# Structure

## **Discovery, structure, and function of filamentous 3methylcrotonyl-CoA carboxylase**

### **Graphical abstract**



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### In brief

Hu et al. serendipitously discover that 3-methylcrotonyl-CoA carboxylase (MCC), an enzyme involved in leucine catabolism, forms filaments using cryogenic electron microscopy. Their work supports a new model of MCC catalysis and casts new light on the unusual structural and functional diversity of enzymatic polymerization.

### **Highlights**

- Native 3-methylcrotonyl-CoA carboxylase from *Leishmania tarentolae* forms filaments
- The structure of filamentous LtMCC was resolved up to 3.4 Å resolution
- Filamentation sequesters LtMCC in an inactive state
- A new model of LtMCC catalysis, termed the dual-swingingdomains model, is proposed







## Article

## Discovery, structure, and function of filamentous 3-methylcrotonyl-CoA carboxylase

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#### SUMMARY

3-methylcrotonyl-CoA carboxylase (MCC) is a biotin-dependent mitochondrial enzyme necessary for leucine catabolism in most organisms. While the crystal structure of recombinant bacterial MCC has been characterized, the structure and potential polymerization of native MCC remain elusive. Here, we discovered that native MCC from *Leishmania tarentolae* (LtMCC) forms filaments, and determined the structures of different filament regions at 3.4, 3.9, and 7.3 Å resolution using cryoEM.  $\alpha_6\beta_6$  LtMCCs assemble in a twisted-stacks architecture, manifesting as supramolecular rods up to 400 nm. Filamentous LtMCCs bind biotin non-covalently and lack coenzyme A. Filaments elongate by stacking  $\alpha_6\beta_6$  LtMCCs onto the exterior  $\alpha$ -trimer of the terminal LtMCC. This stacking immobilizes the biotin carboxylase domains, sequestering the enzyme in an inactive state. Our results support a new model for LtMCC catalysis, termed the dual-swing-ing-domains model, and cast new light on the function of polymerization in the carboxylase superfamily and beyond.

#### INTRODUCTION

3-methylcrotonyl-CoA carboxylase (MCC) is a biotin-dependent enzyme necessary for the catabolism of leucine, <sup>1–3</sup> an essential branched-chain amino acid involved in regulating cellular metabolism, <sup>4,5</sup> protein synthesis, <sup>4,5</sup> and anabolic signaling. <sup>6,7</sup> MCC belongs to a superfamily of biotin-dependent carboxylases with different substrate preferences, such as acetyl-CoA carboxylase (ACC), geranyl-CoA carboxylase (GCC), propionyl-CoA carboxylase<sup>8</sup> (PCC), and pyruvate carboxylase (PC). In eukaryotes, MCC resides in the mitochondrial matrix.<sup>9</sup> Documented in many species, <sup>10–12</sup> MCC shares high cross-species sequence homology (Figure S1). In humans, upregulated MCC expression often correlates with various cancers. <sup>4,13–16</sup> On the other hand, MCC deficiency, one of the most common metabolic disorders in newborns, <sup>17</sup> may cause vomiting, seizures, and other neurological abnormalities. <sup>18,19</sup>

As part of the mitochondrial leucine degradation pathway, MCC catalyzes the conversion of 3-methylcrotonyl-CoA to 3-methylglutaconyl-CoA.<sup>11</sup> Upon covalent biotinylation of the apoenzyme, MCC accelerates two successive reactions: biotin

carboxylation and carboxyl group transfer.<sup>20,21</sup> First, biotin is carboxylated at an  $\alpha$ -subunit active site with bicarbonate as the carbon dioxide donor upon concomitant ATP hydrolysis.<sup>9,20,21</sup> Next, the carboxylated biotin is translocated to the corresponding  $\beta$ -subunit active site, where the carboxyl group is transferred from biotin to 3-methylcrotonyl-CoA.<sup>9,20,21</sup>

Structure

The atomic structure of bacterial MCC has been solved by X-ray crystallography using recombinant *Pseudomonas aeruginosa* MCC (PaMCC<sup>rec</sup>) expressed in *Escherichia coli* with Histags to the  $\beta$ -subunits.<sup>22</sup> PaMCC<sup>rec</sup> subunits oligomerize into a dodecameric complex with a core of six  $\beta$ -subunits sandwiched by two  $\alpha$ -trimers in an  $\alpha_6\beta_6$  architecture.<sup>22</sup> Whether MCCs may exist in other forms is unclear. Albeit, their supramolecular assembly was conjectured based on rod-shaped aggregations of *Achromobacter* IVS MCC observed by negative-stain electron microscopy.<sup>23</sup> Recent advances enabled by cryogenic electron microscopy (cryoEM) have uncovered unexpected modes of enzyme polymerization and elucidated the regulatory roles of such architectural forms.<sup>24–29</sup> For example, high-resolution cryoEM structures illuminated the regulatory functions of several filamentous forms of eukaryotic ACC.<sup>30</sup> In the absence of high-



#### Table 1. CryoEM map statistics

	α-Subunit-centered map	$\beta$ -Subunit-centered map	Filament termini map
Data collection and processing			
Magnification	81,000	81,000	81,000
Voltage (kV)	300	300	300
Electron exposure (e <sup>-</sup> /Ų)	40	40	40
Defocus range (µm)	-1.5 to -2.5	-1.5 to -2.5	-1.5 to -2.5
Pixel size (Å)	1.1	1.1	1.1
Symmetry imposed	D3	D3	C3
Initial particle images (no.)	51,322	51,322	8,992
Final particle images (no.)	7,582	13,250	3,380
Map resolution (Å)	3.9	3.4	7.3
Fourier shell correlation (FSC) threshold	0.143	0.143	0.143
Estimated resolution range (Å)	3–5	3–5	6–8
Map sharpening B factor (Å <sup>2</sup> )	-112	-85	-153

resolution structures of the native MCC enzyme, it remains unsettled whether MCCs can similarly form supramolecular assemblies.

