Expression and Purification of a Cleavable Recombinant Fortilin from *Escherichia coli* for Structure Activity Studies

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Expression and purification of a cleavable recombinant fortilin from *Escherichia coli* for structure activity studies

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ABSTRACT

Complications related to atherosclerosis account for approximately 1 in 4 deaths in the United States and treatment has focused on lowering serum LDL-cholesterol levels with statins. However, approximately 50% of those diagnosed with atherosclerosis have blood cholesterol levels within normal parameters. Human fortilin is an anti-apoptotic protein and a factor in macrophage-mediated atherosclerosis and is hypothesized to protect inflammatory macrophages from apoptosis, leading to subsequent cardiac pathogenesis. Fortilin is unique because it provides a novel drug target for atherosclerosis that goes beyond lowering cholesterol and utilization of a solution nuclear magnetic resonance (NMR) spectroscopy, structure-based drug discovery approach requires milligram quantities of pure, bioactive, recombinant fortilin. Here, we designed expression constructs with different affinity tags and protease cleavage sites to find optimal conditions to obtain the quantity and purity of protein necessary for structure activity relationship studies. Plasmids encoding fortilin with maltose binding protein (MBP), 6-histidine (6His) and glutathione-S-transferase (GST), N-terminal affinity tags were expressed and purified from *Escherichia coli* (E. coli). Cleavage sites with tobacco etch virus (TEV) protease and human rhinovirus (HRV) 3C protease were assessed. Despite high levels of expression of soluble protein, the fusion constructs were resistant to proteinases without the inclusion of amino acids between the cleavage site and N-terminus. We surveyed constructs with increasing lengths of glycine/serine (GGS) linkers between the cleavage site and fortilin and found that inclusion of at least one GGS insert led to successful protease cleavage and pure fortilin with conserved binding to calcium as measured by NMR.

1. Introduction

Atherosclerosis, or the hardening and narrowing of arterial walls, is the leading cause of death worldwide [1]. Atherosclerogenesis occurs when low density lipoproteins (LDL) in plasma diffuse into the intima of arteries and then become oxidized (oxLDL), followed by uptake into macrophages. Macrophages eventually form foam cells, which over time develop into plaques on the arteries, leading to blockages in blood flow that cause cardiac events such as infarction, stroke, hypertension, etc. Presently, medications used to treat atherosclerosis largely depend on inhibiting the formation of cholesterol in *vivo* via the use of statins, which can cause undesirable side effects. For example, HMG-CoA reductase inhibitors (statin drug class) can cause leg cramps in patients at night that lead to decreased compliance with treatment [2]. Additional side effects may include cataracts and short-term memory loss [3]. Moreover, a recent study of 136,905 people showed that nearly half of all patients brought into the emergency room with heart attacks had normal to low LDL levels in their blood [4]. This finding made evident the need to develop drugs that target other mechanisms in the pathology of atherosclerosis than the in *vivo* formation of high cholesterol.

Fortilin is a promising drug target to treat atherosclerosis [5]. Also known as translationally controlled tumor protein (TCTP) or histamine releasing factor (HRF), fortilin is a highly conserved anti-apoptotic protein [6]. Fortilin consists of 172 amino acids and is found throughout the human body; in the cytosol, nucleus, and mitochondria of all cell types, blood and extracellular space [7,8]. Fortilin exerts its...
anti-apoptotic effect through multiple mechanisms including calcium complexation and p53 inactivation, but it contributes to atherosclerosis by keeping macrophages in the M1 pro-inflammatory phenotype [5,9]. Fortilin has been implicated in the progression of atherosclerosis by reducing macrophage apoptosis by binding to and inhibiting p53, a transcription factor that regulates the transcription of BAX, a pro-apoptotic protein. This mechanism promotes foam cell formation, leading to deposits on arterial walls [9]. Fortilin also exhibits anti-apoptotic activity by scavenging free calcium in the cytoplasm of cells, thus exerting a protective effect from calcium-induced apoptosis [10]. Specifically, inhibition of the pro-survival activity of could provide an orthogonal strategy to treat atherosclerosis by shutting down the protective effects of fortillin.

