A Natural Product Hybrid with Potent Antitumor Activity

**Significance:** Bioactive natural products are promising resources in drug development. The authors report a natural product hybrid approach by combining aplyronine A and swinholide A. By synthetically combining the structures of these two marine macrolides, the group has developed an aplyronine A–swinholide A hybrid, which showed comparably potent cytotoxicity and actin depolymerizing activity against HeLa S3 cells.

**Comment:** Synthesis of pyran A involves a Claisen condensation and an asymmetric Evans–Saksena ketone reduction. Pyran A and acetylene C were coupled using a highly diastereoselective reaction governed by a kinetically favored chair-like transition state over a twist-boat conformation. Other key reactions used in the synthesis are Takai olefination, a modified Yamaguchi esterification, and an intramolecular Nozaki–Hiyama–Takai–Kishi coupling.
Breaking the Mirror: Deracemization of α-Branched Aldehydes

Significance: Deracemizations are endergonic processes due to a negative entropy change; therefore, such phenomena demand an exogenous source of energy. Bach (Nature 2018, 564, 240), Knowles and Miller (Science 2019, 366, 364) demonstrated that light could operate as the exogenous driving force to push the racemic system out of equilibrium. In this work, the authors demonstrated a deracemization of α-stereogenic aldehydes based on a photocatalytic E/Z Isomerization of enamines formed in situ and applied this to the enantioselective synthesis of commercial anti-inflammatory drugs.

Comment: In the presence of a chiral primary aminocatalyst, E- and Z-enamines form stereoselectively from each enantiomer of an α-branched aldehyde due to facial selective deprotonation (Angew. Chem. Int. Ed. 2011, 50, 11451). In the presence of visible light (400 nm), an iridium photocatalyst converts the E-enamine into the disfavored Z isomer. The overall effect is the depletion of one of the enantiomers, which affords enantioenriched aldehydes with up to 96% ee within one hour. A modified Pinnick oxidation converts the aldehyde products into α-aryl-propionic acid derived drugs.
Harnessing the Power of Organo- and Biocatalysis to Synthesize STING Agonist MK-1454

**Significance:** The stimulator of interferon gene (STING) protein is part of the innate immune system, which is responsible for the upregulation of interferons and cytokines. There is a huge interest in agonism of STING for cancer treatment. MK-1454, a complex cyclic di-nucleotide, is a potent agonist of STING protein and is currently under clinical development.

**Comment:** The synthesis leveraged conventional organocatalysis and novel biocatalysis reactions, which are well-orchestrated into a concise and highly efficient stereoselective synthesis of MK-1454. Highlights are two stereoselective fluorination reactions using NFSI and the optimization of two biocatalysis reactions by directed evolution.

**SYNFAC T S Contributors:** Dirk Trauner, Ruiyang Bao

**Category:** Chemistry in Medicine and Biology

**Key words:** STING agonist, organocatalysis, biocatalysis, directed evolution

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**Organocatalysis reaction:** conditions are shown in blue

**Biocatalysis reaction:** conditions are shown in green

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**a) Synthesis of modified guanosine A:**

1. TBSCI, py, then TsCl, NMI
2. isobutynil chloride, py

- 63% over 2 steps

**b) Synthesis of modified adenosine B:**

1. L-Leu-NH₂·HCl, NFSI
2. evolved ketoreductase, NADPH

- 64% over 2 steps

**c) Merge of A and B:**

1. kinases, co-factors
2. cobalt-treated cGAS, ZnSO₄

- 75% over 2 steps

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**Key words:**

- STING agonist
- organocatalysis
- biocatalysis
- directed evolution

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SYNFAC T S Contributors: Dirk Trauner, Ruiyang Bao

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Unexpected Hydrolytic Triazoloquinazoline Ring Opening Leads to a Potent HDAC6 Inhibitor

**Significance:** Histone deacetylases (HDACs) play a key role in the epigenetic regulation of gene function, and their overexpression has been found in several tumor types. HDAC6 specifically has been established as a target for anticancer agents, and many structurally diverse inhibitors have been reported. Expanding on known quinazoline-capped HDAC inhibitors, the authors designed 11a and 11b and demonstrated potent in vivo HDAC inhibition of 11a with an original aminotriazoloquinazoline-based scaffold. In attempting to synthesize 11b, an unexpected quinazoline ring opening generated 18, which was also demonstrated to be a novel aminotriazole-based HDAC6 inhibitor.

