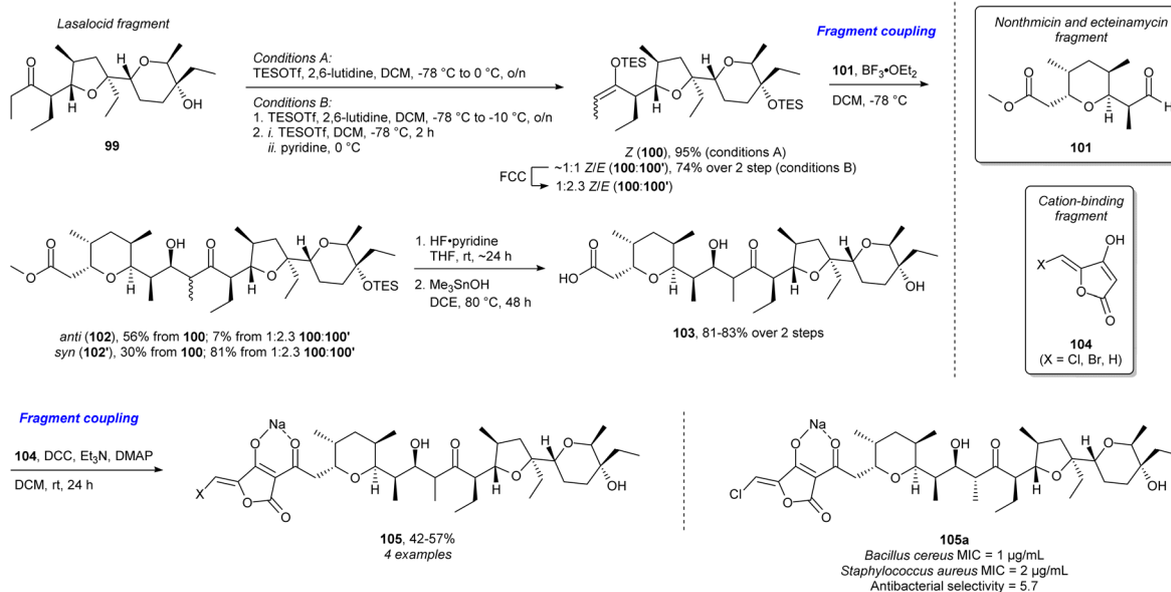


Scheme 9 Synthesis of 26-nor-germanicol (**94**) and 26-nor-lupeol (**95**) and identification of AR antagonists **97** and **98**.¹⁰⁶

diversity and access so-called “hybrid polyethers”.¹⁰⁹ With this structural optimisation they retain the antibacterial activity found in the natural PEIs but achieve improved antibacterial selectivity (Scheme 10). The lasalocid acid fragment **99** was obtained from commercially available lasalocid. Using slightly different conditions, the *Z*-triethylsilyl (TES) enol ether **100** and a mixture of *E*-(**100'**) and *Z*-TES enol ether (2.3 : 1 *E/Z*) were obtained. Then Mukaiyama aldol addition to the synthesised nonthmicin and ecteinamycin fragment **101** yielded the *anti* (**102**) or *syn* (**102'**) product depending on the starting material. Then silyl deprotection and ester hydrolysis afforded **103**. Through DCC-activation of the carboxylic acid, it was coupled to fragment **104** which is proposed to be the group responsible for

cation binding in certain polyether ionophores. This yielded the final hybrid polyethers **105** and the new potent and selective antibiotic **105a**. This is an excellent example of streamlining the synthesis for efficient diversification and investigation of the chemical space and biology related to the NPs, key arguments in FOS, DRA, and PDR. The use of coupling of NP fragments and fragments with a known biological function have strong resemblance to PNP and BIOS.

The described examples show the benefit of rational changes to the retrosynthetic analysis and/or synthesis to reduce synthetic complexity while maintaining structural complexity and biological function (FOS/DRA/PDR) coupled with diversity methodologies to make analogues of the NP (DTS/AOS/DOS) in

Scheme 10 Synthesis of “hybrid polyethers” and identification of the potent and selective antibiotic **105a**.¹⁰⁹ FCC = flash column chromatography, DCC = *N,N'*-dicyclohexylcarbodiimide, DMAP = 4-dimethylaminopyridine, MIC = minimum inhibitory concentration.

the pursuit of “supernatural products”.^{110,111} The examples also illustrate how the boundaries between the different synthetic strategies can be blurred or even broken down, and that this is more likely to be a benefit than a problem. In terms of biological screening, the majority are target-based and influenced by the known bioactivity of the guiding/parent NP since they are screened against the targets known for the guiding NP. However, compound collections may also be made to target a biological phenotype with multiple associated targets or used across various target classes and phenotypes altogether. Therefore, the choice of chemical starting point(s) and synthetic strategies must be carefully considered.

Targeted combination of approaches for a specific project goal?

To conclude the perspective, we have chosen to provide a prospective view to aid researchers seeking to design a strategy for their specific goal, whether this be a target- or phenotype-based screen, or a synthetic chemistry campaign aimed at broad coverage of chemical and biological space. In this context we concur with Shenvi's assertion that the “purpose dictates strategy”.⁷ We have outlined a generalised approach based on a flow chart to ask researchers to consider the project goal and the information available to them at the start, when choosing individual strategies or a combination thereof (Fig. 2).

For example, if the desired outcome is a ligand for a specific target, one must determine whether the target has any known ligands. If that is the case, then the ligand(s) can be broken down into one or two primary scaffolds or fragments with functionalisable handles. Then, the chemical diversity localised around the primary scaffold/fragment should be targeted. This can be achieved by general diversity-generating approaches like (p)DOS and/or fusion with relevant scaffolds as in PNP, as illustrated by the previous work on stemona alkaloids^{85,86} and

our own work on alkaloid- and sterol-inspired compounds.^{35,91,92} This strategy applies to all types of ligands. Additionally, if the ligand is an NP, the NP-driven approaches such as DTS, AOS, DRA, CtD, PDR, and FOS are also very good options. If very specific SAR questions need to be answered by densely populating an area of chemical space, then DTS and AOS are useful. If the NP is in high abundance, CtD is an option to quickly access diverse analogues of the NP. Furthermore, when the pharmacophore is known for the NP, it can be used as a primary fragment in a (p)DOS/BIOS/PNP approach but is also applicable in a FOS/PDR campaign. If the NP is in low abundance, unstable, or hard to access synthetically, breaking it down to useful fragment(s) as above or following a DRA approach may be beneficial as showcased in the work on sala analogues.^{45,46,105}

