# Structural diversity and oligomerization of bacterial ubiquitin-like proteins

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#### Summary

Bacteria possess a variety of operons with homology to eukaryotic ubiquitination pathways that encode predicted E1, E2, E3, deubiquitinase, and ubiquitin-like proteins. Some of these pathways have recently been shown to function in anti-bacteriophage immunity, but the biological functions of others remain unknown. Here, we show that ubiquitin-like proteins in two bacterial operon families show surprising architectural diversity, possessing one to three  $\beta$ -grasp domains preceded by diverse N-terminal domains. We find that a large group of bacterial ubiquitin-like proteins possess three  $\beta$ -grasp domains and form homodimers and helical filaments mediated by conserved Ca<sup>2+</sup> ion binding sites. Our findings highlight a distinctive mode of self-assembly for ubiquitin-like proteins, and suggest that Ca<sup>2+</sup>-mediated ubiquitin-like protein filament assembly and/or disassembly enables cells to sense and respond to stress conditions that alter intracellular metal ion concentration.

## Introduction

Ubiguitination is a multi-step enzymatic cascade in which ubiguitin or a ubiguitin-like protein (Ubl) becomes covalently linked to a target - typically a lysine side chain or other amine group through the sequential action of E1, E2, and (usually) E3 proteins<sup>1</sup>. Ubiquitination and related pathways regulate protein homeostasis and other processes in eukaryotes<sup>2–6</sup>, but related bacterial pathways are usually involved in metabolic pathways and do not mediate protein conjugation. Recently, bioinformatics analysis of bacterial genomes has identified several sparsely distributed families of bacterial operons that encode proteins related to eukaryotic E1, E2, E3, Ubl, and peptidase/deubiquitinase proteins<sup>7-9</sup>. Structural and biochemical studies of three such operon families have now demonstrated that they are related to eukaryotic ubiquitination pathways and perform *bona fide* protein conjugation reactions<sup>10–13</sup>. Type II CBASS anti-bacteriophage (phage) operons encode an E2-E1 fusion protein, Cap2, that conjugates the C-terminus of their cognate CD-NTase to unknown targets to activate CBASS signaling<sup>10,11</sup>. Type I and II Bil (Bacterial ISG15-like) operons encode separate E1, E2, Ubl, and DUB proteins, and when cells are infected with phages, these pathways conjugate their Ubl to a phage tail protein to inhibit virion assembly and infectivity<sup>12,13</sup>. Finally, in an accompanying work, we find that Bacterial ubiguitination-like (Bub) operons, previously termed "6E" or "DUF6527" operons, also encode E1, E2, Ubl, and peptidase proteins and perform protein conjugation<sup>14</sup>. These studies show that ubiguitination pathways evolved and proliferated in bacteria in the contexts of antiphage immunity and stress response.

In eukaryotes, Ubls typically comprise a single small domain with a fold termed " $\beta$ -grasp"<sup>1</sup>. Some Ubls, notably including the innate-immune protein ISG15, possess two  $\beta$ -grasp domains. In prior work, we found that bacterial Ubls in Type II Bil operons show high structural diversity, with up to three predicted  $\beta$ -grasp domains and diverse fused N-terminal domains<sup>12</sup>. Here, we expand on that finding and show that Ubls in Bil and Bub operons possess diverse architectures, and that many such proteins form higher-order oligomers. We find that a large subset of bacterial Ubls contain three  $\beta$ -grasp domains and form filamentous assemblies *in vitro* upon calcium (Ca<sup>2+</sup>) ion binding. We show that Ca<sup>2+</sup>-induced filament formation occurs in diverse Ubls from Type II Bil, Type I Bub, and Type II Bub operons, suggesting that this property plays an important role in Bil/Bub operon function. We propose a mechanism in which assembly and/or disassembly of Ubl filaments enables cells to respond to changes in metal ion concentration during phage infection and/or other stress conditions.

#### Results

#### Bacterial ubiquitin-like proteins are architecturally diverse

To comprehensively characterize the architectural diversity of bacterial Ubls, we took advantage of recent studies that identified hundreds of bacterial operons encoding different combinations of ubiquitination-related proteins, particularly those that encode a predicted Ubl. These operons fall into two major groups termed Bil (<u>B</u>acterial <u>I</u>SG15-<u>like</u>)<sup>15</sup> and Bub (<u>B</u>acterial <u>ub</u>iquitination-like) operons<sup>14</sup>. Bil operons can be divided into two families, Type I and Type II, based on the

sequences of their proteins. All Bil operons encode a Ubl (BilA), an E2 (BilB), a JAB/JAMMfamily peptidase/DUB (BilC), and an E1 protein (**Figure 1a**)<sup>12,15</sup>. Bub operons can also be divided into two families, which encode different but related domains. Type I Bub operons encode a Ubl-E2 fusion (BubAB), a JAB/JAMM peptidase-E1 fusion (BubCD), and a DUF6527 protein with serine protease activity (BubE; **Figure 1a**)<sup>14</sup>. Type II Bub operons encode a Ubl (BubA), an E2-E1 fusion (BubBD), and a DUF6527 protein (BubE), and typically do not encode a JAB/JAMM peptidase (**Figure 1a**).

We used protein sequence alignments and structure predictions from ESMFold<sup>16</sup> and AlphaFold<sup>17,18</sup> to comprehensively outline the predicted domain architectures of Ubls in previously-identified Bil and Bub operons (**Figure 1b-c, Table S1**)<sup>15</sup>. Type I Bil operons overwhelmingly encode Ubls with two predicted  $\beta$ -grasp domains, similar to the structure of ISG15 (ref. <sup>15</sup>), and are not further considered here. Ubls in Type II Bil operons show diverse predicted architectures: 105 out of 134 identified operons (78%) encode a Ubl with a single predicted  $\beta$ -grasp domain, 1 (<1%) contains two  $\beta$ -grasp domains, and 28 (21%) contain three  $\beta$ -grasp domains (**Figure 1b**). Some Ubls with a single predicted  $\beta$ -grasp domain possess a variable N-terminal domain; we identified one Ubl from *Hymenobacter coccineus* with a 151 amino acid N-terminal tail rich in asparagine and glycine (N/G), suggesting that this protein may undergo liquid-liquid phase separation (**Figure S1a-c, Table S1**). We also identified five Ubls with a predicted N-terminal coiled-coil domain, suggesting that these proteins may form homodimers (**Figure S1f-i, Table S1**).