Here, we used a bottom-up structural proteomics cryoEM approach<sup>31,32</sup> to identify and determine the 3.4 Å, 3.9 Å, and 7.3 Å resolution structures (Tables 1 and 2) of native *Leishmania tarentolae* MCC (LtMCC) enriched from mitochondrial fraction by streptavidin affinity pull-down. We discovered that LtMCC filaments consist of stacks of LtMCC  $\alpha_6\beta_6$  dodecamers. The biotin carboxylase (BC) domains of adjacent  $\alpha_6\beta_6$  dodecamers bind

Table 2. LtMCC model statistics				
Model refinement				
Model composition	PDB: 8F3D	PDB: 8F41		
Non-hydrogen atoms	57,846	57,846		
Protein residues	7464	7464		
Ligands	BTI: 6	BTI: 6		
Mean isotropic B factor (Å <sup>2</sup> )				
Protein	70.80	70.80		
Ligand	150.43	150.43		
RMS deviations				
Bond lengths (Å)	0.006	0.005		
Bond angles (°)	1.094	1.089		
Validation				
MolProbity score	2.36	2.34		
Clashscore	21.50	20.46		
Rotamer outliers (%)	0.39	0.39		
Ramachandran statistics				
Outliers (%)	0.00	0.00		
Allowed (%)	9.34	9.35		
Favored (%)	90.66	90.65		
Model-to-map fit, CC (mask)				
α-Subunit-centered map	N/A	0.55		
β-Subunit-centered map	0.66	N/A		

with each other, inhibiting their mobility. In contrast, at the filament termini, the BC domain is flexible, while the linker between the biotin carboxyl carrier protein (BCCP) and the BC-CT mediating (BT) domain is rigid. These observations support a dualswinging-domains model for LtMCC catalysis and suggest a regulatory role of LtMCC filamentation in the mitochondrial matrix.

#### RESULTS

## Discovery and identification of LtMCC filaments from mitochondrial lysate

While examining complexes enriched by sedimentation of mitochondrial lysate from parasitic protist *L. tarentolae* in 10%–30% glycerol gradient (Figure 1A) and affinity pull-down with streptavidin-coated magnetic beads, we serendipitously discovered filamentous structures (Figure 1B). We identified the constituents of these filamentous structures by following the workflow detailed in Figures 1B–1C, which is based on the cryoID bottom-up structural proteomics approach.<sup>31</sup> Using single-particle reconstruction, we obtained two differently centered cryoEM maps of the filament middle segments, one at 3.4 Å and another at 3.9 Å resolution (Figure 1B). Additionally, we processed the termini particles and found that the two termini have the same structure, which was resolved at 7.3 Å resolution (Figure 1B).

To establish the protein identities of the cryoEM maps, we first obtained a partial model prediction from DeepTracer<sup>32</sup> using the 3.9 Å map. The sequence of the output model from DeepTracer<sup>32</sup> was then input into cryoID (Figure 1C). Searching for proteins detected by mass spectrometry (Table S1), cryoID identified LtMCC  $\alpha$ -subunits as a filament component. Upon visual inspection, it was clear that the filaments also contain LtMCC  $\beta$ -subunits. Indeed, LtMCC  $\alpha$ - and  $\beta$ -subunits produced the most peptide spectral counts of all proteins detected by mass spectrometry (Table S1). Furthermore, we ruled out GCCs as the filament components because GCCs are not found in eukaryotes.<sup>36</sup> We built the atomic model for the filament *de novo*, using an Alpha-Fold<sup>34,35</sup> predicted model for the  $\alpha$ -subunit and the  $\beta$ -subunit as references. We found that both the 3.4 Å and 3.9 Å maps contain the same subunits but center on the







#### Figure 1. Discovery of filamentous 3-methylcrotonyl-CoA carboxylase from mitochondria of L. tarentolae

(A) Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) of the glycerol gradient fractions obtained from mitochondrial lysate. The gel was stained by Sypro Ruby.

(B) Workflow for cryoEM imaging and reconstruction. Filaments observed in cryoEM micrographs were visually identified, partitioned into segments of equal length, and classified in 2D and 3D. Refinement of 3D reconstructions yielded an  $\alpha$ -centered and  $\beta$ -centered cryoEM map. The filament ends were picked and reconstructed similarly, converging into a single map for both ends of the filament (i.e., the termini map).

(C) Workflow for protein identification and atomic modeling. From an input cryoEM map, DeepTracer<sup>32</sup> generated an initial model with predicted amino acid sequence, which was input into cryoID<sup>31</sup> together with mass spectrometry data. CryoID predicted that the filaments contained LtMCC. Finally, *de novo* atomic modeling of LtMCC was performed in Coot<sup>33</sup> using an Alpha-Fold predicted model as a reference.<sup>34,35</sup>

 $\beta$ - and  $\alpha$ -subunits, respectively. After translating one map for half of an  $\alpha_6\beta_6$  stack along the filament and a slight rotation around the filament axis, the two maps match well with one another (cross-correlation coefficient of 0.74). The slightly better resolution of the map centered on the  $\beta$ -subunits is probably because the architecture of the  $\beta$ -subunits is more stable.

