What Drives Chorismate Mutase to Top Performance? Insights from a Combined In Silico and In Vitro Study
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Helen V. Thorbjørnsrud,|| Luca Bressan,|| Tamjidmaa Khatanbaatar,|| Manuel Carrer, Kathrin Würth-Roderer, Gabriele Cordara, Peter Kast,* Michele Cascella,* and Ute Krengel*}

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ABSTRACT: Unlike typical chorismate mutases, the enzyme from Mycobacterium tuberculosis (MtCM) has only low activity on its own. Remarkably, its catalytic efficiency $\frac{k_{\text{cat}}}{K_m}$ can be boosted more than 100-fold by complex formation with a partner enzyme. Recently, an autonomously fully active MtCM variant was generated using directed evolution, and its structure was solved by X-ray crystallography. However, key residues were involved in crystal contacts, challenging the functional interpretation of the structural changes. Here, we address these challenges by microsecond molecular dynamics simulations, followed up by additional kinetic and structural analyses of selected sets of specifically engineered enzyme variants. A comparison of wild-type MtCM with naturally and artificially activated MtCMs revealed the overall dynamic profiles of these enzymes as well as key interactions between the C-terminus and the active site loop. In the artificially evolved variant of this model enzyme, this loop is preorganized and stabilized by Pro52 and Asp55, two highly conserved residues in the $\alpha$20$\beta$21 family. The role of Asp55 can be taken over by another acidic residue, if introduced at position 88 close to the C-terminus of MtCM, as suggested by molecular dynamics simulations and confirmed by kinetic investigations of engineered variants.

INTRODUCTION

Pericyclic reactions are common in industrial processes, but very rare in biology.1–4 Chorismate mutase (CM) catalyzes the only known pericyclic process in primary metabolism, the Claisen rearrangement of chorismate (1) to prephenate (2), via a chair-like transition state (Scheme 1).5 This catalytic step at the branch point of the shikimate pathway funnels the key metabolite chorismate toward the synthesis of tyrosine and phenylalanine, as opposed to tryptophan and several aromatic vitamins.6,7 The CM reaction is a concerted unimolecular transformation that is well studied by both experimental and computational means.8 It proceeds ostensibly via the same transition state in both solution and enzyme catalysis.9,10 Due to these factors, CM has long been a model enzyme for computational chemists.11

Natural CMs belong to two main classes with two distinct folds AroH and AroQ, which are equally efficient, with typical $\frac{k_{\text{cat}}}{K_m}$ values in the range of $(1-5) \times 10^7$ M$^{-1}$ s$^{-1}$.12 The AroH fold, exemplified by the Bacillus subtilis CM, has a trimeric pseudo-$\alpha/\beta$-barrel structure,13,14 whereas the structures of AroQ enzymes have all-$\alpha$-helical folds.15–21 The AroQ family is further divided into four subfamilies, $\alpha-9$, $\alpha-20,21$ The AroQ$\delta$ subfamily shows abnormally low catalytic activity compared to prototypical CM enzymes. In fact, the first discovered AroQ$\delta$ enzyme, the intracellular CM from Mycobacterium tuberculosis (MtCM), is on its own only a...
poor catalyst \( (k_{cat}/K_m = 1.8 \times 10^3 \text{ M}^{-1} \text{s}^{-1}) \),\textsuperscript{21} despite its crucial role for producing the aromatic amino acids Tyr and Phe. However, this low activity can be boosted more than 100-fold to a \( k_{cat}/K_m \) of 2.4 \( \times 10^6 \text{ M}^{-1} \text{s}^{-1} \) through formation of a noncovalent complex with the first enzyme of the shikimate pathway, 3-deoxy-D-arabino-heptulosonate 7-phosphate (DAHP) synthase (MtDS) (Figure 1A).\textsuperscript{21} The active site of AroQ CMs is dominated by positive charges, contributed by four arginine residues (Figure 1F). In MtCM, these are Arg18, Arg35, Arg46, and Arg58 (with the prime denoting a different MtCM protomer). Of particular importance for catalysis is Arg46,\textsuperscript{21} or its corresponding cationic residues in other CMs (of both AroH and AroQ families).\textsuperscript{22} However, high catalytic prowess is only achieved when this cationic residue is optimally positioned such that it can stabilize the developing negative charge at the ether oxygen in the transition state (Scheme 1).\textsuperscript{11,14,21,23–26} In MtCM, this is not the case unless MtCM is activated by MtDS.\textsuperscript{21} The MtDS partner repositions residues of the C-terminus of MtCM for interaction with the H1-H2 loop of MtCM that covers its active site, thereby inducing a characteristic kink in this loop (orange circle in Figure 1C). This interaction leads to a rearrangement of active site residues to catalytically more favorable conformations (Figure 1B)\textsuperscript{21} and is likely a key contributing factor for the increase in CM activity, as shown by randomizing mutagenesis of the C-terminal region followed by selection for functional variants.\textsuperscript{26} Complex formation also endows MtCM with feedback regulation by Tyr and Phe through binding of these effectors to the MtDS partner.\textsuperscript{21,27,28} Such inter-enzyme allosteric regulation\textsuperscript{28} allows for dynamic adjustment of the CM activity to meet the changing needs of the cell.

The naturally low activity of MtCM in the absence of its MtDS partner enzyme also provided a unique opportunity for laboratory evolution studies aimed at increasing MtCM efficiency. After four major rounds of directed evolution, the top-performing MtCM variant N-s4.1s emerged,\textsuperscript{17} which is abbreviated as MtCM\textsuperscript{1s} in this manuscript. This variant showed autonomous CM activity \( (k_{cat}/K_m = 4.7 \times 10^5 \text{ M}^{-1} \text{s}^{-1}) \) twice exceeding that of wild-type MtCM in the MtCM-MtDS complex, and can no longer be activated further through the addition of MtDS.\textsuperscript{12} The biggest gains in catalytic activity were due to replacements T52P and V55D in the H1-H2 loop of MtCM that covers the active site, thereby inducing a characteristic kink in this loop (Scheme 1). This conformation adopted by MtCM when in the complex with the MtDS partner enzyme is assumed to be more flexible compared to the \( \alpha \)-helical segments of MtCM.

Here, we used molecular dynamics (MD) simulations to investigate the behavior of MtCM in the absence or presence of ligands and to analyze whether the protein is able to interconvert between activated and nonactivated conformations in the absence of the MtDS partner enzyme. We also compared the wild-type MtCM with the evolved MtCM\textsuperscript{1s}, to see if the acquired amino acid substitutions introduced any new interactions or if they altered the probabilities of existing ones, with potential impact on catalytic activity. From an assessment of the dynamic properties of MtCM and MtCM\textsuperscript{1s}, we proposed a set of single, double, and triple C-terminal variants of the enzyme and subsequently tested these experimentally.

**Materials and Methods**

**Construction of Untagged MtCM Variants.** General cloning was carried out in *E. coli* DH5α or XL1-Blue (both Stratagene, La Jolla, California). All cloning techniques and bacterial culturing were performed according to standard procedures.\textsuperscript{29} Oligonucleotide synthesis and DNA sequencing were performed by Microsynth AG (Balgach, Switzerland).

