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Introduction

The successful total synthesis of natural products depends on orchestrating an array of reactions, careful control of stereochemistry, and protection-deprotection tactics. Based on the principle of retrosynthetic analysis,¹ various strategies have been developed to plan a synthetic route by taking a complex target molecule back to simple building blocks.² Notably, chemical total synthesis has significantly benefited from the advancements of transition metal catalysis and organocatalysis over the years. As one major pillar of asymmetric catalysis, biocatalysis has emerged as a powerful tool for the sustainable production of active pharmaceutical ingredients (APIs), largely due to the development of genome mining and directed evolution.³ Biocatalysis offers some advantages over classical chemical methods, including higher efficiency and selectivity,



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Merging the advantages of biocatalysis and chemocatalysis in retrosynthetic analysis can significantly improve the efficiency and selectivity of natural product synthesis. Here, we describe a unified approach for the synthesis of drimane meroterpenoids by combining heterologous biosynthesis, enzymatic hydroxylation, and transition metal catalysis. In phase one, drimenol was produced by engineering a biosynthetic pathway in *Escherichia coli*. Cytochrome P450_{BM3} from *Bacillus megaterium* was engineered to catalyze the C-3 hydroxylation of drimenol. By means of nickel-catalyzed reductive coupling, six drimane meroterpenoids (+)-hongoquercins A and B, (+)-*ent*-chromazonarol, 8-*epi*-puupehenol, (–)-pelorol, and (–)-mycoleptodiscin A were synthesized in a concise and enantiospecific manner. This strategy offers facile access to the congeners of the drimane meroterpenoid family and lays the foundation for activity optimization.

environmental friendliness, and better safety profiles. In addition, metabolic engineering allows for streamlining multiple enzymatic reactions inside chassis cells, thus providing rapid access to structurally complex molecules.4 Despite these advantages, enzymatic reactions have not been widely used in natural product synthesis largely for two reasons. First, guidelines for merging enzyme catalysis with traditional chemical methods in synthesis design, namely, biocatalytic retrosynthesis,⁵ are underdeveloped. Second, it is highly desirable to enrich the toolbox of biocatalysts for a broad range of transformations and substrates.6 Previously, we have reported the efficient synthesis of bioactive natural products by combining enzymatic reactions with chemical methods.7 Herein, we describe a unified synthetic approach to the drimane meroterpenoids by combining metabolic engineering, enzymatic hydroxylation, and transition metal catalysis.

Meroterpenoids are a diverse family of hybrid natural products that have been isolated from plants, fungi, bacteria, and animals.⁸ Meroterpenoids possess a broad range of biological activities and supply several medicinal drugs or promising lead compounds. For example, hongoquercins A (1) and B (2) (Fig. 1A) were isolated from an unidentified terrestrial fungus by Roll and co-workers and exhibit antibacterial activities toward vancomycin-resistant *Enterococcus faecium* and methicillinresistant *Staphylococcus aureus*.⁹ (+)*-ent*-Chromazonarol (3) was isolated from marine sponge *Disidea pallescens* and shows antitumor activities.¹⁰ These compounds feature a drimane scaffold that is connected to an aromatic ring by C–C and C–O bonds, and belong to the drimane meroterpenoid family. Due to their remarkable biological activities and structural diversity,



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Fig. 1 Structure and chemical synthesis of drimane meroterpenoids. (A) Structure of hongoquercins A (1) and B (2), (+)-*ent*-chromazonarol (3), 8-*epi*-puupehenol (4), (-)-pelorol (5), and (-)-mycoleptodiscin A (6). (B) Synthesis of (-)-chromazonarol (7) was achieved by Yamamoto and co-workers *via* a biomimetic polyene cyclization catalyzed by a Lewis acid-assisted chiral Brønsted acid. (C) Synthesis of (+)-hongoquercin B (2) was achieved by Barrett and co-workers *via* a substrate-controlled epoxy-ene cyclization. (D) The chiral pool approach for the synthesis of drimane meroterpenoids by Baran, Lei, Sau, Wu, Li, and Anderson.

