1	Electronic Supplementary Information		
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3	Modulating stereoselectivity and catalytic efficiency of carbenoid reactions		
4	catalysed by Self-sufficient P450s		
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10 1. Materials and Methods

11 1.1 Materials

12 The racemic standard of ethyl 2-phenylcyclopropane carboxylate and styrene were obtained from

13 MACLIN. Ethyl diazoacetate was provided by Damas-beta. Other required reagents, such as ethanol,

14 acetonitrile, N-hexane, etc., were purchased from Sigma Aldrich.

15 1.2 Strain and amino acid sequence

16 E. coli BL21 (DE3) was employed for the expression of recombinant proteins, six self-sufficient P450s

17 include: P450_{TT} (Genbank number: WP_055423153.1), P450_{JT} (Genbank number: WP_104007096.1),

18 CYP505X (Genbank number: XP 754698), CYP152K6 (GenBank number: 6FYJ A), CviUPO

19 (GenBank number: 7ZCL B), and CYP505A30 (GenBank number: XP 003663647.1). Above

20 nucleotide sequences were synthesized by Tianlin (Wuxi) Technology Co., LTD. and subsequently

21 cloned into pET 28a (+) vector. For the positive control experiments, the nucleotide sequences of

22 P450_{BM3}, myoglobin, and cytochrome C were synthesized according to the references.¹⁻³

23 1.3 Medium

Luria-Bertani (LB) medium (g·L⁻¹): peptone 10.0, yeast extract 5.0, sodium chloride 10.0, pH 7.2,
solid medium with 2% AGAR.

26 Super Broth (TB) medium (g·L⁻¹): yeast extract 24.0, peptone 12.0, K2HPO3 16.4, KH2PO3 2.3,
27 glycerol 5.0.

28 1.4 Mutagenesis of substitution of the key axial residues of P450s

Substitution of key axial residues in P450s was performed through whole plasmid PCR mutagenesis, 29 using six P450 genes as templates. The primers required for introducing the critical residue changes 30 were exclusively synthesized by Sangon Biotech (Shanghai) Co., LTD. (Table S1). PCR amplification 31 system contained complementary primers and used KOD-Plus-Neo DNA polymerase to amplification 32 sequence. PCR program involved an initial pre-denaturation at 94°C for 4 min, followed by 33 amplification for 25 cycles with denaturation at 98°C for 10 s, annealing at 55°C for 15 s and 34 elongation at 72°C for 5 min, and then further elongation at 72°C for 10 min. The plasmids verified 35 by nucleic acid glue and digested with 1 µL of DpnI for 0.5 h at 37°C to remove the methylated 36 templates. Then 10 µL of the digestion mixture were transformed into E. coli BL21 (DE3). Single 37

38 colonies were picked up and cultivated in LB medium (50 μ g/mL Kan). The plasmids of mutations 39 were confirmed by sequencing.

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Table	S1.	List	of	prime	rs
Table	S1 .	List	of	prime	rs

Primer	Sequence		
Р450 _{тт} -С385Н-1	GCCGAGGTGCTGATGGGCGCCGTAACC		
P450 _{TT} -C385H-2	CATCAGCACCTCGGCCGTAACCTCGG		
P450 _{TT} -C385S-1	CATCAGAGCCTCGGCCGTAACCTCGG		
P450 _{TT} -C385S-2	GCCGAGGCTCTGATGGGCGCCGTAACC		
P450 _{JT} -C377S-1	CATCAGAGCATGGGCAAAAACATCGGC		
P450 _{JT} -C377S-2	GCCCATGCTCTGATGGGCACCATAGCCG		
Р450 _{JT} -C377H-1	CATCAGCACATGGGCAAAAACATCGGC		
Р450 _{JT} -C377H-2	GCCCATGTGCTGATGGGCACCATAGCC		
CYP505A30-C411S-1	CGTGCGAGCATCGGGCGTCCTTTCGCG		
CYP505A30-C411S-2	CCCGATGCTCGCACGCATACCAGTACCG		
CYP505A30-C411H-1	CGTGCGCACATCGGGCGTCCTTTCGCG		
CYP505A30-C411H-2	CCCGATGTGCGCACGCATACCAGTACC		
CYP505X-C414H-1	CGCGCACACATTGGTCGTCCGTTTGCG		
CYP505X-C414H-2	ACCAATGTGTGCGCGCATACCGTTGCC		
CYP505X-C414S-1	CGCGCAAGCATTGGTCGTCCGTTTGCG		
CYP505X-C414S-2	ACCAATGCTTGCGCGCATACCGTTGCC		
CYP152K6-C365S-1	CACCGCAGCGCTGGCGAATGGGTTACC		
CYP152K6-C365S-2	GCCAGCGCTGCGGTGGCCCATGAAGTA		
CYP152K6-C365H-1	CACCGCCACGCTGGCGAATGGGTTACC		
CYP152K6-C365H-2	GCCAGCGTGGCGGTGGCCCATGAAGTA		
CviUPO-C20S-1	GCTCCGAGCCCGGCTATGAACAGCCTG		
CviUPO-C20S-2	AGCCGGGCTCGGAGCGCGAAATTCACCC		
CviUPO-C20H-1	GCTCCGCACCCGGCTATGAACAGCCTG		
CviUPO-C20H-2	AGCCGGGTGCGGAGCGCGAAATTCACC		