For the construction of expression plasmids pKTCMM-H-V55D and pKTCMM-H-T52P for the native MtCM single variants, the individual site-directed mutants were first constructed in the pKTNTET background (providing an N-terminal His\textsubscript{6} tag, first 5 residues missing). Parts of the MtCM gene (Gene Rv0948c) were amplified using oligonucleotides 413-MtCM-N-T52P (5'-GTTCGCCTAGCGGAGTTAGTACGTTTGGATCATAGTCGGGAGATGAAGGTCATCGAAC) or 413-MtCM-N-T52P (5'-GTTCGCCTAGCGGAGTTAGTACGTTTGGATCATAGTCGGGAGATGATGAAGGTCATCGAAC) together with oligonucleotides 386-LpLib-N2 (5'-GGTTAAAGCTTTCGGCACGACCTAGTATTTAGTGACCGAGGCGGCCACGGCCCAAT) on template pMG248\textsuperscript{12} to create a 163 bp PCR product. The PCR products were restriction digested with *Xho*I and *Spe*I followed by preparative agarose gel yielding corresponding 260 bp fragments. pKTCMM-H\textsuperscript{1s} was used as acceptor vector and was accordingly restriction digested with *Xho*I and *Spe*I, yielding a 4547 bp acceptor fragment. The fragments were ligated overnight at 16 °C, using T4 DNA ligase (New England Biolabs, Ipswich, Massachusetts) or XL1-Blue (both *E. coli* DH5α and *E. coli* XL1-Blue cells. The cloned PCR’d DNA fragments were confirmed by Sanger sequencing. Subsequently, the genes for MtCM-T52P and MtCM-V55D were isolated by restriction digestion using enzymes *Xho*I and *Spe*I followed by a preparative agarose gel, yielding corresponding 260 bp fragments. pKTCMM-H\textsuperscript{1s} was used as acceptor vector and was accordingly restriction digested with *Xho*I and *Spe*I, yielding a 4547 bp acceptor fragment. The fragments were ligated overnight at 16 °C, using T4 DNA ligase.
The ligation products were transformed into chemically competent *E. coli* KA12 cells and the inserts were analyzed by Sanger sequencing. The gene for variant PHS10-3p3, carrying an N-terminal His$_6$-tag and missing the first five residues, was recloned into the native format provided by plasmid pKTCMM-H. Acceptor vector pKTCMM-H and pKTNTET-PHS10-3p3 were restriction digested with *Xho*I and *Spe*I, and the fragments were isolated from preparative agarose gels. The 4548 bp and 260 bp fragments were ligated overnight at 16 °C with T4 DNA ligase and transformed into chemically competent XL1-Blue cells. The relevant gene sequence was confirmed by Sanger sequencing.

Figure 1. Structural information on *M. tuberculosis* chorismate mutase. (A) Cartoon illustration of the heterooctameric complex of MtCM with DAHP synthase (MtDS) (Protein Data Bank (PDB) ID: 2W1A). MtCM is colored in shades of green and MtDS in shades of gray to emphasize individual subunits; Bartlett’s transition state analogue (TSA) is shown with golden spheres. The location of one of the four active sites of MtCM is marked with a circle. (B) Stereo superimposition of CM active sites of MtCM (shades of violet, with malate bound, white sticks) (PDB ID: 2VKL) and of the MtCM−MtDS complex (with TSA bound, golden sticks) (PDB ID: 2W1A), showing several active site residues as sticks. Shades of violet/green (and prime notation for Arg18′) illustrate separate protomers of MtCM/MtCM−MtDS structures, respectively. An arrow shows the shift in the position of Val55 upon MtDS binding, allowing H-bond formation of its backbone to TSA. (C) Cartoon superimposition of MtCM (PDB ID: 2VKL; violet, with white sticks for malate ligand) and activated MtCM (MtCM DS$^-$) from the MtCM−MtDS complex (PDB ID: 2W1A; green, with golden sticks for TSA). The biggest structural changes upon activation are a kink observed in the H1−H2 loop and interaction of the C-terminus (circled in orange) with the active site of MtCM. (D) Cartoon representation of the artificially evolved MtDS-independent super-active MtCM variant N-s4.15 (PDB ID: 5MPV, cyan), dubbed MtCM$^V$ in this work, having a $k_{cat}/K_m$ typical for the most efficient CMs known to date. Amino acid replacements accumulated after four cycles of directed evolution are emphasized as yellow side-chain sticks (A89 and M90 are not resolved) and labeled for one of the protomers. The H1−H2 loop (shown in red) adopts a kinked conformation similar to that observed for the MtDS-activated MtCM$^DS$ shown in (C). (E) Sequence alignment of wild-type MtCM (MtCM WT) and the highly active variant N-s4.15 (MtCM$^V$). Substituted residues are highlighted in yellow, and the H1−H2 loop is colored red. (F) Schematic representation of the active site of MtCM with bound TSA. Boxed residues refer to the wild-type enzyme, and green font color (Asp55, Ile62) refers to those substituted in MtCM$^V$. Charged residues are highlighted in red and blue.
Different C-terminal variants of the MtCM gene were generated by PCR mutagenesis. DNA fragments were amplified with the same forward primer (containing an NdeI site, underlined) and different reverse primers (containing an SpeI site, underlined) on different DNA templates. The gene encoding MtCM L88D was produced by PCR with primers LB5 (5′-TCGCCATATGAACTTGGAAATG) and LB4 (5′-TAAGCACTAGTTAGTGCAGC) on the template plasmid pKTCMM-H carrying the wild-type gene.21 The gene for the triple variant MtCM (T52P V55D L88D) was assembled with primers LB5 and LB4 on a pKTCMM vector containing MtCM variant 3p3 (T52P V55D).15 The gene for MtCM variant PNAM (D88N) was generated with primers LB5 and LB6 (5′-TAAGCACTAGTTACAGTCCGAAGG) and for the MtCM variant PLAM (D88L) with primers LB5 and LB7 (5′-TAAGCACTAGTTATAGTGACACCGGAGA), in both cases using a version of the template plasmid pKTCMM-H, into which the gene for the top-evolved s4.15 variant had been inserted.12 The resulting 296 bp PCR fragments containing Ndel and SpeI restriction sites at the 5′ and 3′ ends of the MtCM gene, respectively, were digested with the corresponding enzymes to yield 278 bp fragments. These fragments were ligated to the 4529 bp Ndel–SpeI fragment of pKTCMM-H yielding the final 4807 bp plasmids.

**Protein Production and Purification.** E. coli strain KA13,30 carrying an endogenous UV5 P lac-expressed T7 RNA polymerase gene was used to overproduce the (untagged) MtCM variants. KA13 cells were transformed by electroporation with the appropriate pKTCMM-H plasmid derivative that carries the desired MtCM gene variant.