On the other hand, if no ligands are known or an entirely new chemotype is sought, one can employ an X-ray crystal structure or AlphaFold^{112,113} to model a binding site and predict a pharmacophore model which can be targeted by synthesis. The predicted pharmacophore can then be used as a starting point in a (p)DOS/PNP campaign. Alternatively, a fragment-based screen can be a cost-efficient approach to generate new scaffolds and starting points while covering a larger proportion of chemical space, around which a (p)DOS/PNP approach can be centred. Notably, even the fragments themselves can be designed by DOS to increase chemical diversity and Fsp³ content in the fragment collection.¹¹⁴ This has already been used in some cases¹¹⁵ including the reported synthesis of fluorinated Fsp³-rich fragments.¹¹⁶

In addition to a target-based approach, a phenotype of interest can also be selected and screened against. If there are known modulators of the phenotype, then diverse fragments thereof could be picked. The fragments are fused with diverse secondary fragments similar to the case with known ligands. Similarly, the NP-based approaches are also applicable if the modulators are NPs. If no modulators of the phenotype of interest are known,

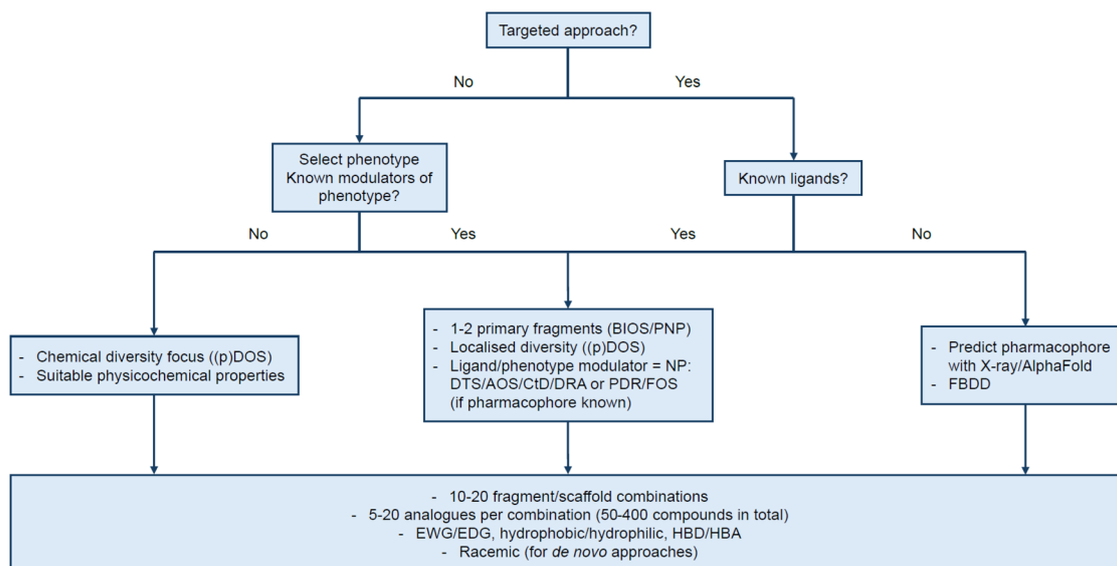


Fig. 2 Flow chart for choosing the optimal strategies based on the project goals.



a chemical diversity-driven approach could be applied, similarly to (p)DOS. Compounds with suitable physicochemical properties to reach the target site should be prioritised. In this regard, using NP-inspired starting points as in BIOS/(d)PNP can be beneficial to reach biologically relevant space. This approach is also useful if no specific phenotype is targeted, but an unbiased phenotypic evaluation such as cell painting is used, as shown with the work on quinalogs.^{79,80} Cell painting is a particularly powerful method to assess the bioactivity of a compound collection with no specific target or phenotype in mind, in an unbiased way. Here, cells are stained with multiple dyes covering different organelles, and changes to staining patterns after compound treatment can be measured. Importantly, with a large and diverse reference set of compounds at hand, it is possible to identify modes of action and even target(s) of new compounds, by comparison. Recent bioactivity clusters that can rapidly be discerned are lysosomotropism¹¹⁷ and cholesterol homeostasis,¹¹⁸ modulation of tubulin polymerisation,¹¹⁹ mitochondrial stress,¹²⁰ modulation of potassium channels,¹²¹ and pyrimidine biosynthesis,¹²² amongst others. It should be noted that in approaches with no known ligands or modulators, high-throughput screening (HTS) is always an option to identify starting points if time and budget allows.

In all library synthesis strategies where bioactive scaffolds or fragments are designed from scratch, the goal is typically to synthesise 10–20 fragment combinations or scaffolds, with approximately 5–20 analogues each for a total of 50–400 compounds. Fragments, or the resulting analogues, should ideally contain diverse substituents including electron-withdrawing groups (EWGs), electron-donating groups (EDGs), hydrophobic and hydrophilic groups, and ideally hydrogen bond donors (HBDs) and hydrogen bond acceptors (HBAs), at different positions. Additionally, substituents with different steric bulk can be explored. This includes flexible or locked and linear or branched groups. Lastly, the compounds should be synthesised as racemic mixtures to reduce bias and obtain twice the number of compounds for biological screening in approaches where compounds are synthesised *de novo*. In strategies where compounds are directly derived from available NPs, this criterion is, in most cases, not possible to fulfil. In our experience the above criteria are both necessary and sufficient to find bioactive compounds for a given target or phenotype of interest, even from a relatively modestly sized library of <100 compounds.

