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Structural insights revealed by crystal structure of B38-CAP, an isoenzyme of carboxypeptidase ACE2, the receptor of SARS-CoV-2

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Abstract

The worldwide pandemic of Coronavirus disease 2019 (COVID-19) is triggered by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) and further worsened by the emergence of a variety of SARS-CoV-2 variants. Angiotensin-converting enzyme 2 (ACE2), a carboxypeptidase of M32 family, serves as the receptor of SARS-CoV-2 and key regulator of host renin-angiotensin system (RAS), both of which are mainly mediated via the carboxypeptidase domain of ACE2 (sACE2) or its activity. sACE2 is thus promising in the treatment of COVID-19 but unfortunately weakened by its unstrigent substrate preference and complex interplay with host RAS. B38-CAP, an isoenzyme of ACE2, partially compensates these defects but still encounters the problem related to carboxypeptidase activity and specificity. In this study, we firstly determined the crystal structure of B38-CAP at a resolution of 2.44 Å which exists in dimeric form with the non-crystallographic two-fold axis being in coincidence with the crystallographic two-fold axis. Further structural analysis revealed the structural conservatism feature among M32 family, particularly the catalytic core and moreover lead us to hypothesize that conformational flexibility might play an pivotal role in the catalysis of B38-CAP and ACE2. The work provided here presents key features of the M32 family carboxypeptidase and provides structural basis for further development of B38-CAP-based anti-SARS-CoV-2 drugs.

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1. Introduction

The worldwide pandemic of Coronavirus disease 2019 (COVID-19), which is triggered by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) [1], a member of genus betacoronavirus within the subfamily Orthocoronavirinae [2], has caused about 435 million infections and more than 59 hundred thousand deaths as of March 3, 2022 [3]. To combat SARS-CoV-2 infection, a series of vaccines, neutralizing antibodies as well as small-molecule drugs have been developed and/or approved [4–8]. However, with the ongoing and repeated spreading of SARS-CoV-2 in the global world, a number of genetic variants of SARS-CoV-2 have emerged, due to its intrinsic property of being prone to genetic mutation while infecting and adapting to human hosts [9]. These variants display considerably different characteristics compared to their ancestral strains, which have been leading to the compromised efficacy of approved vaccines and probably those antibodies and small molecule drugs, hence strengthening the ever-growing demand of developing broad-spectrum drugs against SARS-CoV-2 mutants [10].

Angiotensin-converting enzyme 2 (ACE2), a single transmembrane protein characterized of a carboxypeptidase domain belonging to M32 carboxypeptidase family [11], has been thought to meet the aforementioned need of wide-spectrum anti-SARS-CoV-2 drug development due to its multiple functions in SARS-CoV-2 pathogenesis [12,13]. During the course of SARS-CoV-2 infection, ACE2 (or more specifically, its membrane-bound form) firstly serves as the receptor of SARS-CoV-2. To enter into host cells,
the receptor binding domain (RBD) of SARS-CoV-2 Spike protein needs to first bind to the carboxypeptidase domain of ACE2, immediately followed by the cleavage of Spike protein byTMPRSS2 [12,14]. In the meantime, ACE2, as a key regulator of host renin-angiotensin system (RAS), also employs its carboxypeptidase activity to reduce host inflammatory response and restrict the deterioration of SARS-CoV-2 [13,15]. This is also mediated by the carboxypeptidase domain of ACE2 through the digestion of substrate Ang II to Ang (1–7), despite a dozen of other substrates having been identified for ACE2 [16]. Thus, the soluble carboxypeptidase domain of ACE2 (sACE2) which possesses both RBD-binding (in other way, SARS-CoV-2 neutralizing) and RAS-modulating ability regardless of whether and how SARS-CoV-2 mutates, is a natural drug candidate to treat COVID-19. Currently, a series of sACE2-based anti-SARS-CoV-2 strategy are in investigation [17–19]. However, it’s not until recently that researchers find that sACE2 surprisingly mediate the cell entry of SARS-CoV-2 via interaction with proteins related to RAS [20]. This finding, in addition to indicating the antagonistic role of ACE2 in SARS-CoV-2 pathogenesis, further promotes people to re-examine the clinical complexity of ACE2.

B38-CAP, another member of M32 carboxypeptidase family and in nature the carboxypeptidase domain of ACE2’s carboxypeptidase domain, serves as a better start for developing anti-SARS-CoV-2 drug, due to its non-interacting property with RAS proteins and moreover, an extra ability of digesting Ang(1–9) into Ang(1–8) compared to ACE2 [21]. However, B38-CAP is found to display weaker carboxypeptidase activity compared to ACE2 and retain the shortcoming of unstrict substrate property of ACE2 [21]. To aids our understanding of the carboxypeptidase activity of B38-CAP, we firstly determined the three-dimensional structure of B38-CAP in this study and then performed structural comparison between B38-CAP and other M32-family carboxypeptidase, particularly ACE2. It was shown that B38-CAP which exists in dimeric form shares structural rather than sequence conservatism with ACE2 particularly in the catalytic core, while in the meantime, displays little structural flexibility in dimeric form compared to ACE2, which may have a potential effect and thus contribute to their difference in carboxypeptidase activity.