For the two crystallized MtCM variants T52P (MtCM T52P) and V55D (MtCM V55D), the transformed cells were grown in baffled flasks at 30 °C in LB medium containing 100 μg/mL sodium ampicillin until the OD 600 reached 0.5. Gene expression was induced by centrifugation (6500g for 20 min at 4 °C) and incubation was continued overnight. The cells were harvested by centrifugation (6500g, 15 min) and the supernatant was loaded onto a HiTrap XL SP column (GE Healthcare) and the enzyme concentration ([E]) was determined using the Bradford assay.31

**X-ray Crystallography.** MtCM variants T52P (MtCM T52P) and V55D (MtCM V55D) were crystallized in 96-well two-drop MRC crystallization plates (SWISSCSSC) by the sitting drop vapor diffusion technique. Diffraction-quality crystals of MtCM T52P grew at 20 °C from a 1:1 (375 mL + 375 mL) mixture of protein (28 mg mL−1 in 20 mM BTP, pH 7.5) and reservoir solution containing 0.2 M sodium malonate, 20% PEG 3350 (w/v), and 0.1 M Bis Tris propane buffer, pH 8.5 (PACT premier crystallization screen, condition H12; Molecular Dimensions Ltd.). Crystals of MtCM V55D were obtained from a 1:1 (375 mL + 375 mL) mixture of protein (44 mg mL−1 in 20 mM Bis Tris propane, pH 7.5, 150 mM NaCl) and reservoir solution containing 0.2 M zinc acetate dihydrate, 10% w/v PEG 3000, and 0.1 M sodium acetate, pH 4.5 (JCSG plus crystallization screen, condition C7; Molecular Dimensions Ltd.) at 20 °C.

Diffraction data of MtCM T52P and MtCM V55D crystals were collected at the European Synchrotron Radiation Facility.
(ESRF, Grenoble, France) at the ID30A-3/MASSIF-3 (Dectris Eiger X 4M detector) and ID29 (Pilatus detector) beamlines, respectively, covering 120° with 0.1° oscillation. Diffraction images were integrated and scaled using the XDS software package.\textsuperscript{24} Merging and truncation were performed with AIMLESS\textsuperscript{25} from the CCP4 program suite.\textsuperscript{26} Since data collection statistics of both crystals suggested the presence of anisotropy, the XDS output was reprocessed for anisotropy correction and truncation using the STARANISO server.\textsuperscript{27} The “aniso-merged” output files (merged MTZ file with an anisotropic diffraction cutoff) were subsequently used for structure solution and refinement (Table S1).

The crystal structures of MtCM\textsuperscript{12P} and MtCM\textsuperscript{V55D} were solved by molecular replacement with the program Phaser.\textsuperscript{30} The structure of the top-evolved MtCM variant MtCM\textsuperscript{V} (PDB ID: 5MV)\textsuperscript{12} was used as a search model for solving the structure of MtCM\textsuperscript{12P} since it was expected to be a better match at the Pro52-containing H1–H2 loop compared to wild-type MtCM. For MtCM\textsuperscript{V55D}, we used the MtCM structure from the MtCM–MtDS complex (PDB ID: 2W19)\textsuperscript{21} as a search model, after truncation of the termini and the H1–H2 loop, and removal of the ligand.

The two structures were subsequently refined, alternating between real-space refinement cycles using Coot\textsuperscript{37} and maximum-likelihood refinement with REFMACS.\textsuperscript{38} The models were improved stepwise by first removing ill-defined side chains, and subsequently adding missing structural elements as the quality of the electron density map improved. Water molecules and alternative side-chain conformations were added to the MtCM\textsuperscript{12P} model toward the end of the refinement process, where positive peaks in the $\sigma_A$-weighted $F_o-F_c$ difference map and the chemical surroundings allowed for their unambiguous identification. As a last step, occupancy refinement was carried out with phenix.refine, a tool of the PHENIX software suite.\textsuperscript{39} The final structure of MtCM\textsuperscript{12P} was deposited in the Protein Data Bank (PDB)\textsuperscript{40} with deposition code 6YGT. Data collection and refinement statistics are summarized in Supporting Table S1.

**Determination of Enzyme Kinetic Parameters.** Michaelis–Menten kinetics of the untagged purified MtCM variants were determined by a continuous spectroscopic approach. Data collection and refinement statistics are summarized in Supporting Table S1.

**Molecular Dynamics Simulations.** Molecular dynamics (MD) simulations were carried out on a number of representative structures for CM. They included two independent sets of simulations for apo MtCM, starting either from the X-ray crystal structure of MtCM in complex with malate (after removing malate) (PDB ID: 2VKL)\textsuperscript{22} or from the structure of the CM polyepitide in the apo MtCM–MtDS complex (PDB ID: 2W19, chain D). The malate complex was chosen over ligand-free MtCM (PDB ID: 2QBV)\textsuperscript{41} due to its higher resolution and better refinement statistics. Both simulations gave essentially the same result; therefore, we will not refer to the second data set any further. For the highly active evolved MtCM variant (MtCM\textsuperscript{V}), we used the recent crystal structure (PDB ID: 5MV).\textsuperscript{12} The MtCM–ligand complex (MtCM\textsuperscript{V}) was taken from PDB ID: 2W1A,\textsuperscript{21} excluding the MtDS partner protein, where MtCM was co-crystallized with a transition state analog (TSA) in its active site (Figure 1). Finally, the V55D variant was modeled based on a partially refined experimental structure (Table S1). Residues that were not fully defined were added to the models using (often weak) electron density maps as reference in Coot.\textsuperscript{37} When no interpretable density was visible, geometric restraints (and $\alpha$-helical restraints for residues in helix H1) were applied during model building, to ensure stable starting geometries. The N-termini of all of the models were set at Glu13, corresponding to the first defined residue in almost all of the resolved structures available. Glu13 was capped with an acetyl group to imply the continuation of the H1 helix. CM dimers were generated by 2-fold crystallographic symmetry.

Missing H-atoms were added to the model and the systems were solvated in a periodic box filled with explicit water molecules, retaining neighboring crystallographic waters, and keeping the protein at least 12 Å from the box boundaries. The systems were neutralized through the addition of $\text{Cl}^-$ ions at a minimum distance of 7 Å from the protein and each other. Additional buffering moieties like glycerol or sulfate ions found in the crystals were not considered. MD simulations were run using the Gromacs 5.1.4 package\textsuperscript{42,43} using the AMBER 12 force fields for the protein moieties\textsuperscript{44} and the TIP3P model for water.\textsuperscript{45} The ligand was modeled using the GAFF force field.\textsuperscript{46} The smooth particle mesh Ewald method was used to compute long-range electrostatic interactions,\textsuperscript{47} while a cutoff of 11 Å was used to treat the Lennard–Jones potential.

The systems were minimized using the steepest descent/conjugate gradients algorithms for 500/1500 steps until the maximum force was less than 1000 kJ mol$^{-1}$ nm$^{-1}$. To equilibrate and heat the systems, first we ran 100 ps MD in the NVT ensemble starting from a temperature of 10 K, using the canonical velocity rescaling thermostat\textsuperscript{48} followed by 100 ps in the NpT ensemble with a Parrinello–Rahman barostat\textsuperscript{49} targeting a final temperature of 310 K and a pressure of 1 atm. After initial equilibration, 1 μs of MD simulation was performed for each system. In all MD simulations, the time step size was set to 2 fs.

