Expression and preliminary characterization of human MICU2
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ABSTRACT
MICU2 has been reported to interact with MICU1 and participate in the regulation of mitochondrial Ca2+ uptake, although the molecular determinants underlying the function of MICU2 is unknown. In order to characterize MICU2 we screened a series of N-terminal and C-terminal truncations and obtained constructs which can be expressed in abundance, giving rise to soluble samples to enable subsequent characterizations. Size exclusion chromatography (SEC) expressed in abundance, giving rise to soluble samples to enable C-terminal truncations and obtained constructs which can be characterized. Size exclusion chromatography (SEC) and multi-angle laser light scattering (MALLS) revealed that MICU2 exists as a monomer in Ca2+-free conditions but forms a dimer in Ca2+-bound conditions. Unlike MICU1, the C-helix domain of MICU2 exhibits no influence on protein conformation in both Ca2+-free and Ca2+-bound forms. Furthermore, mutation of the first EF-hand abolishes the ability of MICU2 to switch to a dimer in the presence of Ca2+, indicating that the first EF-hand is not only involved in Ca2+ binding but also in conformational change. Our pull-down and co-immunoprecipitation assays suggest that, in addition to disulfide bonds, salt bridges also contribute to MICU1-MICU2 heterodimer formation.

KEY WORDS: MICU1, MICU2, Calcium, EF-hand, Mitochondria

INTRODUCTION
More than half a century ago it was proposed that Ca2+ could accumulate in mitochondria through the mitochondrial calcium uniporter (DeLuca and Engstrom, 1961; Vasington and Murphy, 1962). Subsequent research demonstrated that Ca2+ flux across the inner mitochondrial membrane (IMM) was driven by the membrane potential (ΔΨ) which was generated by the respiratory chain (Nicholls, 2005; O’Rourke, 2007; Santo-Domingo and Demaurex, 2010; Drago et al., 2011). However, the molecular identities of the uniporter complex were ambiguous until recently. By screening IMMs proteins using selective siRNA, MICU1 was discovered, which heralded the molecular identification of the uniporter complex (Perocchi et al., 2010). Other members of the uniporter complex family (MCU, MCUb, MICU2, EMRE and MCUR1) were identified subsequently (Marchi and Pinton, 2014). A series of evidence has defined MCU as the pore-forming unit of the uniporter (DeLuca and Engstrom, 1961; Vasington and Murphy, 1962). Subsequent research demonstrated that Ca2+ flux across the inner mitochondrial membrane (IMM) was driven by the membrane potential (ΔΨ) which was generated by the respiratory chain (Nicholls, 2005; O’Rourke, 2007; Santo-Domingo and Demaurex, 2010; Drago et al., 2011). However, the molecular identities of the uniporter complex were ambiguous until recently. By screening IMMs proteins using selective siRNA, MICU1 was discovered, which heralded the molecular identification of the uniporter complex (Perocchi et al., 2010). Other members of the uniporter complex family (MCU, MCUb, MICU2, EMRE and MCUR1) were identified subsequently (Marchi and Pinton, 2014). A series of evidence has defined MCU as the pore-forming unit of the uniporter complex (Baughman et al., 2011; De Stefani et al., 2011; Chaudhuri et al., 2013; Kovacs-Bogdan et al., 2014). MCUb is a paralog of MCU, considered to be an endogenous dominant negative isoform since its overexpression reduces the number of channels in the lipid bilayer (Raffaello et al., 2013). MICU2, a paralog of MICU1, interacts with MICU1 to form a heterodimer that regulates the function of MCU (Plovanich et al., 2013; Kamer and Mootha, 2014; Patron et al., 2014). Discovered by mass spectrometry of the MCU proteome, EMRE has been found essential for the interaction of MICU1 and MICU2 with MCU (Sancak et al., 2013). MCUR1 was considered to be a key component of the uniporter complex since its knockout was observed to abolish the activity of the uniporter (Mallilankaraman et al., 2013). Although the molecular identities of the uniporter complex have been revealed, precise understanding of the mitochondrial Ca2+ uptake machinery is still largely lacking.

