A Biocatalytic Platform for Synthesis of Chiral α-Trifluoromethylated Organoborons

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ABSTRACT: There are few biocatalytic transformations that produce fluorine-containing molecules prevalent in modern pharmaceuticals. To expand the scope of biocatalysis for organofluorine synthesis, we have developed an enzymatic platform for highly enantioselective carbene B−H bond insertion to yield versatile α-trifluoromethylated (α-CF3) organoborons, an important class of organofluorine molecules that contain stereogenic centers bearing both CF3 and boron groups. In contrast to current “carbene transferase” enzymes that use a limited set of simple diazo compounds as carbene precursors, this system based on Rhodothermus marinus cytochrome c (Rma cyt c) can accept a broad range of trifluorodiazo alkanes and deliver versatile chiral α-CF3 organoborons with total turnovers up to 2870 and enantiomeric ratios up to 98.5:1.5. Computational modeling reveals that this broad diazo scope is enabled by an active-site environment that directs the alkyl substituent on the heme CF3 carbene intermediate toward the solvent-exposed face, thereby allowing the protein to accommodate diazo compounds with diverse structural features.
the B–H insertion strategy has only been demonstrated with a few aryl-substituted trifluorodiazo compounds.\textsuperscript{25} The deficit of methods for making chiral α-CF\textsubscript{3} organoborons motivated us to develop an enzymatic platform for their synthesis by reprogramming heme proteins to utilize trifluorodiazo alkanes for enantioselective B–H bond insertion reactions (Figure 1a). Harnessing the facile synthetic accessibility of trifluorodiazo alkanes\textsuperscript{26} and the evolvability of heme proteins,\textsuperscript{16,27} this platform can be used to prepare a broad range of chiral α-CF\textsubscript{3} organoborons, many of which are currently unobtainable. The major challenge to developing the enzymatic system is engineering the heme proteins to accommodate structurally diverse heme CF\textsubscript{3}-carbene intermediates for enantioselective B–H bond insertion. Although current carbene transferases can use a variety of substrates for versatile chemical transformations,\textsuperscript{38–43} the scope of diazo compounds in most of these reactions has been very limited (mainly to ethyl diazoacetate (EDA)). To expand the diazo substrate scope, it has usually been necessary to reoptimize the enzyme active site for each new diazo substrate. Furthermore, for all synthetic carbene B–H bond insertion reactions developed to date,\textsuperscript{44–50} high enantioselectivity has only been achieved for diazo compounds containing (hetero)aromatic groups adjacent to the diazo carbon.

We envisioned that these challenges could be met by leveraging the unique structural features of \textit{Rhodothermus marinus} cytochrome \(c\) (Rmcyt \(c\)).\textsuperscript{51} This protein has proven to be a highly versatile platform for developing new enzymatic carbene-transfer reactions.\textsuperscript{33,34} The heme-binding pocket of wild-type Rmcyt \(c\) is surrounded by several α-helices, two of which are connected by a surface loop.\textsuperscript{51} Recent structural characterization of the laboratory-evolved Rmcyt \(c\) with iron porphyrin carbene (IPC) intermediate revealed that the three mutations introduced during directed evolution rendered this surface loop (residues 98–103) highly flexible (Figure 1b, left).\textsuperscript{52} This enhanced flexibility allows the loop to explore more open conformations, not observed in the wild-type structure, that promote interactions between the substrate and the carbene intermediate. We hypothesized that we could engineer an active-site environment that favors a constrained conformation in which the CF\textsubscript{3} group of heme CF\textsubscript{3}-carbene intermediates...
points into the heme pocket, and the bulkier alkyl substituent R faces the solvent-exposed side (Figure 1b, right). In this conformation, the restricted orientation of the CF$_3$ group could ensure highly enantioselective formation of C−B bonds with little or no interference from the alkyl substituent R.