intensive efforts have been made to achieve the chemical synthesis of drimane meroterpenoids. Two general strategies, including the biomimetic cyclization approach and the chiral pool approach, have been developed to this end.^{11,12} For instance, Yamamoto and co-workers have achieved the enantioselective synthesis of (-)-chromazonarol (7) via a biomimetic cyclization induced by a Lewis acid-assisted chiral Brønsted acid (chiral LBA) (Fig. 1B).^{11a} This approach offers the cyclization product with good enantioselectivity (88% ee), albeit with poor diastereoselectivity (69:31 d.r.). Later, Barrett and coworkers developed a highly efficient synthesis of (+)-hongoquercin B (2) via a substrate-controlled cationic epoxy-ene tricyclization (Fig. 1C).11b,c For the chiral pool approach, the groups of Baran, Lei, Sau, Wu, Li, and Anderson respectively used (+)-sclareolide (8) as the starting material in the synthesis of (+)-hongoquercin A (1), (+)-ent-chromazonarol (3), 8-epipuupehenol (4), and (–)-pelorol (5) (Fig. 1D).¹² Notably, Renata and co-workers reported a hybrid synthetic strategy by combining the P450-catalyzed selective hydroxylation of (+)-sclareolide with radical-based transformations, which provides facile access to five C-3 oxidized drimane metroterpenoids.¹³ Despite these advances, a unified and efficient approach for the divergent synthesis of drimane meroterpenoids is highly desirable.

Results and discussion

Synthetic design

Biosynthetically, meroterpenoids are produced via a hybrid machinery that starts with the construction of a non-terpenoid moiety, followed by prenylation, epoxidation, and terpene cyclization.¹⁴ In contrast to the cyclization in the late stage during biosynthesis, we envisioned that the drimane moiety can be prepared in the early stage via heterologous biosynthesis, followed by connecting to the aromatics by transition metalcatalyzed C-C bond formation. Therefore, the target molecules, such as 1-4, are retrosynthetically disconnected from the drimane fragments 9 and 10 with their cognate aromatics (Fig. 2). Compounds 9 and 10 can be traced back to drimenol (11) and 3-(OH)-drimenol (12), respectively. To introduce the C-3 hydroxyl group, we planned to engineer a cytochrome P450 for the selective oxidation of drimenol. Finally, we planned to prepare drimenol by engineering a biosynthetic pathway in Escherichia coli. This unified approach harnesses enzymatic terpene cyclization and C-H hydroxylation to construct the drimane fragments on the gram scale, which otherwise has to be achieved via multi-step chemical synthesis. Moreover, it offers a concise and divergent approach to the congeners in the drimane meroterpenoid family for evaluating their biological activities.

Heterologous synthesis of drimenol

To develop a strain for microbial production of drimenol, we first examined the drimenol synthases from four different species, including PhDS, VoDS, AsDMS, and DrtB.¹⁵ PhDS from *Persicaria hydropiper* and VoDS from *Valeriana officinalis* belong to plant sesquiterpene synthases; AsDMS and DrtB were identified from *Aquimarina spongiae* and *Aspergillus calidoustus*,



Fig. 2 The synthetic strategy for drimane meroterpenoids.

respectively. By using our reported method, the *in vivo* activities of these synthases were evaluated in *E. coli* (Fig. S1†). As expected, both PhDS and VoDS provided drimenol after 72 hours of fermentation at a titer of 276 mg L⁻¹ and 371 mg L⁻¹, respectively (Fig. 3A). To our delight, DrtB afforded the highest titer (1.5 g L^{-1}), which was 5.6-fold relative to PhDS. The specific yield was even 8.6-fold that of PhDS. The products were purified to verify their absolute configuration. We also examined the Cterminal truncated form of DrtB (53 amino acid residues), which was reported to be the only effectively expressible form rather than the full-length protein in *E. coli*. However, it exhibited a lower titer than the full-length DrtB.

To improve the production of drimenol, metabolic engineering was employed to balance host cell growth and carbon flux into terpene production. It has been suggested that isopentenyl pyrophosphate isomerase (IDI) and HMG-CoA reductase (HMGR) in the mevalonate (MVA) pathway are the key ratelimiting enzymes in that isopentenyl pyrophosphate (IPP) and HMG-CoA cause cell growth inhibition.¹⁶ Therefore, we overexpressed a second copy of EcIDI and mvaA gene individually and simultaneously in the DrtB strain. The highest titer and specific yield were achieved by combination of these two rate-limiting enzymes with 30% and 50% increases, respectively.