As a regulator that may interact with the pore-forming unit MCU, the MICU family has attracted more attention and as a result, more is now known about it. However, various experimental results from different groups on the effect of MCU-mediated Ca2+ uptake caused by MICU1 knockdown are in conflict. Earlier studies found that knockdown of MICU1 inhibited the mitochondrial Ca2+ uptake (Perocchi et al., 2010) which was confirmed by subsequent experiments (Alam et al., 2012), whereas other studies discovered that MICU1 knockdown led to constitutive mitochondrial Ca2+ (Ca2+ m) accumulation at low intracellular Ca2+ concentration (Ca2+i) (Mallilankaraman et al., 2012; Patron et al., 2014; Csordas et al., 2013; de la Fuente et al., 2014). Thus, MICU1 is assumed to act as the threshold for Ca2+ uptake and function to inhibit uniporter-mediated Ca2+ uptake at low Ca2+ m (Mallilankaraman et al., 2012; Csordas et al., 2013). In MICU1, the EF-hands act as a Ca2+ sensor, although there are still some controversies about its role. Some studies showed that the EF-hand mutation failed to inhibit Ca2+ flux into mitochondrial which paralleled MICU1 knockdown results (Mallilankaraman et al., 2012). In contrast, others revealed that MICU1 with non-function EF-hands still inhibited Ca2+ uptake (Csordas et al., 2013; Kamer and Mootha, 2014). Moreover, crystal structures of human MICU1 demonstrated that Ca2+ binding by the EF-hand could cause large conformational changes in the interface of the dimer unit (Wang et al., 2014). As a paralog of MICU1, MICU2 shares 27% protein sequence identity and contains two putative EF-hands. It has been shown that MICU2 interacts with MICU1 in a heterodimer through a disulfide bond (Patron et al., 2014; Petrungaro et al., 2015). MICU2 was found to be essential for the MCU-mediated Ca2+ uptake, since both its knockout and EF-hand mutation caused reduced Ca2+ uptake at high Ca2+i, even if the effect was smaller than that of MICU1 (Plovanich et al., 2013; Kamer and Mootha, 2014). Nevertheless, other studies found that MICU2 knockdown could enhance Ca2+ uptake at high Ca2+i, suggesting that it acts as the gatekeeper while MICU1 acts as the activator for the uniporter (Patron et al., 2014). A recent report proposed that MICU2 is the inhibitor of MCU and MICU1 has a double role, acting as an inhibitor at low Ca2+i, and activator at high Ca2+i (Kovacs-Bogdan et al., 2014). In addition, there are studies pointed out that the MICU1-MICU2 heterodimer binds to MCU in order to close the channel at low Ca2+i. With the increase of Ca2+ concentration, EF-hands of the MICU1-MICU2 heterodimer would bind to Ca2+ and result in the release of the dimer from the uniporter (Petrungaro et al., 2015).
Herein, we report the preparation and the characterization of MICU2 using insect and *Escherichia coli* cells. We have constructed and expressed a series of recombinant MICU2 with N-terminal and C-terminal truncations to achieve protein expression, enabling the characterization of MICU2 in Ca\(^{2+}\)-free and Ca\(^{2+}\)-bound states using SEC and MALLS. Through site-specific mutation of the EF-hands, we have studied Ca\(^{2+}\) response of MICU2, which demonstrates a significant difference with MICU1. Furthermore, we have studied MICU1-MICU2 association and revealed new interaction features.

**RESULTS**

**Cloning of full-length MICU2 and truncations**

As shown in Fig. 1, MICU2 possesses an N-terminal targeting sequence, two putative EF-hands and a helix domain at the C-terminus. We designed a series of constructs including full-length and truncated MICU2 with varying lengths. Several constructs of N-terminal deletions were prepared starting at amino acid 66, 84, 105 or 159. The C-helix has been proven crucial in MICU1 (Kamer and Mootha, 2014; Wang et al., 2014), and in order to exploit the role of the C-helix domain in MICU2, the C-terminal clones were constructed by deleting 28 or 36 amino acids in the C-terminus. The details of MICU2 constructs are listed in Table 1. For the pull-down assay of MICU1 and MICU2, MICU1-NA96 and MICU1-NA96 were cloned without any tag. Furthermore, MICU1-NA96 and MICU2-NA84 were cloned into the multiple cloning sites (MCS) of the pETDuet-1 vector respectively, which would allow expression of the two constructs simultaneously. Clones of MICU1 (MICU1-NA96-Myc) and MICU2 (MICU2-NA84-Flag and MICU2-NA84-CA36-Flag) constructs used for co-immunoprecipitation were designed with Myc-tag and Flag-tag, respectively. The details of MICU1 constructs are listed in Table S1.