2. Materials and methods

2.1. Expression and purification

The coding sequence of B38-CAP was synthetized and cloned into the vector pGEX-6P-1 (GE Healthcare) using the BamH I and XhoI restriction sites. After verified by sequencing, the recombinant plasmid was transformed into Escherichia coli strain BL21 (DE3) for protein expression. Cultures were grown in LB medium containing 0.1 mg/mL ampicillin at 310 K until the optical density at 600 nm reached 0.8. Isopropyl-ß-D-thiogalactopyranoside (IPTG) was added to the above buffer containing 10 mM Tris-HCl, pH 8.0, 100 mM NaCl. The purifying GST-tagged B38-CAP protein. The fusion protein, after concentration of 0.5 m

The crystal structure of B38-CAP was solved by molecular replacement method in Phaser using the structure of BsuCP (PDB entry 3HQ2) as the search model [22]. The initial model was then subjected to manual rebuilt in COOT [23] and automatic refinement in PHENIX [24] iteratively until the stereochemical parameters suit the criteria of Molprobity [25]. The refinement strategy includes individual_sites refinement in real-space and reciprocal-space, atomic displacement refinement and TLS refinement. In the final stage of refinement, solvent molecules were added at a contour

2.2. Crystallization

The purified B38-CAP protein was concentrated to 20 mg/mL in the above buffer containing 10 mM Tris-HCl, pH 8.0, 100 mM NaCl. In the primary stage, commercial kits including were used to screen for preliminary crystallization conditions for B38-CAP. Crystallization trials were set in 48-well crystallization plates at 291 K using the sitting-drop vapour diffusion method. Crystallization drops were carefully set by mixing 1.0 µL protein solution with 1.0 µL of the reservoir solution and then left to equilibrate with 80 µL reservoir solution. Initial crystals of B38-CAP protein were obtained after 24 h under multiple conditions. The crystals used for diffraction data collection grew in the optimized solution containing 0.1 M Tris–HCl pH8.5, 1.0 M Sodium citrate.

2.3. Data collection and processing

B38-CAP crystals were cryoprotected in a solution containing 1.0 M Sodium citrate, 0.1 M Tris–HCl, pH 8.5 supplemented with 10% glycerol and then mounted in a nylon loop and flash-cooled in a nitrogen stream at 100 K. The X-ray diffraction data sets were collected using an R-AXIS IV++ image-plate system and an in-house rotating-anode Cu generator (Rigaku, USA) at a wavelength of 1.5418 Å. The crystals showed high-quality diffraction patterns. All intensity data were indexed, integrated and scaled with the HKL-3000 package. A complete data set diffracting to 2.0 Å was collected and the related data collection and processing statistics are summarized in Table 1.

2.4. Structure solution, refinement and analysis

The crystal structure of B38-CAP was solved by molecular replacement method in Phaser using the structure of BsuCP (PDB entry 3HQ2) as the search model [22]. The initial model was then subjected to manual rebuilt in COOT [23] and automatic refinement in PHENIX [24] iteratively until the stereochemical parameters suit the criteria of Molprobity [25]. The refinement strategy includes individual_sites refinement in real-space and reciprocal-space, atomic displacement refinement and TLS refinement. In the final stage of refinement, solvent molecules were added at a contour

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level of 3.0 σ. Structural comparison were performed using the DALI program [26]. The figures were prepared in molecular graphics system PyMOL [27].

2.5. *In vitro* carboxypeptidase activity measurements

The real-time measurement of the carboxypeptidase activity of B38-CAP and ACE2 carboxypeptidase domain was investigated using Nma-His-Pro-Lys(Dnp) as the fluorescent substrate using the EnSpire Multilabel Plate Reader (PerkinElmer, USA). The reaction mixture contained 40 μL of 0.1 M HEPES pH 7.5, containing 0.3 M NaCl, 0.01% Triton X-100, 80 μM fluorescent substrate, and 0.0125 μg of B38-CAP protein or ACE2 carboxypeptidase domain in a total volume of 50 μL. The reaction mixture was incubated at 37 °C for 1 min and then measured spectrophotometrically at an emission wavelength 440 nm upon excitation wavelength 340 nm on an EnSpire multilabel plate reader [28].

3. Results and discussion

3.1. Characterization of the oligomeric state and carboxypeptidase activity of B38-CAP

B38-CAP and ACE2 carboxypeptidase domain were expressed using *E. coli* and Bac-to-Bac baculovirus system, respectively. To investigate their oligomeric state, gel-filtration characterization using a Superdex™ 200 Increase 10/300 GL column was performed and the elution volume for B38-CAP and ACE2 carboxypeptidase domain was calibrated as 14.2 and 13.2 mL, respectively, indicating that B38-CAP, unlike ACE2 carboxypeptidase domain, exists in dimeric instead of monomeric form in solution (Fig. 1A). Their carboxypeptidase activity was then measured by the fluorescent method. It's noteworthy that an real-time mode is employed here rather than the average mode reported in earlier literature [21]. It turns out that B38-CAP displays considerably weak carboxypeptidase activity compared to ACE2 carboxypeptidase domain (Fig. 1B), raising the concern that how this can be linked to their respective structural differences.