**RESULTS**

The fact that MtCM exhibits only low natural catalytic activity provided us with a perfect opportunity to probe features that optimize CM catalysis by directed evolution.\textsuperscript{12} Since the biggest gains in catalytic activity were contributed by exchanging the H1–H2 loop residues S2 (TS2P) and S5 (V55D), we set out to determine the crystal structures of these two enzyme variants. Together, these two substitutions led to an increase in $k_{cat}/K_m$ by 22-fold compared to the parent enzyme.\textsuperscript{12}

**Crystal Structures of MtCM\textsuperscript{12P} and MtCM\textsuperscript{V55D}**. Whereas MtCM\textsuperscript{12P} crystals had the same space group (P4$_1$2$_1$2) and similar cell parameters as the wild-type enzyme (PDB IDs: 2VKL,\textsuperscript{21} and 2QBV,\textsuperscript{41}) with one protomer in the asymmetric unit, MtCM\textsuperscript{V55D} crystallized in a different space group
where the asymmetric unit contained the biological dimer. The MtCM<sup>T52P</sup> structure was refined to 1.6 Å and <i>R</i><sub>work</sub>/<i>R</i><sub>free</sub> values of 24.0/26.5% (Table S1 and Figure S1B), whereas MtCM<sup>V55D</sup> diffraction data yielded lower-quality electron density, particularly for the H1−H2 loop (Figure S1C,D). Consistent with this, the Wilson B-factor of MtCM<sup>V55D</sup> is high (57.8 Å<sup>2</sup>), indicating structural disorder. Refinement of the 2.1 Å MtCM<sup>V55D</sup> model stalled at <i>R</i><sub>work</sub>/<i>R</i><sub>free</sub> values of 27.6/34.9%, with very high B-factors for H1−H2 loop residues, especially for protomer B. For both structures, residues preceding residue Glu13 and C-terminal to Leu88 showed poorly defined electron density. Therefore, the terminal residues were not included in the final model.

Overall, the crystal structures of both MtCM<sup>T52P</sup> (PDB ID: 6YGT, this work; yellow), MtCM<sup>V55D</sup> (this work; orange), and top-evolved MtCM<sup>Y</sup> (PDB ID: 5MPV; 12 cyan) are very similar to the structure of substrate-free wild-type MtCM (PDB ID: 2QBV),<sup>41</sup> with RMSD = 0.3 and 0.4 Å, respectively. However, the H1−H2 loops (<sup>47</sup>MASGGPRLDHS<sup>57</sup>) of both protomers of MtCM<sup>V55D</sup> adopt a different conformation (RMSD = 2.3 Å compared to PDB ID: 2QBV), which most closely resembles the kinked conformation in the MtCM−MtDS complex (PDB ID: 2W19; RMSD = 0.8 Å) (Figure 2A). In the crystal structure

![Figure 2. Comparison of MtCM crystal structures, with focus on Arg46 and H1−H2 loop.](https://doi.org/10.1021/acs.biochem.2c00635)

![Figure 3. Secondary structure changes of MtCM during MD simulations.](https://doi.org/10.1021/acs.biochem.2c00635)
of MtCM\textsuperscript{VSSD}, Asp55 in the H1–H2 loop forms a salt bridge with Arg46, similar to the one in MtCM\textsuperscript{V} (compare Figure S1E,G,H). This interaction reorganizes the active sites of both MtCM variants for catalytic activity, mimicking MtCM in the complex with MtDS (Figure 2B). However, the overall conformation of the active site loop, which is involved in extensive crystal contacts that are highly distinct for the different crystal forms (Figure S2), differs significantly between the structures (Figures S1 and 2A).

**MD Simulations.** To evaluate the behavior of MtCM in the absence of crystal contacts, we probed the MtCM structures by MD simulations. We used four model systems: low-activity apo wild-type MtCM, MtCM\textsuperscript{V} (“ligand complex”: wild-type MtCM from the MtCM–MtDS structure in complex with TSA, the transition state analog of the CM reaction; \textsuperscript{11} Scheme 1 and Figure 1), MtCM\textsuperscript{V}, corresponding to the highly active evolved variant N-s4.1S,\textsuperscript{12} and MtCM\textsuperscript{VSSD}, which shows the highest catalytic activity among the single-substitution MtCM variants.\textsuperscript{12} We compared the overall dynamic profiles of these models and inspected the interactions formed between the C-termini and the H1–H2 loops covering the active sites, to find general features that could be associated with increased catalytic competence.

**Apo Structures of MtCM Are Characterized by Significant Flexibility.** We anticipated that the model systems would more or less retain the same fold as observed in the crystal structures, but that regions associated with crystal contacts, like the C-termini and the H1–H2 loop, would rapidly move away from their starting positions. Instead, the MD simulations revealed large changes from the initial crystal geometries in the apo protein structures, causing a rather high root-mean-square deviation (RMSD) from the original crystal structure geometry for the CM core regions (RMSD = 2.8 ± 1.2 Å (MtCM) or 3.4 ± 1.5 Å (MtCM\textsuperscript{V})). In particular, helix H2 showed a tendency to unravel (Figure S3). Due to the large flexibility observed, the two protomers making up the biological dimer instantaneously broke their symmetry, independently exploring different conformations in two chains. In contrast, the ligand-bound structure MtCM\textsuperscript{VLC} retained the secondary structure throughout the 1 μs simulation (Figure 3), with a lower RMSD (1.7 ± 0.6 Å) than the two apo structures. Intriguingly, a similar stabilization was observed for the unliganded variant MtCM\textsuperscript{VSSD} (Figure 3).

**Kinked Conformation of the H1–H2 Loop.** One of the biggest conformational changes in the crystal structure upon formation of the MtCM–MtDS complex occurs in the H1–H2 loop (Figures 1C and 2A).\textsuperscript{21} Whereas in the X-ray structure of the MtDS-activated MtCM, the H1–H2 loop is strongly kinked, this is not the case in nonactivated MtCM. We investigated the conformational landscape of this loop by simulations, using Arg53 from the loop as reporter residue. As shown in Figure 4, in one of the two protomers of MtCM, Arg53 remained in an extended conformation for the entire 1 μs MD simulation. In contrast, the same amino acid in the other protomer oscillated between the extended and the helical region of the Ramachandran plot (Figure 4), the latter being characteristic of the catalytically active conformation of the loop. Statistically averaging the two distributions, it appears that the apo form of MtCM is preferentially found in its inactive conformation, whereas in MtCM\textsuperscript{V} both protomers assumed the kinked active loop conformation, and retained it for the whole length of the simulation. However, TSA binding promoted the active conformation also in wild-type MtCM (represented by MtCM\textsuperscript{VLC}). The fact that the fluctuations of the MtCM\textsuperscript{V} H1–H2 loop are contained within the conformational basin of the catalytically competent geometry (Figure 4 and Table 1) is an indication that MtCM\textsuperscript{V} has an intrinsically preorganized loop, a condition that helps to minimize the entropy loss during substrate binding and consequently favors catalysis.

To test the effect of ligand binding, we repeated simulations of MtCM loaded with only one TSA ligand (MtCM\textsuperscript{VCL}). Interestingly, ligand presence in one of the two binding pockets was sufficient to stabilize the structure of the whole

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**Figure 4.** Conformation of Arg53 in the H1–H2 loop. Ramachandran plot showing backbone dihedral angles $\Phi$ and $\psi$ for Arg53. Red dots mark starting conformations. For MtCM, the H1–H2 loops from the two protomers (left and middle plots in the top row) assume different ensembles of conformations; overall, the catalytically favored conformation ($\psi \sim 0$) is observed less frequently than the nonproductive one. In contrast, in MtCM\textsuperscript{V}, both loops retained the active conformation during the entire course of the simulation, similarly to that observed for MtCM\textsuperscript{VLC}. When only one ligand was bound (MtCM\textsuperscript{VCL}), the TSA-loaded site (holo) retained the active conformation, while the loop in the other protomer (apo) remained flexible.
evolved sequence. Interestingly, also MtCM rigidification as a direct consequence of its promoted the elongation of helix H2, resulting in a significant MtCM retain considerable independence.