As in Type II Bil, Ubls from Type I and II Bub operons Ubls show diverse predicted architectures: 218 (40%) contain a single predicted  $\beta$ -grasp domain, 38 (7%) contain two  $\beta$ grasp domains, and 286 (53%) contain three  $\beta$ -grasp domains (Figure 1b). Among those Ubls with a single predicted  $\beta$ -grasp domain, we identified two with long N/G-rich N-terminal tails, and three with predicted N-terminal coiled-coils (Table S1). We also identified 7 Ubls with tandem DUF1508 domains followed by a single  $\beta$ -grasp domain (Figure 1c, Figure S1d-e). DUF1508 domains are found in stress-induced genes including *E. coli* YegP<sup>19,20</sup> and appear either as single domains that form homodimers (PDB ID 6Q2Z)<sup>20</sup>, tandem domains that form symmetric pseudo-dimers (PDB 2K8E), or as extended arrays (e.g. Pfam PF07411). Overall, our sequence analyses show that bacterial Ubls from Type II Bil and Type I/Type II Bub operons possess diverse architectures with variable numbers of  $\beta$ -grasp domains, plus other domains that could mediate self-association or oligomerization (Figure 1c). Because all of these proteins are named either "BilA" or "BubA" but these names do not capture their structural diversity, in the following sections we use a standard nomenclature with "Ubl" followed by a subscript number indicating the number of  $\beta$ -grasp domains in each protein (e.g. "Ubl<sub>3</sub>" for a protein with three predicted  $\beta$ -grasp domains).

We performed size exclusion chromatography coupled to multi-angle light scattering (SEC-MALS) on six Ubl proteins with different predicted architectures: two encoding two predicted  $\beta$ -grasp domains (Ubl<sub>2</sub>), two with a predicted coiled-coil domain followed by one  $\beta$ -grasp domain (CC-Ubl<sub>1</sub>), and two with N-terminal tandem DUF1508 domains followed by one  $\beta$ -grasp domain (DUF1508-Ubl<sub>1</sub>). The Ubl<sub>2</sub> proteins and the DUF1508-Ubl<sub>1</sub> proteins all formed monomers in

solution (**Figure S2a-d**). We observed that one CC-Ubl<sub>1</sub> formed a homodimer, while the other was monomeric (**Figure S2e-f**).

## Structures of Ubl<sub>3</sub> proteins reveal a conserved homodimeric assembly

Bacterial Ubls predicted to contain three  $\beta$ -grasp domains (Ubl<sub>3</sub>) are nearly as common as those encoding one  $\beta$ -grasp domain. We expressed and purified three diverse Ubl<sub>3</sub> proteins from a Type II Bil operon (*Methylobacterium brachiatum* DSM 19569 Ubl<sub>3</sub>), a Type I Bub operon (Citrobacter sp. RHBSTW-00271 Ubl<sub>3</sub>), and a Type II Bub operon (E. coli ZDHYS365 Ubl<sub>3</sub>). We first crystallized and determined a 1.9 Å resolution crystal structure of E. coli Ubl<sub>3</sub>. The structure revealed the expected array of three  $\beta$ -grasp domains, with the first two domains tightly associated with one another through a predominantly hydrophobic interface burying ~500 Å<sup>2</sup> of surface area per domain. The interface between the second and third  $\beta$ -grasp domains is much smaller (280 Å<sup>2</sup> per domain) and polar, implying flexibility (Figure 2a-b). All three domains adopt the  $\beta$ -grasp fold, with five antiparallel  $\beta$ -strands embracing a single short  $\alpha$ -helix (Figure 2c-d). In β-grasp domains 2 and 3, the short g-helix is buttressed by an additional two-stranded antiparallel  $\beta$ -sheet comprising one strand from the loop linking  $\beta$ -strands S4 and S5, and a second strand just upstream of the α-helix (which is itself inserted between β-strands S2 and S3). Within the crystallographic asymmetric unit, we observed a dimeric assembly mediated by an antiparallel arrangement of the first two  $\beta$ -grasp domains, burying ~1,100 Å of mostly polar surface area per protomer (Figure 2b). The dimer is apparently stabilized by two symmetricallybound divalent cations (built as Ca<sup>2+</sup> based on coordination geometry and refined difference electron density), which are each coordinated by three main-chain carbonyl groups and a glutamate side-chain from one protomer (E136), plus an aspartate side-chain from the second protomer (D73; Figure 2e). Sequence alignments show that these residues are highly conserved across Ubl<sub>3</sub> proteins (Figure S3a-b).

We next crystallized and determined a 3.0 Å resolution crystal structure of *M. brachiatum* Ubl<sub>3</sub>. In the structure, we observed only the first two of three predicted  $\beta$ -grasp domains; the third domain is likely to be flexible and disordered (**Figure 3a-b**, **Figure S4a-d**). Like *E. coli* Ubl<sub>3</sub>, *M. brachiatum* Ubl<sub>3</sub> forms a symmetric homodimer through association of  $\beta$ -grasp domains 1 and 2, and the dimer symmetrically binds two ions (again built as Ca<sup>2+</sup>), which are coordinated identically to those bound by *E. coli* Ubl<sub>3</sub>, by E148 and D85 (**Figure 3b** *inset*).

Within the crystallographic asymmetric unit, two *M. brachiatum* Ubl<sub>3</sub> dimers pack against one another and are related by a 60° right-handed rotation and an ~18 Å translation along the crystallographic c axis. Further, these crystals adopt space group P3<sub>1</sub>21, which contains a screw axis along the crystallographic c axis with a right-handed twist of ~120° and a rise of 35.2 Å (105.5 Å c axis divided by three) per asymmetric unit. Combining non-crystallographic and crystallographic symmetry gives an apparent helical filament of *M. brachiatum* Ubl<sub>3</sub> dimers, with a right-handed helical twist of 60° and a rise of 17.6 Å (105.5 Å divided by six) per dimer (Figure **3c**). We observe an additional ion symmetrically bound between each Ubl<sub>3</sub> dimer pair, coordinated by two main-chain carbonyl groups and an aspartate side-chain (D131) from each of the two protomers involved (Figure **3c** *inset*). Sequence alignments show that the acidic

residues involved in both intra-dimer and inter-dimer Ca<sup>2+</sup> ion coordination are highly conserved across Ubl<sub>3</sub> proteins, implying that these sites are functionally relevant (**Figure S3a-b**).