Outlook and conclusions

In addition to the considerations outlined thus far, we see great benefits in adopting recent developments in the field, including C–H and late-stage functionalisation, as well as single-atom and skeletal editing strategies, to streamline analogue synthesis and further expand the diversity of a given collection, but also for the medicinal chemistry optimisation of the hit compounds into leads. Efficiently combining library synthesis with C–H/late-stage functionalisation has proven effective as shown from the work on oxazatwistane-derived analogues.¹²³ It can provide access to new analogues from a vector/diversification

point that is not obvious. Several strategies for C–H functionalisation are known including free-radical, metal-catalysed, photochemical, electrochemical, and chemoenzymatic reactions.¹²⁴ Chemoenzymatic synthesis has proven to be a very useful strategy for C–H functionalisation. Very recent work combined chemoenzymatic synthesis with DOS and CtD, which was termed chemoenzymatic DOS (CeDOS).¹²⁵ Chemoenzymatic C–H functionalisation of the NP parthenolide enabled the synthesis of a diverse parthenolide-based library using divergent chemical routes. With chemoenzymatic synthesis applied successfully to TS of NPs,^{126–133} this is a powerful example of the combination of chemoenzymatic synthesis with a diversifying strategy such as DOS. Two main strategies can be considered when utilising chemoenzymatic synthesis in library synthesis and medicinal chemistry.¹³⁴ The first is the early-stage synthesis of novel building blocks combined with a general diversifying strategy to access new analogues. The second is the late-stage diversification of NP-inspired advanced key intermediates or analogues. Late-stage single-atom and skeletal editing is an attractive approach in medicinal chemistry to quickly elucidate SAR of hit and lead compounds.¹³⁵ Removal, addition, or exchange of a single atom in a molecule is often achieved by modifying the synthesis of the compound from an early point in the route or by a totally different route. However, recent advantages in direct atom deletion, insertion, and exchange^{136–153} can in some cases remove the need for new retrosynthetic analysis and provide new diverse compounds more efficiently. Additionally, subtle changes to the overall molecular shape and not just single atoms can have a large impact on the function and properties of compounds. Efficient methods to access isomeric chemical space by “shapeshifting” have also been reported.¹⁵⁴ This example bears some resemblance to CtD strategies, which may also be considered skeletal editing, in a broad sense. Even combining chemoenzymatic methods with skeletal editing can be a powerful tool as showcased by the recent example which allowed for ring expansion at aliphatic C–H sites.¹⁵⁵ With the current and continuously growing number of methods in the area of late-stage C–H functionalisation and skeletal editing, we see these becoming more integrated into the synthesis of NP-inspired compounds and lead optimisation and in accessing new chemically and biologically relevant space.

In summary, we have outlined how a large variety of strategies are available to access diverse NP-derived and -inspired compound collections. We emphasise how combining several strategies or elements thereof, depending on the specific need and purpose, can be beneficial in the pursuit of new bioactive molecules in an efficient manner. We hope that the ideas outlined here will serve to help chemists push the boundaries in the synthesis of natural-product inspired, biologically relevant compound collections.

Abbreviations

5-HT _{1A}	5-Hydroxytryptamine 1A
5-HT _{2B}	5-Hydroxytryptamine 2B receptor
5-HT _{2C}	5-Hydroxytryptamine 2C receptor



σ_1R	Sigma-1 receptor	MEM	2-Methoxyethoxymethyl
σ_2R	Sigma-2 receptor	Mes	Mesityl
Ac	Acetyl	MIC	Minimum inhibitory concentration
ADS	Activity-directed synthesis	MLCK1	Myosin light chain kinase 1
AOS	Analogue-oriented synthesis	mol.	Molecular sieves
AR	Androgen receptor	sieves	
Ar	Aryl	MWI	Microwave irradiation
B/C/P	Build/couple/pair	NBS	<i>N</i> -Bromosuccinimide
BenzP*	1,2-Bis(<i>tert</i> -butylmethylphosphino)benzene	<i>n</i> -Bu	<i>n</i> -Butyl
BINAP	2,2'-Bis(diphenylphosphino)-1,1'-binaphthyl	NCS	<i>N</i> -Chlorosuccinimide
BIOS	Biology-oriented synthesis	NP	Natural product
Bn	Benzyl	o/n	Overnight
Boc	<i>tert</i> -Butyloxycarbonyl	pDOS	Privileged-substructure-based diversity-oriented synthesis
CeDOS	Chemoenzymatic diversity-oriented synthesis	PDR	Pharmacophore-directed retrosynthesis
CFL	Compact fluorescent lamp	PEI	Polyether ionophore
CLS	Combinatorial library synthesis	Ph	Phenyl
COD	1,5-Cyclooctadiene	PMP	<i>para</i> -Methoxyphenyl
conc.	Concentrated	PNP	Pseudo-natural product
CS	Conventional synthesis	PPTS	Pyridinium <i>para</i> -toluenesulfonate
CtD	Complexity-to-diversity	PS	Polystyrene
Cy	Cyclohexyl	<i>p</i> -Ts	<i>para</i> -Tosyl
DABCO	1,4-Diazabicyclo[2.2.2]octane	QuinoxP*	2,3-bis(<i>tert</i> -Butylmethylphosphino)quinoxaline
DCC	<i>N,N'</i> -Dicyclohexylcarbodiimide	RCM	Ring-closing metathesis
DCE	Dichloroethane	rt	Room temperature
DCM	Dichloromethane	sala	Salvinorin A
DIPEA	<i>N,N</i> -Diisopropylethylamine	SAR	Structure-activity relationship
DMA	Dimethylacetamide	(<i>S</i>)-Trifer	1,1'-bis{1-[(<i>R</i>)-Ferrocenyl-2- (<i>S</i>)-ethyl-1-(diethylamino)phenyl]-(<i>R</i>)-phosphino}ferrocene
DMAP	4-Dimethylaminopyridine	TBTU	<i>N,N,N',N'</i> -Tetramethyl- <i>O</i> -(benzotriazol-1-yl)uronium tetrafluoroborate
DMEDA	<i>N,N'</i> -Dimethylethylenediamine	<i>t</i> -Bu	<i>tert</i> -Butyl
DMF	Dimethylformamide	TES	Triethylsilyl
DMSO	Dimethyl sulfoxide	Tf	Triflyl
DOS	Diversity-oriented synthesis	TFA	Trifluoroacetic acid
DPEPhos	bis[(2-Diphenylphosphino)phenyl]ether	THF	Tetrahydrofuran
dPNP	Diverse pseudo-natural product	TM	Target molecule
dr	Diastereoisomeric ratio	TMS	Trimethylsilyl
DRA	Dynamic retrosynthetic analysis	TOS	Target-oriented synthesis
DTS	Diverted total synthesis	TPS	Two-phase synthesis
DTU	Technical University of Denmark	TS	Total synthesis
EC ₅₀	Half maximal effective concentration	Xantphos	(9,9-Dimethyl-9 <i>H</i> -xanthene-4,5-diyl)bis(diphenylphosphane)
EDG	Electron-donating group	XPhos	Dicyclohexyl[2',4',6'-tris(propan-2-yl)[1,1'-biphenyl]-2-yl]phosphane
ee	Enantiomeric excess		
ESCRT	Endosomal sorting complexes required for transport		
Et	Ethyl		
EWG	Electron-withdrawing group		
FBDD	Fragment-based drug-discovery		
FCC	Flash column chromatography		
FLS	Focussed library synthesis		
FOS	Function-oriented synthesis		
Fsp ³	Fraction of sp ³ -hybridised carbons		
HBA	Hydrogen bond acceptor		
HBD	Hydrogen bond donor		
HFIP	Hexafluoroisopropanol		
Hh	Hedgehog		
HTS	High-throughput screening		
IC ₅₀	Half maximal inhibitory concentration		
K _i	Inhibitory constant		
LC3	Microtubule-associated protein 1A/1B light chain 3		
MCR	Multicomponent reaction		
Me	Methyl		

Data availability

The supplementary figures with graphical illustrations, explanations, and examples of the individual strategies are available in the ESI.†

Author contributions

F. S. B. and L. L. conceived the perspective. F. S. B. produced all figures. F. S. B. and L. L. conducted the literature search and wrote the manuscript.



Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

We thank the Novo Nordisk Foundation (NNF21OC0067188), the European Research Council (ERC, ChemBioChol, 101041783) and DTU for funding. We thank Prof. Herbert Waldmann and Dr Daniel Foley for critical reading of the manuscript and helpful comments.

References

- L. Laraia and H. Waldmann, *Drug Discovery Today: Technol.*, 2017, **23**, 75–82.
- B. J. Huffman and R. A. Shenvi, *J. Am. Chem. Soc.*, 2019, **141**, 3332–3346.
- M. Grigalunas, A. Burhop, A. Christoforow and H. Waldmann, *Curr. Opin. Chem. Biol.*, 2020, **56**, 111–118.
- M. Grigalunas, S. Brakmann and H. Waldmann, *J. Am. Chem. Soc.*, 2022, **144**, 3314–3329.
- R. J. Young, S. L. Flitsch, M. Grigalunas, P. D. Leeson, R. J. Quinn, N. J. Turner and H. Waldmann, *JACS Au*, 2022, **2**, 2400–2416.
- J. Liu, M. Grigalunas and H. Waldmann, in *Annual Reports in Medicinal Chemistry*, Academic Press, 2023, vol. 61, pp. 1–53.
- R. A. Shenvi, *ACS Cent. Sci.*, 2024, **10**, 519–528.
- S. B. Bharate and C. W. Lindsley, *J. Med. Chem.*, 2024, **67**, 20723–20730.
- D. J. Newman and G. M. Cragg, *J. Nat. Prod.*, 2020, **83**, 770–803.
- C. R. Pye, M. J. Bertin, R. S. Lokey, W. H. Gerwick and R. G. Linington, *Proc. Natl. Acad. Sci. U. S. A.*, 2017, **114**, 5601–5606.
- M. D. Burke and S. L. Schreiber, *Angew. Chem., Int. Ed.*, 2004, **43**, 46–58.
- W. R. J. D. Galloway, A. Isidro-Llobet and D. R. Spring, *Nat. Commun.*, 2010, **1**, 80.
- S. Oh and S. B. Park, *Chem. Commun.*, 2011, **47**, 12754–12761.
- J. Kim, H. Kim and S. B. Park, *J. Am. Chem. Soc.*, 2014, **136**, 14629–14638.
- G. Karageorgis, S. Warriner and A. Nelson, *Nat. Chem.*, 2014, **6**, 872–876.
- G. Karageorgis, S. Liver and A. Nelson, *ChemMedChem*, 2020, **15**, 1776–1782.
- G. Karageorgis, D. J. Foley, L. Laraia and H. Waldmann, *Nat. Chem.*, 2020, **12**, 227–235.
- G. Karageorgis, D. J. Foley, L. Laraia, S. Brakmann and H. Waldmann, *Angew. Chem., Int. Ed.*, 2021, **60**, 15705–15723.
- S. Wetzler, R. S. Bon, K. Kumar and H. Waldmann, *Angew. Chem., Int. Ed.*, 2011, **50**, 10800–10826.
- H. van Hattum and H. Waldmann, *J. Am. Chem. Soc.*, 2014, **136**, 11853–11859.
- P. A. Wender, V. A. Verma, T. J. Paxton and T. H. Pillow, *Acc. Chem. Res.*, 2008, **41**, 40–49.
- M. E. Abbasov, R. Alvarino, C. M. Chaheine, E. Alonso, J. A. Sánchez, M. L. Conner, A. Alfonso, M. Jaspars, L. M. Botana and D. Romo, *Nat. Chem.*, 2019, **11**, 342–350.
- R. W. Huigens III, K. C. Morrison, R. W. Hicklin, T. A. Flood Jr, M. F. Richter and P. J. Hergenrother, *Nat. Chem.*, 2013, **5**, 195–202.
- S. Woo and R. A. Shenvi, *Acc. Chem. Res.*, 2021, **54**, 1157–1167.
- J. T. Njardarson, C. Gaul, D. Shan, X.-Y. Huang and S. J. Danishefsky, *J. Am. Chem. Soc.*, 2004, **126**, 1038–1040.
- R. M. Wilson and S. J. Danishefsky, *J. Org. Chem.*, 2006, **71**, 8329–8351.
- Y. Ishihara and P. S. Baran, *Synlett*, 2010, **12**, 1733–1745.
- Z. A. Könst, A. R. Szklarski, S. Pellegrino, S. E. Michalak, M. Meyer, C. Zanette, R. Cencic, S. Nam, V. K. Voora, D. A. Horne, J. Pelletier, D. L. Mobley, G. Yusupova, M. Yusupov and C. D. Vanderwal, *Nat. Chem.*, 2017, **9**, 1140–1149.
- S. Bag, J. Liu, S. Patil, J. Bonowski, S. Koska, B. Schölermann, R. Zhang, L. Wang, A. Pahl, S. Sievers, L. Brieger, C. Strohmann, S. Ziegler, M. Grigalunas and H. Waldmann, *Nat. Chem.*, 2024, **16**, 945–958.
- A. Schuffenhauer, P. Ertl, S. Roggo, S. Wetzler, M. A. Koch and H. Waldmann, *J. Chem. Inf. Model.*, 2007, **47**, 47–58.
- V. Pogacic, A. N. Bullock, O. Fedorov, P. Filippakopoulos, C. Gasser, A. Biondi, S. Meyer-Monard, S. Knapp and J. Schwaller, *Cancer Res.*, 2007, **67**, 6916–6924.
- S. D. Larsen, C. F. Stachew, P. M. Clare, J. W. Cubbage and K. L. Leach, *Bioorg. Med. Chem. Lett.*, 2003, **13**, 3491–3495.
- B. M. Ibbeson, L. Laraia, E. Alza, C. J. O' Connor, Y. S. Tan, H. M. L. Davies, G. McKenzie, A. R. Venkitaraman and D. R. Spring, *Nat. Commun.*, 2014, **5**, 3155.
- J. Kim, J. Jung, J. Koo, W. Cho, W. S. Lee, C. Kim, W. Park and S. B. Park, *Nat. Commun.*, 2016, **7**, 13196.
- F. S. Bro, L. Depta, N. J. Dekker, H. P. Bryce-Rogers, M. L. Madsen, K. F. Præstegaard, T. Petersson, T. Whitmarsh-Everiss, M. Kubus and L. Laraia, *ACS Cent. Sci.*, 2025, **11**, 136–146.
- J. Ceballos, M. Schwalfenberg, G. Karageorgis, E. S. Reckzeh, S. Sievers, C. Ostermann, A. Pahl, M. Sellstedt, J. Nowacki, M. A. Carnero Corrales, J. Wilke, L. Laraia, K. Tschapalda, M. Metz, D. A. Sehr, S. Brand, K. Winklhofer, P. Janning, S. Ziegler and H. Waldmann, *Angew. Chem., Int. Ed.*, 2019, **58**, 17016–17025.
- A. Nören-Müller, I. Reis-Corrêa, H. Prinz, C. Rosenbaum, K. Saxena, H. J. Schwalbe, D. Vestweber, G. Cagna, S. Schunk, O. Schwarz, H. Schiewe and H. Waldmann, *Proc. Natl. Acad. Sci. U. S. A.*, 2006, **103**, 10606–10611.
- I. R. Corrêa Jr, A. Nören-Müller, H.-D. Ambrosi, S. Jakupovic, K. Saxena, H. Schwalbe, M. Kaiser and H. Waldmann, *Chem.-Asian J.*, 2007, **2**, 1109–1126.
- P. A. Wender, J. L. Sloane, Q. H. Luu-Nguyen, Y. Ogawa, A. J. Shimizu, S. M. Ryckbosch, J. H. Tyler and C. Hardman, *J. Org. Chem.*, 2020, **85**, 15116–15128.