To identify mutations that can stabilize the conformation proposed in Figure 1b, we began directed evolution of R$_{\text{ma cyt}}$ with N-heterocyclic carbene (NHC) borane 1 as the model borane substrate and (3-diazo-4,4,4-trifluorobutyl)benzene (2 in Figure 2) as the model diazo carbene precursor. We expected that the large phenylethyl group in 2 would facilitate its positioning toward the solvent-exposed face. While wild-type R$_{\text{ma cyt}}$ barely catalyzed carbene B−H insertion with 2, site-saturation mutagenesis (SSM) at residue V75 led to the discovery of the V75S variant, which afforded the desired product 2a with 220 total turnovers (TTN) and 80:20 enantiomeric ratio (e.r.). Residue V75 is located on an α-helix that is close to the heme cofactor and was previously shown to be a key site for expanding the scope of diazo compounds.33

With R$_{\text{ma cyt}}$ V75S as the parent, we performed additional rounds of SSM on residues M100, M103, T101, and M99 to further improve the catalytic performance of this borylation catalyst. These residues reside on the front loop and are important for controlling the structure of the heme pocket.52 Through this engineering, we obtained a quadruple mutant, R$_{\text{ma cyt}}$ V75S M100L M103D M99A (SLDA), which produced product 2a with 1960 TTN and 96.5:3.5 e.r.

Y44 on the A helix of R$_{\text{ma cyt}}$ is another residue that might affect catalysis of the target borylation reaction. In an R$_{\text{ma cyt}}$ c-catalyzed carbene Si−H insertion reaction, this residue was expected to interact with a silane substrate approaching the enzyme active site, as indicated by the crystal structure of an R$_{\text{ma cyt}}$ c-bound heme-carbene intermediate.52 SSM of the quadruple mutant SLDA at Y44 and screening yielded the Y44I mutation that further improved borylation activity to 2460 TTN and enantioselectivity to 97.5:2.5 e.r.

With optimized variant R$_{\text{ma cyt}}$ Y44I V75S M99A M100L M103D (denoted BOR-CF$_3$) in hand, we next probed its activity toward a panel of structurally diverse trifluorodiazo alkanes. If BOR-CF$_3$ stabilizes the “solvent-exposed” conformation of the R group as we expect, then it should accept a wide range of trifluorodiazo alkanes for enantioselective carbene B−H insertion. Indeed, as shown in Figure 3a, BOR-CF$_3$ could be used to synthesize a spectrum of α-CF$_3$ organoborons with diverse structural features. High enantioselectivity and activity were obtained for diazo compounds bearing different phenyl substitution patterns (3a−6a). Extending the chain length of the alkyl substituent R has a more profound impact on the performance of BOR-CF$_3$ (7a, 8a). Strikingly, reducing the chain length to one carbon in 9a abolishes borylation activity. As indicated by computational modeling, the iron-carbene intermediate generated from diazo 9 with a shorter alkyl chain can adopt conformations in which the phenyl moiety is close to the carbene reaction center. Unfavorable steric interactions with the phenyl group in these
conformations would inhibit the borane substrate’s approach to the carbene (see Figure S6 in the Supporting Information). We further challenged BOR-CF₃ with trifluorodiazooalkanes containing alkyl chains without aromatic groups (10−13). Although these diazo compounds are structurally distinct from the model compound 2 used for directed evolution, BOR-CF₃ can still effectively convert them into α-CF₃ organoborons with 730−1630 TTN and 96:4−98:2 e.r. (10a−13a). This result shows that the high enantioselectivity of BOR-CF₃ does not require specific recognition of the aromatic substituent. One synthetic advantage of this method is that the trifluorodiazooalkane starting compounds can be synthesized easily from a variety of starting materials such as alkyl halides, aldehydes, and carboxylic acids.²⁶,⁵³,⁵⁴ As these compounds are widely present in nature, this method could provide a facile way to obtain chiral organoborons that contain motifs of complex natural products. To demonstrate this, we synthesized trifluorodiazoo compound 13 bearing a geranyl structural motif. Subjecting 13 to the standard conditions described in Figure 3 with BOR-CF₃ as the catalyst afforded organoboron product 13a with 1630 TTN and 98:2 e.r. Given the prevalence of the geranyl structural motif in bioactive
molecules,55 this organoboron compound may find applications in syntheses of fluorinated analogues of natural products such as pheromones. To further demonstrate the synthetic utility of our method, we performed this biocatalytic transformation on preparative scale and obtained borane product 2a in 52% isolated yield and 97:3 e.r. The NHC borane 2a can be readily converted to the boronic acid 2c while retaining the stereochemistry of the boron-substituted chiral center, which would facilitate its further derivatization to other chiral trifluoromethyl-containing molecules.