**Protein expression and purification**

Our expression results of the MICU2 constructs are summarized in Table 1. For full length constructs, MICU2-I was expressed in Sf21 insect cells. MICU2-N (N-terminal His-tag), MICU2-C (C-terminal His-tag) and maltose-binding protein (MBP)-MICU2 were expressed in *E. coli* BL21 (DE3) cells. Though we were able to detect MICU2-I expression in the insect cells by western blotting (Fig. 2A), the SDS-PAGE results indicated that most of the protein was accumulated in the precipitant, which was further confirmed by western blot (Fig. S1). When fused with a MBP tag, MICU2 was also largely expressed in the precipitant in *E. coli* (Fig. 2B); therefore, we concluded that the constructs with the targeting sequence were all insoluble, even in the presence of a MBP tag. After truncating the targeting sequence, we obtained soluble MICU2-NA66 and MICU2-NA84 which were purified by affinity chromatography (Fig. S2A,B), however large amounts of the proteins were still present in the precipitant (Fig. S2A,B). We next screened the C-terminal truncations. Several constructs (MICU2-NA66-CA28, MICU2-NA66-CA36, MICU2-NA84-CA28 and MICU2-NA84-CA36) had good protein expression with high purity (Fig. S2C-F). However, for those constructs with more N-terminal amino acids eliminated, such as MICU2-NA105-CA28 and MICU2-NA159-CA28, the protein would express in inclusion bodies in *E. coli*.

**Protein purification by SEC**

The eluted samples were concentrated and purified by SEC. Considering the EF-hands of MICU2, the buffer used for SEC either contained EGTA to provide a Ca\(^{2+}\)-free condition, or CaCl\(_2\) to maintain the protein in a Ca\(^{2+}\)-bound state. The results are shown in Fig. 3 and the retention volume (V\(_R\)) values of the peaks are shown in Table S2. In the Ca\(^{2+}\)-free condition, peak integration revealed that the V\(_R\) values of MICU2-NA66 and MICU2-NA84 were ~83-85 ml (Fig. 3A). Similarly, for the other constructs (MICU2-NA66-CA28, MICU2-NA66-CA36, MICU2-NA84-CA28, MICU2-NA84-CA36), the V\(_R\) values were ~84 ml (Fig. 3C). Therefore, it seems that the C-helix domain has no significant influence on the protein conformation in Ca\(^{2+}\)-free condition. In Ca\(^{2+}\)-bound state, the V\(_R\) values were ~80 ml for all aforementioned six MICU2 constructs, indicating that the C-helix domain has no influence on the protein conformation in the Ca\(^{2+}\)-binding state as well (Fig. 3B,D). However, comparing the V\(_R\) values of all tested constructs between EGTA and Ca\(^{2+}\) conditions, consistently there was a 3-5 ml shift which is not insignificant, suggesting that there may be conformational changes after binding Ca\(^{2+}\). SDS-PAGE results of various constructs are shown in Fig. 4, Figs S2 and S3, in which the construct MICU2-NA84-CA28 exhibited the best purity.
To further investigate the conformation of MICU2 in the Ca²⁺-free condition, we performed Multi-angle laser light scattering (MALLS) analysis of ΔMICU2-NΔ84-C28Δ28EF1mut and ΔMICU2-NΔ84-C28Δ28EF2mut (Fig. 5). Similar molecular weights for ΔMICU2-NΔ84-C28Δ28EF1mut and ΔMICU2-NΔ84-C28Δ28EF2mut were detected in the Ca²⁺-free condition, as the MALLS results were 48.2 kDa, 44.3 kDa and 50.2 kDa, respectively. However, when Ca²⁺ was added, the MALLS results of ΔMICU2-NΔ84-C28Δ28EF1mut and ΔMICU2-NΔ84-C28Δ28EF2mut were 87.4 kDa, 55.9 kDa and 78.4 kDa, respectively. The predicted molecular weight of ΔMICU2-NΔ84-C28Δ28 was ~40 kDa. Considering the influence of protein purity, it is not surprising that there may be a deviation between the experimental result and the actual molecular weight. However, our MALLS results are sufficiently clear that ΔMICU2-NΔ84-C28Δ28EF1mut would not be influenced by Ca²⁺, and the conformational change is controlled by the first EF-hand. Moreover, our SEC results in Fig. 3 show that the addition of Ca²⁺ would cause a retention volume shift for ΔMICU2-NΔ84-C28Δ28EF1mut, which did not happen for ΔMICU2-NΔ84-C28Δ28EF2mut. These results are consistent with the MALLS analysis.