- 40 E. Llabani, R. W. Hicklin, H. Y. Lee, S. E. Motika, L. A. Crawford, E. Weerapana and P. J. Hergenrother, *Nat. Chem.*, 2019, **11**, 521–532.
- 41 K. Yamamoto, R. M. Garbaccio, S. J. Stachel, D. B. Solit, G. Chiosis, N. Rosen and S. J. Danishefsky, *Angew. Chem., Int. Ed.*, 2003, **42**, 1280–1284.
- 42 X. Lei and S. J. Danishefsky, *Adv. Synth. Catal.*, 2008, **350**, 1677–1681.
- 43 R. B. Woodward, M. P. Cava, W. D. Ollis, A. Hunger, H. U. Daeniker and K. Schenker, *J. Am. Chem. Soc.*, 1954, **76**, 4749–4751.
- 44 R. B. Woodward, M. P. Cava, W. D. Ollis, A. Hunger, H. U. Daeniker and K. Schenker, *Tetrahedron*, 1963, **19**, 247–288.
- 45 S. Hirasawa, M. Cho, T. F. Brust, J. J. Roach, L. M. Bohn and R. A. Shenvi, *Bioorg. Med. Chem. Lett.*, 2018, **28**, 2770–2772.
- 46 S. J. Hill, N. Dao, V. Q. Dang, E. L. Stahl, L. M. Bohn and R. A. Shenvi, *ACS Cent. Sci.*, 2023, **9**, 1567–1574.
- 47 P. Ertl, S. Roggo and A. Schuffenhauer, *J. Chem. Inf. Model.*, 2008, **48**, 68–74.
- 48 S. L. Schreiber, *Science*, 2000, **287**, 1964–1969.
- 49 D. R. Spring, *Org. Biomol. Chem.*, 2003, **1**, 3867–3870.
- 50 D. L. Boger and C. E. Brotherton, *J. Org. Chem.*, 1984, **49**, 4050–4055.
- 51 R. J. Spandl, A. Bender and D. R. Spring, *Org. Biomol. Chem.*, 2008, **6**, 1149–1158.
- 52 A. Krzyzanowski, A. Pahl, M. Grigalunas and H. Waldmann, *J. Med. Chem.*, 2023, **66**, 12739–12750.
- 53 T. I. Oprea and C. Bologa, *J. Med. Chem.*, 2023, **66**, 12710–12714.
- 54 P. A. Clemons, N. E. Bodycombe, H. A. Carrinski, J. A. Wilson, A. F. Shamji, B. K. Wagner, A. N. Koehler and S. L. Schreiber, *Proc. Natl. Acad. Sci. U. S. A.*, 2010, **107**, 18787–18792.
- 55 M. P. Gleeson, A. Hersey, D. Montanari and J. Overington, *Nat. Rev. Drug Discovery*, 2011, **10**, 197–208.
- 56 A. R. Leach and M. M. Hann, *Curr. Opin. Chem. Biol.*, 2011, **15**, 489–496.
- 57 T. Böttcher, *J. Chem. Inf. Model.*, 2016, **56**, 462–470.
- 58 R. M. Demoret, M. A. Baker, M. Ohtawa, S. Chen, C. C. Lam, S. Khom, M. Roberto, S. Forli, K. N. Houk and R. A. Shenvi, *J. Am. Chem. Soc.*, 2020, **142**, 18599–18618.
- 59 P. A. Wender, *Nat. Prod. Rep.*, 2014, **31**, 433–440.
- 60 P. A. Wender, *Tetrahedron*, 2013, **69**, 7529–7550.
- 61 S. Ichikawa, *Chem. Rec.*, 2016, **16**, 1106–1115.
- 62 I. Pavlinov, E. M. Gerlach and L. N. Aldrich, *Org. Biomol. Chem.*, 2019, **17**, 1608–1623.
- 63 S. E. Motika and P. J. Hergenrother, *Nat. Prod. Rep.*, 2020, **37**, 1395–1403.
- 64 N. J. Truax and D. Romo, *Nat. Prod. Rep.*, 2020, **37**, 1436–1453.
- 65 A. Nelson and G. Karageorgis, *RSC Med. Chem.*, 2021, **12**, 353–362.
- 66 B. O. Alkubaisi, A. I. Shahin, R. A. Zenati, A. Ravi, R. Alchami, M. Alkalla, R. Khaled, M. I. El-Gamal and T. H. Al-Tel, *ChemMedChem*, 2023, **18**, e202300117.
- 67 X.-F. Cheng, M. Grigalunas and H. Waldmann, *Arkivoc*, 2024, **5**, 202312153.
- 68 P. Craven, A. Aimon, M. Dow, N. Fleury-Bregeot, R. Guilleux, R. Morgentin, D. Roche, T. Kalliokoski, R. Foster, S. P. Marsden and A. Nelson, *Bioorg. Med. Chem.*, 2015, **23**, 2629–2635.
- 69 G. Li Petri, M. V. Raimondi, V. Spanò, R. Holl, P. Barraja and A. Montalbano, *Top. Curr. Chem.*, 2021, **379**, 34.
- 70 Y. Choi, S. Lee, H. Kim and S. B. Park, *Front. Chem.*, 2022, **10**, 841250.
- 71 F. P. Player and D. J. Foley, *Asian J. Org. Chem.*, 2024, **13**, e202400397.
- 72 A. Lawer, F. P. Player, V. M. Avery and D. J. Foley, *Adv. Synth. Catal.*, 2024, **366**, 2090–2100.
- 73 J. J. Ciardiello, H. L. Stewart, H. F. Sore, W. R. J. D. Galloway and D. R. Spring, *Bioorg. Med. Chem.*, 2017, **25**, 2825–2843.
- 74 K. T. Mortensen, T. J. Osberger, T. A. King, H. F. Sore and D. R. Spring, *Chem. Rev.*, 2019, **119**, 10288–10317.
- 75 A. K. Yudin, *Chem. Sci.*, 2014, **6**, 30–49.
- 76 D. Garcia Jimenez, V. Poongavanam and J. Kihlberg, *J. Med. Chem.*, 2023, **66**, 5377–5396.
- 77 G. Niggemeyer, A. Knyazeva, R. Gasper, D. Corkery, P. Bodenbinder, J. J. Holstein, S. Sievers, Y.-W. Wu and H. Waldmann, *Angew. Chem., Int. Ed.*, 2022, **61**, e202114328.
- 78 A. Knyazeva, S. Li, D. P. Corkery, K. Shankar, L. K. Herzog, X. Zhang, B. Singh, G. Niggemeyer, D. Grill, J. D. Gilthorpe, M. Gaetani, L.-A. Carlson, H. Waldmann and Y.-W. Wu, *Proc. Natl. Acad. Sci. U. S. A.*, 2024, **121**, e2317680121.
- 79 D. J. Foley, S. Zinken, D. Corkery, L. Laraia, A. Pahl, Y.-W. Wu and H. Waldmann, *Angew. Chem., Int. Ed.*, 2020, **59**, 12470–12476.
- 80 M. Grigalunas, A. Burhop, S. Zinken, A. Pahl, J.-M. Gally, N. Wild, Y. Mantel, S. Sievers, D. J. Foley, R. Scheel, C. Strohmman, A. P. Antonchick and H. Waldmann, *Nat. Commun.*, 2021, **12**, 1883.
- 81 D. J. Foley and H. Waldmann, *Chem. Soc. Rev.*, 2022, **51**, 4094–4120.
- 82 R. Narayan, J. O. Bauer, C. Strohmman, A. P. Antonchick and H. Waldmann, *Angew. Chem., Int. Ed.*, 2013, **52**, 12892–12896.
- 83 R. Narayan, M. Potowski, Z.-J. Jia, A. P. Antonchick and H. Waldmann, *Acc. Chem. Res.*, 2014, **47**, 1296–1310.
- 84 T. Schneidewind, S. Kapoor, G. Garivet, G. Karageorgis, R. Narayan, G. Vendrell-Navarro, A. P. Antonchick, S. Ziegler and H. Waldmann, *Cell Chem. Biol.*, 2019, **26**, 512–523.
- 85 K. J. Frankowski, B. Neuenswander and J. Aubé, *J. Comb. Chem.*, 2008, **10**, 721–725.
- 86 K. J. Frankowski, V. Setola, J. M. Evans, B. Neuenswander, B. L. Roth and J. Aubé, *Proc. Natl. Acad. Sci. U. S. A.*, 2011, **108**, 6727–6732.
- 87 K. J. Frankowski, V. Setola, J. M. Evans, B. Neuenswander, B. L. Roth and J. Aubé, *Proc. Natl. Acad. Sci. U. S. A.*, 2012, **109**, 15526–15527.
- 88 A. Shaabani, A. Maleki, H. Mofakham and H. R. Khavasi, *J. Comb. Chem.*, 2008, **10**, 323–326.



