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# Directed Evolution of a Cytochrome P450 Carbene Transferase for Selective Functionalization of Cyclic Compounds

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KEYWORDS Biocatalysis, Cytochrome P450, Carbene Transfer, Indole Alkylation, Cyclopropanation.

**ABSTRACT:** Transfers of carbene moieties to heterocycles or cyclic alkenes to obtain  $C(sp^2)$ –H alkylation or cyclopropane products are valuable transformations for synthesis of pharmacophores and chemical building blocks. Through their readily tunable active site geometries, hemoprotein "carbene transferases" could provide an alternative to traditional transition metal catalysts by enabling heterocycle functionalizations with high chemo-, regio- and stereocontrol. However, carbene transferases accepting heterocyclic substrates are scarce; the few enzymes capable of heterocycle or cyclic internal alkene functionalization described to date are characterized by low turnovers or depend on artificially introduced, costly iridium–porphyrin cofactors. We addressed this challenge by evolving a cytochrome P450 for highly efficient carbene transfer to indoles, pyrroles, and cyclic alkenes. We first developed a spectrophotometric high-throughput screening assay based on 1-methylindole  $C_3$ -alkylation that enabled rapid analysis of thousands of P450 variants and comprehensive directed evolution via random and targeted mutagenesis. This effort yielded a P450 variant with 11 amino acid substitutions and a large deletion of the non-catalytic P450 reductase domain, which chemoselectively  $C_3$ -alkylates indoles with up to 470 turnovers per minute and 18,000 total turnovers. We subsequently used this optimized alkylation variant for parallel evolution towards more challenging heterocycle carbene functionalizations, including  $C_2/C_3$  regioselective pyrrole alkylation, enantioselective indole alkylation with ethyl-2-diazopropanoate, and cyclic internal alkene cyclopropanation. The resulting set of efficient biocatalysts showcases the tunability of hemoproteins for highly selective functionalization of cyclic targets and the power of directed evolution to enhance the scope of new-to-nature enzyme catalysts.

## Introduction

Cyclic compounds, particularly heterocycles, are key components in biologically active compounds, from the building blocks of life (e.g. nucleic acids, amino acids and carbohydrates) to natural products with pharmacologically relevant activities (e.g. vitamins, alkaloids and antibiotics). Accordingly, heterocycles have been playing an increasingly significant role in the modern repertoire of medicinal chemistry,<sup>1</sup> with molecules containing at least one heterocyclic fragment comprising nearly 60 % of FDA-approved pharmaceuticals.<sup>2</sup> Because heterocycles are powerful handles to regulate polarity, lipophilicity and hydrogen bonding properties, extensive efforts are directed at tuning these structures to manipulate the pharmacological, pharmacokinetic and toxicological features of drug candidates.<sup>3-5</sup> Therefore, development of selective and efficient methods for functionalization of heterocycles will streamline synthesis of medicinal compounds and pharmacophores.

51 One major way to derivatize heterocyclic fragments is to di-52 rectly functionalize C-H bonds with new chemical entities.<sup>6</sup> Al-53 ternatively, one can derivatize ring structures by functionalizing existing C-C bonds<sup>7</sup>. Nature employs different enzymatic ma-54 chineries to accomplish these two processes. For instance, me-55 thyltransferases install methyl moieties on the C-H bonds of 56 DNA bases;8 flavin-dependent halogenases are responsible for 57 installing halogens on heterocycle-based natural products, such 58

as indoles, pyrroles and azoles;<sup>9</sup> and (mono)oxygenases like cytochrome P450s or flavoenzymes are involved in oxidation of heterocyclic substrates to induce or lose aromaticity.<sup>10–12</sup>

To expand nature's capacity for functionalization of (hetero)cyclic molecules, we wished to use recently developed "carbene transferase" enzymes to functionalize carbon–hydrogen and carbon–carbon double bonds in heterocycles. Carbene transferases are engineered heme-dependent proteins capable of transferring carbene moieties to different target compounds.<sup>13</sup> Reactivities developed in recent years include carbene transfer to olefins,<sup>14–16</sup> alkynes,<sup>17</sup> and X–H bonds (X = C, N, S, Si, B)<sup>18–23</sup> to generate chemical structures often not found in nature. These enzymatic transformations grant biocatalytic access to important pharmaceuticals including levomilnacipran, ticagrelor, tasimelteon, tranylcypromine and others.<sup>24–26</sup>