We next attempted to further improve the titer of drimenol by the multienzyme assembly strategy.¹⁷ Recently, Xia and coworkers have utilized orthogonal protein reactions (SnoopTag/ SnoopCatcher and SpyTag/SpyCatcher pair) to covalently assemble three key enzymes in the mevalonate/terpene biosynthetic pathway.¹⁸ The Jiang group have also developed a modified Tag/Catcher system (NGTag/NGCatcher pair) by integrating covalent bonds and noncovalent interactions for highly efficient protein self-assembly.¹⁹ Both methods have been successfully used to improve terpenoid production in *E. coli*. Thus, we designed (1) a top MVA pathway assembly (AtoB– mvaS–mvaA three enzymes assembly, namely T-ab) by the SnoopTag/SnoopCatcher and SpyTag/SpyCatcher pair; and (2) cross-linking of the MVA-drimenol pathway (ERG20–EcIDI– DrtB three enzyme assembly, namely C-ab) by the NGTag/ NGCatcher pair and peptide interaction *via* a caveolar membrane scaffold (Fig. S2†). Unfortunately, neither assembly strategy afforded a higher titer of drimenol. In contrast, the titer decreased dramatically by means of the second approach. Last, we compared the impact of the *E. coli* strain on the drimenol titer.

As shown in Fig. 3B, the MG1655(DE3) strain harboring DrtB afforded the highest titer (2.1 g L^{-1}), which was 1.3-fold relative to BL21(DE3). In contrast, the titer and specific yield of drimenol from MG1655(DE3) harboring DrtB with another copy of EcIDI and mvaA are lower than those of BL21(DE3). Therefore, we used the MG1655(DE3) strain harboring DrtB for drimenol production.

Engineering of P450_{BM3} for selective oxidation of drimenol

With a few grams of drimenol in hand, we carried out biocatalytic oxidation of drimenol by engineering $P450_{BM3}$ (CYP102A1) from *Bacillus megaterium*. $P450_{BM3}$ is highly expressible in *E. coli* and has one of the highest CYP turnover rates among the reported cytochrome $P450s.^{20}$ Several groups hence have engineered $P450_{BM3}$ to catalyze the regio- and stereoselective C–H hydroxylation of cyclic substrates by means of directed evolution.²¹ However, to the best of our knowledge, selective oxidation of drimenol by this enzyme has not been



Fig. 3 Heterologous biosynthesis of drimenol in *E. coli*. (A) The BL21(DE3) strain harboring MVA pathway genes, FPPS gene (ERG20), and drimenol synthase gene from 4 species (yellow background); additional copy of EcIDI and mvaA (blue background) and multienzyme assembly containing the AtoB-mvaS-mvaA assembly (T-ab) and ERG20-EcIDI-DrtB assembly (C-ab) (purple background) based on a DrtB over-expressing strain. (B) Comparison of drimenol production between BL21(DE3) and MG1655(DE3) strain which harbored comparative good construct in terms of species of drimenol synthase (DrtB), rate-limiting enzyme of metabolic pathway (EcIDI & mvaA) and multienzyme assembly (T-ab), respectively, and PhDS as control. The data represent the averages \pm standard deviations of three independent colonies.



Fig. 4 C3-hydroxylation of drimenol was catalyzed by P450_{BM3} variants. (A) Purified P450_{BM3} (F87A) catalyzed C–H hydroxylation and C–C epoxidation of 1 mM and 2 mM drimenol (yellow background) to produce compounds **12** (blue background) and **13** (purple background). (B) X-ray crystallography of 3-(OH)-drimenol (**12**). (C) Docking model of drimenol (cyan) in the active site of P450_{BM3} (F87A) (green). Several residues (dark blue) show hydrophobic interactions (gray line) with drimenol. (D) Heat map of the yield of compound **12** by screening the P450_{BM3} L75/F87 mutant library. The yield was determined by GC-MS analysis. (E) Whole cell catalysis (yellow background) and cell lysate catalysis (blue background) by P450_{BM3} (F87A, L75A/F87I, L75G/F87I). (F) The gram-scale oxidation of drimenol with P450_{BM3} (L75A/F87I).

reported thus far. To this end, we first investigated this reaction using the purified $P450_{BM3}$ (F87A) (Fig. 4A). To our delight, the desired C3-hydroxylation product **12** was obtained as the main product (Fig. 4B). Its structure was confirmed by 2D NMR and Xray crystallography (CCDC 2307775, Table S3†). Interestingly, a side product **13** was detected and structurally characterized. We reasoned that **12** was produced through selective epoxidation of compound **12**, which was proved by an *in vitro* enzymatic assay of P450_{BM3} (F87A) using compound **12** as the substrate (Fig. S3†).