- 89 A. K. Ghosh and M. Brindisi, *J. Med. Chem.*, 2015, **58**, 2895–2940.
- 90 K. J. Frankowski, J. E. Golden, Y. Zeng, Y. Lei and J. Aubé, *J. Am. Chem. Soc.*, 2008, **130**, 6018–6024.
- 91 R. Yao, A. A. Jensen, H. P. Bryce-Rogers, K. Schultz-Knudsen, L. Zhou, N. P. Hovendal, H. Pedersen, M. Kubus, T. Ulven and L. Laraia, *J. Med. Chem.*, 2023, **66**, 11536–11554.
- 92 T. Whitmarsh-Everiss, A. H. Olsen and L. Laraia, *Angew. Chem., Int. Ed.*, 2021, **60**, 26755–26761.
- 93 B. Melillo, J. Zoller, B. K. Hua, O. Verho, J. C. Borghs, S. D. Jr. Nelson, M. Maetani, M. J. Wawer, P. A. Clemons and S. L. Schreiber, *J. Am. Chem. Soc.*, 2018, **140**, 11784–11790.
- 94 K. A. Scott, N. Ropek, B. Melillo, S. L. Schreiber, B. F. Cravatt and E. V. Vinogradova, *Curr. Res. Chem. Biol.*, 2022, **2**, 100028.
- 95 D. J. Foley, P. G. E. Craven, P. M. Collins, R. G. Doveston, A. Aimon, R. Talon, I. Churcher, F. von Delft, S. P. Marsden and A. Nelson, *Chem.–Eur. J.*, 2017, **23**, 15227–15232.
- 96 C. Townley, L. McMurray, S. P. Marsden and A. Nelson, *Bioorg. Med. Chem. Lett.*, 2022, **62**, 128631.
- 97 E. A. Okolo, A. Pahl, S. Sievers, C. M. Pask, A. Nelson and S. P. Marsden, *Chem.–Eur. J.*, 2023, **29**, e202203992.
- 98 S. Liu, J. S. Scotti and S. A. Kozmin, *J. Org. Chem.*, 2013, **78**, 8645–8654.
- 99 N. G. Paciaroni, R. Ratnayake, J. H. Matthews, V. M. Norwood IV, A. C. Arnold, L. H. Dang, H. Luesch and R. W. Huigens III, *Chem.–Eur. J.*, 2017, **23**, 4327–4335.
- 100 V. Srinivasulu, P. Schilf, S. Ibrahim, M. A. Khanfar, S. M. Sieburth, H. Omar, A. Sebastian, R. A. AlQawasmeh, M. J. O'Connor and T. H. Al-Tel, *Nat. Commun.*, 2018, **9**, 4989.
- 101 V. Srinivasulu, S. M. Sieburth, M. A. Khanfar, I. A. Abu-Yousef, A. Majdalawieh, M. Ramanathan, A. Sebastian and T. H. Al-Tel, *J. Org. Chem.*, 2021, **86**, 12872–12885.
- 102 G. Srikanth, A. Ravi, A. Sebastian, J. Joseph, M. A. Khanfar, M. I. El-Gamal, R. A. Al-Qawasmeh, I. A. Shehadi, S. McN. Sieburth, I. A. Abu-Yousef, A. F. Majdalawieh and T. H. Al-Tel, *Eur. J. Org. Chem.*, 2023, **26**, e202300080.
- 103 J. Liu, J. Flegel, F. Otte, A. Pahl, S. Sievers, C. Strohmam and H. Waldmann, *Angew. Chem., Int. Ed.*, 2021, **60**, 21384–21395.
- 104 V. Porte, B. C. van Veen, H. Zhang, P. Piacentini, S. A. Matheu, S. Woolford, K. R. Sokol, S. Shaaban, H. Weinstabl and N. Maulide, *Org. Lett.*, 2024, **26**, 4873–4876.
- 105 J. J. Roach, Y. Sasano, C. L. Schmid, S. Zaidi, V. Katritch, R. C. Stevens, L. M. Bohn and R. A. Shenvi, *ACS Cent. Sci.*, 2017, **3**, 1329–1336.
- 106 Z. D. Stempel, H. S. Radomska, C. C. Coss and G. C. Micalizio, *Org. Lett.*, 2024, **26**, 3054–3059.
- 107 A. Fürstner, D. Kirk, M. D. B. Fenster, C. Aïssa, D. De Souza and O. Müller, *Proc. Natl. Acad. Sci. U. S. A.*, 2005, **102**, 8103–8108.
- 108 N. J. Truax, S. Ayinde, K. Van, J. O. Liu and D. Romo, *Org. Lett.*, 2019, **21**, 7394–7399.