However, enzymatic carbene transfer for heterocycle derivatization faces several challenges. Carbene transfer has been achieved mostly for sterically unhindered functional groups, such as terminal olefins. Thus it remains to be determined whether sterically more demanding, internal carbon–carbon double bonds in heterocycles can be efficiently functionalized. A second challenge for enzymatic carbene transfer to heterocycles is selectivity. Synthetic chemists have used transition metal catalysts based on Rh, Ru, Pd, Cu and others to selectively transfer carbenes to heteroarenes, such as indoles and pyrroles, yielding either cyclopropanes or  $C(sp^2)$ –H alkylation products.<sup>27-32</sup> Mimicking this activity would require enzymes to have exquisite control over chemo-, regio- and stereo-selectivity. Here, we set out to direct the evolution of enzyme catalysts with high activity and selectivity in heterocycle carbene functionalization reactions.

Towards this goal, we used a two-stage engineering approach. We first evolved a cytochrome P450 variant for  $C_3$  alkylation of indoles, using a high-throughput spectrophotometric screen that allowed us to evaluate several thousand enzyme variants per generation. This enabled a mutagenesis strategy targeting the entire P450 protein sequence, which delivered a variant containing 11 mutations as well as a truncation of a large part of the non-catalytic reductase domain and which chemoselectively and efficiently alkylates indoles. Hypothesizing that this enzyme, evolved for alkylation of a heterocyclic substrate, would provide a launch pad for further exploration of challenging heterocycle functionalizations, in the second stage we performed parallel directed evolution to engineer a set of P450 catalysts capable of regio-divergent pyrrole alkylations, enantioselective indole alkylations with  $\alpha$ -disubstituted carbenes, and stereoselective cyclic alkene cyclopropanations.

#### **Results and Discussion**

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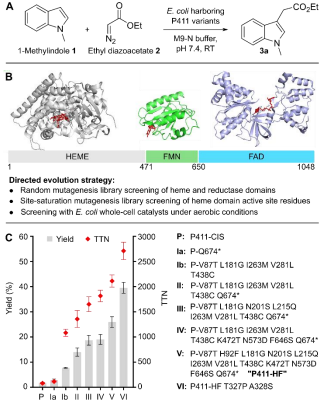
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# Leveraging a high-throughput screening assay to evolve a cytochrome P450 for indole alkylation

We began by testing a range of hemoproteins for carbene transfer (using ethyl diazoacetate (EDA) as carbene precursor) to a selection of heterocyclic substrates (Supplementary Table 1). Cytochrome P450-BM3 variant P411-CIS, a serine-ligated variant previously evolved for styrene cyclopropanation,<sup>14</sup> stood out as highly active for  $C_3$ -alkylation of 1-methylindole, catalyzing this reaction with 14% yield and 690 TTN. To our delight, we found that the resulting alkylated product **3a** showed a pronounced red-shift in UV absorbance compared to 1-methylindole. This difference in absorbance properties allowed us to develop a simple UV-Vis spectrophotometric screening assay, in which the catalytic activities of enzyme variants for the alkylation reaction (performed in whole *Escherichia coli* (*E. coli*) cells) could be measured by UV absorbance at 305 nm in a 96-well plate reader (Figure 1A, Supplementary Figure 1).

39 P411-CIS is a 1048-residue protein consisting of a heme domain containing the active site (residues 1-470) and a reductase 40 domain that shuttles electrons from NADPH to the heme do-41 main and is essential for P450-BM3's native oxo-transfer chem-42 istry (residues 471-1048, subdivided into FMN and FAD do-43 mains; Figure 1B).<sup>35</sup> Having a screen that enables analysis of 44 several thousand enzyme variants, we introduced mutations 45 throughout the heme and reductase domains using both random 46 mutagenesis and site-saturation mutagenesis and screened the 47 resulting libraries for improved 1-methylindole alkylation ac-48 tivity (Supplementary Table 2).