#### Structure of native eukaryotic LtMCC filaments

LtMCC filaments are predominantly straight (Figure 2A), although some are curved (Figure 2B), with a maximum observed curvature reaching 3.1  $\mu$ m<sup>-1</sup>. The length distribution resembles a Poisson distribution (Figure 2C), similar to that of ACC filaments isolated from animal tissues and observed by

negative-stain electron microscopy.<sup>37</sup> Most filaments consist of four to six  $\alpha_6\beta_6$  dodecamers, or stacks (Figure 2C), with a mean length of 993 Å. With an architecture similar to that of PaMCC<sup>rec</sup>, each  $\alpha_6\beta_6$  stack in the LtMCC filament is characterized by an  $\alpha_6\beta_6$  cylindrical dodecamer exhibiting D3 symmetry (Figure 2D). Measuring to a height of 216 Å and width of 148 Å, the dodecamer is composed of a core of six  $\beta$ -subunits ( $\beta$ -core) sandwiched by two layers of trimeric  $\alpha$ -subunits (Figure 2D and Video S1). Modeling the atomic structures of LtMCC  $\alpha_6\beta_6$  stacks into our cryoEM reconstruction reveals that the filament has a twisted-stacks architecture; that is, neighboring  $\alpha_6\beta_6$  stacks are related by a rotation of 23° and a translation of 216 Å (Figure 2D). In the filament, the active sites are

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Figure 2. Structure of the filamentous LtMCC See also Figures S2, S3, and S4

(A-C) CryoEM images of a straight (A) and curved (B) LtMCC filament and distribution of filament lengths (C).

(D and E) CryoEM map and atomic models of the LtMCC filament. The composite cryoEM map of the filament (left column in D) is composed of three copies of the  $\alpha$ -centered map, three copies of the  $\beta$ -centered map, and one copy of the termini map, with overlapping subunits. The atomic models (middle and right columns in D) of an  $\alpha_6\beta_6$  dodecamer and its trimeric stacks are shown in various views as ribbons, colored by domains according to the scheme in (E). The view directions are indicated by arrows, planes, and the camera. The dashed lines in the right column demarcate the boundaries between neighboring subunits ( $\beta$ -core is two lavered).

(F and G) Atomic model of an α-subunit (F) and a β-subunit (G) of LtMCC shown as ribbons colored by domains with active sites labeled. Insets: example cryoEM densities superimposed with the atomic model.

exposed in both  $\alpha$ - and  $\beta$ -subunits (indicated by stars in the middle panel of Figure 2D).

Each  $\alpha$ -subunit contains the BC, BT, and BCCP domains and two linkers: the BC-BT linker and the BT-BCCP linker, of which the latter was unmodeled in PaMCC<sup>rec</sup> (Figure 2E). A prominent feature of the BC domain is the presence of two large central  $\beta$ -sheets surrounded by  $\alpha$ -helices. The active site of each  $\alpha$ -subunit is located inside the central pocket of the BC domain (Figure 2F) and is responsible for carboxylating BCCP-bound biotin. In contrast to the BT domain of PaMCC<sup>rec</sup>, which contains a central  $\alpha$ -helix accompanied by a seven-stranded antiparallel  $\beta$ -barrel,<sup>22</sup> the LtMCC BT domain features eight  $\beta$ -strands (Figure S2). When the dodecameric PaMCC<sup>rec</sup> and LtMCC are superimposed globally, the BC, BT, and BCCP domains in LtMCC are rotated with respect to those in PaMCC<sup>rec</sup> (Figures S3A and S3B). However, when individual domains are





Figure 3. Presence of non-covalently bound biotin and lack of CoA in the LtMCC filament See also Figure S5.

(A) Close-up view of the middle column of Figure 2D, showing the biotin-containing site surrounded by the CT-linker (purple), the BCCP (red), N-CT (turquoise), and C-CT (green) domains. The conserved Lys651 (red) and the non-covalently bound biotin (black) are shown as sticks, superimposed with their cryoEM densities (semi-transparent gray). The distance between the carboxyl group of biotin and the e-amino group of Lys651 is about 10 Å. Notably, the CoA binding site (pink circle region) lacks cryoEM density, indicating absence of CoA.

(B) Comparison of biotin and CoA orientations documented in various carboxylases.<sup>22,38-41</sup> The red star indicates the location of the N1' atom of biotin.

superimposed independently, they fit well with each other (Figures S3C-S3E), suggesting that the above noted structural differences between the LtMCC filament and PaMCCrec are likely due to filamentation, although we cannot rule out genuine species-specific differences.

Like in PaMCC<sup>rec</sup>, each LtMCC β-subunit possesses an N-carboxyltransferase (N-CT) domain and a C-carboxyltransferase (C-CT) domain. We have designated two new domains in LtMCC: the N-dock domain and CT-linker (Figure 2E), located before and after the N-CT domain, respectively. The N-dock domain was introduced to facilitate description of interactions between  $\alpha$ - and  $\beta$ -subunits in the filament (see below). Designating the helix-loop-helix-loop fragment between the N-CT and C-CT domains as a separate domain (CT-linker) renders the N-CT and C-CT domains to have near-identical backbone folds (Figure S4), despite sharing only 19% sequence identity. Each of the six  $\beta$ -core active sites is wedged at the N-CT and C-CT interface between a top and a bottom  $\beta$ -subunit (Figures 2G and 3A) and is responsible for carboxylating 3-methylcrotonyl-CoA. Similar to that in PaMCCrec, each LtMCC β-core active site is roughly 80 Å away from its corresponding  $\alpha$ -subunit active site.