- 109 S. Lin, H. Liu, E. B. Svenningsen, M. Wollesen, K. M. Jacobsen, F. D. Andersen, J. Moyano-Villameriel, C. N. Pedersen, P. Nørby, T. Tørring and T. B. Poulsen, *Nat. Chem.*, 2021, **13**, 47–55.
- 110 K. K. Wan and R. A. Shenvi, *Synlett*, 2016, **27**, 1145–1164.
- 111 Z.-C. Wu and D. L. Boger, *Nat. Prod. Rep.*, 2020, **37**, 1511–1531.
- 112 J. Jumper, R. Evans, A. Pritzel, T. Green, M. Figurnov, O. Ronneberger, K. Tunyasuvunakool, R. Bates, A. Židek, A. Potapenko, A. Bridgland, C. Meyer, S. A. A. Kohl, A. J. Ballard, A. Cowie, B. Romera-Paredes, S. Nikolov, R. Jain, J. Adler, T. Back, S. Petersen, D. Reiman, E. Clancy, M. Zielinski, M. Steinegger, M. Pacholska, T. Berghammer, S. Bodenstein, D. Silver, O. Vinyals, A. W. Senior, K. Kavukcuoglu, P. Kohli and D. Hassabis, *Nature*, 2021, **596**, 583–589.
- 113 J. Abramson, J. Adler, J. Dunger, R. Evans, T. Green, A. Pritzel, O. Ronneberger, L. Willmore, A. J. Ballard, J. Bambrick, S. W. Bodenstein, D. A. Evans, C.-C. Hung, M. O'Neill, D. Reiman, K. Tunyasuvunakool, Z. Wu, A. Žemgulytė, E. Arvaniti, C. Beattie, O. Bertolli, A. Bridgland, A. Cherepanov, M. Congreve, A. I. Cowen-Rivers, A. Cowie, M. Figurnov, F. B. Fuchs, H. Gladman, R. Jain, Y. A. Khan, C. M. R. Low, K. Perlin, A. Potapenko, P. Savy, S. Singh, A. Stecula, A. Thillaisundaram, C. Tong, S. Yakneen, E. D. Zhong, M. Zielinski, A. Židek, V. Bapst, P. Kohli, M. Jaderberg, D. Hassabis and J. M. Jumper, *Nature*, 2024, **630**, 493–500.
- 114 H. F. Klein, D. J. Hamilton, I. J. P. de Esch, M. Wijtman and P. O'Brien, *Drug Discovery Today*, 2022, **27**, 2484–2496.
- 115 S. L. Kidd, T. J. Osberger, N. Mateu, H. F. Sore and D. R. Spring, *Front. Chem.*, 2018, **6**, 460.
- 116 N. S. Troelsen, E. Shanina, D. Gonzalez-Romero, D. Danková, I. S. A. Jensen, K. J. Śniady, F. Nami, H. Zhang, C. Rademacher, A. Cuenda, C. H. Gotfredsen and M. H. Clausen, *Angew. Chem., Int. Ed.*, 2020, **59**, 2204–2210.
- 117 L. Laraia, G. Garivet, D. J. Foley, N. Kaiser, S. Müller, S. Zinken, T. Pinkert, J. Wilke, D. Corkery, A. Pahl, S. Sievers, P. Janning, C. Arenz, Y. Wu, R. Rodriguez and H. Waldmann, *Angew. Chem., Int. Ed.*, 2020, **59**, 5721–5729.
- 118 T. Schneidewind, A. Brause, B. Schölermann, S. Sievers, A. Pahl, M. G. Sankar, M. Winzker, P. Janning, K. Kumar, S. Ziegler and H. Waldmann, *Cell Chem. Biol.*, 2021, **28**, 1780–1794.
- 119 M. Akbarzadeh, I. Deipenwisch, B. Schoelermann, A. Pahl, S. Sievers, S. Ziegler and H. Waldmann, *Cell Chem. Biol.*, 2022, **29**, 1053–1064.
- 120 S. Rezaei Adariani, D. Agne, S. Koska, A. Burhop, C. Seitz, J. Warmers, P. Janning, M. Metz, A. Pahl, S. Sievers, H. Waldmann and S. Ziegler, *J. Med. Chem.*, 2024, **67**, 13252–13270.
- 121 T. Whitmarsh-Everiss, Z. Wang, C. Hauberg Hansen, L. Depta, E. Sasseti, O. Rafn Dan, A. Pahl, S. Sievers and L. Laraia, *ChemBioChem*, 2023, **24**, e202200555.