**Pull-down and co-immunoprecipitation assays of MICU1 and MICU2**

It has been reported that MICU1 and MICU2 form a heterodimer via a disulfide bond through the cysteine residues in the C-helix domain (Cys463 in MICU1 and Cys413 in MICU2) (Patron et al., 2014; Petrungaro et al., 2015). In order to characterize this bonding, we performed in vitro pull-down assays and the results are shown in Fig. 6 and Fig. S5. In this experiment, we first confirmed the disulfide bond between MICU1 and MICU2 through the co-expression experiment (Fig. S5). The western blot shows that the two constructs could form a complex with a molecular weight similar to the disulfide bond.
and MICU2-NΔ84-CA36 which lack Cys413 and would therefore not be expected to form the heterodimer, there are still two bands (Fig. 6B). This surprising result demonstrates that both MICU2-NΔ66-CA36 and MICU2-NΔ84-CA36 retain interaction with MICU1-NΔ96. To further substantiate these unexpected observations, we carried out co-immunoprecipitation and western blot analysis, which clearly indicates that MICU2 (MICU2-NΔ84-Flag and MICU2-NΔ84-CA36-Flag) interacts with MICU1-NΔ96-Myc either with or without Cys413 (Fig. 6E). Thus, it seems that there are other factors contributing to MICU1-MICU2 interaction in addition to the disulfide bond. In order to determine whether the EF-hands are involved in the interaction of the heterodimer, pull-down assays were next performed using MICU1-NΔ96 and MICU2 EF-hand mutants. Interestingly, both the two MICU2 EF-hand mutants which lack Cys413 and Ca\(^{2+}\) binding residues still interacted with MICU1, indicating that the EF-hands do not participate in MICU1-MICU2 interaction (Fig. 6C). According to the structure model of MICU1 (Wang et al., 2014), we designed the mutants MICU1-NΔ54R221A and MICU2-NΔ84-CΔ28D330N. Neither mutants interacted with the corresponding wild-type MICU2-NΔ84-CΔ28 or MICU1-NΔ96 as there was only one band present (Fig. 6D,E). Therefore, MICU1 associates with MICU2 through the salt bridge created by Arg221 in MICU1 and Asp330 in MICU2, in addition to the disulfide bond.

Fig. 4. Fraction analysis of MICU2-NΔ84-CΔ28 in Ca\(^{2+}\)-free or Ca\(^{2+}\)-bound conditions. SDS-PAGE analysis of SEC fractions revealed that MICU2-NΔ84-CΔ28 exhibited the best purity among all the MICU2 truncations. The white vertical line refers to the fact that MW markers were loaded on the same SDS-PAGE, but not in the adjacent lanes.

Fig. 5. MALLS results for MICU2-NΔ84-CΔ28 and EF-hand mutations. (A,C,E) MALLS results for MICU2-NΔ84-CΔ28 and EF-hand mutations in Ca\(^{2+}\)-free (2 mM EGTA) condition. (B,D,F) MALLS results for MICU2-NΔ84-CΔ28 and EF-hand mutations in Ca\(^{2+}\)-bound (5 mM CaCl\(_2\)) condition. Ca\(^{2+}\) could change MICU2-NΔ84-CΔ28 and MICU2-NΔ84-CΔ28EF2mut from a monomer to a dimer which is not the case for MICU2-NΔ84-CΔ28EF1mut. Therefore, the conformational change would take place in parallel with Ca\(^{2+}\) binding with the first EF-hand. The red line indicates the laser light scattering data and the blue line indicates the differential refractive index data.
DISCUSSION