## Ubl<sub>3</sub> proteins oligomerize in the presence of Ca<sup>2+</sup>

Our structures of *E. coli* Ubl<sub>3</sub> and *M. brachiatum* Ubl<sub>3</sub> suggest that these proteins bind ions at conserved sites within and between Ubl<sub>3</sub> dimers, and that ion binding could mediate assembly of dimers and helical filaments. When we purified *Citrobacter* Ubl<sub>3</sub>, we found that the protein eluted from a size exclusion column in two peaks: one at the void volume (suggestive of a large oligomer), and one at the position expected for a monomer (**Figure S5a**). We pooled and concentrated fractions for the oligomer peak and subjected the protein to inductively coupled plasma mass spectrometry (ICP-MS) to measure metal ion content. We measured ~0.2 molar equivalents of calcium, ~0.05 molar equivalents of magnesium, and vanishingly small amounts of manganese and iron (see **Materials and Methods**). These data support our assignment of ion densities in the crystal structures of *M. brachiatum* and *E. coli* Ubl<sub>3</sub> as Ca<sup>2+</sup>.

To further explore the role of divalent cations in Ubl<sub>3</sub> oligomerization, we performed size exclusion chromatography coupled to multi-angle light scattering (SEC-MALS) on three Ubl<sub>3</sub> proteins. We incubated purified E. coli Ubl<sub>3</sub>, M. brachiatum Ubl<sub>3</sub>, and Citrobacter Ubl<sub>3</sub> in a buffer containing EDTA to chelate any bound cations, then performed SEC-MALS (Figure 4a-c). In all three cases, we observed predominantly monomeric Ubl<sub>3</sub>, with *Citrobacter* Ubl<sub>3</sub> migrating as two peaks: one representing a monomer and a second between the expected molecular weight of a monomer and a dimer (Figure 4c). We next removed EDTA by buffer-exchange and performed SEC-MALS in a buffer containing  $Mg^{2+}$  (Figure 4d-f). In this condition, *E. coli* Ubl<sub>3</sub> and *M.* brachiatum Ubl<sub>3</sub> migrated as monomers (Figure 4d-e), while Citrobacter Ubl<sub>3</sub> shifted to a predominantly dimeric form (Figure 4f). Finally, we performed SEC-MALS in a buffer containing  $Ca^{2+}$  (Figure 4g-i). Both *M. brachiatum* Ubl<sub>3</sub> and *Citrobacter* Ubl<sub>3</sub> migrated in the void volume of the size exclusion column, forming large assemblies in the MDa size range (Figure 4h-i). E. coli Ubl<sub>3</sub> precipitated from solution in the presence of 5 mM Ca<sup>2+</sup>, but we could perform SEC-MALS on the protein after incubation with 50  $\mu$ M Ca<sup>2+</sup>. This analysis showed a mix of large oligomers in the MDa size range, plus apparent monomers and dimers (Figure 4g). Finally, we performed isothermal titration calorimetry (ITC) on Citrobacter Ubl<sub>3</sub> to measure the affinity and stoichiometry of  $Ca^{2+}$  binding. This analysis showed that *Citrobacter* Ubl<sub>3</sub> binds ~4 Ca<sup>2+</sup> ions per Ubl<sub>3</sub> protomer, with the best fit to the data suggesting one high-affinity binding site ( $K_d \sim 100$ nM), and three lower-affinity binding sites ( $K_d \sim 3 \mu$ M; Figure S3c).

## Ubl<sub>3</sub> proteins form helical filaments in the presence of Ca<sup>2+</sup> ions

To determine the molecular basis for Ca<sup>2+</sup>-mediated Ubl<sub>3</sub> oligomerization, we prepared cryoelectron microscopy (cryoEM) grids of oligomerized *Citrobacter* Ubl<sub>3</sub> and *M. brachiatum* Ubl<sub>3</sub>. For *Citrobacter* Ubl<sub>3</sub>, we used a fraction eluting in the void volume of a size exclusion column upon initial purification of the protein from *E. coli* expression without EDTA addition (**Figure S5a**). For *M. brachiatum* Ubl<sub>3</sub>, we added 5 mM CaCl<sub>2</sub> to purified monomeric Ubl<sub>3</sub>, incubated 60 minutes at 4°C, then prepared cryoEM grids. Initial images of both samples showed filaments ~20 nm in diameter (**Figure 5a, Figure S6a**). We collected a full cryoEM dataset for each sample and determined the structures using helical reconstruction methods,

resulting in a 2.4 Å resolution structure of *Citrobacter* Ubl<sub>3</sub> (**Figure S5**) and a 3.1 Å resolution structure of *M. brachiatum* Ubl<sub>3</sub> (**Figure S6**).

The cryoEM structure of *Citrobacter* Ubl<sub>3</sub> reveals a helical filament with Ubl<sub>3</sub> homodimers assembled identically to the dimers observed in our crystal structures of both *E. coli* and *M. brachiatum* Ubl<sub>3</sub> proteins (**Figure 5b, S3e-g**), stacked into a helical filament with a right-handed helical twist of 55° and a rise of 28 Å per Ubl<sub>3</sub> dimer (**Figure 5c**). Ubl<sub>3</sub> dimer-dimer interactions are nearly identical to that observed in the crystal structure of *M. brachiatum* Ubl<sub>3</sub>, with β-grasp domains 1 and 2 highly ordered and mediating filament formation, and β-grasp domain 3 extending outward from the filament. Despite the flexibility of β-grasp domain 3, we could use 3D variability analysis and local refinement to generate a 2.7 Å resolution map showing all three β-grasp domains, enabling us build and refine a model of the full-length protein (**Figure 5b**, **Figure S3e-f, Figure S5h-i**).

The cryoEM structure of *M. brachiatum* Ubl<sub>3</sub> reveals a helical filament with a right-handed helical twist of 63° and a rise of 30 Å per Ubl<sub>3</sub> dimer, distinct from the parameters in our crystal structure of the same protein (right-handed twist of 60° and rise of 17.6 Å per Ubl<sub>3</sub> dimer) (**Figure S6f**). The packing of Ubl<sub>3</sub> dimers in both structures is equivalent, with small conformational changes at dimer-dimer interfaces resulting in a significantly more extended filament as imaged by cryoEM versus x-ray crystallography. As in our x-ray crystal structure, our cryoEM structure of *M. brachiatum* Ubl<sub>3</sub> revealed no evidence of  $\beta$ -grasp domain 3, likely due to flexibility of the linker between  $\beta$ -grasp domains 2 and 3.