- 122 B. Schölermann, J. Bonowski, M. Grigalunas, A. Burhop, Y. Xie, J. G. F. Hoock, J. Liu, M. Dow, A. Nelson, C. Nowak, A. Pahl, S. Sievers and S. Ziegler, *ChemBioChem*, 2022, **23**, e202200475.
- 123 L. Laraia, K. Ohsawa, G. Konstantinidis, L. Robke, Y.-W. Wu, K. Kumar and H. Waldmann, *Angew. Chem., Int. Ed.*, 2017, **56**, 2145–2150.
- 124 H. M. L. Davies and D. Morton, *J. Org. Chem.*, 2016, **81**, 343–350.
- 125 A. R. Bortz, J. M. Bennett and R. Fasan, *Chem*, 2024, **10**, 3488–3502.
- 126 X. Zhang, E. King-Smith and H. Renata, *Angew. Chem., Int. Ed.*, 2018, **57**, 5037–5041.
- 127 J. Li, F. Li, E. King-Smith and H. Renata, *Nat. Chem.*, 2020, **12**, 173–179.
- 128 X. Zhang, E. King-Smith, L.-B. Dong, L.-C. Yang, J. D. Rudolf, B. Shen and H. Renata, *Science*, 2020, **369**, 799–806.
- 129 F. Li and H. Renata, *J. Am. Chem. Soc.*, 2021, **143**, 18280–18286.
- 130 F. Li, H. Deng and H. Renata, *J. Am. Chem. Soc.*, 2022, **144**, 7616–7621.
- 131 J. Li, F. Chen and H. Renata, *J. Am. Chem. Soc.*, 2022, **144**, 19238–19242.
- 132 Y. Jiang and H. Renata, *Nat. Chem.*, 2024, **16**, 1531–1538.
- 133 C. N. Stout and H. Renata, *J. Am. Chem. Soc.*, 2024, **146**, 21815–21823.
- 134 F. Li, H. Deng and H. Renata, *Nat. Synth.*, 2023, **2**, 708–718.
- 135 C. Ma, C. W. Lindsley, J. Chang and B. Yu, *J. Med. Chem.*, 2024, **67**, 11459–11466.
- 136 B. D. Dherange, P. Q. Kelly, J. P. Liles, M. S. Sigman and M. D. Levin, *J. Am. Chem. Soc.*, 2021, **143**, 11337–11344.
- 137 S. Liu and X. Cheng, *Nat. Commun.*, 2022, **13**, 425.
- 138 J. C. Reisenbauer, O. Green, A. Franchino, P. Finkelstein and B. Morandi, *Science*, 2022, **377**, 1104–1109.
- 139 S. C. Patel and N. Z. Burns, *J. Am. Chem. Soc.*, 2022, **144**, 17797–17802.
- 140 E. E. Hyland, P. Q. Kelly, A. M. McKillop, B. D. Dherange and M. D. Levin, *J. Am. Chem. Soc.*, 2022, **144**, 19258–19264.
- 141 G. L. Bartholomew, F. Carpaneto and R. Sarpong, *J. Am. Chem. Soc.*, 2022, **144**, 22309–22315.
- 142 J. Wang, H. Lu, Y. He, C. Jing and H. Wei, *J. Am. Chem. Soc.*, 2022, **144**, 22433–22439.
- 143 T. J. Pearson, R. Shimazumi, J. L. Driscoll, B. D. Dherange, D.-I. Park and M. D. Levin, *Science*, 2023, **381**, 1474–1479.
- 144 J. Woo, C. Stein, A. H. Christian and M. D. Levin, *Nature*, 2023, **623**, 77–82.
- 145 Q. Cheng, D. Bhattacharya, M. Haring, H. Cao, C. Mück-Lichtenfeld and A. Studer, *Nat. Chem.*, 2024, **16**, 741–748.
- 146 J. Luo, Q. Zhou, Z. Xu, K. N. Houk and K. Zheng, *J. Am. Chem. Soc.*, 2024, **146**, 21389–21400.
- 147 F.-P. Wu, J. L. Tyler, C. G. Daniliuc and F. Glorius, *ACS Catal.*, 2024, **14**, 13343–13351.
- 148 F.-P. Wu, M. Lenz, A. Suresh, A. R. Gogoi, J. L. Tyler, C. G. Daniliuc, O. Gutierrez and F. Glorius, *Chem. Sci.*, 2024, **15**, 15205–15211.
- 149 D. Kim, J. You, D. H. Lee, H. Hong, D. Kim and Y. Park, *Science*, 2024, **386**, 99–105.
- 150 A. Conboy and M. F. Greaney, *Chem*, 2024, **10**, 1940–1949.
- 151 S. Liu, Y. Yang, Q. Song, Z. Liu, P. Sivaguru, Y. Zhang, G. de Ruiter, E. A. Anderson and X. Bi, *Nat. Commun.*, 2024, **15**, 9998.
- 152 A.-S. Paschke, Y. Brägger, B. Botlik, E. Staudinger, O. Green and B. Morandi, *ChemRxiv*, 2024, preprint, DOI: [10.26434/chemrxiv-2024-prwm8](https://doi.org/10.26434/chemrxiv-2024-prwm8).
- 153 B. Ghosh, P. Kafle, R. Mukherjee, R. Welles, D. Herndon, K. M. Nicholas, Y. Shao and I. Sharma, *Science*, 2025, **387**, 102–107.
- 154 A. Sanchez, A. Gurajapu, W. Guo, W.-Y. Kong, C. J. Laconsay, N. S. Settineri, D. J. Tantillo and T. J. Maimone, *J. Am. Chem. Soc.*, 2023, **145**, 13452–13461.
- 155 R. Fasan, J. Bennett and A. Bortz, *ChemRxiv*, 2024, preprint, DOI: [10.26434/chemrxiv-2024-9shgl](https://doi.org/10.26434/chemrxiv-2024-9shgl).