Nonetheless, the H1 dimer. Nonetheless, the H1–H2 active site loop of the apo protomer retained its intrinsic flexibility (Figure 4). The fact that the active site loop of the unloaded protomer behaved like the apo MtCM system suggests that the two active sites in MtCM retain considerable independence.

Contrary to MtCM and MtCMV (Figure 1C,D), the presence of the additional carboxylate group in MtCMVSSD promoted the elongation of helix H2, resulting in a significant shortening of the active H1–H2 loop (Figure 5A). This structural rearrangement is associated with the formation of persistent salt bridges between Asp55, now localized in the first turn of H2, and active site residues Arg18′ and Arg46 that were retained for the entire length of the simulation. Noticeably, in MtCMWT, where such stabilizing electrostatic interactions are absent, no such contacts were observed, with the side chain of Val55 keeping a distance of more than 10 Å from the side chains of both Arg18′ and Arg46 for the whole duration of the simulation.

Overall, MtCMVWT shows a noisier RMSF profile over the whole amino acid sequence compared to MtCMV and to the ligand complex MtCMVLC (Figure S3). This result reflects the expected rigidification occurring upon substrate binding due to additional protein–ligand interactions in MtCMVLC. MtCMV accomplishes rigidification as a direct consequence of its evolved sequence. Interestingly, also MtCMVSSD shows generally dampened fluctuations, possibly due to the extended helical motif observed in that structure.

Positioning of Active Site Residues. The MtCM active site contains four arginine residues (Figure 1F), among them the key catalytic residue Arg46. In contrast to the observation in the two MtCM–MtDS crystal structures, the conformation of Arg46 was not strictly maintained during MD simulations. In the absence of a ligand, Arg18′, Arg46, and Arg58 repelled each other, and at least one of the residues was pushed out of the active site in the majority of the simulations. Only one of the four arginine residues (Arg35) maintained its position (Table 1), appropriately placed for substrate binding by wild-type MtCM, with an RMSF below 1 Å, while RMSF values >2 Å for the other Arg residues signal a substantial increase in the conformational freedom. This changes upon complex formation with MtDS, guiding also the important Arg46 into a catalytically competent conformation.

In contrast to MtCMVWT, the two variants MtCMV and MtCMVSSD exhibited lower RMSF values for all active site Arg residues (Table 1 and Figure 1F) and maintained their catalytically competent conformation during the MD simulations even in the apo forms (Figure 5). The more stable positioning of Arg18′ and Arg46 appears to be a direct consequence of the replacement of Val55 with Asp, which introduces a negative charge, mitigating the surplus positive charges in the active site.

Interactions between C-Terminal Residues and H1–H2 Loop. A crucial factor for the enhanced activity of MtCM in the MtCM–MtDS complex is an MtDS-induced interaction between MtCM’s H1–H2 loop and its C-terminus. The interaction can be divided into two contributions: a salt bridge between the C-terminal carboxylate and the side chain of Arg53, and a hydrophobic contact between Leu54 and Leu88 (Figure S4A).

Our 1 μs-long simulations detected persistent, multiple interactions involving the C-terminal carboxylate. In contrast, the hydrophobic contacts between Leu54 and Leu88 were disrupted in the first nanoseconds, and almost never observed again during the rest of the simulation time (Figure S4B,C).

Table 1. Root-Mean-Square Fluctuation (RMSF) Values of Selected Active Site Residues

|          | RMSF (Å) |
|----------|----------|
|          | MtCM<sup>WT</sup> | MtCMV | MtCM<sup>VLC</sup> | MtCM<sup>VSSD</sup> |
| Arg18′   | 2.2 ± 0.5 | 1.5 ± 1.0 | 0.7 ± 0.2 | 1.1 ± 0.5 |
| Arg35    | 0.8 ± 0.4 | 0.7 ± 0.4 | 0.4 ± 0.1 | 0.6 ± 0.2 |
| Arg46    | 2.0 ± 0.6 | 1.7 ± 0.7 | 0.7 ± 0.5 | 0.9 ± 0.5 |
| Arg58    | 2.3 ± 1.0 | 1.9 ± 2.0 | 2.0 ± 0.9 | 1.5 ± 0.6 |

“RMSF values were calculated as an average over all nonhydrogen atoms for each residue compared to the average structure of the simulation. The reported σ values reflect the different relative fluctuations of the individual atoms composing the residues in the two symmetric protomers.

Figure 5. Role of MtCMV residue Asp55 in positioning active site residues. (A) Extension of H2 and stabilization of the H1–H2 loop by residue Asp55. Substitution of Val55 by Asp stabilizes helix H2 through interactions with Arg18′ and Arg46 across the active site (the image shows the structure of VSSD after 1 μs of MD simulations). Note that Arg46 is a catalytically essential residue for MtCM and its correct orientation is critical for catalytic proficiency. (B) Distance plotted between MtCMV Arg46 (black, chain A) or Arg18′ (red, from chain B), and Asp55 (chain A) observed during the simulation. In both cases, the distance measured is between Asp Cγ and Arg Cζ using PDB nomenclature.
Salt Bridges with C-Terminus. In our MD simulations, the C-terminal carboxylate formed interchangeable contacts with Arg53 and the catalytically important Arg46, which is located in the last turn of helix H1 (Figure 6A). Notably, the presence of a salt bridge between Arg46 and the C-terminus correlated with the apparently active conformation of the H1–H2 loop (Figure 6A).

The observed fluctuations suggest that the catalytically competent conformation of the binding site is malleable in wild-type MtCM and that additional interactions, i.e., with the substrate, are required to stabilize it. This is in line with studies of a topologically redesigned monomeric CM from *Methanococcus jannaschii*. This artificial enzyme was found to be catalytically active in the presence of the substrate despite showing extensive structural disorder without a ligand, reminiscent of a molten globule.

**MtCM** V Exhibits Strengthened Interactions between C-Terminus and H1–H2 Loop. In MtCM V, the four C-terminal residues Arg–Leu–Gly–His (RLGH) are substituted with Pro–Asp–Ala–Met (PDAM) at positions 87–90, which include another carboxylate, introduced through Asp88. Our MD simulations show that the Asp88 carboxylate in the evolved variant MtCM V offers an alternative mode of interaction with Arg53 of the H1–H2 loop (Figures 6B and S5), which is not possible for wild-type MtCM. This allows for a persistent interaction of C-terminal residues with the H1–H2 loop throughout the simulation, while maintaining a highly flexible C-terminus. Moreover, in MtCM V, Arg46 is topologically displaced from its original position with respect to the loop and no longer able to engage in a catalytically unproductive salt bridge with the C-terminus.