49 After six rounds of directed evolution for formation of **3a**, of 50 which the last four rounds were conducted under aerobic con-51 ditions, we obtained a P411 variant we refer to as P411-HF 52 (Heterocycle Functionalization) (Figure 1C). This variant contains eight mutations in the heme domain, three in the FMN do-53 main and a stop codon at the end of the FMN domain  $(Q674^*)$ , 54 which deletes the entire FAD domain (Figure 1B, Supplemen-55 tary Figure 2). Of these mutations, five were introduced by site-56 saturation mutagenesis (V87T, H92F, L181G, I263M, T438C), 57



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**Figure 1: Evolution of P411-CIS for 1-methylindole** *C*<sup>3</sup> **alkylation.** (A) Screening reaction used to evolve P411 variants for higher indole alkylation activity. (B) Domain structure of P450-BM3. As a crystal structure of the entire P450-BM3 protein is lacking, individual domain structures with approximate domain boundaries are shown (pdb entries: heme domain 4h23<sup>14</sup>; FMN domain 1bvy<sup>33</sup>; FAD domain 4dqk<sup>34</sup>); co-factors are shown in red. (C) Evolutionary lineage from P411-CIS to P411-HF T327P A328S. Data were obtained from two independent assays with six replicates per variant, run in 96-well plates under aerobic conditions with 10 mM **1** and 15 mM **2** at room temperature (RT) for 16h.

and seven (N201S, L215Q, V281L, K472T, N573D, F646S, Q674\*) were introduced by random mutagenesis. A final round of double site-saturation mutagenesis library screening on P411-HF revealed mutations T327P and A328S. This enzyme delivered the alkylated product with up to 2700 TTN in aerobic 96-well plate screening reactions (Figure 1C).

#### P411-HF: A catalyst for chemoselective alkylation of unprotected indoles

P411-HF is a highly active alkylation enzyme: analytical whole-cell reactions with P411-HF under anaerobic conditions with 30 mM 1-methylindole and 1.5 equivalents of EDA delivered alkylation product **3a** in 95% yield with almost 12,000 TTN (Figure 2A, entry 2). Under catalyst-limiting conditions, P411-HF delivered more than 18,000 TTN (Figure 2A, entry 4). Under aerobic conditions, variant P411-HF T327P A328S proved superior to P411-HF and delivered the alkylated product in 74% yield and with 1670 TTN (Figure 2A, entry 6; P411-HF delivered **3a** in 46% yield under these conditions). Thus, even with four rounds of directed evolution under aerobic conditions,

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P411-HF is about an order of magnitude more effective under anaerobic conditions. Furthermore, P411-HF delivers almost two orders of magnitude higher TTN than recently reported variants of the hemoproteins Mb or YfeX catalyzing indole alkylations (170 to 240 TTN),<sup>36,37</sup> and up to three orders of magnitude higher TTN than Rh, Cu, Pd or Fe-based transition metal catalysts reported for the same type of reaction. 30,31,38-40

To investigate catalyst lifetime, we tested sequential additions of 10 mM 1 and 2 substrate equivalents to whole-cell reactions in 30-min intervals. This did not increase product formation, which indicates that the enzyme is inactivated during the reac-10 tion (Supplementary Figure 3A). Furthermore, a 10-minute 11 EDA pre-incubation of P411-HF whole-cell catalyst resulted in 12 a 26% loss in catalytic activity (Supplementary Figure 3B). Pre-13 sumably, alkylation of enzyme residues and/or the porphyrin 14 cofactor limits enzyme lifetime, which may possibly be ad-15 dressed by further directed evolution.41

16 P411-HF anaerobic whole-cell reactions reached completion af-17 ter 2 h. The high initial turnover frequency (TOF) of 470 turn-18 overs min<sup>-1</sup> represents a 35-fold increase over the parent variant P411-CIS (Figure 2B). Under aerobic conditions, P411-HF 19 T327P A328S showed an initial rate of 21 turnovers min<sup>-1</sup>. com-20 pared to 3 turnovers min<sup>-1</sup> for P411-CIS (Figure 2C). The initial 21 rates of P411-HF compare well to previously disclosed Mb var-22 iants, which exhibited ~10 turnovers min<sup>-1</sup> in the alkylation of 23 indole with EDA under anaerobic conditions.36 24