Fortilin is a 19 kDa protein that has 11 beta-strands and an extended flexible loop between residues Arg36 and Ile67 (Fig. 1A). The amino- and carboxy-termini are in proximity to one another, creating a cavity that could be interrogated for small molecule binding. The calcium binding site is coordinated by six oxygen atoms within the beta barrel among residues N131, Q133, L149 and D150 (Fig. 1B) [11]. While a direct connection between fortillin, calcium scavenging, and atherosclerosis has not yet been made, calcium-dependent apoptosis does occur during ox-LDL-induced apoptosis and it is hypothesized that fortillin could play a role through its calcium scavenging activity [12].

A structure-based approach to targeting fortillin will provide important information for drug screening and defining drug binding sites. Structure-based drug design (SBDD) is a technique commonly used to create small molecule therapeutics that specifically target a given protein [13,14]. Solution nuclear magnetic resonance spectroscopy (NMR) is an excellent tool for SBDD, for example, in screening small molecule inhibitors (SMIs), mapping binding hot spots, and determining binding affinities and kinetics [14]. A solution structure of recombinant fortillin has been reported using NMR [11]. Feng et al., used a construct with a 6His affinity tag and required high salt (200 mM) conditions for high quality NMR spectra.

NMR experiments require milligram quantities of pure, isotopically enriched, and bioactive protein. E. coli expression platforms are routinely used as they allow for straightforward, high yield and affordable isotope enrichment. Affinity tags are often used for protein purification [15]. Ideally, the affinity tags should be removed by enzymatic cleavage to ensure no pseudo binding effects due to the presence of the affinity tag. The N- and C-termini of fortillin form a beta strand that forms a pocket in the native fortillin structure. Affinity tags attached at either the N- or C-terminus complicate the potential to target this site. Functional studies of fortillin with an affinity tag are inconclusive due to the presence of the affinity tag at the suspected site of small molecule binding site [16,17]. Here, we describe the design and assessment of several fortillin constructs for the purpose of expressing and purifying a recombinant fortillin molecule with minimal non-native residues. Inclusion of a short GGS linker between an HRV 3C protease cleavage site and the N-terminus of fortillin resulted in milligram quantity of isotopically enriched 15N GGS-fortillin. Calcium binding was observed by NMR titrations, demonstrating the structure and activity of this construct were maintained. The work presented here shows that pure recombinant fortillin can be proteolytically cleaved from an affinity tag and retain biological activity by the insertion of a GGS linker sequence.

2. Results and discussion

Initial efforts to produce recombinant tag-less fortillin were hindered by low or no cleavage with either TEV or HRV 3C proteases directed at
N-terminal cleavage sites with either 6His, maltose binding protein (MBP), glutathione-S-transferase, or 6His-MBP affinity tag constructs (Fig. 2). Observed cleavage efficiencies for these constructs led to development of a systematic approach to define a minimal linker length required for effective and efficient affinity tag removal.

2.1. Expression and purification of recombinant human fortilin

Each fortilin construct was made using the ATUM pD441 T5 IPTG inducible promoter system. For each of the constructs used, soluble and high purity quantities from 100 mL LB cultures grown at 37 °C were successful. After affinity chromatography using glutathione agarose to purify the (GGS)_n constructs, GST-(GGS)_n-fortilin was purified typically with some GST protein made during the protein expression process (SI. Figs. 1–3, 5). For cleavage analysis, the cleaved fortilin construct was compared to the un-cleaved construct, so we decided not to purify further than affinity chromatography for this reason. The same occurred with the MBP-containing constructs where some MBP protein was eluted during affinity chromatography with the desired purified protein (SI. Figs. 4 and 6). Samples containing pure MBP construct were combined and concentrated to workable small volumes for cleavage with either TEV or 3C protease. For the initial cleavage analysis, all constructs were grown at smaller culture volumes (100 mL) to save on materials. The construct we decided to use for NMR titration analysis was the GST-GGS-fortilin construct because cleavage with 3C protease provided the fewest amino acids remaining at the N-terminus of the protein, leaving only GGS-fortilin (SI. Fig. 7). The yield from a 1 L, 15N-labeled culture was 2.08 mg by a reading of the A$_{280}$ and an ε value of 0.595 (mL/mg)$^{-1}$ cm$^{-1}$. This was an ideal amount for an approximately 200 μM sample for NMR titration analysis.