Another interesting substitution, which emerged within the four C-terminal residues during the laboratory evolution toward variant MtCM V, is a proline residue (RLGH to PDAM). However, in contrast to Pro52, Pro87 did not appear to have a major influence on the simulations. While
Pro52 is likely contributing to H1–H2 loop rigidity, with an average RMSF of 1.6 Å in MtCMV compared to 2.5 Å (MtCM) for this region, the C-termini showed similarly high RMSF values in the two models (>3 Å). Although Pro87 induced a kink at the C-terminus, this did not affect to appear the flexibility of the three terminal residues Asp88—Ala89—Met90.

**Kinetic Analysis to Probe Predicted Key Interactions of Engineered MtCM Variants.** In the course of the directed evolution of MtCMV, the L88D replacement was only acquired after the H1–H2 loop-stabilizing substitutions TS2P and V55D were already introduced. Guided by the outcome of the MD simulations, we therefore probed the kinetic impact of the innocuous single L88D exchange in the context of three different sets of MtCM variants to experimentally assess the benefit of the introduced negative charge for fine-tuning and optimizing catalytic efficiency. We looked at (i) changing Asp88 in the MtCMV sequence 87PD/PADM90 into Asn88 or Leu88, (ii) directly introducing Asp88 into the MtCM wild-type sequence, and (iii) the triple variant TS2P V55D L88D (MtCM Triple). All variants were obtained in their native format, i.e., with their native N-terminus and without a His-tag, to allow for optimal comparison with the structural and computational results. The variants were purified by ion-exchange and size-exclusion chromatography from the E. coli host strain KA13, which is devoid of CM genes to rule out contamination by endogenous CMs.18,30 Subsequently, the enzymes’ kinetic parameters were characterized by a spectrophotometric chorismate dephosphorylation assay.

As shown in Table 2, removing the negative charge at residue 88 by replacing Asp with Asn in the top-evolved variant MtCMV leads to a 2.5-fold drop in the catalytic efficiency $k_{cat}/K_m$ to $1.7 \times 10^7$ M$^{-1}$ s$^{-1}$. This decrease is due both to a slightly lower catalytic rate constant ($k_{cat}$) as well as a reduced substrate affinity (doubled $K_m$). When residue 88 is further changed to the similarly sized but nonpolar wild-type residue Leu88 in variant MtCM PLAM, the catalytic parameters essentially remain the same as for the Asn88 variant (Table 2), independently confirming the catalytic advantage of the negative charge introduced through Asp88.

For the second set of variants that directly started out from the sluggish MtCM wild-type enzyme (MtCMV), a trend for an increase in catalytic activity upon replacing Leu88 by Asp88 was observed (1.6-fold higher $k_{cat}/K_m$ reaching $2.7 \times 10^7$ M$^{-1}$ s$^{-1}$; Table 2). This is mainly caused by an increase in $k_{cat}$ rather than an altered substrate affinity. Interestingly, the L88D exchange together with TS2P and V55D in the MtCM triple variant does not lead to a significant increase in $k_{cat}/K_m$ compared to MtCM 3p3,12 which just carries the two loop substitutions TS2P and V55D.

Thus, the substitution of Leu88 with Asp88 indeed results in a beneficial effect on the performance of MtCM. However, this effect is only prominent in combination with other selected exchanges, such as those present in MtCMV. As a single amino acid replacement in the wild-type enzyme or on top of the two substitutions in the H1–H2 loop, the effect of L88D is less noticeable, if present at all.

In summary, a comparison of the dynamic behavior of wild-type MtCM in its apo and ligand-bound states with MtCMV and MtCMV33D revealed that the catalytically favorable conformation of the active site is achieved by the interplay of several interactions, which balance charges and entropic disorder of the H1–H2 loop. Structuring is promoted, in particular, by increasing the number of the negatively charged carboxylate groups that can both shield the electrostatic charge of the various arginine side chains within or next to the active site and orient catalytically important residues by hydrogen bonding and salt bridge formation. Simulations of MtCMV revealed the special importance of Asp55 in the V55D variant for coordinating Arg18 and Arg46, thus promoting the preorganization of the active site region. These results echo the conclusions from directed evolution, which also identified the V55D substitution as the most important contributor for catalytic enhancement, causing a 12-fold increase in $k_{cat}/K_m$.

At the same time, we determined and rationalized the more subtle and context-dependent effect of the L88D replacement that introduced an additional negative charge for electrostatic preorganization of the active site. Overall, the high catalytic activity of MtCMV clearly results from many individual larger and smaller contributions mediated by substitutions at diverse locations within the enzyme structure.

## DISCUSSION

### Important Activating Factors in MtCMV and MtCMV

MtCM has intrinsically low activity but can be activated to rival the performance of the best CMs known to date through the formation of a heteroocotameric complex with MtDS,21 which aligns crucial active site residues to catalytically competent conformations. Most importantly, binding to MtDS induces preorganization of Arg46 into a catalytically favorable
conformation (Figure 2B), via H-bonding to the carbonyl oxygens of Thr52 and Arg53.21 Arg46 is the crucial catalytic residue interacting with the ether oxygen of Bartlett’s transition state analogue (TSA)31 in the complex with MtDS (PDB ID: 2W1A)21 (Figures 1B,F and 2B); upon replacing Arg with Lys, the enzyme’s efficiency drops 50-fold.21

Both MtCM<sup>DS</sup> and MtCM<sup>W</sup> exhibit a kinked H1—H2 loop conformation (Figures 1C,D and 2A), which was hypothesized to be important for increased catalytic efficiency.2 However, in MtCM<sup>W</sup> and MtCM<sup>DS</sup>, the kink is exacerbated by crystal contacts, which are different in the two crystal forms (Figure S2). This kink is much less prominent in wild-type MtCM, or even MtCM<sup>T52P</sup> (Figures 2A and S1B), and completely lost during the simulations of MtCM<sup>WT</sup> (we did not carry out simulations on the single variant MtCM<sup>T52P</sup>). Thus, this conformation may well be a crystallization artifact rather than a prerequisite for an active MtCM.

Nevertheless, preorganization and prestabilization appear to be of crucial importance for the catalytic prowess of MtCM. The largest boost in catalytic efficiency (12-fold enhancement) by a single substitution was observed for the V55D mutant of MtCM<sup>DS</sup> (Figure 1D,E) and forms a salt bridge to the catalytically important Arg46 at the top of helix H1 (Figure 5A), an interaction that is also observed in the crystal structure of MtCM<sup>W</sup> (Figure S1G,H). During the MD simulations of MtCM<sup>W</sup> and the single variant MtCM<sup>VS5D</sup>, the presence of Asp55 reduced the mobility of active site residues. By interacting with Arg18 and Arg46, this residue helps to preorganize the active site for catalysis and reduce unfavorable conformational fluctuations caused by electrostatic repulsion in the absence of a substrate. This is supported by the lower RMSF values of MtCM<sup>W</sup> compared to uncomplexed wild-type MtCM (Table 1) and by a slightly higher melting temperature of MtCM<sup>VS5D</sup> (ΔT = 3 °C from differential scanning fluorimetry (DSF) measurements; preliminary data). By decreasing thermal fluctuations in the active site, Asp55 likely also reduces the entropic penalty associated with substrate binding. Pro52 appears to exert a similar stabilizing effect on the protein, despite the rather small structural changes, as suggested by a 2 °C increase in melting temperature of MtCM<sup>T52D</sup> in DSF experiments compared to MtCM (preliminary data). This single substitution alone raises the k<sub>cat</sub>/K<sub>m</sub> value of the enzyme by a factor of six.12 It is worth noting that the simultaneous substitution of T52P and VS5D increased the melting temperature by 6 °C (monitored by circular dichroism spectroscopy) and boosted k<sub>cat</sub>/K<sub>m</sub> by 22-fold.12 The top-evolved MtCM<sup>W</sup> even showed a melting temperature of 83 °C compared to 74 °C for the parent MtCM.12

Importance of the C-Terminus. MtCM activation by MtDS involves a change in conformation of the C-terminus of MtCM and its active site H1—H2 loop.21 Specifically, a salt bridge is formed between the C-terminal carboxylate of MtCM (which is repositioned upon MtDS binding) and loop residue Arg53, possibly bolstered by a newly formed hydrophobic interaction between Leu88 and Leu54 (Figure S4A). The 1 μs simulations suggest that salt bridge formation with Arg53 occurs in solution in all tested cases, whereas the hydrophobic contact is less important.