#### LtMCC filaments bind biotin but lack coenzyme A

PaMCCrec was crystalized as a holoenzyme (with bound CoA and covalently linked biotin) and an apoenzyme (without CoA and biotin). By contrast,  $\alpha_6\beta_6$  stacks in the LtMCC filament contain non-covalently-bound biotin but lack CoA (Figures 3A and S5). For LtMCC to perform its two-step catalytic process, covalent biotinylation must occur first. Although biotin is

observed at the expected  $\beta$ -core active sites, it is untethered to Lvs651 in the highly conserved Ala-Met-Lvs-Met sequence in the BCCP domain. Indeed, the carboxyl group of biotin is about 10 Å apart from the  $\varepsilon$ -amino group of Lys651 (Figure 3A), which is too far to form a prosthetic group.<sup>21</sup> The existence of non-biotinylated BCCP domains in LtMCC filaments suggests that the filamentous LtMCC is inactive. Similarly, inactive forms of PCC also bind biotin without forming a covalent adduct.<sup>42,43</sup>

The bicyclic ring system in biotin consists of an ureido ring and a thiophane ring.44 Carboxylation requires the transfer of the carboxyl group attached to the N1' atom of the biotin ureido ring to acyl-CoA.45 Therefore, the orientation of the N11 atom of biotin is important for carboxylation. In LtMCCs, the N1' atom in the ureido ring of biotin is proximal to the CoA binding site, but not in PaMCCrec (Figure 3A). This is because the biotin bicyclic ring in LtMCC is approximately 180° rotated relative to that of PaMCCrec (Figure 3B). The spatial arrangement of LtMCC biotin is consistent with PCC biotin38,39 and PC biotin<sup>40</sup> (Figure 3B). In contrast, the arrangement of PaMCC<sup>rec</sup> biotin is consistent with ACC biotin<sup>22,41</sup> (Figure 3B). Therefore, biotin orientation is variable at the  $\beta$ -core active site for different carboxylases and carboxylases in different catalytic states.

#### $\alpha$ -Subunits associate more with $\beta$ -subunits than with intra-trimer *a*-subunits

Multiple interactions are observed between an a-subunit and the  $\beta$ -core, with 1,515 Å<sup>2</sup> of buried surface area, as calculated by ChimeraX,<sup>46</sup> and a binding affinity of -11.7 kcal mol<sup>-1</sup>, as calculated by PRODIGY.<sup>47,48</sup> Interactions between each α-subunit and the

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Figure 4. Subunit interactions in the filamentous LtMCC See also Figure S6.

(A–D) Interacting regions are boxed and examined in close-up views with their interfacial residues labeled. Interactions between  $\alpha$ - and  $\beta$ -subunits (A and B) are more extensive than interactions among  $\alpha$ -subunits (C and D).

β-core are mainly mediated by a hook between the central α-helix and β1 strand of one BT domain that curls into a groove of a β-subunit (yellow segment from Asp494–Thr506 in Figure 4A). This groove is formed by the N-CT, C-CT, and N-dock domains (Figure S6). Due to its shorter sequence, the LtMCC hook contains only one hairpin in contrast to two hairpins in the PaMCC<sup>rec</sup> hook (Figure S2). In both structures, the hook is crucial for interactions with the β-subunit to reinforce inter-subunit associations.<sup>22</sup> The additional structures resolved in our cryoEM map also unveil previously unrecognized associations among domains of three neighboring subunits: the BT-BCCP linker of an α-subunit, N-dock domain of a β-subunit, and N-CT domain of another β-subunit (Figure 4B). Here, the BT-BCCP linker "docks" onto the crescent N-dock domain (Figure 4B).

Fewer interactions are observed between intra-trimer  $\alpha$ -subunits, contributing just 579 Å<sup>2</sup> of buried surface area and a binding affinity of -6.1 kcal mol<sup>-1</sup>. Within an  $\alpha$ -trimer, each  $\alpha$ -subunit interacts with its neighboring  $\alpha$ -subunits through the BC-BC and BC-BT interfaces. At the BC-BC interface, a loop from one BC domain (Glu34–His36) interacts with a loop (Phe379–Pro385) and a helix (Arg423–Gln427) from a neighboring BC domain (Figure 4C). At the BC-BT interface,  $\beta 6$  strand (Gly572–Ala581) from the  $\beta$ -barrel in the BT domain interacts with a short  $\beta$ -strand (Ala59–Cys61) in the BC domain through  $\beta$ -sheet augmentation<sup>49</sup> (Figure 4D).

#### Structure of filament termini reveals mechanism of LtMCC polymerization

To investigate the mechanism of LtMCC polymerization, we reconstructed the filament termini at 7.3 Å resolution (Figure 5A). We found that both ends of the filament terminate at  $\alpha$ -trimers, with partially resolved densities attributed to the BC domains (Figure 5A). When displayed at a high threshold, cryoEM densities for the BC, BCCP domain, and the BC-BT linker disappear, while density for the BT domain and BT-BCCP linker of the same  $\alpha$ -subunit remain visible (Figure 5A). Therefore, the weaker densities of the BC and BCCP domains are due to their flexibility, rather than low occupancy of the  $\alpha$ -subunit at the termini, which would make all domains of the  $\alpha$ -subunit disappear. The flexibility of the BC and BCCP domains is further supported by limited intra-trimer  $\alpha$ -subunit associations (Figures 4C and 4D). In contrast, the BT-BCCP linker is mostly rigid as it interacts extensively with the  $\beta$ -subunits (Figure 4B).