2.2. Protease cleavage sites distal to fortilin increases MBP removal effectiveness

After treatment of up to 48 h with TEV protease, there was no evidence by SDS-PAGE that the 6His-TEV-fortilin exhibited any cleavage (Fig. 4, lanes 7–10). Next, MBP-TEV-fortilin was treated TEV protease for up to 48 h at room temperature, with again no evidence of cleavage (Fig. 3, lanes 9–12). A construct consisting of 6His-MBP-10N–3C-TEV-fortilin treated with TEV again showed no evidence of cleavage (Fig. 3, lanes 4 and 7). TEV activity was validated with a positive control of MBP-TEV-OSM, or Oncostatin M, a construct known to cleave successfully upon addition of TEV protease [18]. Interestingly, the 6His-MBP-10N–3C-TEV-fortilin construct contained an 3C HRV protease cleavage site distal to the N-terminal site that was left intact after cloning in the TEV site. Treatment with 3C resulted in 72% cleavage after 24 h with no further cleavage after 48 h (Fig. 3, lanes 5 & 8).

2.3. Protease cleavage sites distal to fortilin increases GST removal effectiveness

The GST-3C-TEV-fortilin construct showed no cleavage after two days of incubation with TEV protease (day 2 data not shown). However, upon incubation with HRV 3C protease, cleavage was observed after 24 h. TEV-fortilin is indicated by an arrow in Fig. 4. No cleavage was observed for the 6His-TEV-fortilin construct upon addition of TEV protease after 48 h of incubation at room temperature. The 6His-TEV-fortilin band was confirmed by mass spectrometry, demonstrating no cleavage.
cleavage site is N-terminal to the TEV protease cleavage site. The arrow indicates successfully cleaved TEV-fortilin, where the 3C indicate molecular weight markers on a 4% bis-acrylamide gradient SDS gel. The arrow indicates successfully cleaved TEV-fortilin, where the 3C indicate molecular weight markers on a 4% bis-acrylamide gradient SDS gel.

Fig. 4. Protease cleavage sites located proximal to the N-terminus are resistant to cleavage. Cleavage reactions of GST and 6His constructs against a positive control lane of MBP-TEV-OSM. Protein ladder is shown in the first lane to indicate molecular weight markers on a 4–12% bis-acrylamide gradient SDS gel. The arrow indicates successfully cleaved TEV-fortilin, where the 3C indicate molecular weight markers on a 4% bis-acrylamide gradient SDS gel.

2.4. A minimum GGS tripeptide is required for efficient protease cleavage

It is well-documented that affinity tags can introduce steric hindrance, making the enzymatic cleavage less efficient. Further, the 6His-MBP-10N–3C-TEV-fortilin construct has an additional Human 3C rhinovirus protease cleavage site N-terminal to the TEV cleavage site. Treatment of the 6His-MBP-10N–3C-TEV-fortilin with 3C protease after 24 h at 4 °C resulted in 72% cleaved fortilin. However, this construct was undesirable because it left additional 3C (Leu-Glu-Val-Leu-Phe-Gln-Gly-Pro) and TEV (Glu-Asn-Leu-Tyr-Phe-Gln-Gly-Ser) amino acids; further, there was no additional cleavage observed after 48 h. These observations led us to hypothesize that for efficient cleavage, a linker was needed between the protease cleave site and the fortilin protein.

We systematically investigated what minimum linker length was required for efficient protease cleavage. We used (GGS)ₙ linker sequences where N = 0, 1, 2 or 3. We found that the inclusion of just one GGS linker sequence (GGS)₁ led to cleaved fortilin with the sequence GGS-fortilin; this result was verified against a negative control of HRV 3C protease. While additional linker sequences proved successful, we moved forward with (GGS)₁-fortilin for NMR analysis to minimize N-terminal binding interference by tag amino acids.