The most striking mutation found during the directed evolution 25 was the truncation of the FAD domain; introduction of the stop 26 codon by Q674\* reduces the size of the protein catalyst by 35% 27 (from 1048 to 673 residues) and increases reaction yield by ca. 28 1.7-fold, driven primarily by an increase in protein expression 29 level (ca. 1.6-fold) with a slight increase in TTN (Figure 1B, 30 Supplementary Figure 4A). Comparable positive effects of de-31 leting the FAD domain have subsequently been observed for other P411-catalyzed carbene transfer reactions: FAD trunca-32 tion discovered in the course of this study proved beneficial for 33 carbene C(sp<sup>3</sup>)-H insertion<sup>18</sup> as well as cyclopropanation reac-34 tions (Supplementary Figure 4B).<sup>42</sup> Notably, FAD domain de-35 letion significantly destabilized the protein, as whole-cell alkyl-36 ation assays performed with a protein denaturant pre-treatment 37 revealed up to 33% lower activity of P411-CIS Q674\* com-38 pared to P411-CIS (Supplementary Figure 4C).43,44 Thus, FAD 39 domain deletion exerts a destabilizing effect, but contributes to 40 higher P411 expression levels and carbene transfer activity. 41 Mounting evidence suggests that native P450-BM3 adopts a ho-42 modimeric quaternary structure in solution via FAD domain interactions, and that the dimeric form is required for oxo-transfer 43 activity.<sup>34,45–48</sup> The data presented here suggest that for non-nat-44 ural carbene-transfer reactions a different, possibly monomeric, 45 catalyst structure may be suitable or even beneficial. Attempts 46 to obtain crystal structures of the FAD-truncated P411-HF var-47 iant have unfortunately not been successful. 48

Next, we compared 1-methylindole alkylation activities of 49 P411-CIS and P411-HF as purified proteins and with different 50 reducing agents (Supplementary Figure 5). We found that puri-51 fied P411-CIS catalyzes 1-methylindole alkylation with 165 52 TTN with sodium dithionite as reductant, 135 TTN with 53 NADPH as reductant, but only 8 TTN in the absence of reduct-54 ant. To our surprise, P411-HF showed 1890 TTN with NADPH (corresponding to 94% yield), 1750 TTN with sodium dithio-55 nite, and an almost equally high 1670 TTN without any reduct-56 ant present. Thus, P411-HF maintains a catalytically competent 57

Α	Entry	Catalyst	Conditions	1 (mM)	<b>2</b> (mM)	Yield <b>3a</b> (%)	TTN
	1	P411-CIS	Anaerobic OD <sub>600</sub> =30	30	45	13	1660
	2	P411-HF	Anaerobic OD <sub>600</sub> =30	30	45	95	11900
	3	P411-CIS	Anaerobic OD <sub>600</sub> =10	30	45	5	2010
	4	P411-HF	Anaerobic OD <sub>600</sub> =10	30	45	51	18700
	5	P411-CIS	Aerobic OD <sub>600</sub> =30	10	15	4	140
	6	P411-HF T327P A328S	Aerobic OD <sub>600</sub> =30	10	15	74	1670

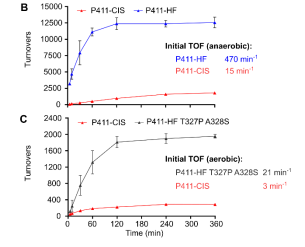
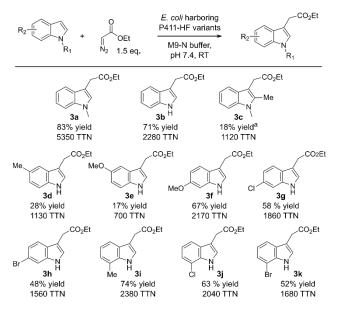


Figure 2: Characterization of P411-HF. (A) Comparison of P411-CIS and P411-HF in analytical (400 µL) biocatalytic wholecell reactions for formation of 3a under aerobic or anaerobic conditions and different substrate and catalyst loadings. (B,C) Kinetics of whole cell-reactions for formation of 3a under anaerobic (B) and aerobic (C) conditions. Reactions were run with cells at OD<sub>600</sub>=30 with 30 mM 1 and 45 mM 2 (anaerobic, B), or 10 mM 1 and 15 mM 2 (aerobic, C). Initial turnover frequencies were calculated from samples analyzed 10 minutes after starting the reaction. Data in (A) to (C) were obtained from two to three independent assays with two replicates per variant.