42 1.5 Protein expression and purification

Recombinant E. coli BL21 (DE3) including six P450s and variants were cultivated at 37°C and 180 43 rpm in LB medium supplemented with 1% glucose. The cultures were grown until OD₆₀₀ reached 44 0.6~0.8. Subsequently, they were induced with 0.2 mM isopropyl β-D-thiogalactopyranoside (IPTG) 45 for gene expression. 2 mM FeSO₄ and 1 mM 5-aminolevulinic acid hydrochloride (ALA) were added 46 for synthesis of iron porphyrin in the catalytic center of P450s. And further culture was conducted at 47 16 °C, 120 rpm for 24 h. After culture, the culture medium was removed by centrifugation at 4000 rpm 48 for 10 min. The induced cells were harvested by centrifugation (4°C, 8,000×g for 10 min) to be freeze-49 dried. When protein purification was required, the cells were resuspended in phosphate buffer A (25 50 mM Tris-HCl, 300 mM NaCl, 20 mM imidazole, 5 mM β-mercaptoethanol, pH 7.4), then disrupted 51 with ultra-sonication (work for 2 s, pause for 3 s, 150 W). The resulting supernatant was loaded onto 52 a His-Trap HP nickel affinity column pre-equilibrated with buffer A using AKTA Avant System (GE 53 Healthcare, USA), P450s were eluted off by a 20-300 mM imidazole gradient in buffer B (25 mM 54 Tris-HCl, 300 mM NaCl, 300 mM imidazole, 5 mM β-mercaptoethanol, pH 7.4). The collected 55 proteins were verified by SDS-PAGE analysis and then concentrated by an ultrafiltration tube. The 56 target pure enzyme is subjected to the next experiment. 57

58 1.6 Small-scale of carbenoid model reactions

The reactions were initiated by adding 10 µL of styrene (1.2 M stock solution in methanol), followed 59 by the addition of 10 µL of EDA (0.4 M stock solution in methanol) using a syringe under anaerobic 60 conditions. The reaction mixture was stirred for 12 hours at 16°C. Upon completion of the reaction, it 61 was terminated by adding an equal volume of 3 M HCl. The resulting solution was then centrifuged at 62 12,000 rpm for 10 min to separate the phases. After centrifugation for 5 min, the solution was filtered 63 using a 0.22 µm organic filter membrane. Subsequently, the analysis was performed using an 64 ACQUITY UPC2 system by Waters Corporation, with a Trefoil CEL2 Supercritical Column (3.0 \times 65 150 mm, 2.5 µm) employed. The mobile phase A consisted of a mixture of ethanol and acetonitrile. 66 UV detection was carried out at a wavelength of 210 nm, with the column temperature maintained at 67 30 °C. The flow rate was set at 1.2 mL/min, and an injection volume of 3 µL was used. It was 68 recommended to conduct at least two parallel experiments for each group of reactions. The enzymatic 69 cyclopropane reaction system consists of crude enzymes (Table S2) and purified enzymes (Table S3). 70

Table S2. Reaction system for crude enzymes

Ingredients	Volume (µ		
Crude enzyme	3444		
Styrene (1.2 M)	33		
EDA (400 mM)	50		
NaS_2O_4	33		

Table S3. Reaction system for purified enzymes

Ingredients	Volume (μ		
Purified Enzyme	10 µM		
Styrene (1.2 M)	33		
EDA (400 mM)	50		
NaS_2O_4	33		
Tris HCl buffer (100 mM	Filled up to 2000		

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75 1.7 Phylogenetic tree analysis, Sequence analysis, Moleclular modeling and Simulation

Phylogenetic tree was constructed using is the maximum likelihood (ML) approach, and the 76 software used is RAxML 8.2.12 and PhyloSuite.^{4, 5} Using P450_{BM3}, myoglobin, and six self-sufficient 77 homologous sequences was P450s as templates, searched on the NCBI website 78 (https://www.ncbi.nlm.nih.gov/), incomplete and redundant sequences were removed, and the 79 sequences based on sequence consistency were rename. Sequence alignment was performed using the 80 multiple sequence alignment software MAFFT,⁶ with specific parameters: FFT-NS-2 (default) 81 strategy, Gap opening penalty, default: 1.53, Offset (works like gap extension penalty), default: 0.0. 82 Before the tree constructed, it is essential to determine evolutionary model for the sequence using 83 Model Finder.⁷ Bayesian Information Criterion (BIC) was employed to identify the best-fitting 84 substitution model among various evolutionary models: LG+G4. Finally, online phylogenetic tree 85 editing tool ITOL v5 (https://itol.embl.de/) was employed to beautify the tree.8 86

87 Sequence alignments and conserved analysis were conducted using online tool Clustal

(https://www.genome.jp/tools-bin/clustalw) and NCBI service (https://www.ncbi.nlm.nih.gov/). To
further enhance the obtained results, online website ENDscript/ESPrip was utilized for beautification.⁹
With the exception of certain protein structures like P450_{TT} and P450_{BM3}, all molecular models
were generated using AlphaFold 2. The analysis of interactions around the heme group was performed
through Discovery Studio software. The enzyme-substrate complexes were further optimized using
the CHARM force field in the Discovery Studio software package. The structures were visualized and
generated using Pymol.