In order to gain further functional insights, it is necessary to establish a reliable method of producing recombinant MICU2 for in-depth characterization. Our experiments show that constructs containing an N-terminal targeting sequence did not express in E. coli. Subsequently, we attempted the insect expression system and MBP fusion construct. Although the expression was successful, unfortunately the protein was mostly insoluble. Lacking the N-terminal target sequence, constructs MICU2-NΔ66 and MICU2-NΔ84 improved both protein solubility and expression level (Figs S1 and S2). The SEC results revealed that there was significant shift between the retention volume for MICU2-NΔ66 and MICU2-NΔ84 in Ca\(^{2+}\)-free and Ca\(^{2+}\)-bound condition, indicating that there was conformational change induced by Ca\(^{2+}\) binding.

The helix domain of MICU1 in the C-terminus is considered to be essential for protein conformation. It is reported that in the wild-type MICU1 C-helices bundle together to form Ca\(^{2+}\)-free hexamer, while the C-helix truncation behaves as a dimer in both Ca\(^{2+}\)-free and Ca\(^{2+}\)-bound conditions (Wang et al., 2014). Considering the similarity of the functional domain between MICU1 and MICU2, we wondered whether the C-helix domain would play a similarly important role in MICU2 as well. Our results revealed that when the C-terminal 28 or 36 amino acids were truncated, the constructs exhibited similar properties as the N-terminal constructs in either Ca\(^{2+}\)-free or Ca\(^{2+}\)-bound condition. Thus, the C-helix in MICU2 seems to have no influence on protein conformation as observed for MICU1. The MALLS results further revealed that Ca\(^{2+}\) triggers MICU2-NΔ84-CA28 to switch from a monomer to a dimer. Therefore, the conformational change of MICU2 is largely controlled through the combined effect of EF-hands and Ca\(^{2+}\) binding. A recent paper reports that MICU1 would rearrange in a Ca\(^{2+}\)-free hexamer, while our results reveal that MICU2 aggregates switch from monomer to dimer after binding Ca\(^{2+}\). This difference may be related to their functional role in mitochondrial Ca\(^{2+}\) uptake as MICU1 and MICU2 exert opposite effects on MCU activity (Patron et al., 2014; Matesanz-Isabel et al., 2016). Therefore, conformational change of MICU1 and MICU2 after binding Ca\(^{2+}\) may be able to disable MICU2’s inhibitory effect and evoke MICU1’s activating function for mitochondrial Ca\(^{2+}\) uptake.

To discern the influence of each EF-hand on conformational change, we probed the two EF-hands by site-directed mutagenesis. It has been reported that EF-hand mutants of MICU1 with a C-terminus truncation behave as dimers and would not be influenced by Ca\(^{2+}\) (Wang et al., 2014). Unexpectedly, in our SEC experiment, Ca\(^{2+}\) caused a retention volume shift for MICU2-NΔ84-CA28 and MICU2-NΔ84-CA28\(\text{EF2mut}\) but not for MICU2-NΔ84-CA28\(\text{EF1mut}\). In order to substantiate these observations, we carried out further MALLS experiments and the results revealed that MICU2-NΔ84-CA28\(\text{EF2mut}\) changed from a monomer to a dimer after binding Ca\(^{2+}\) while MICU2-NΔ84-CA28\(\text{EF1mut}\) did not. Therefore, it is the first EF-hand, not the second, which induces the conformational change upon Ca\(^{2+}\) binding.

Our observed conformational change of MICU2 caused by the first EF-hand is different from that of MICU1. Ca\(^{2+}\)-free structure of MICU1 suggested that the dimer interaction is via a salt bridge between Asp376 and Arg221. Upon binding Ca\(^{2+}\), hydrogen bonds between Arg221 and Gln224 from one monomer with Ser382 and His385 from another, which provide the interaction between dimers (Wang et al., 2014). Since MICU2 structure has not been determined, our conjecture of the conformation change of MICU2 is described as follows (Fig. 7). Alignment of MICU1 and MICU2 shows that only the aspartic acid is conserved (Asp376 in MICU1 and Asp330 in MICU2). The corresponding residue of MICU1 Arg221 in MICU2 is Ser175; the salt bridge would not form for MICU2 and this may be the reason why MICU2 is a monomer without Ca\(^{2+}\). The switch of MICU2 from monomer to dimer induced by Ca\(^{2+}\) binding in the first EF-hand may be considerably different from MICU1 since all the residues contributing to MICU1 dimer formation are not conserved in MICU2. The first EF-hand may experience large conformational change after binding Ca\(^{2+}\) and result in potential new interactions between the first EF-hand and the other MICU2 monomer.