Incorporating a Gly-Gly-Ser (GGS) linker between the protease cleavage site and fortilin increased enzymatic protease cleavage efficiency. We chose to incorporate a repeating GGS motif between the protease cleavage site and fortilin. GGS motifs were chosen for their flexibility, experimentally determined random coil behavior, and charge neutrality [19,20]. GGS linker units were either not included in the construct (GGS)₀ or included in increasing repeats of GGS from (GGS)₁ to (GGS)₃. Addition of just one GGS linker to the N-terminus of fortilin was sufficient to produce a successfully cleaved construct, with approximately 86.5% cleavage GGS-fortilin after treatment with 3C protease (Table 1). Addition of two GGS linker sequences provided 81.5% cleavage, and addition of three GGS linkers resulted in 85% cleavage efficiency. Confirmation of cleavage reactions was analyzed by SDS-PAGE (Fig. 5) and compared to a positive control of GST-3C-Syk protein to ensure protease activity. All samples were also compared against negative controls wherein no protease was added. Addition of extra (GGS)ₙ linkers showed no increase in cleavage efficiency. Addition of one GGS linker sequence at the N-terminus of fortilin appeared to have provided sufficient access for a protease to cleave the affinity tag from fortilin for biological studies. Recovery yields after protease cleavage are available in Table 1. Yields were quantified in comparison to the respective non-cleaved construct on the same SDS gel.

2.5. Calcium titration of (GGS)₁-fortilin

It is well established that fortilin binds to calcium and the binding site has been characterized (see Fig. 1) [11,22,23]. For the present study, we reproduced CaCl₂ titration analysis with the ¹⁵N-labeled (GGS)₁-fortilin protein using NMR spectroscopy and obtained a Kᵣ value of 13 μM, which is within reason to previously reported affinities of fortilin for Ca²⁺ [11]. Multiple literature reports provide the Kᵣ values for calcium binding to fortilin that range from 10 μM to 17.5 μM to 22 mM and 25 mM [10,11,24]. The present study shows that recombinant fortilin, in the absence of an affinity tag, still binds calcium. Biologically, fortilin binds calcium to mediate histamine release from basophils, which play a critical role in immune system function [6].

<table>
<thead>
<tr>
<th>Construct</th>
<th>Cleaved construct</th>
<th>Percent (%) recovery of cleaved fortilin</th>
</tr>
</thead>
<tbody>
<tr>
<td>6His-TEV-fortilin</td>
<td>fortilin</td>
<td>0</td>
</tr>
<tr>
<td>6His-MBP-10N–3C-TEV-fortilin</td>
<td>TEV-fortilin</td>
<td>72</td>
</tr>
<tr>
<td>GST-3C-TEV-fortilin</td>
<td>TEV-fortilin</td>
<td>28</td>
</tr>
<tr>
<td>GST-3C-(GGS)₀-fortilin</td>
<td>fortilin</td>
<td>0.5</td>
</tr>
<tr>
<td>GST-3C-(GGS)₁-fortilin</td>
<td>GGS-fortilin</td>
<td>86.5</td>
</tr>
<tr>
<td>GST-3C-(GGS)₂-fortilin</td>
<td>(GGS)₂-fortilin</td>
<td>81.5</td>
</tr>
<tr>
<td>GST-3C-(GGS)₃-fortilin</td>
<td>(GGS)₃-fortilin</td>
<td>85</td>
</tr>
</tbody>
</table>

Table 1

Percent yield of tag free fortilin after one day to cleave constructs. Yields quantified using SDS PAGE image analysis using GelAnalyzer [21].

Fig. 5. Cleavage reactions of (GGS)ₙ constructs where N = 0, 1, 2 or 3 against a positive control lane of cleaved GST-3C-Syk. Protein ladder is shown in the first lane to indicate molecular weight markers on a 4–12% polyacrylamide SDS gradient gel. Arrow indicates successfully cleaved (GGS)ₙ-fortilin that was confirmed by mass spectrometry.
A 1 M aqueous solution of calcium chloride (CaCl$_2$) was titrated into a solution of 176 μM (GGS)$_2$-fortilin. Aliquots of 1 μL CaCl$_2$ were used for each titration point for a total of 6 points. Notable peak shifts in the titration can be seen in Fig. 6 where arrows indicate increasing concentrations of CaCl$_2$. The combined chemical shifts were calculated for Y132 and a quadratic expansion of the isotherm binding model binding isotherm was fit to the data to yield a binding affinity of $K_d$ of 14 mM ± 3.6 mM.