97 2 Standard curve of product (EPC) using ultra-performance convergence chromatography

99 Figure S1. Analysis of ethyl 2-phenyl cyclopropane methane carboxylate (EPC) using ultra-100 performance convergence chromatography. (a) The ultra-performance convergence chromatography 101 results for different configurations of EPC were as follows: (R, R)-EPC eluted at 1.216 min, (S, S)-102 EPC eluted at 1.315 min, (S, R)-EPC eluted at 1.470 min, and (R, S)-EPC eluted at 1.649 min. (b) 103 Standard curve of different configurations of EPC analyzed by ultra-performance convergence 104 chromatography.



106 4. Expression and purification of recombinant P450s

Figure S2. SDS-PAGE analysis of six recombinant P450s. The results showed that six different candidate genes for self-sufficient P450 skeleton enzymes were all soluble and in line with the expected protein size. M: protein Marker, lane $1\sim2$: P450_{TT} supernatant, precipitate; Lane $3\sim4$: P450_{JT} supernatant, precipitate; Lane $5\sim6$: CviUPO supernatant, precipitate; Lane $7\sim8$: CYP505X supernatant, precipitation; Lane $9\sim10$: CYP505A30 supernatant, precipitate; Lane $11\sim12$: CYP152K6 supernatant, precipitate





116 Fig S3. SDS-PAGE analysis of three purified self-sufficient P450 enzymes. (a) P450_{JT}-WT. (b)

117 P450_{TT}-C385S. (c) CYP505X-WT.







Fig S6. SDS-PAGE analysis of purified $P450_{BM3}$



129 5. Comparison of carbenoid model reactions catalyzed by different self-sufficient P450s

Fig S7. Fingerprint of carbenoid model reactions catalysed by different self-sufficient P450s.
Experiments were performed using crude extract of recombinant P450s. Catalyst were normalized
such that all lyophilized cells added to buffer in the same amount for ultrasonication. Product was
determined by ultra-performance convergence chromatography.





Fig S8. Ultra-performance convergence chromatography analysis of reactions catalyzed by crude
 enzymes of P450_{TT} and its variants.





Fig S9. Ultra-performance convergence chromatography analysis of reactions catalyzed by crude
 enzymes of P450_{JT} and its variants.





Fig S10. Ultra-performance convergence chromatography analysis of reactions catalyzed by crude
 enzymes of CYP505A30 and its variants.

151 152	Fig S11 . Ultra-performance convergence chromatography analysis of reactions catalyzed by crude
152	enzymes of CYP505X and its variants.
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Fig S12. Ultra-performance convergence chromatography analysis of reactions catalyzed by crude
 enzymes of CYP152K6 and its variants.



Fig S13. Ultra-performance convergence chromatography analysis of reactions catalyzed by crude
 enzymes of CviUPO and its variants.

167 6. Comparison of turnover frequencies (TOFs) and stereoselectivity of P450s

Purified Enzymes	TOF (min ⁻	$de_E(\%)$	ee_{cis} (%)	ee_{trans} (%)
	1)			
P450 _{TT} -WT	0.75	94.0	99.9	79.8
P450 _{TT} -C285H	0.80	95.4	99.9	83.3
P450 _{TT} -C285S	1.50	31.5	58.0	20.9
P450 _{JT} -WT	1.20	50.7	68.0	33.4
P450 _{JT} -C377H	0.45	84.0	99.9	56.7
P450 _{JT} -C377S	0.70	83.3	93.5	51.9
CYP505X	0.95	91.6	99.9	75.8
P411-CIS	0.40	99.9	99.9	n.d.
P450-CIS	0.30	99.9	99.9	n.d.
Myoglobin	0.50	81.1	99.9	99.9
Cytochrome C	0.30	84.4	69.2	99.9

Table S4. Comparison of turnover frequencies (TOFs) and stereoselectivity of carbenoid model
reaction catalyzed by different purified P450s

171 8. Residues interacted with the heme cofactors of different P450s



Fig S14. Residues interacted with the heme cofactors of different P450s. Molecular simulations were conducted on the heme pocket of the following proteins: (a) $P450_{TT}$ (PDB: 6GII), (b) $P450_{JT}$ (homology model built using AlphaFold 2), (c) CYP505A30 (homology model built using AlphaFold 2), (d) CYP505X (homology model built using AlphaFold 2), (e) CYP152K6 (homology model built using AlphaFold 2), and (f) CviUPO (homology model built using AlphaFold 2). The heme cofactor (represented in red) and the surrounding residues were visualized as sticks.

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