Directed evolution experiments carried out by randomizing the final four C-terminal positions 87—90 of MtCM had previously revealed that a great variety of residues with quite distinct physico-chemical properties are compatible with a functional catalytic machinery.20 Conserved positions emerged only when probing for an intact activation mechanism by MtDS.26 Still, when residues 87—90 of MtCM<sup>V</sup> were evolved from Arg—Leu—Gly—His to Pro—Asp—Ala—Met (Figure 1E), an increase in k<sub>cat</sub>/K<sub>m</sub> by roughly a factor of four was achieved.12 Here, we resolved this apparent paradox by investigating C-terminal factors important for the fine-tuned optimization of CM function. Even though the replacement R87P induced a kink in the structure, the presence of the proline did not appear to have a major influence in the simulations. Notably, the C-terminal substitutions together result in a change in net charge from +1 to −2, including the terminal carboxylate, providing the basis for more extensive electrostatic interactions with the positively charged Arg53 than is possible for wild-type MtCM. Indeed, our kinetic analysis of Asp88-containing MtCM variants demonstrates that this residue increases CM’s catalytic efficiency (Table 2). The fact that Asp88 did not significantly augment k<sub>cat</sub>/K<sub>m</sub> in the context of the MtCM double variant T52P VS5D (i.e., MtCM Triple; Table 2) suggests that the extent of catalytic improvement by L88D depends on the particular structural context.

Our simulations indicate that in free wild-type MtCM, an interaction of the C-terminal carboxylate with the key active site residue Arg46 is possible but infrequent due to fluctuations (Figure 6A and Table 1). In contrast, in MtCM<sup>W</sup> and MtCM<sup>DS</sup> the side chain of Arg46 points toward the catalytic pocket (Figures 5 and 2B), and any unproductive reorientation of Arg46 toward the C-terminus would easily result in a clash with the H1—H2 loop. Thus, an additional feature of this loop may be to act as a conditional shield (illustrated for MtCM<sup>W</sup> in Figure 7). In the conformation assumed in MtCM<sup>V</sup> and MtCM<sup>DS</sup>, this loop blocks the reorientation of Arg46 toward the C-terminus and hence prevents an unproductive conformation accessible for free wild-type MtCM. MtCM<sup>DS</sup> and MtCM<sup>V</sup> use different means to correctly position active site residues, which correlates with a bent H1—H2 loop in both cases. This is either achieved through conformational changes imposed upon MtCM<sup>DS</sup> by MtDS binding, or by establishing a salt bridge across the active site, between Arg46 and Asp55, as seen for MtCM<sup>V</sup> and also for the single variant MtCM<sup>V55D</sup> (Figures 5 and S1E,G,H).

General Implications for CM Catalysis. It is obviously impossible to directly transfer our findings of critical detailed molecular contacts from the AroQ<sub>γ</sub> subclass CM of <i>M. tuberculosis</i> to the evolutionary distinct AroH class CMs, or even to the structurally and functionally divergent AroQ<sub>γ</sub> and AroQ<sub>ρ</sub> subclasses.53 Neither of those groups of CMs have evolved to be deliberately poor catalysts that become proficient upon regulatory interaction with a partner protein. Nevertheless, they have malleable and allow for conformational switching between a poorly and a highly active form. In contrast, this region is rigidified in a catalytically competent conformation in the overwhelming majority of CMs from other subclasses. This is exemplified by the prototypic EcCM (AroQ<sub>ρ</sub> subclass) and the secreted *MtCM (AroQ<sub>γ</sub>), which possess the sequence 45PVRD<sup>48</sup> and 66PIED<sup>69</sup>, respectively, at the position corresponding to the malleable H1—H2 loop sequence 32TRLV<sup>55</sup> of wild-type MtCM.12 Remarkably, the two most impactful substitutions T52P and VS5D occurring during the
evolution of MtCM\textsuperscript{V} have led to the tetrapeptide sequence \textsuperscript{5}PRLD, with both Pro and Asp being conserved in naturally highly active CMs.\textsuperscript{12}

The AroQ\textsubscript{δ} subclass CM from \textit{Corynebacterium glutamicum} is another structurally well-characterized poorly active CM ($k_{\text{cat}}/K_m = 110 \text{ M}^{-1} \text{s}^{-1}$) that requires complex formation with its cognate DAHP synthase for an impressive 180-fold boost in catalytic efficiency.\textsuperscript{14} In that case, inter-enzyme allosteric regulation involves a conformational change of a different malleable segment between helices H1 and H2. Thus, while the molecular details important for the activation of a particular AroQ\textsubscript{δ} CM cannot be transferred directly from one system to another, our findings suggest as a general regulatory principle the deliberate and reversible destabilization of a catalytically critical loop conformation.

In both the \textit{M. tuberculosis}\textsuperscript{12} and the \textit{C. glutamicum} systems,\textsuperscript{54} crystal contacts in the H1–H2 loop region impede the structural interpretation of the activity switching. The MD simulations shown here represent an interesting alternative approach to dynamic high-resolution structure determination methods for sampling the conformational space adopted by malleable peptide segments with and without ligands.

\section*{CONCLUSIONS}

MD greatly aided the analysis of crystal structures that were compromised or biased by extensive crystal contacts at the most interesting structural sites. Our aim was to obtain insight into the crucial factors underlying CM activity by comparing the structure and dynamics of the poorly active wild-type MtCM ($k_{\text{cat}}/K_m = 1.7 \times 10^3 \text{ M}^{-1} \text{s}^{-1}$) with the top-performing MtCM variant MtCM\textsuperscript{V} ($k_{\text{cat}}/K_m = 4.3 \times 10^5 \text{ M}^{-1} \text{s}^{-1}$), which emerged from directed evolution experiments. Both in MtDS-activated wild-type MtCM and in MtCM\textsuperscript{V}, high activity correlated with a kinked H1–H2 loop conformation and an interaction of this region with the C-terminus of MtCM. The autonomously fully active variant MtCM\textsuperscript{V} had amino acid changes in both of these regions that augment these structural features. In this report, we focussed on substitutions T52P, V55D, and L88D.