To avoid overoxidation and improve the yield of compound 12, we designed and constructed a focused mutagenesis library of P450_{BM3}. Based on the docking model of drimenol in the active site of P450_{BM3} (F87A), we selected the positions of F87 and L75 for site-directed mutagenesis (Fig. 4C). NNK codon degeneracy that encodes all 20 amino acids has been widely used for saturation mutagenesis. The screening of ~ 3000 transformants is required to achieve 95% library coverage of a two-residue mutagenesis library. Reetz and co-workers reported the use of NDC codon degeneracy encoding only 12 amino acids to reduce the transformant number to 430.21b Based on the structure of drimenol, we envisaged that mutation of the key residues in the active site to hydrophobic amino acids or those capable of forming a hydrogen bond would improve enzyme-substrate interactions. Therefore, F87 and L75 were mutated to nine amino acids (Ala, Phe, Gly, Ile, Leu, Met, Ser, Thr, Val) to generate a focused library containing 81 mutants (Fig. 4D). After screening, 6 mutants showed higher activity than P450_{BM3} (F87A). Two mutants, P450_{BM3} (L75A/F87I) and P450_{BM3} (L75G/F87I), were further validated with whole-cell catalysis and cell lysate. As shown in Fig. 4E, both variants showed higher conversions of drimenol and higher yields of compound 12 than P450_{BM3} (F87A). Previously, Renata and coworkers reported that an engineered P450_{BM3} variant MERO1 could efficiently transform sclareolide into 3-hydroxysclareolide.¹³ MERO1 contains 10 mutations at positions that do not interact directly with the substrate. Considering these mutations might have an impact on the catalytic activity, we introduced them into P450_{BM3} (L75A/F87I). Unfortunately, no increase in the yield of compound **12** was observed (Fig. S4A†). We next optimized the reaction conditions by changing the percentage of the co-solvent. We discovered that in the presence of 5% DMF, a GC yield of 80% was afforded at an analytical scale (Fig. S4B and C†). The gram-scale oxidation of drimenol with P450_{BM3} (L75A/F87I) was performed under this condition, and afforded compound **12** in an isolated yield of 68% (Fig. 4F).

Synthesis of meroterpenoid natural products

We firstly targeted natural products (+)-ent-chromazonarol (3), 8-epi-puupehenol (4), (-)-pelorol (5), and (-)-mycoleptodiscin (6) (Scheme 1). As shown in Table 1, Ni-catalyzed reductive cross-coupling between drimane halides and aryl halides was evaluated.22 According to Weix's seminal contribution of Nicatalyzed reductive coupling of alkyl halides with aryl halides, we initially tested the cross-coupling of two iodides under similar conditions, with pyridine as an effective additive to inhibit β -H elimination (Table 1, entry 1). However, the desired product 15 was only obtained in 6% yield. The elimination product B was found to be the major side-product, accompanied with the reduced aryl compound C and the homocoupling product D. We then used drimanyl bromide which is less prone to elimination in the reductive coupling (Table 1, entry 2). To our delight, the yield of the elimination product B was significantly decreased, while the yield of the desired product 15 was increased to 40%. As a comparison, the cross-coupling of drimanyl iodide with the aryl bromide afforded compound 15 only