Previous experiments indicate that MICU1 and MICU2 form a heterodimer through disulfide bonds between Cys463 in MICU1 and Cys413 in MICU2. MICU1 can also form a homodimer when it is overexpressed, which is consistent with its crystal structures (Patron et al., 2014; Petrungaro et al., 2015). By using non-reducing
SDS-PAGE analysis, we confirmed that there is a disulfide bond between MICU1 and MICU2. However, our pull-down results using constructs lacking Cys413 suggest that the disulfide bond is not the only interaction between MICU1 and MICU2 which is further confirmed by co-immunoprecipitation in situ. Since the aspartic acid is conserved (Asp376 in MICU1 and Asp330 in MICU2), based on the structural analysis we speculate that in the heterodimer, Arg221 in MICU1 may interact with Asp330 in MICU2 to form a salt bridge similar to the Asp376-Arg221 salt bridge in the MICU1 homodimer. Pull-down results using Arg221 or Asp330 mutants provide strong support for our hypothesis.

**Conclusion**

Through construct screening by truncating the N-terminus or C-terminus of MICU2, we have successfully obtained soluble MICU2 truncations which enabled subsequent characterization. The SEC and MALLS experiments demonstrate that MICU2 exists as a monomer without Ca\(^{2+}\) and changes to a dimer after binding Ca\(^{2+}\). Mutation of the first EF-hand, but not the second, abolishes the Ca\(^{2+}\)-induced conformational change of MICU2. Compared with MICU1, our results indicate a different role of the first EF-hand in MICU2 which binds Ca\(^{2+}\) and induces conformational change. In addition to the disulfide bond, our results suggest a salt bridge also contributes to the MICU1-MICU2 interaction. A full understanding of the molecular determinants underlying these new observations would have to wait for structural characterization.

**MATERIALS AND METHODS**

**Cloning of MICU1 and MICU2**

The details of various clones for MICU1 and MICU2 are given in Table 1 and Table S1. Generally, human MICU1 (BC004190.2) or MICU2 (BC031089.1) genes were amplified by PCR with the desired restriction sites and cleaved with the corresponding restriction enzymes (Thermo Scientific). The digested PCR products were ligated into vectors, namely pFastBac HT A for insect expression, pET-28a(+) and pETDuet-1 for E. coli expression, and pcDNA3.1 for HEK293T expression, respectively. It should be mentioned that for the construction of MBP-MICU2, MBP tag was first cloned into pET-28a(+) between the Ncol and NdeI restriction sites (a Prescision protease cleavage site was inserted before the NdeI restriction site). Next the cleaved MICU2 PCR products (5′ NdeI, 3′ XhoI) were ligated into the modified vector which was digested using the same enzymes. For MICU1, the reverse primers were designed with a termination codon to ensure lacking of His-tag. For MICU2, clones were designed with His-tag in the N-terminus. In order to co-express MICU1 and MICU2, we introduced both genes into the pETDuet-1 vector which contains two MCSs. Therefore, the first protein inserted into the first MCS contains His-tag and the second protein does not have any tag. After that, the recombinant vectors were transformed into TOP10 cells and cultured on the agar LB medium plate overnight at 37°C. A single clone was picked up, cultivated and confirmed by DNA sequencing. The recombinant plasmid was extracted (Plasmid Miniprep Kit, Tiangen) and stored at −40°C for further use.