Close inspection of the cryoEM maps of both *Citrobacter* and *M. brachiatum* Ubl<sub>3</sub> revealed the presence of multiple ordered Ca<sup>2+</sup> ions bound to each Ubl<sub>3</sub> protomer (**Figure 5d, Figure S3**, **Figure S4e**). We observed five Ca<sup>2+</sup> ions bound to each *Citrobacter* Ubl<sub>3</sub> protomer, three of which were also observed in the *M. brachiatum* Ubl<sub>3</sub> cryoEM structure. Site #1 is equivalent to the intra-dimer Ca<sup>2+</sup> ion we observed in our crystal structures of *E. coli* and *M. brachiatum* Ubl<sub>3</sub>. In *Citrobacter* Ubl<sub>3</sub>, this Ca<sup>2+</sup> ion is coordinated by two main-chain carbonyl groups (residues 143 and 145) and the side-chain of a glutamate (E147) of one protomer, and the side-chain of an aspartate (D85) on the second protomer (**Figure 5d**). In the *M. brachiatum* Ubl<sub>3</sub> cryoEM structure, site #1 involves three main-chain carbonyl groups and the side-chain of E148 from one protomer, and D85 from the second protomer is nearby (**Figure S6g**). A second intra-dimer Ca<sup>2+</sup> ion (site #2) in *Citrobacter* Ubl<sub>3</sub> is coordinated by the side chains of two glutamate residues on opposite protomers (E62 and E68). The residues at site #2 are less well conserved in Ubl<sub>3</sub> proteins than the site #1 residues (**Figure S3**), and we did not observe this ion in either the *E. coli* or *M. brachiatum* Ubl<sub>3</sub> cryoEM structure.

We observe two inter-dimer  $Ca^{2+}$  ions in the *Citrobacter* Ubl<sub>3</sub> helical filament (**Figure 5d**). The  $Ca^{2+}$  ion at site #3 is symmetrically coordinated by an aspartate side chain (D30) and both the main-chain carbonyl and side-chain hydroxyl of a threonine (T32) on Ubl<sub>3</sub> protomers from two different dimers. Site #4 is also symmetric, and involves a main-chain carbonyl (residue 125) and an aspartate side chain (D130) side chain from two Ubl<sub>3</sub> protomers from two dimers. The residues involved in both sites #3 and #4 are conserved in Ubl<sub>3</sub> proteins (**Figure S3**), and we observe  $Ca^{2+}$  ions bound to both sites in our cryoEM structure of *M. brachiatum* Ubl<sub>3</sub>. (**Figure** 

**S6g**). Overall, our structural data closely match our ITC results that showed *Citrobacter* Ubl<sub>3</sub> binding ~4 molar equivalents of  $Ca^{2+}$  ions (**Figure S3c**): we identify five distinct  $Ca^{2+}$  binding sites in *Citrobacter* Ubl<sub>3</sub>, with the two inter-dimer sites symmetrically bridging two protomers each, thereby yielding a total of four molar equivalents of  $Ca^{2+}$  binding this protein.

To directly link  $Ca^{2+}$  binding to Ubl<sub>3</sub> oligomerization, we designed two multi-site mutants of *Citrobacter* Ubl<sub>3</sub>: mutant 1 (E62A/E68A/D85A/E147A) was designed to disrupt  $Ca^{2+}$  binding to intra-dimer sites #1 and #2, and mutant 2 (D30A/D130A) was designed to disrupt inter-dimer  $Ca^{2+}$  binding to sites #3 and #4. By SEC-MALS, we found that mutant 1 is monomeric in solution in both the absence and presence of  $Ca^{2+}$  ions, suggesting that  $Ca^{2+}$  binding at the intra-dimer sites #1 and #2 is critical for higher-order assembly (**Figure 5e**). Like wild-type *Citrobacter* Ubl<sub>3</sub>, mutant 2 migrated as two peaks in the absence of  $Ca^{2+}$ , one representing a monomer and a second between the expected molecular weight of a monomer and a dimer (**Figure 5f**). In the presence of  $Ca^{2+}$  ions at the intra-dimer sites #1 and #2 enables dimer assembly, but its inability to bind  $Ca^{2+}$  ions at the inter-dimer sites #3 and #4 disrupts filament assembly. Overall, these data strongly implicate  $Ca^{2+}$  binding in filament formation by *Citrobacter* Ubl<sub>3</sub>.

# Discussion

It is now understood that bacteria possess a range of biochemical pathways related to eukaryotic ubiquitination, which mediate protein conjugation in a variety of contexts including antiphage immunity<sup>7,8,10–13,15</sup>. To date, Type II CBASS operons were demonstrated to bear homology to a noncanonical eukaryotic ubiquitination pathway acting in autophagy<sup>10,11</sup>, and Type I/II Bil (bacterial ISG15-like) operons were shown to be related to canonical eukaryotic ubiquitination pathways <sup>12</sup>. Finally, two groups of Bub operons, previously termed 6E or DUF6527 operons<sup>7,8,10</sup>, encode predicted E1, E2, peptidase, and DUF6527 proteins in addition to Ubls with diverse predicted architectures<sup>14</sup>.

While bacterial Bil and Bub pathways encode Ubls with high structural similarity to eukaryotic ubiquitin, Ubl-target conjugation in bacteria is unlikely to constitute a degradation signal as it does in eukaryotic cells. Therefore, the functional consequences of Ubl-target conjugation in bacteria remains a key unanswered question. For one Type I Bil system, specific Ubl conjugation to phage central tail fiber proteins was shown to disrupt assembly of virions and also impair infectivity of fully-assembled virions<sup>13</sup>. The specific targets and the biological consequences of Ubl-target conjugation in Type II Bil and in Type I/II Bub are thus far unknown. These pathways may similarly function in antiphage immunity, or they may constitute stress response pathways that provide a selective advantage to their host in specific stress or pathogenic contexts.