3.2. The overall structure of B38-CAP

The crystal structure of B38-CAP which belongs to C2 space group was solved to a resolution of 2.44 Å by molecular replacement method using the program Phaser and finally refined an Rwork/Rfree factor of 15.6% and 20.0% respectively. The final structure consists of 30 α/β-helices and 3 β-sheets, with one molecule per asymmetric unit, seeming contradictory to gel-filtration profile of B38-CAP. A fine inspection of crystal packing reveals that the non-crystallographic two-fold axis between the two molecules of B38-CAP dimer is in perfect coincidence to the crystallographic two-fold axis of the C2 space group (Fig. 2A). The interface area between the two molecules constituting B38-CAP dimer is as high as 984.5 Å². An extensive hydrogen bond, salt bridge and hydrophobic interaction network exists between these two B38-CAP molecules, contributing to the stabilization of B38-CAP dimer (Fig. 2B). Inside of the structure, there is a zinc ion, which, by coordination with residues H269, H273, E273 and water molecules, forms the catalytic core (Fig. 2A). To assess the conservative property of B38-CAP, we then performed DALI structural similarity search and identified eight M32 carboxypeptidase in PDB database, all bacteria-derived, with r.m.s. deviation between them ranging from 1.1 to 2.9 Å (Table S1). Structural alignment clearly shows that B38-CAP adopts a typical M32 carboxypeptidase domain fold (Fig. 2C). This is in good accordance with the sequence comparison result, which means B38-CAP and the eight carboxypeptidases share both structural and sequence conservation (Table S1).

3.3. The structural conservation and diversity between B38-CAP and ACE2

Sequence comparison displays little, if there is, identity between B38-CAP and ACE2 carboxypeptidase domain (Fig. 3A), raising the question that whether ACE2 has evolved a brand-new model for carboxypeptidase catalysis. Considering that structural conservatry is more reliable, a DALI analysis was performed using the structure of B38-CAP monomer and ACE2 carboxypeptidase domain, which however gives an r.m.s. deviation of 5 Å with 437 Ca atoms aligned. It's surprising to find that further inspection of their fine structure reveals that the catalytic core of B38-CAP and ACE2 are completely conservative between each other and so is the spatial configuration of the structural elements adjacent to catalytic core (Fig 3B). This strongly indicates that the catalytic core of M32 family carboxypeptidases was strictly reserved during their evolution process, which however poses a confusion for us to understand the differences displayed in their carboxypeptidase activity, until we surprising find that all and only bacteria-derived carboxypeptidase adopts dimeric form. As shown in Fig. 4, the dimeric property is strictly conservative among B38-CAP and the above eight bacteria-derived carboxypeptidases, albeit a little bit flexibility is revealed in distal end compared to the tight state in the dimeric interface. On contrast, ACE2 has evolved at least two states, namely open and closed state which are indispensable for efficient catalysis [29]. This marked difference promotes us to reasonably infer that liberation of ACE2 into monomeric state during its evolution process might contribute to its higher efficiency during...
Fig. 1. Characterization of the oligomeric state and carboxypeptidase activity of B38-CAP and ACE2 carboxypeptidase domain. (A) The gel-filtration profile of B38-CAP and ACE2 carboxypeptidase domain in a Superdex™ 200 Increase 10/300 GL column. (B) Real-time characterization of the carboxypeptidase activity of B38-CAP and ACE2.

Fig. 2. The overall structure of B38-CAP. (A) The dimeric structure of B38-CAP in cartoon representation. (B) A close-up view of the interaction on the dimeric interface of B38-CAP. (C) Structural alignments between B38-CAP (palegreen) monomer and other carboxypeptidase monomers. PDB entry 3HQ2, 3HOA, 5WVV, 7A03, 5E3X, 1K9X, 5GIV, 3DWC are colored in red, marine, yellow, magenta, lime green, olive, gray and orange, respectively.
carboxypeptidase catalysis. The presented high-resolution structure of B38-CAP and corresponding structural comparison with ACE2 help elucidate their structural conservation feature and also their structural diversity and is believed to aid further understanding of the catalytic machinery.

4. Conclusions

In this study, we characterized the dimeric state of B38-CAP which is in accordance with its solved crystal structure. We also characterized the real-time carboxypeptidase activity of B38-CAP which is weaker compared to ACE2. Structural comparison between B38-CAP and other M32-family carboxypeptidase showed highly sequential and structural similarity while only structural conservatism was retained between B38-CAP and ACE2 carboxypeptidase domain, particularly in their catalytic core. Moreover, the dimeric architecture was shown to be strictly conservative among bacteria-derived carboxypeptidases while ACE2 did not obey this rule. This marked difference is considerably worthy of in-depth study and might contribute to a new understanding of the diversity of the catalytic machinery within M32 family carboxypeptidases and also the ongoing development of B38-CAP-based anti-SARS-CoV-2 drugs [30].

Declaration of competing interest

The authors declare that they have no conflicts of interest.

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Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.bbrc.2022.03.077.

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