**Site-specific mutation of MICU2 constructs**

To disable Ca\(^{2+}\) binding, mutation of the EF-hand was performed according to literature by mutating the first and the last residues of the predicted 12 residues in Ca\(^{2+}\) binding loop of each EF-hand to alanine and to lysine, respectively (Perec et al., 2010). To investigate the interaction of MICU1 and MICU2, we also designed MICU1-R221A and MICU2-D330N mutants. The mutation was operated by PCR using KOD polymerase (TOYOBO). After purification by agarose gel electrophoresis and DNA purification kit (Tiangen), the PCR product was incubated with DpnI for 30 min. The digested product was next transformed into TOP10 and cultured on the agar LB medium plate overnight. A single clone was picked and cultured and the mutated sites were confirmed by DNA sequencing.

**Protein expression**

The expression of the full-length MICU2 in insect cells was done using the Bac-to-Bac baculovirus expression system (Invitrogen). The recombinant donor plasmid containing MICU2 gene was first transformed into competent DH10Bac E. coli cells and incubated on the agar LB medium plate (50 µg/ml kanamycin, 7 µg/ml gentamicin, 10 µg/ml tetracycline, 100 µg/ml X-gal and 40 µg/ml IPTG) for 48 h at 37°C. A white clone was picked and cultured in 15 ml LB medium (50 µg/ml kanamycin, 7 µg/ml gentamicin, 10 µg/ml tetracycline) overnight. The recombinant bacmid was isolated (Plasmid Mini Kit, Qiagen) and analyzed by PCR using pUC/M13 primers (pUC/M13 forward: 5′-CCCCAAGTCAGCTACGTTGTGAACACCCG-3′; pUC/M13 reverse: 5′-AGCGGATAACAATTTCACACAGG-3′) to confirm that the target gene was inserted into the bacmid. Purified recombinant bacmid (~1 µg) was used to transfect Sf21 insect cells (~1×10^6 cells) for 3 days using the Cellfectin reagent (Cellfectin II reagent, Invitrogen) to produce P1 viral stock. The high-titer P2 viral stock was generated by using the P1 viral stock to transfect the Sf21 insect cells (~1.2×10^6 cells in 25 ml medium). P3 viral stock was generated using the P2 viral stock with the same method. To express the protein, Sf21 cells (1 litre) were first suspension cultured to a density of 1×10^7 to 2×10^7 cells/ml and infected with the P3 viral stock (20 ml) for 48 h before the cells were harvested. The expression of MICU2 was determined by western blotting (anti-His-tag mouse monoclonal antibody, CWbio). For the expression of MICU1, MICU2 and the coexpression of MICU1-MICU2 construct in E. coli, the recombinant plasmids were transformed into BL21 (DE3) E. coli strain. A single clone was picked and cultured in 5 ml LB medium for 12 h, and 1 ml cells were then transferred into 100 ml LB medium to amplify overnight. 30 ml of the amplified cells were
Lipofectamine 3000 (Invitrogen, Cat No. 1756121) and cultured sample of each fraction was assessed by 12% SDS-PAGE.

The cells were harvested by centrifugation and stored at −4°C until further use.

**Protein purification**

The general protocol for the purification of the full-length and truncated MICU2 is as follows. The harvested cells were first re-suspended in buffer I (20 mM Tris pH 7.0, 300 mM NaCl, 20 mM imidazole, 0.3% Triton X-100). For the insect cells, cells were lysed by freezing and thawing. For *E. coli* cells, cells were lysed by sonication. Both the cell lysates were centrifuged at 18,000 rpm (39,000 × g) for 30 min. The column with 2 ml of Ni-NTA resin (Qiagen) was first equilibrated with buffer I and precooled at 4°C and the supernatant was slowly loaded onto the resin. Afterwards, the protein was washed with 150-200 ml buffer II (20 mM Tris pH 7.0, 300 mM NaCl, 50 mM imidazole) and eluted with 20 ml buffer III (20 mM Tris pH 7.0, 300 mM NaCl, 400 mM imidazole). Protein purity in each step was monitored by 12% SDS-PAGE. The eluted protein was concentrated by a 10-kDa molecular weight cut-off centrifugal filter (Merck Millipore) and samples were further analyzed by SEC using a 120 ml Hidal Superdex 20 10/300 GL column (GE Healthcare Biosciences AB) on AKTA purifier system. The column was equilibrated with buffer IV (20 mM MES pH 6.8, 300 mM NaCl, 2 mM EGTA or 5 mM CaCl₂) before protein analysis. The protein was detected by a UV detector at a wavelength of 280 nm and the sample of each fraction was assessed by 12% SDS-PAGE.