A widespread group of bacterial Ubl proteins found in both Type II Bil and Type I/II Bub pathways is the Ubl<sub>3</sub> family, whose members possess three ubiquitin-like  $\beta$ -grasp domains. We show here that Ubl<sub>3</sub> proteins form helical filaments upon binding Ca<sup>2+</sup> ions, and that the Ca<sup>2+</sup> binding sites and the property of filament formation are conserved across diverse Ubl<sub>3</sub> proteins.

We envision two possible scenarios for these filaments' function in the context of stress response. In the first scenario, unperturbed cells possess Ubl<sub>3</sub> filaments that cannot be used as a substrate for the pathways' E1 and E2 proteins for target conjugation. Upon stress that reduces intracellular Ca<sup>2+</sup>, Ubl<sub>3</sub> filaments would disassemble, freeing Ubl<sub>3</sub> to be conjugated to target proteins. In a second scenario, Ubl<sub>3</sub> proteins are monomeric in unperturbed cells, possibly pre-conjugated to target proteins. Upon stress that increases intracellular Ca<sup>2+</sup>, Ubl<sub>3</sub> filaments would form and thereby sequester the conjugated target proteins. This scenario is conceptually similar to the known tendency of some metabolic proteins to form filaments in stress conditions that reversibly alter their activity<sup>21</sup>; in at least one case, filamentation of bacterial glutamine synthetase was shown to be regulated by metal ion binding<sup>22</sup>. In either scenario, we hypothesize that filament formation by Ubl<sub>3</sub> proteins represents a mechanism to respond to stress conditions that alter intracellular Ca<sup>2+</sup> concentration.

Bacterial Ubls show high diversity, possessing one to three β-grasp domains and N-terminal domains with unknown functions, some of which are predicted to mediate oligomerization. Just as bacterial ubiquitination-related pathways show high diversity and likely act in many stress-response contexts, bacterial Ubls likely have many roles that depend on their diverse architectures. Notably, a large proportion of bacterial Bub operons are associated with known DNA damage-responsive transcriptional regulators, including the WYL domain protein CapW/BrxR<sup>23–25</sup> and the two-protein CapH+CapP system<sup>26</sup> (Table S1). Future studies will likely need to move beyond antiphage immunity to explore the full range of adaptive benefits provided by these diverse, and as-yet mysterious, pathways.

# **Materials and Methods**

# **Bioinformatics**

To analyze bacterial Ubl diversity, Type II Bil operons identified by Chambers et al. <sup>12</sup> and DUF6527-containing operons identified by Millman et al. <sup>15</sup> were visually inspected in the IMG (Integrated Microbial Genomes & Microbiomes) database (https://img.jgi.doe.gov) for operon architecture and neighboring transcription factors. Structure predictions of Ubl proteins were performed by AlphaFold 2 <sup>17</sup>, AlphaFold 3 <sup>18</sup>, or ESMFold <sup>16</sup>. Disorder/LLPS predictions were performed using catGRANULE (http://service.tartaglialab.com/new\_submission/catGRANULE) <sup>27</sup>, and coiled-coil predictions were performed using PairCoil2 (https://cb.csail.mit.edu/cb/paircoil2/) <sup>28</sup>.

# Protein expression, purification, and characterization

All proteins used in this study are listed in Table S2. Codon-optimized genes encoding Ubl proteins without their C-terminally fused E2 domains (when present) into E. coli expression vectors encoding an N-terminal TEV protease-cleavable His6-tag (UC Berkeley Macrolab vector 2B-T, Addgene ID 29666) or an N-terminal TEV protease-cleavable His<sub>6</sub>-maltose binding protein tag (UC Berkeley Macrolab vector 2C-T, Addgene ID 29706). Vectors were transformed into E. coli Rosetta 2 pLysS (EMD Millipore), and 1 L cultures were grown at 37°C in 2XYT media to an OD<sub>600</sub> of 0.6 before induction with 0.25 mM IPTG at 20°C for 16-18 hours. Cells were harvested by centrifugation, resuspended in buffer A (25 mM Tris-HCl pH 8.5, 300 mM NaCl, 5 mM MgCl<sub>2</sub>, 10% glycerol and 5 mM mercaptoethanol) containing 5 mM imidazole, then lysed by sonication (Branson Sonifier). Lysates were clarified by centrifugation, then supernatants were passed over a Ni-NTA Superflow column (Qiagen) in buffer A. The column was washed in wash buffer (buffer A containing 20 mM imidazole), then eluted in elution buffer (buffer A containing 400 mM imidazole). Eluates were concentrated by ultrafiltration (Amicon Ultra; EMD Millipore). For crystallography, His<sub>6</sub>-tagged proteins were cleaved with TEV protease (expressed and purified in-house from plasmid pRK793 (Addgene #8827))<sup>29</sup> then passed over a HisTrap HP column (Cytiva) to remove His6-tagged TEV protease and uncleaved proteins. All proteins were finally passed over a Superdex 200 Increase size exclusion column (Cytiva) in size exclusion buffer (25 mM Tris-HCl pH 8.5, 300 mM NaCl, 5 mM MgCl<sub>2</sub>, 10% glycerol and 1mM DTT). Peak fractions were concentrated by ultrafiltration and stored at 4°C. For purification with EDTA treatment, eluates from the first Ni-NTA column were concentrated and 5 mM EDTA was added. After 16 hours at 4°C, the EDTA was removed by buffer exchange in centrifugal concentrators and the sample was passed over the size exclusion column in size exclusion buffer.

For characterization of oligomeric state by size exclusion chromatography coupled to multiangle light scattering (SEC-MALS), 100 µl of purified proteins at a concentration of 5 mg/ml were injected onto a size exclusion column (Superdex 200 Increase 10/300 GL, Cytiva) in size exclusion buffer, then light scattering and differential refractive index (dRI) profiles were collected using miniDAWN TREOS and Optilab T-rEX detectors (Wyatt Technology). SEC-MALS data were analyzed using ASTRA software version 8 and visualized with Prism version 10 (GraphPad Software). For analysis of divalent cations' role in oligomerization, proteins were pre-incubated in size exclusion buffer plus supplemented with 5 mM EDTA, 5 mM MgCl<sub>2</sub>, or 5 mM CaCl<sub>2</sub> for 30 minutes at  $4^{\circ}$ C, then passed over the size exclusion column in the same buffer.