Our observation that LtMCC filaments terminate at a-subunits suggests that the building block for the filament is an  $\alpha_6\beta_6$  stack. Filaments plausibly elongate one stack at a time, attaching through BC domains to form an  $\alpha$ - $\alpha$  interface (Video S2), which is defined as the associations between two  $\alpha$ -trimers from adjacent stacks (Figure 5B). The  $\alpha$ - $\alpha$  interface has a total buried surface area of 1,689 Å<sup>2</sup> and a binding affinity of -11.6 kcal mol<sup>-1</sup>. At the  $\alpha$ - $\alpha$  interface, the BC domain of an  $\alpha$ -subunit from the lower  $\alpha$ -trimer interacts with BC domains of two  $\alpha$ -subunits from the upper  $\alpha$ -trimer (Figure 5B). Interactions contributing to the  $\alpha$ - $\alpha$  interface occur primarily between loops, along a local 2-fold axis (right panel of Figure 5B): the first site of interaction is between two loops (Leu338-Asp344), with its neighboring residues Lys319 and Arg231 also participating in  $\alpha$ - $\alpha$  interactions; the second site of interaction occurs between two loops connected to two a-helices (His257-Glu261). These interactions at the  $\alpha$ - $\alpha$  interface stabilize the BC domains between neighboring stacks in the LtMCC filament (Figure 5B).

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#### DISCUSSION

This study reveals a previously unrecognized role of the BC domain in MCC catalysis. During catalysis, the BCCP domain must shuttle biotin between the CT active site ( $\beta$ -subunit) and the BC active site (α-subunit), located approximately 80 Å apart (Figure 5A). However, we observed that the BT-BCCP linker is fixed (Figure 5A). Therefore, the maximum distance that the BCCP domain can translocate by itself is about 60 Å, which is insufficient for the BCCP domain to reach the BC active site (right panel of Figure 5A). As such, for catalysis to occur, both the BCCP and active-site-containing BC domain must move so that BCCP-bound biotin can reach the BC active site. Indeed, at the filament termini, the BC domain is flexible, thus permitting BC domain movement. We term this the dual-swinging-domains model to highlight the difference from the original swingingdomain model,<sup>9</sup> in which only the BCCP domain swings. Beyond MCC, both the BCCP and BC domains have been documented to move in PC.<sup>50,51</sup> These observations suggest that carboxylase catalysis, in general, may involve a dual-swinging-domains mechanism.

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#### Figure 5. Filament formation stabilizes flexible BCCP and BC domains in LtMCC

(A) Structure of the filament termini. The left and middle panels show the 7.3 Å cryoEM map of the filament termini at a low and high density threshold, respectively. The  $\alpha$ -subunit is colored by domain according to the scheme in Figure 2E, and the entire  $\beta$ -core is turquoise. Note, while visible at low threshold, the BCCP and BC domain densities disappear at high threshold. The right panel shows the atomic model of the filament terminus with the rigid and flexible domains in harder and softer shades, respectively.

(B) Structure of the  $\alpha$ - $\alpha$  interface in the LtMCC filament. The cryoEM map (left panel) is colored as in (A), but the atomic models (middle panel) of the interfacing  $\alpha$ -subunits are colored differently from each other to delineate individual  $\alpha$ -subunits. Close-up views (right panel) of the two boxed areas in the middle panel show interfacing residues at the  $\alpha$ - $\alpha$  interface. The ellipse in the top right panel denotes the local 2-fold symmetry axis.

(C) Proposed model of filament formation and stabilization. Blurry domains indicate their flexibility.

In LtMCC filaments, the  $\alpha$ - $\alpha$  interface fixes the BC domain, thus preventing its swinging. According to our dual-swinging-domains model, immobilization of the BC domain should inhibit catalysis. Granted, the BC domain and the BT-BCCP linker in LtMCC filaments may be more flexible in a cellular environment, and cryoEM reconstructions did not capture flexible states. However, as biotin is non-covalently bound to filamentous LtMCC, LtMCC must be inactive (Figure 3A).

Therefore, filamentation likely sequesters LtMCCs in a quiescent state in the mitochondrial matrix (Figure 5C), which may provide a capacity to reactivate the enzyme in response to environmental signals, such as an influx of nutrients, which may trigger covalent attachment of biotin to the BCCP domain through biotin protein ligase. Polymerization appears to be a readily deployable and economic mechanism for regulating the activity of abundant mitochondrial matrix enzymes in response to rapid changes in cellular metabolism requirements. Such capacity is particularly relevant for parasitic protists cycling between the amino acid-rich gut environment of insect vectors and the high glucose concentration in the bloodstream of their hosts.

Our inquiries into MCC polymerization further reflect the diversity of polymerization in the carboxylase superfamily. It is known that both isoforms of ACC, ACC1 and ACC2, form filaments.<sup>30,37,52-56</sup> Recently, high-resolution structures of filamentous ACC1 have been reported.<sup>30</sup> In contrast to LtMCC, ACC1 assembles into two filamentous forms: an active single-stranded and inactive double-stranded helical structure. Eukaryotic MCC and ACC filamentation exhibit different



architectures and regulatory functions, supporting the distinct evolutionary lineages of these enzymes.<sup>22</sup> Nonetheless, filament formation may be a general way to regulate enzymatic activity in the carboxylase superfamily. We suspect that MCC filamentation is not unique to LtMCC, since the BCfo domain of LtMCC, which includes the  $\alpha$ - $\alpha$  interface, generally shares a high (>50%) sequence identity with BC domains from other organisms.

Evolutionarily, employing the same gene costs less than "inventing" a new gene for regulation. Polymerization is a remarkable example of such a cost-effective strategy. In the case of LtMCC, stacking of active  $\alpha_6\beta_6$  dodecamers can inhibit biotin shuttling (Figure 5C), thus sequestering the enzyme by polymerization. In other enzymes, polymerization can alter specificity<sup>57</sup>, decrease<sup>28</sup> or increase activity<sup>58</sup>, and even introduce non-enzymatic functions such as conducting electrons between bacterial cells<sup>27</sup> and regulating cell curvature.<sup>59</sup> Just like how complex life forms arose by the aggregation and polymerization of unicellular forms, the diverse functions arising from enzyme polymerization can be considered as a rudimentary form of evolution.