**Multi-angle laser light scattering**

The multi-scale field and polydispersity of MICU2-NΔ84-CΔ28 and its EF-hand mutants were analyzed by SEC equipped with a MALS detector. Protein samples were prepared by concentrating to 0.5-2.5 mg/ml and filtered through a 0.22-µm filter. 200 µl of the protein sample was injected into the loop and chromatographic analysis was done using KW-G and KW-803 columns (Shimadzu) at a flow rate of 0.5 ml/min. The protein signals were detected by a DAWN HELOS-II light scattering detector and an Optilab re refractive index detector (Wyatt Technology). The molecular weight was calculated by peak integrations (ASTRA software version 5.3.4.13). Each sample was carried out independently and repeated at least three times.

**Pull-down assay for MICU1 and MICU2**

Pull-down assays for MICU1 and MICU2 were operated by co-expression or by mixing cells individually expressing MICU1 or MICU2 together. The harvested cells of MICU1 constructs (MICU1-NΔ54, MICU1-NΔ54P221A, MICU1-NΔ96) with MICU2 construct (MICU2-NΔ66-CΔ36, MICU2-NΔ84-CΔ28, MICU2-NΔ84-CΔ28, MICU2-NΔ84-CΔ36, MICU2-NΔ84-CΔ28EF1mut, MICU2-NΔ84-CΔ28EF2mut, MICU2-NΔ84-CΔ28D330N) were re-suspended in buffer I, and the mixed cells were sonicated together. Cell lysates were centrifuged and the supernatant was applied to the Ni-NTA resin that was precooled and equilibrated with the buffer I. The resin was washed with 100 ml buffer II and the proteins were eluted together with buffer III. The controls for the pull-down assay were performed by purifying MICU1-NΔ54, MICU1-NΔ96 and MICU1-NΔ54P221A as aforementioned individually. All eluted samples were analyzed by reducing SDS-PAGE. The co-expression cells (MICU1-NΔ96-MICU2-NΔ84 or MICU2-NΔ84-MICU1-NΔ96) were purified by the protocols aforementioned. The eluted samples were exposed to air for oxidation and ensuring disulfide formation and then analyzed by western blot with non-reducing SDS-PAGE. The assays were carried out independently and repeated at least three times.

**Transfection and co-immunoprecipitation of MICU1 and MICU2**

Transfected HEK293T cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM, Hyclone GE Healthcare Life Sciences) supplemented with 10% v/v fetal bovine serum (FBS, Gibco) and 1% v/v penicillin-streptomycin (PS, Caisson). The cells were cultured in a 95% air and 5% CO₂ environment at 37°C. MICU1-NΔ96-Myc with MICU2-NΔ84-Flag or MICU2-NΔ84-CΔ36-Flag genes were transfected into cells using Lipofectamine 3000 (Invitrogen, Cat No. 1756121) and cultured sequentially in a 10-cm Petri dish for 48 h. Transfected HEK293T cells were harvested by PBS buffer (Hyclone, GE Healthcare Life Sciences) and then lysed in 300 µl of TBS buffer (20 mM Tris pH 7.4, 150 mM NaCl) containing 0.2% n-dodecyl-β-D-maltoside (DDM, Sigma) and protease inhibitor cocktail (Roche). After being lysed on ice for 30 min, the lysate was centrifuged at 13,000× g for 30 min and 40 µl of the cell supernatant was removed and used as positive control. Protein A/G agarose beads (GE Healthcare) were washed three times with TBS buffer and then incubated with anti-Flag antibody (Transgen, Cat No. HT201) and cell supernatant for 4 h. The beads were washed three times with TBS and eluted by adding 50 µl of SDS-PAGE loading buffer. Both the control and eluted samples were analyzed by western blotting using anti-Flag antibody and anti-c-Myc antibody (Transgen, Cat No. HT101).

**Competing interests**

The authors declare no competing or financial interests.

**Author contributions**

D.L., W.W., H.P. and Q.Y. performed the experiments; D.L., Q.W., J.Z. and Z.J. analyzed the data; D.L., J.Z. and Z.J. wrote the manuscript.

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**Supplementary information**

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**References**