### 3. Conclusion

The successful cleavage of the 6His-MBP-10N-3C-TEV-fortilin and GST-3C-TEV-fortilin constructs led us to the hypothesis that, at the N-terminus, space is required for a protease searching for a cleavage recognition sequence to gain access to the cleavage site. Insertion of just one GGS linker sequence, at the N-terminus and before the cleavage site, provided 78% recovery of GGS-fortilin (Fig. 5). Further characterization of this activity needs to be done to determine the mechanism by which addition of a GGS sequence facilitates cleavage, but to also characterize the interaction of small molecules at the N-terminus, such as artemisinin and DHA, which bind but have not been characterized in human recombinant fortilin [16,17]. Additionally, NMR studies need to be performed on fortilin in other expression systems, such as yeast, to explore the possibility of post-translational modifications (PTMs), and to determine the differences in binding between recombinant fortilin products in E. coli, P. pastoris, and human cell lines.

### 4. Materials and methods

#### 4.1. Reagents

Unless otherwise mentioned, chemical reagents were purchased from Fisher Scientific (Waltham, MA). Isotopically enriched reagents were purchased from MilliPoreSigma (St Louis, MO).

**4.1.1. Expression and purification of S219V TEV protease from E. coli**

Recombinant TEV protease was expressed and purified using pRK793 [28] (a gift from David Waugh (Addgene plasmid # 8827; http://n2t.net/addgene:8827; RRID:Addgene_8827)) in BL21 (DE3) strain E. coli. From a glycerol stock, 20 mL of Luria Bertani (LB) broth containing 100 μg/mL ampicillin, 25 μg/mL chloramphenicol was inoculated, and incubated overnight at 37 °C with shaking at 200 rpm. This overnight culture was then used to inoculate 1L LB containing 100 μg/mL ampicillin, 25 μg/mL chloramphenicol with shaking at 37 °C with 200 rpm. Once the OD$_{600}$ reached between 0.5 and 0.7, a pre-induction sample equivalent to 1 mL of cells at OD$_{600}$ of 0.8 was collected, and the culture flask placed on ice and gently rotated every few minutes to cool the cultures. The incubator was then set to 30 °C and the cultures were reintroduced at the elevated temperature and expression was induced with 1 mM IPTG, with shaking at 200 rpm. This culture was incubated overnight. On the third day, a post-induction sample equivalent to 1 mL of cells at OD$_{600}$ of 0.8. The pre- and post-induction samples were used to analyze expression using SDS-PAGE. Lysis was halted by pouring the cultures into pre-weighed 500 mL centrifuge bottles and the cultures centrifuged at 6000 × g for 30 min at 4 °C. After centrifugation, the supernatants were removed, and the mass of the pellets was determined. The pellet was resuspended in 40 mL of lysis buffer consisting of 50 mM Tris-Cl pH 8.0, 300 mM NaCl, 20 mM imidazole, 10% (w/v) sorbitol, 20 mM β-mercaptoethanol, 0.02 mg/mL lysozyme, and 1X HALT protease inhibitors (Thermofisher, USA). The suspension was sonicated on ice water at 60% power, 30 s on/30 s off for 4.5 min of total pulse time and a 75 μl sample collected for SDS-PAGE to check post sonication pellet and supernatant. The sonicated suspension was centrifuged at 16,000 × g for 30 min at 4 °C. The supernatant was collected and filtered using a 0.45 μm PES filter. To the filtered lysate, 5 mL of Ni-NTA superflow slurry (Qiagen Cat #30410/30,430) was added and the mixture incubated overnight at 4 °C with rocking. On the fourth day, the lysate/resin mix was loaded into a column and the flow through collected. Purification was carried out with a BioLogic LP System using EconoColumn 1.0 × 10 cm chromatography.