#### **STAR\*METHODS**

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- QUANTIFICATION AND STATISTICAL ANALYSIS

#### SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.str. 2022.11.015.

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#### **AUTHOR CONTRIBUTIONS**

Z.H.Z. and R.A. initiated and supervised the project. I.A. prepared the sample. C.Y. and L.H. conducted mass spectrometry. Y-T.L. carried out cryoEM imaging and data processing with assistance from J.K.J.L. and J.J.H. J.K.J.L. and J.J.H. built the atomic models. J.K.J.L., J.J.H., and Y-T.L. analyzed the data. J.J.H., J.K.J.L., and Y-T.L. made illustrations and wrote the original draft. Z.H.Z., Y-T.L., J.J.H., and J.K.J.L. finalized the manuscript. All authors reviewed and approved the paper.

#### **DECLARATION OF INTERESTS**

The authors declare no competing interests.

#### **INCLUSION AND DIVERSITY**

One or more of the authors of this paper self-identifies as a gender minority in their field of research. While citing references scientifically relevant for this work, we also actively worked to promote gender balance in our reference list. We avoided "helicopter science" practices by including the participating local contributors from the region where we conducted the research as authors on the paper. We support inclusive, diverse, and equitable conduct of research.

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#### **STAR**\***METHODS**

#### **KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, peptides, and recombinant proteins		
Complete protease cocktail tablet	Roche	Sigma product no: 11836170001
Deposited data		
Crystal structure of PaMCC <sup>rec</sup> apoenzyme	Huang et al. <sup>22</sup>	PDB: 3U9T
Crystal structure of PaMCC <sup>rec</sup> holoenzyme	Huang et al. <sup>22</sup>	PDB: 3U9S
Crystal structure of PCC D422-mutant $\beta$ -subunit	Arabolaza et al. <sup>42</sup>	PDB: 3IB9
Crystal structure of PCC wild-type β-subunit	Diacovich et al. <sup>43</sup>	PDB: 1XNY
CryoEM structure of ACC-Citrate filament	Hunkeler et al. <sup>30</sup>	PDB: 6G2D
CryoEM structure of ACC–BRCT filament	Hunkeler et al. <sup>30</sup>	PDB: 6G21
Crystal structure of ACC holoenzyme	Wei et al. <sup>41</sup>	PDB: 5CSL
Crystal structure of PCC holoenzyme	Huang et al. <sup>38</sup>	PDB: 3N6R
CryoEM map of α-subunit centered LtMCC	This paper	EMDB: EMD-28849
CryoEM map of β-subunit centered LtMCC	This paper	EMDB: EMD-28846
CryoEM map of LtMCC termini	This paper	EMDB: EMD-28847
CryoEM structure of the LtMCC filament centered on $\alpha$ -subunits	This paper	PDB: 8F41
CryoEM structure of the LtMCC filament centered on $\beta$ -subunits	This paper	PDB: 8F3D
Experimental models: Organisms/strains		
in axenic culture in various laboratories. Parasite culture used in this study was received from Prof. Larry Simpson laboratory at UCLA in 2004 and transfected with pX based plasmid expressing RNA editing ligase 1 (KREL1).		
Software and algorithms		
MotionCorr2	Zheng et al. <sup>61</sup>	https://emcore.ucsf.edu/ucsf-software
Gctf	Zhang, <sup>62</sup>	https://github.com/JackZhang-Lab/GCTF
RELION 3.1	Zivanov et al. <sup>63</sup>	https://www3.mrc-lmb.cam.ac.uk/ relion/index.php/Main_Page
ChimeraX	Goddard et al. <sup>46</sup>	https://www.cgl.ucsf.edu/chimerax/
Coot	Emsley et al. <sup>33</sup>	https://www2.mrc-Imb.cam.ac.uk/ personal/pemsley/coot/
Phenix	Liebschner et al. <sup>64</sup>	https://phenix-online.org/
DeepTracer	Pfab et al. <sup>32</sup>	https://deeptracer.uw.edu/
cryoID	Ho et al. <sup>31</sup>	https://github.com/EICN-UCLA/cryoID
ESPript 3	Robert and Gouet, <sup>65</sup>	https://espript.ibcp.fr/ESPript/ESPript/
AlphaFold	Jumper et al., Mirdita et al. <sup>34,35</sup>	https://github.com/sokrypton/ColabFold
IMOD	Kremer et al. <sup>66</sup>	https://bio3d.colorado.edu/imod/
SerialEM 3.8	Mastronarde, <sup>67</sup>	https://bio3d.colorado.edu/SerialEM/
ExPASy Translate	Gasteiger et al.68	https://web.expasy.org/translate/
Clustal Omega	Sievers and Higgins <sup>69</sup>	https://www.ebi.ac.uk/Tools/msa/clustalo/
PRODIGY	Vangone and Bonvin, Xue et al. <sup>47,48</sup>	https://wenmr.science.uu.nl/prodigy/
ResMap	Kucukelbir et al. <sup>70</sup>	http://resmap.sourceforge.net/

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Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Other		
MagStrep XT beads	IBA Lifesciences	2-4090-002
Zeba™ Spin Desalting Columns	Thermo Fisher Scientific	Catalog number: 89882

#### **RESOURCE AVAILABILITY**

#### Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Z. Hong Zhou (hong.zhou@ucla.edu).

#### **Materials availability**

This study did not generate new unique reagents.