# Crystallography

For crystallization of E. coli ZDHYS365 Ubl<sub>3</sub>, purified protein was subjected to surface lysine methylation by treating with borane (50 mM final concentration) and formaldehyde (100 mM final concentration) for 1 hour at 4°C, followed by addition of glycine (25 mM final concentration) to guench the reaction for 30 minutes, followed by buffer exchange by centrifugal concentration. Methylated protein at 10 mg/mL in crystallization buffer (25 mM Tris-HCl pH 8.5, 200 mM NaCl, 5 mM MgCl<sub>2</sub>, and 1 mM TCEP (tris(2-carboxyethyl)phosphine)) was mixed 1:1 with well solution containing 1 M LiCl, 0.1 M sodium acetate, and 30% PEG 6000 in sitting drop format. Crystals were harvested directly from the crystallization drop and frozen in liquid nitrogen. Diffraction data were collected at the Advanced Light Source (Lawrence Berkeley National Lab) beamline 5.0.1 on August 30, 2022 (collection temperature 100 K; x-ray wavelength 0.97741) (Table S3). Data were processed with XDS <sup>30</sup> and converted to structure factors with TRUNCATE <sup>31</sup>. The structure was determined by molecular replacement in PHASER <sup>32</sup> using a predicted structure from AlphaFold 2<sup>17</sup> as a search model. The model containing two protomers was manually rebuilt in COOT <sup>33</sup> and refined in phenix refine <sup>34</sup> using positional and individual isotropic Bfactor refinement. The final model has good geometry, with 98.77% of residues in favored Ramachandran space, 1.23% allowed, and 0% outliers. The overall MolProbity score is 1.07, and the MolProbity clash score is 2.12.

For crystallization of *M. brachiatum* Ubl<sub>3</sub>, purified protein was subjected to surface lysine methylation as above. Methylated protein at 12 mg/mL in the crystallization buffer was mixed 1:1 with a well solution containing 100 mM CAPS pH 10.5, 0.2 M Li<sub>2</sub>SO<sub>4</sub>, 1.2 M NaH<sub>2</sub>PO<sub>4</sub>, and 0.8 M K<sub>2</sub>HPO<sub>4</sub> in sitting drop format. Crystals were harvested into a cryoprotectant solution containing 30% glycerol and frozen in liquid nitrogen. Diffraction data were collected at the Advanced Photon Source (Argonne National Lab) NE-CAT beamline 24ID-C on March 2, 2023 (collection temperature 100 K; x-ray wavelength 0.97911 Å) (Table S3). Data were processed with the RAPD data-processing pipeline, which uses XDS for data indexing and reduction, POINTLESS <sup>31</sup> for space group assignment, and AIMLESS <sup>31</sup> for scaling. The structure was determined by molecular replacement in PHASER using a predicted structure from AlphaFold 2 as a search model. Despite crystallizing full-length protein, only  $\beta$ -grasp domains 1 and 2 were visible in the model. The model containing four protomers was manually rebuilt in COOT and refined in phenix.refine using positional and individual isotropic B-factor refinement. The final model has good geometry, with 99.49% of residues in favored Ramachandran space, 0.51% allowed, and 0% outliers. The overall MolProbity score is 1.11, and the MolProbity clash score is 3.22.

# **ICP-MS**

For ICP-MS, we purified *Citrobacter* Ubl<sub>3</sub> in size exclusion buffer lacking divalent cations, and concentrated fractions eluting from a size exclusion column at the void volume for analysis, to a final concentration of 25 mg/mL (0.956 mM). We diluted the protein 100-fold into 1% nitric acid in ultrapure water, then analyzed metal content using an iCAP RQ ICP-MS instrument (Thermo

Fisher Scientific). We measured manganese at 0.102 ppb (parts per billion), iron at 0.415 ppb, magnesium at 11.120 ppb, and calcium at 84.458 ppb. After accounting for the 100X dilution, these measurements correspond to 0.000185 mM manganese, 0.000727 mM iron, 0.0463 mM magnesium, and 0.192 mM calcium.

## **Isothermal Titration Calorimetry**

For ITC, *Citrobacter* Ubl<sub>3</sub> was purified with EDTA treatment, then passed over a size exclusion column in a buffer containing 25 mM Tris-HCl pH 8.5, 300 mM NaCl, and 1 mM NaN<sub>3</sub>, and fractions representing monomeric protein were pooled and concentrated. ITC was performed using an ITC-200 instrument (Malvern Panalytical), and four independent runs were performed at 25°C with 20-33  $\mu$ M protein in the analysis cell and 300  $\mu$ M CaCl<sub>2</sub> in the syringe (40 x 3  $\mu$ L injections with 120 seconds delay between each injection). Data were fit using a two-site binding model.

## Cryoelectron Microscopy of Citrobacter Ubl<sub>3</sub>

For structure determination of *Citrobacter* sp. RHBSTW-00271 Ubl<sub>3</sub>, fractions eluting in the void of a Superdex 200 Increase size exclusion chromatography columns (Cytiva) were pooled and concentrated to 1 mg/mL. Prior to use, Quantifoil copper 1.2/1.3 300 mesh grids were plasma cleaned for 12 seconds using a preset program in the Solarus II plasma cleaner (Gatan). A 3.5  $\mu$ L protein sample was applied to the grid within the environmental chamber adjusted to 4°C temperature and approximately 95% humidity in a Vitrobot Mark IV (Thermo Fisher Scientific). After a 5-second incubation, grids were blotted with a blot force of 4 for 4 seconds, the sample was then plunge-frozen into liquid nitrogen-cooled liquid ethane.

Grids were mounteds into standard AutoGrids (Thermo Fisher Scientific) for imaging. An initial dataset was collected on a Talos Arctica transmission electron microscope (Thermo Fisher Scientific) operated at 200 kV and equipped with a Gatan K3 direct electron detector. Movies were collected at a magnification of 130,000x and a pixel size of 0.95 Å, with a total dose of 55 e-/Å<sup>2</sup>. A defocus range of -0.5 to -2.2 was used during the data collection. In total, 396 movies were used in the final data processing after applying CTF fitting and excessive motion criteria.