Fig. 6. $^1$H,$^15$N HSQC spectral overlay of GGS-fortilin titrated with calcium. Y132, Y151, N131, Q133, and H77 are labeled with arrows indicating increasing concentrations of calcium.
columns (Bio-Rad, Hercules, CA, USA). To line A, the lysis buffer described previously was attached. To line B, the same lysis buffer with the only difference being a 300 mM imidazole concentration was attached. Buffer A was run over the column at a flow rate of 1.0 mL/min until the A280 stabilized, all while collecting the flow through. The program for protein purification was as follows: Line A for 10 min, Line B (0%-100% B, 1.0 mL/min) for 60 min, hold Line B for 10 min, and back to 100% A for 10 min. Results were evaluated by SDS-PAGE. Fractions containing purified TEV protease were combined, and protein concentration determined by absorption at 280 nm. Pooled fractions were diluted to a final concentration of 1 mg/mL in a 50% glycerol solution, and 0.5 mL aliquots stored at −80 °C.

4.1.2. Expression plasmid constructs
Plasmids were purchased from ATUM (Newark, CA) using their pD441-SR vector. These constructs consisted of glutathione S-transferase (GST), maltose binding protein (MBP), and 6-histidine (6His) affinity tags, human rhinovirus (HRV) 3C and tobacco etch virus (TEV) protease recognition sites, and glycine-glycine-serine (GGS) linkers of varying length (GGS)n, where N = 0, 1, 2 or 3 at the N-terminus of fortilin in the pD441-SR expression vector.

4.1.3. Expression of recombinant fusion proteins
Plasmids were transformed into 50 μL of competent E. coli cells using the heat-shock method at 42 °C. Single colonies were inoculated into 5 mL of LB medium containing 50 μg/mL kanamycin and grown at 37 °C overnight with shaking at 250 rpm. One mL of each overnight culture was inoculated into 100 mL LB medium containing 50 μg/mL kanamycin and grown at 37 °C with shaking at 250 rpm until the OD600 reached between 0.60 and 0.80. Once reached, IPTG was added to 0.1 mM to induce protein expression. Expression continued at these conditions for 6 h. Cells were harvested by centrifugation at 4,000 × g for 20 min and expression was assessed by SDS-PAGE on 4–12% bis-acrylamide gradient gels.

4.1.4. Expression of 15N-labeled recombinant fusion proteins for NMR spectroscopy
Plasmids were transformed into 50 μL of competent E. coli cells using the heat-shock method at 42 °C. Single colonies were inoculated into 5 mL of LB medium containing 50 μg/mL kanamycin and grown at 37 °C and grown for approximately 8 h. One mL of each culture was inoculated into 50 mL of 15N-labeled M9 minimal media containing 50 μg/mL kanamycin and grown at 37 °C overnight.

4.1.5. Cell lysis
Cell pellets (approximately 0.13 g each) from 100 mL cultures were resuspended in 2 mL lysis buffer containing 1X phosphate buffered sal- line (PBS), pH 7.4 and 0.1% Triton X-100. Lysomyze was added to a final concentration of 0.25 mg/mL as well as ½ tablet of HALT protease inhi- bitor cocktail (Pierce, Rockford, IL). The lysate was then subjected to sonication at 70% power for 30 s on/30 s off for a total of 2 min. Samples were then centrifuged at 17,000 × g for 45 min at 4 °C. The resulting lysate was filtered through a 0.45 μm PES filter and subjected to affinity purification.

4.1.6. Purification of GST-3C-(GGS)n-fortilin and GST-3C-TEV-fortilin from BL21(DE3) cells
GSTrap sepharose 4B 1 mL pre-packed columns were used for puri- fication (GE Healthcare, Marlborough, MA). Separate columns were used for each linker sequence to avoid contamination of samples. Samples were loaded onto the column with a flow rate of 1.0 mL/min and washed with 1X PBS with a flow rate of 1.0 mL/min for 20 mL, followed by a 10 mL elution with GSTrap elution buffer containing 50 mM Tris-HCl pH 8.0 and 10 mM reduced glutathione. Elution was fol- lowed by a 10 mL wash at 1.0 mL/min with 1X PBS (Supp. Figs 1-3 and 5). All solutions used for affinity chromatography were filtered through 0.22 μm nylon filters and degassed under vacuum with rigorous stirring for at least 1 h.