#### Data and code availability

CryoEM maps of the  $\alpha$ -subunit centered,  $\beta$ -subunit centered, and filament termini map have been deposited in the Electron Microscopy Data Bank under accession numbers EMD-28849, EMD-28846, and EMD-28847, respectively. The coordinates of LtMCC have been deposited in the Protein Data Bank under accession number 8F3D and 8F41. All aforementioned deposited data are publicly available as of the date of publication and accession numbers are also listed in the key resources table. This paper does not report original code. DOIs are listed in the key resources table. This paper analyzes existing, publicly available data. These accession numbers for the datasets are listed in the key resources table. Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

#### **EXPERIMENTAL MODEL AND SUBJECT DETAILS**

*L. tarentolae* cells were grown at 27°C in brain heart infusion media supplemented with 5 mg/L of hemin and harvested during late-exponential growth phase at  $\sim 2 \times 10^8$  cell/mL. The cells are from *L. tarentolae* TATII/UC strain.

#### **METHOD DETAILS**

#### **Preparation of mitochondrial lysate**

Mitochondrial fraction was enriched by hypotonic cell lysis and sequential separation of membrane-containing fraction on RenoCal76 density gradients.<sup>71</sup> Mitochondrial pellets (0.5–0.8 g wet weight) were lysed in 1 mL of pH 7.3, 50 mM HEPES, 150 mM KCl, 2 mM EDTA, 1% NP40, and 50  $\mu$ L of 20x Complete protease cocktail by sonication at 24W for 15 seconds and centrifuged at 30,000 RPM in an SW55 rotor for 15 minutes. The supernatant was recovered and separated on a continuous 10–30% gradient glycerol in pH 7.3, with 20 mM HEPES, 100 mM KCl, and 1 mM EDTA and prepared in SW28/32 Setton clear tubes for 15 hours at 72,000 g. Glycerol gradient fractions of 1.5 mL were collected from the top and those corresponding to 20S-40S region were combined.

#### Purification of LtMCC by streptavidin affinity pulldown

Glycerol gradient fractions were supplemented octylglucoside to 2 mM and incubated on Strep-Tactin®XT magnetic beads in a Binding Buffer (50 mM Tris-HCl, pH 8.0, 100 mM KCl, 1 mM EDTA, 2 mM OG) for 1 hour at 4°C on a nutating mixer. Beads were washed with 5 mL of Binding Buffer twice. For elution, the beads were incubated in 0.2 mL of Elution Buffer (20 mM Tris-HCl, pH 8.0, 100 mM KCl, 1 mM EDTA, 2 mM OG, 100 mM biotin) at 4°C for 10 minutes. The 130 µL of purified material was exchanged into Sample Buffer (20 mM Tris-HCl, pH 7.5, 60 mM KCl, 5 mM MgCl<sub>2</sub>, 1 mM DTT, 5 mM OG) using Zeba<sup>™</sup> Spin Desalting Columns, 7K MWCO (0.5 mL). The sample was then centrifuged at 21,000g for 10 minutes and the supernatant was stored on ice before grid preparation.

#### Protein identification by LC-MS/MS

Affinity-purified complexes were sequentially digested with LysC peptidase and trypsin. LC-MS/MS was carried out by nanoflow reversed phase liquid chromatography (RPLC) using an UltiMate 3000 RSLC (Thermo Fisher Scientific) coupled on-line to an Orbitrap Fusion Lumos mass spectrometer (Thermo Fisher Scientific). A cycle of full FT scan mass spectrum (m/z 375–1500, resolution of 60,000 at m/z 400) was followed by MS/MS spectra acquired in the linear ion trap for 3 seconds at top speed with normalized collision energy (HCD, 30%). Following data extraction to MGF format using MSConvert from ProteoWizard,<sup>72</sup> resultant peak lists for each LC-MS/MS experiment were submitted to Protein Prospector (UCSF) for database searching.<sup>73</sup> Each project was searched against a normal form concatenated with the random form of the *L. tarentolae* Parrot Tar II from TriTrypDB.<sup>74</sup> The mass accuracy for parent



ions and fragment ions were set as  $\pm$  10 ppm and 0.6 Da, respectively. Trypsin was set as the enzyme, with a maximum of two missed cleavages allowed. Cysteine carbamidomethylation was set as a fixed modification, and protein N-terminal acetylation, methionine oxidation, and N-terminal conversion of glutamine to pyroglutamic acid were selected as variable modifications.

#### **CryoEM sample preparation**

Lacey carbon cryoEM grids with a 2 nm continuous carbon film (Ted Pella) were first glow discharged for 45 seconds with a target current of 15 mA, using PELCO easiGlowTM. Then a 2.5 µL sample was applied to the grids. After waiting for 5 seconds, the grids were blotted for 4 seconds with blot force 0, at 100% humidity and 4°C temperature; after blotting, the grids were plunge-frozen into liquid ethane using an FEI Mark IV Vitrobot (Thermo Fisher Scientific). The grids were stored in a liquid nitrogen dewar before imaging.

#### **CryoEM** image acquisition

The cryoEM grids were loaded and imaged using a Titan Krios (Thermo Fisher Scientific) operated at 300 kV, with a Gatan K3 camera and a Gatan Imaging Filter Quantum LS. Movies were recorded with SerialEM<sup>67</sup> by electron counting in super-resolution mode at a pixel size of 0.55 Å/pixel. The exposure time was 2 seconds and fractionated to 40 frames. The defocus range was between -1.5 to  $-2.5 \mu m$ . The total dosage was approximately 40 electrons/Å<sup>2</sup>. A total of 3,328 movies were collected.

#### **CryoEM** image processing

The movies were processed with MotionCor2,<sup>61</sup> leading to dose-weighted and drift-corrected electron micrographs with a calibrated pixel size of 1.1 Å. We discarded the first frame due to severe drift of this frame. The defocus of the micrographs was determined by Gctf.<sup>62</sup> Using IMOD,<sup>66</sup> the micrographs were binned by a factor of 10 to make a stack for particle picking. After directly observing filaments on the micrographs, we manually picked the start and end of each filament using the 3dmod command in IMOD.<sup>66</sup> A total of 4,496 filaments were picked. To derive the constituents of the filaments, we coded a Python script to subdivide the filaments into equally spaced segments of 87 Å. The coordinates were then converted through a Bash script to RELION format. A total of 51,322 particles were extracted in total through RELION 3.1, with a box size of 384.