All data processing was performed using cryoSPARC version 4 <sup>35</sup>. From the initial dataset, particle picking was performed using the blob picker with an estimated particle diameter of 200 Å. 189,755 initial particles were subjected to multiple rounds to 2D classification, yielding seven high-quality 2D classes from 53,925 particles that were used for template-based particle picking using the cryoSPARC Filament Tracer job. 207,260 particles were subjected to 2D classification and *ab initio* reconstruction, yielding an initial 3D volume from 101,204 particles. This volume was used to generate templates for a second round of particle picking using the Filament Tracer job. 189,327 particles were subjected to 2D classification, and helical refinement, yielding a final 3.4 Å resolution map.

The final dataset was collected on a Titan Krios G4 transmission electron microscope (Thermo Fisher Scientific) operated at 300 kV and configured for fringe-free illumination, equipped with a Falcon 4 direct electron detector with Selectris X energy filter. The microscope was operated in

EFTEM mode with a slit width of 20 eV. Automated data acquisition was performed using EPU (Thermo Fisher Scientific). Movies were collected at a magnification of 130,000x and a pixel size of 0.935 Å, with a total dose of 45 e-/Å<sup>2</sup>. A defocus range of -0.5 to -2.2 was used during the data collection. In total, 1,851 movies were used in the final data processing after applying CTF fitting and excessive motion criteria (**Table S4**).

Particle picking in the final dataset was performed using templates generated from the 3.4 Å map from the initial dataset, in the Filament Tracer job. The filament diameter was set to 200 Å and the separation distance between picks was set to 0.14 diameters, enabling picking of adjacent asymmetric units along the helical filament. Particles were picked with a 320x320 pixel box size and Fourier cropped to 80x80 pixels for initial processing. 1,634,973 particles were subjected to four successive rounds of 2D classification (in rounds 1 and 2, the data were split into three batches), yielding a final set of 892,949 particles. Ab initio reconstruction was performed with three models, and the best model contained 640,938 particles. Particles were re-extracted at 320x320 pixels with re-centering (yielding 629,161 particles) and used for helical refinement with D1 symmetry (two-fold rotational symmetry oriented 90° from the helical axis). Helical symmetry was not applied during refinement as initial particle picking used a singleasymmetric-unit step. Final helical parameters are a right-handed helical twist of 55.78° per asymmetric unit, and a helical rise of 28.4 Å per asymmetric unit. The final refinement using the Helical Refinement job type resulted in a 2.6 Å resolution map revealing  $\beta$ -grasp domains 1 and 2, which participate in helical packing. To resolve the flexible third  $\beta$ -grasp domain, particles were symmetry-expanded (using D1 symmetry) then subjected to local refinement using a mask covering four protomers. The resulting 2.4 Å resolution map was subjected to 3D variability analysis and clustering, resulting in one class with strong density for the third  $\beta$ -grasp domain. This class (containing 148,697 particles) was refined to 2.7 Å resolution (Table S4).

For modeling, initial models of each  $\beta$ -grasp domain generated by AlphaFold 2 were manually placed, then the model was manually rebuilt in COOT with manual addition of ordered Ca<sup>2+</sup> ions. The model was refined in phenix.refine using positional and individual isotropic B-factor refinement. For global refinement with four Ubl<sub>3</sub> protomers, strict non-crystallographic symmetry restraints were applied during refinement for all protein atoms (Table S4).

# Cryoelectron Microscopy of *M. brachiatum* Ubl<sub>3</sub>

For structure determination of *M. brachiatum* Ubl<sub>3</sub>, pooled fractions from the monomeric peak of EDTA-treated protein at 0.5 mg/mL were mixed with 5 mM CaCl<sub>2</sub>, incubated 60 minutes at 4°C, then a 3.5  $\mu$ L protein sample was applied to a plasma-cleaned Quantifoil grid within the environmental chamber adjusted to 4°C temperature and approximately 95% humidity in a Vitrobot Mark IV (Thermo Fisher Scientific). After a 5-second incubation, grids were blotted with a blot force of 4 for 4 seconds, the sample was then plunge-frozen into liquid nitrogen-cooled liquid ethane.

Grids were mounteds into standard AutoGrids (Thermo Fisher Scientific) for imaging, and a 1074 movie dataset was collected on a Titan Krios G3 transmission electron microscope (Thermo Fisher Scientific) operated at 300 kV and equipped with a Gatan K3 direct electron detector. Movies were collected at a magnification of 130,000x and a pixel size of 0.95 Å, with a

total dose of 50 e-/Å<sup>2</sup>. A defocus range of -1.2 to -2 was used during the data collection. In total, 1039 movies were used in the final data processing after applying CTF fitting and excessive motion criteria (**Table S4**).

All data processing was performed using cryoSPARC version 4. From the initial dataset, particle picking was performed using the Filament Tracer job with an estimated filament diameter of 120 Å. 1,303,971 initial particles were extracted with a box size of 400 pixels and Fourier cropped to 100 pixels. These particles were subjected to multiple rounds to 2D classification, yielding a dataset of 216,012 particles that were used for *ab initio* 3D reconstruction. 117,271 particles segregating to the best class were re-extracted at full resolution and refined using the Helical Refinement job with D1 symmetry, yielding a 3.22 Å map. This map was used to generate templates for a second round of particle picking using the Filament Tracer type, yielding a set of 495,217 particles. These were subjected to multiple rounds of 2D classification and 3D heterogeneous refinement before a final set of 151,090 particles were refined using the Helical Refinement type with D1 symmetry, yielding a final 3.08 Å resolution map (**Table S4**).

An initial model from the x-ray crystal structure was manually placed, then the model was manually rebuilt in COOT with manual addition of ordered Ca<sup>2+</sup> ions. The model was refined in phenix.refine using positional and individual isotropic B-factor refinement. For global refinement with four Ubl<sub>3</sub> protomers, strict non-crystallographic symmetry restraints were applied during refinement for all protein atoms (**Table S4**).

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**Figures** 



**Figure 1. Bacterial ubiquitin-like proteins show diverse architectures. (a)** Operon schematics of representative Type I and Type II Bil (Bacterial ISG15-like) operons, and Type I and Type II Bub (Bacterial ubiquitination-like) operons. Ubiquitin-like genes (*bilA/bubA*) are shown in orange; E2-like genes (*bilB/bubB*) in dark blue, JAB/JAMM peptidase genes (*bilC/bubC*) in magenta, E1-like genes (*bilD/bubD*) in yellow, and DUF6527 genes (*bubE*) in light blue. See **Table S1** for a catalog of Type II Bil, Type I Bub, and Type II Bub operons. (b) Ubl architecture in Type I Bub, Type II Bub, and Type II Bil operons. Ubls with one β-grasp domain (Ubl<sub>1</sub>) are shown in dark orange, those with two β-grasp domains (Ubl<sub>2</sub>) light orange, and three β-grasp domains (Ubl<sub>3</sub>) in light yellow. See **Table S1**. (c) Summary of Ubl architecture in Type II Bub, and Type II Bil operons. Ubl<sub>1</sub> proteins show diverse N-terminal domains. See **Figure S1** for structural predictions of Ubl<sub>1</sub> proteins with predicted disordered, coiled-coil, or DUF1508 N-terminal domains.