We next used molecular dynamics (MD) simulations to obtain further insights into the stereocontrol imposed by the enzyme and the roles of the introduced mutations. Analysis of the active-site shape of BOR-CF3 in the absence of both substrates revealed that the hydroxyl group in the 7S side chain forms a hydrogen bond with the Y71 amide backbone, which directs the Cα hydrogen of serine to face toward the heme (Figure 4a). Modeling of the active-site structure of BOR-CF3 with a bound diazo compound 2 showed that this serine arrangement generates more space at the heme distal side to better accommodate the CF3 group. As a result, diazo compound 2 can mainly be docked in one specific conformation in the BOR-CF3 active site (Figure 4b). Intriguingly, this conformation is structurally analogous to the transition state for iron-carbene generation via N2 loss (Figure 4c). This result suggests that the active-site environment of BOR-CF3 not only constrains the conformation of the trifluorodiazo alkane, but also promotes the formation of the iron-carbene intermediate by facilitating the interaction between the diazo compound and the heme iron center.

We further modeled the BOR-CF3-bound trifluoroalkyl-carbene intermediate 2b formed from diazo compound 2. MD simulations revealed that 2b can adopt two major conformations as shown in Figure 5a,b. The main difference between the two conformers is the slight rotation of the Fe–C bond, which is possible because no specific contacts between the CF3 group and the protein stabilize one conformation over the other. Nonetheless, in both conformations, the CF3 group is pointing into the active site, and the bulkier phenyl group lies between the M103D and Y44I side chains and is exposed to the solvent, in line with our original hypothesis. More interestingly, both conformers of 2b expose the same pro-R face of the carbene intermediate to the empty volume generated in the active site, consistent with the observed R absolute configuration of product 2a. To better understand the interaction between the borane substrate 1 and the carbene intermediate 2b, we used DFT calculations to describe the transition state (TS) for B–H carbene insertion using a model iron porphyrin system (see the SI, section VIII, for computational details). A structural comparison between this TS geometry and the structure of protein-bound carbene intermediate 2b revealed that borane 1 can be effectively accommodated in the empty volume of 2b where it can adopt a catalytically competent pose to reach the TS for C–B bond formation (TS-CB). In this reaction scheme, the newly introduced M99A and M100L mutations alter the conformational dynamics of the front loop and generate an appropriate binding pocket for borane substrate 1 in the carbene-bound enzyme, while the major role of M103D and Y44I is to facilitate binding of the diazo substrate and stabilize both the diazo and the carbene intermediate in this binding pose. Notably, in this interaction mode, the alkyl substituent R on the CF3-carbene intermediate should have little influence on how the borane substrate approaches the carbene intermediate, which explains the general high enantioselectivity exhibited by BOR-CF3 for most trifluorodiazo alkanes tested. The generality of this rationale for other trifluorodiazo alkanes is suggested by the catalytic performance of BOR-CF3 and its earlier variants on geranyl-containing diazo compound 13, where the trend in activity and enantioselectivity was similar to that with model diazo compound 2 (Scheme S1 in the Supporting Information).

In conclusion, we have developed a biocatalytic platform for synthesis of chiral α-CF3 organoborons. Using directed evolution, we created an active site in Rma cyt c that can structurally diverse trifluorodiazo alkanes for highly enantioselective carbene B–H insertion reactions. The effects of beneficial mutations on the catalytic activities and conformational dynamics have been rationalized by computational modeling. These efforts have expanded the scope of carbene intermediates accessible to heme proteins and provided new mechanistic insights into enzymatic carbene-transfer reactions.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscentsci.8b00679.

Detailed experimental procedures, spectroscopic data for all new compounds, details for molecular dynamics (MD) simulations, and DFT calculations (PDF)

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**Notes**

The authors declare no competing financial interest.

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