**Comment:** Anthranilic anilide was converted into a benzoazainone by an intramolecular annulation, and the resulting fused heterocycle was condensed with aminoguandine A to obtain the desired tricyclic aminotriazoloquinazoline. Changing the substitution of the benzoate group to generate 11b led to hydrolytic opening of the pyrimidine ring upon ester hydrolysis, which is a reactivity that is unreported for triazoloquinazolines in the absence of carbon nucleophiles. Interestingly, the unanticipated product 18 demonstrated five-fold more potent inhibition compared to the designed product 11a, offering an additional scaffold for further HDAC inhibitor design.

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Expanding the E3 Ligase Toolbox by Co-opting Methyl Readers

**Significance:** Proteolysis targeting chimeras (PROTACs) are tools for targeted protein degradation. The authors use L3MBTL3, a methyl reader protein that binds to the E3 ligase complex, to induce nuclear-specific protein degradation of FKBP12 and BRD2.

**Comment:** The authors use UNC1215, a potent antagonist of L3MBTL3 synthesized via Buchwald–Hartwig cross-coupling, as a handle to recruit the E3 ligase complex. The L3MBTL3-recruiting PROTACs promote nuclear-specific protein degradation.
Novel Macrocycle-Based Type-II TRK Inhibitors

**Significance:** Tropomyosin receptor kinases (TRK) are considered targets for anticancer drug discovery, but mutations at the solvent-front (SF) and the DFG motif render type-I inhibitors sensitive to drug resistance. A highly selective macrocyclic TRK inhibitor with a unique type II binding mode was synthesized that was able to overcome any DFG mutation related resistance.

**Comment:** The synthesis started from the commercially available 3-iodo-2-methyl-5-nitrobenzoic acid. Conversion of the nitro group into a phenol ether, followed by two consecutive Sonogashira reactions, formed the precursor for the macrocyclic core. Acidic deprotection followed by macrolactamization completed the macrocycle.
A Rapid Approach to Cyclopropane-Fused Lactams

Significance: The medicinally attractive scaffold of cyclopropane-fused N-heterocycles finds application in many biologically active compounds. Facilitating their synthetic access, the authors developed a mild, rapid, and modular approach implementing novel bench- and thermally stable $\alpha$-diazo acyl chloride reagents. Reaction with readily available allyl amines via an acylation, intramolecular [3+2] cycloaddition, fragmentation sequence afforded cyclopropane-fused lactams.

Comment: The developed methodology tolerates a broad scope of functional groups in various positions and is suitable for the construction of complex fused ring systems. The usefulness of this transformation has been demonstrated with the efficient syntheses of the two antidepressants, amitifadine and milnacipran.
**Tuning Colors of the Drug: A Systematic Study of the Emissive Features of the Deferasirox Scaffold**

**Significance:** Deferasirox, an FDA-approved iron chelator, is shown to have anticancer and antimicrobial activities. Its core has been identified as a novel scaffold for an easy-to-visualize aggregation-induced emission platform with similar therapeutic effects as deferasirox (J. Am. Chem. Soc. 2021, 143, 1278). A broad library of deferasirox derivatives is synthesized with various fluorescent emission profiles, which shows promise in bioimaging applications.

**Comment:** The fluorescent deferasirox derivatives retain the iron-dependent antibiotic activity while permitting bioimaging in the presence of commonly used imaging agents. ExNMe₂, emitting in the blue channel with no overlap emission with green and red channels, proves useful for in vivo and in vitro imaging. The simultaneous use of several fluorescent derivatives maximizes the information gained during a single imaging experiment.
A Reversible Covalent Inhibitor Targeting the BCR-ABL Active-Site Lysine

**Significance:** Permanent activity of the BCR-ABL kinase causes certain types of leukemia such as chronic myeloid leukemia. Various noncovalent inhibitors targeting the ATP-binding site of this protein have been clinically investigated and they greatly reduce the tumor burden in patients. The leukemia cells however often develop a resistance against the treatment, which is, in many cases, caused by point mutations in the kinase. Targeted covalent inhibitors represent a possibility to overcome the resistance problem. In the highlighted publication the development of the first cell-active covalent BCR-ABL inhibitor is described based on previous work (Angew. Chem. Int. Ed. 2021, 60, 17131).

**Comment:** The synthesis of inhibitor D proceeded via functionalization of azaindole A featuring two Suzuki cross-coupling reactions with fragments B and C. Upon binding of D to the BCR-ABL protein, the salicylaldehyde moiety forms an imine (E) with the active site lysine 271, which is stabilized by a hydrogen-bond with the neighboring hydroxyl group. In vitro inhibition experiments revealed a high potency of D against the wild-type kinase as well as against various drug-resistant mutants. In comparison, the inhibitory activity of compound F, which lacks the salicylaldehyde moiety, was significantly lower. Compound D showed 50 nM cellular activities in a chronic myeloid leukemia cell line but low metabolic stability in mouse liver microsomes.
Fluorogenic Cyclopropenones for Cellular Imaging

**Significance:** Fluorogenic bioorthogonal reactions that can be used to image cellular features are highly sought after. These reactions are attractive because they ‘light up’ only in the presence of the required reagents and remove the need to wash-out unreactive probe and reagents. In the highlighted paper, the authors describe first-in-class cyclopropenone-based probes that cyclize to form fluorescent coumarins upon reaction with a bio-orthogonal phosphine. The pentafluorophenyl ester probes can be conjugated to primary amines in biomolecules. Additionally, the authors show that this reaction is compatible with strain-promoted click chemistry in cellular imaging experiments.