4.1.7. Purification of 6His-MBP-10N–3C-TEV-fortilin and MBP-TEV-fortilin
Amylose resin (New England Biolabs, Ipswich, MA) was used for purification of MBP-tagged proteins. An approximately 50% slurry of affinity beads to 20% ethanol were used to pack a column by gravity until the column volume reached 3 mL. The column was equilibrated with 25 mL of column buffer containing 20 mM Tris-HCl pH 7.4, 0.2 M NaCl and 1 mM EDTA. The clarified lysate was added to the column and the flow through collected. This step was repeated once to improve protein binding. Next, the column was washed with 10 column volumes (CV) of the column buffer described above. This was followed by elution by flowing 5 CV of column buffer containing 10 mM maltose, collected over three elution steps. A final wash step of 10 CV column buffer was performed to remove any remaining protein (Supp. Figs. 4 and 6).

4.1.8. Purification of 6His-TEV-fortilin
Nickel agarose resin (HisPur™ Ni-NTA Superflow agarose, (Ther- moFisher Scientific, Waltham, MA) was used for purification of 6His-TEV-fortilin. An approximately 50% slurry of affinity beads in 20% ethanol were used to pack a column by gravity until the column volume reached 3 mL. The column was equilibrated with 25 mL of buffer containing 50 mM Tris-Cl pH 8.0, 300 mM NaCl, 20 mM imidazole. Next, the clarified lysate was added to the column and the flow through collected. This step was repeated once to improve binding to the column. The column was then washed with 10 CV of buffer containing 50 mM Tris-Cl pH 8.0, 300 mM NaCl, 20 mM imidazole. This was followed by elution with 5 CV of buffer containing 50 mM Tris-Cl pH 8.0, 300 mM NaCl, 200 mM imidazole (Supp. Fig. 5).

4.1.9. HRV 3C cleavage of affinity tags from fusion proteins and removal of proteases
Samples containing purified fusion protein containing the 3C protease site (LEVLFQ/GP) underwent cleavage with either human rhino- virus 3C (HRV 3C) using the Pierce HRV 3C protease solution kit (Pierce, Rockford, IL) according to manufacturer protocols. Protease removal was also done according to manufacturer protocols by dialysis using 0.5–3.0 mL capacity dialysis cassettes (ThermoFisher Scientific, Waltham, MA) into 3C cleavage buffer containing 50 mM Tris-Cl pH 7.0, 150 mM NaCl and 1 mM EDTA. Samples were dialyzed and cleaved with 23 μL of the 3C protease at 4 °C; 25 μL of sample was checked for cleavage by SDS-PAGE after 24 and 48 h. For the (GGS)1 construct, the cleaved protein (GGS-fortilin) was buffer exchanged from protease buffer by size exclusion chromatography using an S75 column equilibrated within 100 mM NH4HCO3 (Cytiva, Marlborough, MA). Fractions containing pure protein were combined and concentrated using 5 MWCOS PES Vivaspin 20 columns (Sartorius, Stonehouse, UK) by centrifugation at 4000 × g until the volume reached approximately 10 mL. These were then aliquoted such that microcentrifuge tubes contained 2 mg protein each and lyophilized at 2 atm and −80 °C overnight using a LabConco FreeZone 2.5 Plus lyophilizer (LabConco, Kansas City, MO). Samples were stored at −20 °C.

4.1.10. TEV protease cleavage of affinity tags from fusion proteins
Samples containing purified fusion protein containing the tobacco etchvirus (TEV) protease site (ENLYFQ/GS) using in-house grown TEV protease (expression and purification protocol above) using 2 μL pro- tease solution for every 200 μg of protein in the elution buffer according to the corresponding construct. Samples were rotated at room temper- ature and 25 μL aliquots checked at 24 and 48 h by SDS-PAGE.