Scheme 1 Synthesis of (+)-*ent*-chromazonarol (3), (+)-8-*epi*-puupehenol (4), (–)-pelorol (5), and (–)-mycoleptodiscin A (6). Reagents and conditions: (a) CBr₄, PPh₃, THF, 0 °C, 80%; (b) Nil₂ (10 mol%), dtbpy (5 mol%), dppbe (5 mol%), Co^{II}(Pc) (5 mol%), Mn, pyridine, DMPU, 55 °C, 53%; (c) HCl, iPrOH, 55 °C, 95%; (d) Pt-catalyst (3 mol%), AgOTf (6 mol%), DCE, rt, 90%; (e) Nil₂ (10 mol%), dtbpy (5 mol%), Co^{II}(Pc) (5 mol%), Co^{II}(Pc) (5 mol%), Mn, pyridine, DMPU, 48%; (f) K₂CO₃, MeOH, rt, 95%; (g) Pt-catalyst (3 mol%), AgOTf (6 mol%), DCE, rt, 90%; (h) Pd/C, H₂, MeOH, rt, 95%. (i) Nil₂ (10 mol%), dtbpy (5 mol%), dppbe (5 mol%), Co^{II}(Pc) (5 mol%), Mn, pyridine, DMPU, 55 °C, 57%; (j) TMSOTf, DCM, 0 °C, 73%; (k) Nil₂ (10 mol%), dtbpy (5 mol%), Co^{II}(Pc) (5 mol%), Mn, pyridine, DMPU, 55 °C, 45%; (l) TMSOTf, DCM, 0 °C, 70%. MOM = methoxymethyl, DCE = 1,2-dichloroethane, TMSOTf = trimethylsilyl triflate.

in 3% yield with a 75% yield of compound **B** (Table 1, entry 3). Interestingly, when both the drimanyl bromide and the aryl bromide were used, compound 15 was obtained in 49% yield (Table 1, entry 4). To further increase the yield of 15, $Co^{II}(Pc)$ was added as a cocatalyst to promote alkyl radical formation for the subsequent reductive cross-coupling.22d-g Pleasingly, the yield of 15 was increased to 57% (Table 1, entry 5). The aryl iodide was also tested under the Ni/Co cocatalytic system, and the yield of 15 was further increased to 62%. The use of other reducing agents in place of Mn(0), such as Zn(0) or 1,1,2,2,tetrakis(dimethylamino)ethylene (TDAE), still produces an appreciable amount of the eliminated product B, respectively (Table 1, entries 7 and 8). Thus, under the optimized conditions in Table 1, entry 5, compounds 15, 18, 22, and 25 (Scheme 1) were obtained successfully in 57%, 48%, 57% and 45% yields, respectively. Deprotection of the MOM group of 15 and the acetyl group of 18 afforded intermediates 16 and 19, respectively

(Scheme 1A). Under the catalysis of the platinum catalyst and silver triflate,²³ **16** was transformed into (+)-*ent*-chromazonarol (**3**) in 90% yield as a single diastereomer. **19** was subjected to the same conditions and compound **20** was provided in 90% yield. After deprotection of the benzyl group of **20**, (+)-8-*epi*-puupehenol (**4**) was obtained in 95% yield. Hence, (+)-*ent*-chromazonarol (**3**) and (+)-8-*epi*-puupehenol (**4**) were efficiently synthesized over 4 steps (36% overall yield) and 5 steps (31% overall yield), respectively. For the synthesis of (-)-pelorol (**5**) and (-)-mycoleptodiscin A (**6**), the corresponding reductive coupling products **22** and **25** respectively underwent TMSOTf-mediated hydroarylation reactions to provide the core structures **23** and **26**, which can be converted to the natural products (-)-pelorol (**5**) and (-)-mycoleptodiscin A (**6**) according to the literature.^{12f;24}

We next investigated the synthesis of hongoquercins A (1) and B (2) (Scheme 2). We prepared compound 10 from 3-(OH)-

Table 1 Optimized reaction conditions for the Ni-catalyzed reductive coupling to assemble compound 15^a



^{*a*} dtbpy = 4,4-di-*tert*-butyl bipyridine, dppbe = 1,2-bis(diphenylphosphino)benzene, DMPU = 1,3-dimethyl-3,4,5,6-tetrahydro-2(1*H*)-pyrimidinone, TDAE = tetrakis(dimethylamino)ethylene. The yields of isolated products are given. ^{*b*} Yields of **15** and **B** were calculated based on the drimane moiety. ^{*c*} Yields of **C** and **D** were calculated based on the aryl halide moiety.