**Comment:** Upon reaction with a bioorthogonal phosphine, the cyclopropenone forms a ketene ylide that reacts with an intramolecular nucleophile to yield a fluorescent coumarin. Significant synthetic and design efforts were made to favor the formation of the fluorogenic coumarin product over the non-fluorescent furanone and to determine the best phosphine for the reaction. Reactions with the optimized probe molecule provided a >1600-fold increased signal turn-on response compared with the unreacted probe. The optimized probe synthesis for bioconjugation features a Sonogashira cross-coupling, followed by the installation of the cyclopropenone via a difluorocarbene insertion, hydrolysis and finally, acidic deprotection to yield the desired probe.
An Improved Orthosteric Antagonist of the M₅ Muscarinic Acetylcholine Receptor

Significance: The M₅ muscarinic acetylcholine receptor is a Gq-coupled GPCR expressed in the mesolimbic reward pathway. One hypothesis is that M₅ antagonists could mitigate the rewarding effects of opioids and thus serve as a potential treatment for opioid use disorder. A selective and potent M₅ orthosteric antagonist, ML375, was previously developed but suffered from poor pharmacokinetic properties. The authors report a new M₅ orthosteric antagonist, VU6019650, that displays favorable brain penetration to serve as a tool compound for investigating M₅ antagonism.

Comment: An efficient synthesis of VU6019650 starts with mesylation of primary alcohol A, followed by substitution with imidazole B. Deprotection of the Boc group and substitution with sulfonyl chloride C yields VU6019650. Antagonist VU6019650 has an IC₅₀ of 36 nM against human M₅ receptor and was demonstrated to inhibit oxycodone self-administration in rat models. Efforts are underway to improve the poor metabolic stability of the antagonist, likely arising from the thioether linkage.
A Diastereoselective Cyclopropanation Strategy

Significance: While cyclopropanols and cyclopropylamines are interesting synthetic equivalents, they are also important pharmacophores in medicinal chemistry. Despite the existing cyclopropane synthesis, diastereoselective cyclopropanation strategies remained less studied. Here, the authors report a Ti-catalyzed diastereoselective cyclopropanation method that couples alkenes with carboxylic acids or amides. This method has been shown to be widely compatible across a broad range of substrates.

Comment: Unlike Kulinkovich cyclopropanation, this protocol does not require stoichiometric Grignard reagent and the catalyst is regenerated by using dichlorodimethylsilane. The proposed mechanism involves the formation of a titanacyclopropane intermediate. In case of carboxylic acid esters, the diastereoselectivity is governed by the steric hindrance between the alkene substituent and the methyl groups of Cp*.

Proposed mechanism and stereodetermining T.S.:

in case of carboxylic acid/esters

in case of amides

From lithocholic acid 62% yield, dr = 10:1
From arachidic acid 55% yield, dr = 17:1
Late-stage functionalization of estrone 45% yield, dr = 5:1
Intramolecular 45% yield, dr = 20:1
From tamoxifen 81% yield, dr = 4:1
From paroxetine 69% yield, dr = 8:1
From atomoxetine 72% yield, dr = 5:1

Key words
cyclopropanol
cyclopropylamine
titanium-catalyzed cyclopropanation
titanacyclopropane

Category
Chemistry in Medicine and Biology

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The First Total Synthesis of Bryostatin 7

**Significance:** The bryostatins are a family of 21 marine natural products boasting a wide range of biological activities. They have been explored as anticancer agents, treatments for Alzheimer’s disease, and as antivirals against HIV. Of this family, the first to succumb to total synthesis was bryostatin 7 in a heroic synthesis by the Masamune group requiring a total of 79 steps with a 41 step longest linear sequence.

**Comment:** Retrosynthetically, bryostatin 7 was broken into three fragments (A, B, and D). A and B were used for a matched aldol reaction followed by Hg-mediated cyclization. Further manipulations yielded C. Fragment D (prepared in 22 steps) underwent a Julia–Lythgoe olefination with C to complete the unification of all fragments. A DCC-mediated macrocyclization was used to close the lactone.