The active site of all natural CMs contains a high density of positive charges. In MtCM, four arginine residues (Arg18', Arg35, Arg46, and Arg58, of which Arg18' is contributed by a different MtCM protomer) are responsible for binding and rearranging the doubly negatively charged substrate chorismate. Only one of these residues (Arg35) is firmly in position before the substrate enters the active site. Of critical importance for catalysis is Arg46. During the MD simulations, Arg46 competes with another arginine residue (Arg53) for binding to the C-terminal carboxylate (Figure 6A) and adopts a catalytically unproductive conformation unless an aspartate residue (Asp55 or Asp88) comes to its rescue. As shown here, Asp55 not only properly orients Arg46 for catalysis but additionally stabilizes the active site. Together with T52P, which preorders the H1–H2 loop, the V55D exchange results in reduced mobility of residues in the active site through stabilizing interactions, thereby preorganizing it for efficient catalysis and lowering the entropic cost of substrate binding. Another aspartate residue (Asp88), also acquired in the top-evolved MtCM\textsuperscript{V},\textsuperscript{12} helps to balance charges, and—by interacting with Arg53—imposes a steric block that prevents nonoptimal positioning of Arg46 (Figure 6B), explaining why the L88D exchange can increase $k_{\text{cat}}/K_m$ by about 2- to 3-fold.

In summary, we tested our hypotheses on the specific importance of critical substitutions acquired during the directed evolution of MtCM\textsuperscript{V}, namely, T52P, V55D, and L88D by investigating single variants as well as combinations with other residue replacements that were found to augment catalysis. The variants were characterized by crystallography, MD simulations, and enzyme kinetics. The two residues Pro52 and Asp55 exert a major impact by prestabilization and preorganization of catalytically competent conformations of active site residues, while Asp88 contributes to fine-tuning and optimizing the catalytic process. By expanding on the previous directed evolution studies, we have shown here how the accumulated set of amino acid substitutions found in MtCM\textsuperscript{V} has resulted in an activity level matching that of the most active CMs known to date.\textsuperscript{12}

Figure 7. Shielding interaction mediated by the H1–H2 loop. (A) Conformation of important Arg residues in chain A of MtCM\textsuperscript{V} (cyan) after 31.7 ns of MD simulations. The key active site residue Arg46 is positioned on the opposite side of the H1–H2 loop, which in turn is bolted to the C-terminus by a salt bridge between Arg53 and Asp88 (cartoon representation, with side chains shown as sticks). (B) Cartoon summarizing the important stabilizing interactions in the top-evolved variant MtCM\textsuperscript{V} depicted in (A) that properly position Arg46 for catalysis. Asp55 stabilizes the stretched-out conformation of Arg46, whereas alternating salt bridges accessible for Arg53 with the negatively charged groups present in the C-terminal region hinder Arg46 from adopting an unfavorable interaction with the C-terminal carboxylate. One example of an alternative backbone conformation that allows for interactions between the C-terminal carboxylate and Arg53 is depicted with dashed outlines.
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.biochem.2c00635.

Crystal structures and electron density maps shown for catalytically important regions (Figure S1); crystal contacts of the H1−H2 loop (Figure S2); root-mean-square fluctuations of MtCM during MD simulations (Figure S3); interactions in MtCM between its C-terminus and H1−H2 loop (Figure S4); MD snapshots of interactions between C-terminus and H1-H2 loop of MtCMV (Figure S5); and data collection and refinement statistics (Table S1) (PDF)

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AUTHOR INFORMATION

Corresponding Authors
Peter Kast — Department of Organic Chemistry, ETH Zürich, CH-8093 Zürich, Switzerland; orcid.org/0000-0002-0209-8975; Email: kast@org.chem.ethz.ch

Michele Cascella — Department of Chemistry and Hylleraas Centre for Quantum Molecular Sciences, University of Oslo, Oslo 0315 NO, Norway; orcid.org/0000-0003-2266-5399; Email: michele.cascella@kjemi.uio.no

Ute Krengel — Department of Chemistry and Hylleraas Centre for Quantum Molecular Sciences, University of Oslo, Oslo 0315 NO, Norway; orcid.org/0000-0001-6688-8151; Email: ute.krengel@kjemi.uio.no

Authors
Helen V. Thorbjørnsrud — Department of Chemistry and Hylleraas Centre for Quantum Molecular Sciences, University of Oslo, Oslo 0315 NO, Norway
Luca Bressan — Laboratory of Organic Chemistry, ETH Zürich, CH-8093 Zürich, Switzerland
Tamjidmaa Khatanbaatar — Department of Chemistry and Hylleraas Centre for Quantum Molecular Sciences, University of Oslo, Oslo 0315 NO, Norway
Manuel Carrer — Department of Chemistry and Hylleraas Centre for Quantum Molecular Sciences, University of Oslo, Oslo 0315 NO, Norway
Kathrin Würth-Roderer — Laboratory of Organic Chemistry, ETH Zürich, CH-8093 Zürich, Switzerland; orcid.org/0000-0002-0233-9863
Gabriele Cordara — Department of Chemistry and Hylleraas Centre for Quantum Molecular Sciences, University of Oslo, Oslo 0315 NO, Norway; orcid.org/0000-0001-8029-8043

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.biochem.2c00635

Author Contributions
H.V.T., L.B., and T.K. contributed equally to this work.

Author Contributions
U.K. conceived the study. H.V.T., P.K., and Mi.C. were additionally involved in the planning of the experiments. H.V.T. performed most of the calculations, transformed, produced, purified, and crystallized the two single MtCM variants, and solved the crystal structure of MtCMV (PDF) supervised by Mi.C. and U.K., respectively. Ma.C. contributed with additional simulations, supervised by Mi.C. T.K. solved the crystal structure of MtCMV (PDF) and refined the crystal structures of both MtCM variants, supervised by G.C. and U.K., who also validated the structures. L.B. constructed, produced, and purified additional sets of MtCM variants and characterized their kinetic parameters to validate computational results, and K.W.-R. designed and constructed the MtCM variants T52P and V55D and prepared the final figures; both were supervised by P.K. The initial version of the manuscript was written by H.V.T. and U.K., which was complemented with contributions from all authors and revised by P.K., Mi.C., and U.K.

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Notes
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ABBREVIATIONS
BTP, 1,3-bis[tris(hydroxymethyl)methylamino] propane; CM, chorismate mutase; DAHP, 3-deoxy-d-arabinose-7-phosphate; IPTG, isopropyl-β-D-thiogalactopyranoside; LC, ligand complex (i.e., complex with TSA); MD, molecular dynamics; MES, 2-(N-morpholino)ethanesulfonic acid; MtCM, chorismate mutase from M. tuberculosis; MtCMV, MtCM from MtCM–MtDS complex; MtCMs, TSA-bound MtCM from MtCM–MtDS complex (LC1 refers to only one of the protomers containing TSA); MtCMV (PDF), MtCM variant V55D, top-performing MtCM variant N-s4.15 from directed evolution study; MtCM Triple, MtCM variant with the three substitutions T52P, V55D, and L88D; MtCMV (PDF), MtCM variant V55D; MtCMV (PDF), wild-type MtCM; MtDS, DAHP synthase from M. tuberculosis; MWCO, molecular weight cutoff; pI, isoelectric point; PMSF, phenylmethanesulfonyl fluoride; RMSD, root-mean-square deviation (or root-mean-square difference, if concerning structural comparisons); RMSF, root-mean-square fluctuation; SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis; Tris, tris(hydroxymethyl)aminomethane; TSA, transition state analogue.

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