After extraction, we performed reference-free 2D class averaging on the particles with RELION 3.1.<sup>66</sup> We chose the best classes and proceeded to 3D classification. We made an initial 3D model from the best 3D class averages through RELION 3.1. The 3D classification was done with D3 symmetry and yielded  $\alpha$ - and  $\beta$ -centered 3D class averages. After removing duplicates, the  $\alpha$ - and  $\beta$ -centered classes contained 7,582 and 13,250 particles, respectively. We did 3D refinement for the  $\alpha$ - and  $\beta$ -centered map independently with D3 symmetry using RELION 3D refinement. We then performed CTF refinement for both maps independently using RELION 3.1. We used CTF refinement<sup>63</sup> to improve the accuracy of the defocus for each particle and to correct beam tilt for the dataset. After performing CTF refinement, we performed 3D refinement for both maps again, independently. At the end of all processing, based on RELION's gold-standard Fourier shell correlation (FSC) at the 0.143 criterion,<sup>75</sup> we obtained  $\alpha$ - and  $\beta$ -centered maps at 3.9 and 3.4 Å resolution, respectively (Figure S7). The resolution towards the center of the map is better than the edge of the map (Figure S7).

For the filament termini map, we used RELION 3.1 to extract the previously picked filament start-and-end points at a box size of 384 pixels. We calculated the in-plane rotation angle using the direction of vectors from the start to end of the filament. We used a Python code to include the in-plane rotation angle information in the RELION star file for the particles. We performed 2D classification for these particles without local search around the precalculated in-plane rotation angle. To generate an initial 3D model, we used the IMOD clip function<sup>66</sup> to change the center of the  $\beta$ -centered map. The best particles were selected for 3D reconstruction with C3 symmetry. We performed 3D classification and removed bad particles. During 3D classification, we set the "limit\_tilt" parameter to 85 to keep side-view particles. We proceeded to 3D refinement with 3,380 particles. We obtained a final reconstruction with C3 symmetry at 7.3 Å resolution based on the gold-standard FSC at the 0.143 criterion.<sup>75</sup>

#### **Sequence determination**

The amino acid sequence for the LtMCC  $\alpha$ -subunit was obtained from gene LtaP31.3620 from TriTrypDB.<sup>74</sup> The amino acid sequence for the LtMCC  $\beta$ -subunit was obtained by analyzing the genome sequencing data of *L. tarentolae* Parrot Tar II,<sup>76</sup> since the putative, shotgun LtMCC  $\beta$ -subunit sequence on the NCBI protein database was notably shorter than MCCs from other organisms, deviating from the high sequence homogeneity observed across MCC sequences. A frameshift occurs near the middle of the sequence, likely caused by a sequencing error. The frameshift is not present in the previous genome sequencing data of *L. tarentolae* Parrot Tar II.<sup>77</sup> Since MCC plays an essential role in the catabolism of leucine across many species, it is unlikely that MCC exhibits a highly different sequence in *L. tarentolae*.

#### Atomic modeling and model refinement

An initial AlphaFold<sup>34</sup> model prediction was generated from ColabFold<sup>35</sup> using the LtMCC  $\alpha$ -subunit and  $\beta$ -subunit sequences. Atomic model building was conducted *de novo* in regions of low sequence conservation. First, an individual  $\alpha$ -subunit and  $\beta$ -subunit were modeled into the cryoEM map in Coot<sup>33</sup> and then real-space refined in Phenix.<sup>64</sup> The  $\alpha$ -subunit was modeled using the  $\alpha$ -subunit centered map, and the  $\beta$ -subunit was modeled using the  $\beta$ -subunit centered map. Biotin was modeled *de novo* into its observed density. Then, the subunits were duplicated according to D3 symmetry and fitted into a composite cryoEM map where  $\alpha$ -centered maps were fitted with  $\beta$ -centered maps as illustrated in Figure 2D. This singular atomic model, which was built using both the  $\alpha$ - and





 $\beta$ -centered maps, was refined iteratively using Phenix<sup>64</sup> and Coot.<sup>33</sup> In Coot, the atomic model was checked for rotamer outliers and Ramachandran outliers. All outliers were eliminated through the "Real Space Refine Zone" tool in Coot, with the  $\alpha$ -subunit refined against the  $\alpha$ -subunit centered map, and the  $\beta$ -subunit refined against the  $\beta$ -subunit centered map. Then, the model, along with the  $\alpha$ -subunit centered maps, were submitted to Phenix for automatic real space refinement using the "phenix.real\_space\_refine" tool. This process was performed iteratively to reduce rotamer outliers, geometric outliers, Ramachandran outliers, and clashscore. Note, we did not use the filament termini map for *de novo* atomic model building or refinement because its resolution was too low.

#### **Model analysis**

Buried surface area and map cross-correlation coefficient were calculated in ChimeraX.<sup>46</sup> Binding affinity calculations were done using the PRODIGY web server.<sup>47,48</sup> Sequence alignments and similarity calculations were performed using Clustal Omega<sup>69</sup> from the EMBL-EBI web server and visualized with ESPript 3.<sup>74</sup>

#### **QUANTIFICATION AND STATISTICAL ANALYSIS**

Filament lengths from micrographs were calculated by the IMOD model2point function.<sup>66</sup> The lengths were then counted and compiled together in a spreadsheet for histogram analysis (see Figure 2C).