Scheme 2 Synthesis of hongoquercins A (1) and B (2). Reagents and conditions: (a) TBSCl, imidazole, DMF, rt, 90%; (b) NaH, BnBr, DMF, rt; (c) TBAF, THF, 60 °C, 85% over 2 steps; (d) CBr₄, PPh₃, THF, 0 °C, 62%; (e) Nil₂ (10 mol%), dtbpy (5 mol%), dppbe (5 mol%), Co^{II}(Pc) (5 mol%), Mn, pyridine, DMPU, 55 °C, 48% for **31**, 48% for **32**; (f) HCl, iPrOH, 65 °C, 89%; (g) Pt-catalyst (3 mol%), AgOTf (6 mol%), DCE, rt, 80%; (h) LiOH, MeOH/ H₂O, reflux, 95%; (i) Pd(OAc)₂, L, CsOAc, H₂O₂ (35% aq.), DMA, 60 °C, 65%; (j) Pd/C, H₂, MeOH, rt; (k) Ac₂O, pyridine, rt, 80% over 2 steps; (l) K₂CO₃, MeOH, rt, 90%. TBS = *tert*-butyldimethylsilyl, TBAF = tetrabutylammonium fluoride, DMA = *N*,*N*-dimethylacetamide.

drimenol (12) via four steps (Scheme 2A). The primary hydroxyl group was selectively protected with a TBS group in 90% yield. After benzylation of the C-3 hydroxyl, TBS was deprotected to give intermediate 28 in 85% yield over two steps. In the presence of CBr₄ and PPh₃, compound 28 was converted into bromide 10 in 62% yield. With compounds 9 and 10 in hand, we first tested the cross-coupling reaction with iodide 29. Unfortunately, no desired cross-coupling product was obtained under the same conditions, possibly due to the steric hindrance of the aromatic fragment. Therefore, we decided to introduce the hydroxyl group in the later stage by the palladium-catalyzed C-H oxidation developed by Yu and co-workers.25 Towards this end, compounds 9 and 10 were coupled with iodide 30 to afford intermediates 31 and 32 in 48% and 48% yields, respectively (Scheme 2B). After deprotection of the MOM group under acidic conditions, both intermediates were subjected to platinumcatalyzed hydroalkoxylation.23 Compounds 33 and 34 were obtained in 80% and 85% yield, respectively. The methyl esters of 33 and 34 were hydrolyzed by lithium hydroxide to provide 35 and 36. The resulting products underwent palladium-catalyzed C-H oxidation to provide (+)-hongoquercin A (1) and compound 37 successfully. To complete the synthesis of (+)-hongoquercin B (2), compound 37 was debenzylated with Pd/C and hydrogen. After protection and selective deprotection, (+)-hongoquercin B (2) was afforded.

Conclusion

Remarkable progress has been achieved in the fields of chemoand biocatalysis over the past few decades. However, harnessing the benefits of both technologies in natural product synthesis is underexplored. To this end, biocatalytic retrosynthesis has been proposed to aid synthetic chemists in designing synthetic routes with enzyme-catalyzed reactions.5 Based on this concept, we demonstrated that metabolic engineering, enzymatic hydroxylation, and transition metal catalysis can be orchestrated in natural product synthesis. Our strategy harnesses the biosynthetic machinery of terpenoids for rapid construction of the drimane scaffold. By using an engineered P450_{BM3} variant, the hydroxyl group was site- and stereoselectively introduced at the C-3 position of drimenol. The drimane and aromatic fragments were assembled by transition metal-catalyzed C-C and C-O bond formation reactions. Six drimane meroterpenoids, (+)-hongoquercins A (1) and B (2), (+)-ent-chromazonarol (3), 8epi-puupehenol (4), (-)-pelorol (5) and (-)-mycoleptodiscin A (6), were synthesized in 4-12 chemical steps. We envisioned that this hybrid approach can also be applied to the synthesis of other bioactive terpenoid natural products and their analogues.

Data availability

All experimental procedures and associated data are provided in the ESI.† Crystallographic data for compound **12** has been deposited at the CCDC under 2307775. For ESI and crystallographic data in CIF see DOI: https://doi.org/10.1039/ d4sc06060a.

Author contributions

C. X. and Z. X. supervised the project and wrote the manuscript; Y. Y. performed the chemical synthesis experiments; X. Z., W. X. and T. K. performed the biosynthesis experiments.

Conflicts of interest

There are no conflicts to declare.

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