4.1.11. SDS-PAGE analysis of protein expression, purification and cleavage
Protein expression, purification and cleavage steps were analyzed using sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-
PAGE). Samples were prepared such that equal loading was observed in each gel. Gels were prepared according to standard protocol and used for protein separation with (4–12%) resolving and stacking (6%) gels (Invitrogen, Carlsbad, CA). Quantification was done using GelAnalyzer 19.1 [21].

4.1.12. NMR spectroscopy for activity assay via calcium titration of (GGS)$_2$fortilin

Lyophilized (GGS)$_2$fortilin (1.76 mg) was resuspended in 500 µL of a 20 mM tris-Cl buffer, pH 8.0, containing 200 mM NaCl and 10% D$_2$O. An aqueous 1 M CaCl$_2$ stock solution was used for each titration. Spectra were recorded at 298 K on a Bruker AVANCE III 600 MHz spectrometer equipped with a 5 mm TCI cryprobe with z-axis gradients. $^{1}H$, $^{15}N$ HSQC spectra were collected with the Bruker pulse program hsqcfp3gpphwq [29]. Spectra were processed with Topspin 3.2 (Bruker, Billerica, MA), NMRPipe [30] and plotted using NMRglue [31]. For each titrant, 1–2 µL stock CaCl$_2$ was added to a microcentrifuge test tube containing the NMR sample. The sample was then placed into a glass NMR tube, with care taken to avoid air bubbles. This was then placed in the NMR spectrometer and measurements taken. Chemical shift analysis was done using CcpNmr Analysis V3 [32,33]. Binding analysis was performed on the largest chemical shift perturbations (CSPs) using the combined chemical shift difference in both $^{15}$N and $^1H$ [34]. CSPs were then plotted against the concentration of Ca$^{2+}$ delivered and fit to a saturation binding model, given by:

$$
\Delta \delta = \frac{Kd + [L] + [P]}{2[P]} - \sqrt{\left( Kd + [L] + [P]\right)^2 - 4[L][P]} \Delta \delta_{max}
$$

where $\Delta \delta$ is the change in chemical shift upon addition of ligand, [P] is the concentration of fortilin, [L] is the concentration of ligand (or Ca$^{2+}$), and $\Delta \delta_{max}$ is the maximum chemical change upon saturation of fortilin with Ca$^{2+}$. Curve fitting and error analysis was performed using Wolfram Mathematica.

4.1.13. LC-MS based proteomics analysis

Proteomic analysis was performed using a method established previously with modifications [35]. Excised gel pieces were de-stained in 50 mM ammonium bicarbonate/50% acetonitrile, reduced in 10 mM cysteine carboxymethylated (NCS), alkylated in 55 mM iodoacetamide for 60 min at room temperature in the dark. Gel pieces were then digested with proteases grade trypsin (Thermo Fisher Scientific, Waltham, MA) overnight at 37 °C. Resulting peptide mixtures were chromatographically separated on a reverse-phase C$_{18}$ column (10 cm × 75 mm, 3 µm, 120 Å) and analyzed on a Velos Pro Dual-Pressure Linear Ion Trap mass spectrometer (Thermo Fisher Scientific, Waltham, MA) as described previously [35].

Peptide spectral matching and protein identification were achieved by database search using Sequest HT algorithms in Proteome Discoverer 2.2 (ThermoFisher Scientific, Waltham, MA). Raw spectral data were searched against a protein sequence database containing the anticipated sequences and UniProtKB/Swiss-Prot protein database for E. coli (acquired from www.uniprot.org on November 3, 2020). Main search parameters included: trypsin, maximum missed cleavage site of two, precursor mass tolerance of 1.5 Da, fragment mass tolerance of 0.8 Da, fixed modification of cysteine carbamidomethylation (+57.021 Da), and variable modification of methionine oxidation (+15.995 Da). Decoy database search was performed to calculate false discovery rate (FDR). Proteins containing one or more peptides with FDR <0.01 were considered positively identified and reported.

Data availability

All mass spectrometry data and results are available upon request. Please contact Dr. Shin Pu at shinpu@boisestate.edu and Dr. Maranda Cantrell at marandacantrell@u.boisestate.edu.

Author contributions


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Appendix A. Supplementary data

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References