state of the heme cofactor even in the absence of reductant, in stark contrast to P411-CIS. While native P450 oxo-transfer chemistry requires a stoichiometric supply of electrons, P411 carbene transfer catalysis functions without an electron supply as long as the iron cofactor is in the ferrous state.<sup>14</sup> Therefore. truncating the reductase domain can be expected to be less consequential for carbene transfer catalysis. The extent to which FAD domain truncation contributes to maintaining the enzyme in the catalytically active ferrous state remains to be determined.

Lastly, we tested the substrate scope of P411-HF and its suitability for synthesis of alkylated indoles. With 1-methylindole, preparative-scale reactions (0.6 mmol) delivered 3a with 83 % isolated yield (108 mg) (Figure 3). To our delight, we found that P411-HF selectively alkylates unprotected indoles at the C<sub>3</sub> position. No N-H insertion products were observed, and alkylated products were isolated in good yields across a range of substituted, unprotected indoles (Figure 3). We observed a clear reactivity trend, with only trace amount of alkylation product obtained for  $C_4$ -substituted indoles, comparatively low yields for  $C_5$ -substituted indoles, and good yields for indoles with substitutions at  $C_6$  or  $C_7$ . Presumably, substitutions at  $C_4$  or  $C_5$  interfere with productive indole binding in the active site. Also, only low activity was observed for 1,2-dimethylindole; alkylation of this substrate was improved by the M263A mutation, which likely creates more space in the active site above the heme co-factor and enables productive binding of this sterically hindered 1,2-disubstituted indole.



**Figure 3: P411-HF indole alkylation substrate scope.** Preparative-scale reactions were performed at 0.6 mmol scale (1-methylindole) or 0.3 mmol scale (all other substrates) to yield the *C*<sub>3</sub>-alkylated products. Isolated yields are given; TTN are based on the isolated yield. <sup>a</sup>For 1,2-dimethylindole, reactions were performed with variant P411-HF M263A and HPLC yield and TTN are given.

# Utilizing P411-HF as a starting point for additional cyclic alkene functionalization reactions

While evolving for new functions, enzymes may gain activities for additional reactions that were not actively selected for. However, such promiscuous activities may in turn serve as evolutionary starting points to evolve novel functions. 49-51 For example, we found that variant P411-CIS, previously evolved from P450-BM3 for styrene cyclopropanation,14,15 showed more than 30-fold higher activity for 1-methylindole alkylation than P450-BM3 (Supplementary Table 1). Thus, P411-CIS was chosen as starting point for evolving indole alkylation activity, ultimately yielding variant P411-HF (Figure 4A). In turn, we speculated that P411-HF, engineered to accommodate a heterocyclic structure near the catalytic iron center, may provide a good starting point for further evolution to unlock additional carbene transfer reactions to (hetero)cyclic alkenes. To test this hypothesis, we performed three evolutionary "case studies", focusing on regioselective pyrrole alkylation, enantioselective indole alkylation with an α-disubstituted carbene precursor, and cyclic alkene cyclopropanation.

First, we attempted functionalizing pyrroles at either the  $C_2$  or  $C_3$  position, a considerable challenge for small-molecule catalysts.<sup>27</sup> Transition metal catalysis typically relies on electronic bias between the two positions and predominantly leads to functionalization of  $C_2$ –H bonds.<sup>52</sup> Alternatively, bulky substituents on the nitrogen were shown to shift regioselectivity towards the

 $C_3$  position.<sup>53</sup> Enzyme catalysis can take advantage of precise substrate orientation within the active site, and thus potentially drive site selectivity independent of electronic or steric bias.<sup>54</sup> Accordingly, we speculated that the P411-HF active site could be tuned by directed evolution to favor alkylation at either the  $C_2$  or  $C_3$  position.