**Figure 2. Crystal structure of** *E. coli* Ubl<sub>3</sub>. (a) Domain structure of *E. coli* Ubl<sub>3</sub>. (b) Crystal structure of an *E. coli* Ubl<sub>3</sub> dimer, with one protomer colored dark orange/light orange/light yellow and the second protomer colored dark blue/medium blue/light blue. Bound Ca<sup>2+</sup> ions are shown as gray spheres. (c) Structure of *H. sapiens* ubiquitin (PDB ID 1UBQ) <sup>36</sup>, colored as a rainbow from N-terminus (blue) to C-terminus (red) and with  $\beta$ -strands 1-5 labeled. (d) Structure of *E. coli* Ubl<sub>3</sub>  $\beta$ -grasp domains 1-3, colored as in panel (b) and aligned with the structure of *H. sapiens* ubiquitin. (e) Close-up of the intra-dimer Ca<sup>2+</sup> ion bound by *E. coli* Ubl<sub>3</sub>  $\beta$ -grasp domains 1-2 showing conservation of this site.



**Figure 3. Crystal structure of** *M. brachiatum* Ubl<sub>3</sub>. (a) Domain structure of *M. brachiatum* Ubl<sub>3</sub>. (b) Crystal structure of an *M. brachiatum* Ubl<sub>3</sub> dimer, with one protomer colored dark orange/light orange and the second protomer colored dark blue/medium blue ( $\beta$ -grasp domain 3 is disordered and not shown). Bound Ca<sup>2+</sup> ions are shown as gray spheres. *Inset:* Close-up of the intra-dimer Ca<sup>2+</sup> ion. See **Figure S4** for detailed views of individual  $\beta$ -grasp domains in each bacterial Ubl<sub>3</sub> protein. (c) Structure of a *M. brachiatum* Ubl<sub>3</sub> filament assembled by a combination of crystallographic and non-crystallographic symmetry. Six Ubl<sub>3</sub> dimers (three asymmetric units related by a P3<sub>1</sub> screw axis) are depicted in alternating blue and orange. Bound Ca<sup>2+</sup> ions are shown as spheres in the cartoon depiction at right and labeled according to whether they link protomers within a dimer (intra-dimer) or link protomers in different dimers (inter-dimer). *Inset:* Close-up of the inter-dimer Ca<sup>2+</sup> ion.

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**Figure 4. Ubl**<sub>3</sub> **proteins oligomerize in the presence of Ca**<sup>2+</sup>**. (a-c)** SEC-MALS analysis of *E. coli* Ubl<sub>3</sub> (a), *M. brachiatum* Ubl<sub>3</sub> (b), and *Citrobacter* Ubl<sub>3</sub> (d) in the absence of divalent cations. Black lines indicate dRI (change in refractive index; left Y axis) and red dots indicate measured molar mass (right Y axis; shown in logarithmic scale). The expected molar mass of a monomer species is shown as a dotted line. **(d-f)** SEC-MALS analysis of *E. coli* Ubl<sub>3</sub> (d), *M. brachiatum* Ubl<sub>3</sub> (e), and *Citrobacter* Ubl<sub>3</sub> (f) in the presence of 5 mM MgCl<sub>2</sub>. The expected molar mass of monomer and dimer species are shown as dotted lines. **(g-i)** SEC-MALS analysis of *E. coli* Ubl<sub>3</sub> in the presence of 50 µM CaCl<sub>2</sub> (g), *M. brachiatum* Ubl<sub>3</sub> in the presence of 5 mM CaCl<sub>2</sub> (h), and *Citrobacter* Ubl<sub>3</sub> in the presence of 5 mM CaCl<sub>2</sub> (i).



Figure 5. CryoEM structure of a Citrobacter Ubl<sub>3</sub> filament. (a) CryoEM micrograph of Citrobacter Ubl<sub>3</sub> filaments (see Figure S5a for size exclusion chromatography). (b) Top: Domain structure of Citrobacter Ubl<sub>3</sub>. Bottom: CryoEM structure of a Citrobacter Ubl<sub>3</sub> dimer (see Figure **S5**) with one protomer colored dark orange/light orange/light yellow and the second protomer colored dark blue/medium blue/light blue. Bound Ca<sup>2+</sup> ions are shown as gray spheres and labeled 1-5. (c) Closeup views of  $Ca^{2+}$  sites 1-5. Sites 1 and 2 link Ubl<sub>3</sub> protomers within a dimer (intra-dimer), and sites 3 and 4 link Ubl<sub>3</sub> protomers in different dimers (inter-dimer). Site 5 is coordinated entirely by residues in β-grasp domain 3. (d) CryoEM structure of the *Citrobacter* Ubl<sub>3</sub> filament (see Figure S5), showing six Ubl<sub>3</sub> dimers (alternating blue and orange). Bound Ca<sup>2+</sup> ions are shown as spheres in the cartoon depiction at right. (e) SEC-MALS analysis of *Citrobacter* Ubl<sub>3</sub> mutant 1 (E62A/E68A/D85A/E147A), designed to disrupt intra-dimer Ca<sup>2+</sup> sites 1 and 2. Top panel shows analysis in the absence of divalent cations, and bottom panel shows analysis in the presence of 5 mM CaCl<sub>2</sub>. The expected molar mass of a monomer species is shown as a dotted line. (f) SEC-MALS analysis of *Citrobacter* Ubl<sub>3</sub> mutant 2 (D30A/D130A), designed to disrupt inter-dimer Ca<sup>2+</sup> sites 3 and 4. Top panel shows analysis in the absence of divalent cations, and bottom panel shows analysis in the presence of 5 mM CaCl<sub>2</sub>. The expected molar masses of monomer and dimer species are shown as a dotted line.

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