We found that P411-HF catalyzes carbene transfer to 1methylpyrrole with 180 TTN, showing a 3:1 selectivity for the electronically favored C2-alkylated product (Figure 4A, Supplementary Figure 6, Supplementary Table 3). Subsequent site-saturation mutagenesis and screening revealed a new variant with mutation M263T, providing a 1:1 ratio of both products and confirming that active site mutations can indeed shift the selectivity. Further evolution yielded two divergent P411 lineages that selectively yield the  $C_2$  or  $C_3$  alkylation products: variant P411-HF M263T G181E A82T makes the C<sub>3</sub>-alkylated product with 705 TTN and 9:91 C2:C3 ratio, whereas variant P411-HF M263T T87W G181K yields the C2-alkylated product with 485 TTN and 98:2  $C_2:C_3$  ratio (Figure 4A). Thus, the P411-HF active site can be sculpted with only three rounds of mutagenesis and screening to exert precise control over pyrrole heterocycle alkylation regioselectivity. The resulting variants may provide starting points for evolving enzymes to selectively alkylate other pyrrole derivatives.

Next, we asked whether P411-HF could serve as a starting point to evolve for enantioselective indole alkylation using an  $\alpha$ -disubstituted carbene precursor (e.g., ethyl-2-diazopropanoate, Me-EDA), thereby delivering chiral alkylated indole core 5 found in bioactive natural products.<sup>56</sup> Alkylation of 1-methylindole with Me-EDA represented a difficult biocatalytic reaction: no activity was found for either free hemin or for P411-CIS, and variant P411-HF M263Y delivered the product with only 2 TTN. This low activity, however, was sufficient for GC-MS based screening, and we evolved this variant over seven rounds of site-saturation mutagenesis and screening to obtain variant P411-HF T87A Y263E A328Y L437M A268G T327P, which delivers 5 with 775 TTN (Figure 4B, Supplementary Figure 7, Supplementary Table 4). While this represents a >300-fold increase in catalytic activity, enantioselectivity remained at ca. 50% over the course of evolution. The limited enantioselectivity may be grounded in the enzymatic reaction mechanism: nucleophilic attack by the indole  $C_3$  carbon at the iron-carbone center followed by protonation is generally considered the mechanistic pathway of indole alkylation.36,38,57 This process would require the enzyme to have perfect stereocontrol over the protonation step in order to yield high enantioselectivity. We speculate that further engineering of the P411-HF active site could possibly fine-tune hydrogen- bonding networks between the reaction intermediate and the protein, which might allow better control of the protonation step and lead to higher stereoselectivity. The evolved catalyst is nonetheless among the few P411's efficiently catalyzing transfer of an α-disubstituted carbene.

Finally, we reasoned that P411-HF may provide a good starting point to unlock cyclopropanation reactions of (hetero)cyclic internal alkenes. We first tested cyclopropanation using indene and EDA and found that P411-HF yielded the *cis*-configured cyclopropane **6a** with 1880 TTN, 95:5 dr and 94% *ee* (Figure 4C). This was significantly higher than P411-CIS, showing 305 TTN, 69:31 *dr* and 78% *ee* (Supplementary Figure 8, Supplementary Table 5), supporting the hypothesis that evolution for

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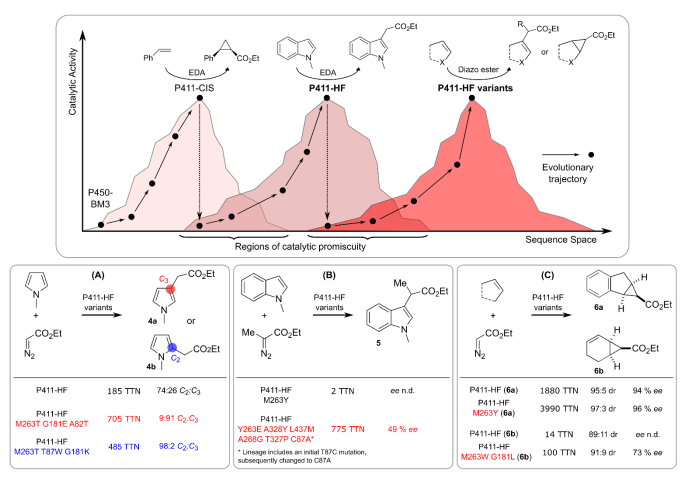
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**Figure 4: Evolving P411-HF for selective (hetero)cycle functionalization reactions.** Top: Schematic representation of the relation between protein sequence and catalytic function. A given enzyme sequence may exhibit additional 'promiscuous' functions that can be optimized by directed evolution.<sup>55</sup> We set out to test whether promiscuous activities of P411-HF, gained during evolution for 1-methylindole alkylation, could serve as starting points to evolve further (hetero)cycle functionalization reactions. (A) Evolution of P411-HF for regiose-lective 1-methylpyrrole alkylation. (B) Evolution of P411-HF for enantioselective 1-methylindole alkylation with ethyl-2-diazopropanoate. (C) Evolution of P411-HF for stereoselective cyclic alkene cyclopropanation. Data in (A) to (C) were derived from two independent experiments performed in duplicate; see Supplementary Tables 3 to 6 and Supplementary Figures 6 to 8 for evolutionary lineages and experimental details.

indole alkylation in parallel raised activity for cyclopropanation of sterically similar substrates. One additional round of site-saturation mutagenesis and screening delivered P411-HF M263Y, providing **6a** with almost 4000 TTN, 97:3 dr and 96% ee (Figure 4C). Comparable hemoprotein cyclopropanation activity on internal alkene substrates was achieved previously only with iridium porphyrin-substituted hemoprotein variants,<sup>58</sup> and this enzymatic reaction delivers a potentially useful cyclopropane core recently explored as pharmacophore.<sup>59</sup>

Similar results were obtained on a less-activated cyclic alkene,
1,3-cyclohexadiene (Figure 4C, Supplementary Figure 8, Supplementary Table 6). Again, P411-HF showed enhanced cyclopropanation activity over its parent P411-CIS, and two additional rounds of site-saturation mutagenesis and screening
yielded P411-HF M263W G181L, delivering the *cis*-configured
cyclopropane **6b** with 100 TTN, 91:1 *dr* and 73% *ee*. Lower
TTNs (< 100) were observed on heterocyclic substrates, including 2,3-benzofuran, 2,3-dihydrofuran and benzothiophene,</li>
while no activity was observed with aliphatic internal alkenes
2,4-hexadiene, 2-octene, cyclopentene, cyclohexene or cycloheptene, indicating that these substrates still present a challenge

for the current P411-HF carbene transferase lineage. Nonetheless, the results presented here demonstrate that enzymes containing the native iron-heme cofactor are capable of internal alkene cyclopropanation, with activity and selectivity tunable by directed evolution.

# **Concluding Remarks**

Enzyme catalysis represents a potentially sustainable alternative to chemical synthesis. Unlocking its promise, however, requires that enzymes perform equal or better than state-of-theart synthetic methods. In this study, we engineered a set of cytochrome P411 variants for carbene transfer to heterocycles and internal alkenes, synthetically important substrate classes largely absent from the repertoire of carbene transferases described to date. We tackled this problem with a two-tiered engineering strategy, first subjecting a cytochrome P411 variant to a comprehensive directed evolution effort, which delivered the highly active indole  $C_3$ -alkylation variant P411-HF. Further engineering of P411-HF towards more challenging heterocycle and internal alkene functionalizations then delivered a suite of P411 variants for a diverse set of heterocycle functionalization reactions. Only a few rounds of site-saturation mutagenesis and screening were required to achieve regioselective pyrrole alkylation, enantioselective indole alkylation, and stereoselective cyclic alkene cyclopropanation. The evolved enzymes catalyze hundreds to thousands of turnovers, are fully genetically encoded, and function with an earth-abundant iron cofactor. These variants complement existing transition metal catalysts for selective heterocycle functionalizations, provide potential starting points for future directed evolution endeavors, and expand the catalytic repertoire of the carbene transferase family of enzymes.

# ASSOCIATED CONTENT

**Supporting Information**. Experimental procedures, supplementary data, calibration curves, and characterization of alkylation and cyclopropanation products. This material is available free of charge via the Internet at http://pubs.acs.org.

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## Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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1 2 3 4 5 6 7 8 9 10 11 12	P450 directed evolution strategy: • Random mutagenesis of entire enzyme and site-saturation mutagenesis of active site residues • Spectrophotometric high-throughput and LC/GC-based reaction screening $\downarrow \downarrow $
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