Fate of Pup inside the Mycobacterium Proteasome Studied by in-Cell NMR

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Abstract

The Mycobacterium tuberculosis proteasome is required for maximum virulence and to resist killing by the host immune system. The prokaryotic ubiquitin-like protein, Pup-GGE, targets proteins for proteasome-mediated degradation. We demonstrate that Pup-GQQ, a precursor of Pup-GGE, is not a substrate for proteasomal degradation. Using STINT-NMR, an in-cell NMR technique, we studied the interactions between Pup-GQQ, mycobacterial proteasomal ATPase, Mpa, and Mtb proteasome core particle (CP) inside a living cell at amino acid residue resolution. We showed that under in-cell conditions, in the absence of the proteasome CP, Pup-GQQ interacts with Mpa only weakly, primarily through its C-terminal region. When Mpa and non-stoichiometric amounts of proteasome CP are present, both the N-terminal and C-terminal regions of Pup-GQQ bind strongly to Mpa. This suggests a mechanism by which transient binding of Mpa to the proteasome CP controls the fate of Pup.

Introduction

Inside a cell, macromolecular complexes are assembled along specific pathways necessary to carry out biological functions in the presence of a crowded cytosol [1–3]. Often, during assembly, effector molecules such as ligands or substrates are also present. The presence of these molecules prior to or following the expression of components of the complex can play a regulatory role in the assembly of that complex. Furthermore, the binding of these effector molecules may alter the pathway through which proper, biologically active conformations are achieved. It is not clear a priori that the final conformation and commensurate activity of the complex will be different due to this temporal control [4].

One method to study macromolecular complexes inside a cell that affords temporal control over assembly is STINT-NMR [5–7]. STINT-NMR is used to elucidate STructural INTerations between proteins within their native environment by using in-cell NMR [8]. In STINT-NMR, protein over-expression is induced in labeling medium to produce a uniformly labeled [U-13C] target protein containing NMR-active nuclei; cells are then transferred to non-labeling medium to induce over-expression of the interactor protein. 15N- or 13C- edited heteronuclear single quantum coherence (1H–15N-HSQC or 1H–13C-HSQC) NMR experiments are performed to monitor the chemical shifts of target backbone amide or side chain 13C–1H nuclei as the concentration of interactor is increased. Monitoring the chemical shift changes of the labeled target delineates the intermolecular interaction surface between the target and the interactor(s) [5]. Most importantly, the order of sequential over-expression of target and interactor proteins can be reversed, allowing temporal control over the assembly of the complex.

In this work STINT-NMR was used to examine the interactions between the Ubiquitin-like protein, Pup, in the presence of the Mycobacterium proteasome ATPase, Mpa, and the active 1.2 megadalton proteasome complex, consisting of Mpa and the Mycobacterial proteasome core particle (CP) [9,10]. The importance of this macromolecular complex is underlined by the fact that Mycobacterium tuberculosis is particularly resistant to reactive nitrogen intermediates generated by host immune system, and this resistance is related to the proteasome [11] and mpa [12]. The Mtb proteasome CP consists of 14 copies each of two distinct but related polypeptides, α and β [13]. The overall architecture of the CP is conserved: α- and β-type subunits segregate into four juxtaposed rings of β-type subunits are flanked on top and bottom by a ring of α-type subunits to form the barrel-shaped complex. The main function of the α-rings is to form a gated channel that controls the passage of unfolded substrates into and cleaved particles out of the proteolytic chamber. Studies
have shown that *Mtb* Mpa forms 404 kDa hexameric rings similar to AAA+ ATPases found in the eukaryotes [10,12]. Structural analysis predicts that *Mtb* Mpa physically interacts with the α-rings of *Mtb* proteasome CP and plays a role in binding, unfolding and translocating substrates into the proteasome complex [10,15,16].

Proteins that are targeted for degradation in eukaryotes are generally tagged with the Ubiquitin, a small (76 aa) highly conserved regulatory protein [17]. By using *Mtb* Mpa as bait in a bacterial two-hybrid screen of an *Mtb* genomic library searching for potential binding partners of *Mtb* Mpa, the first prokaryotic Ubiquitin-like protein, Pup was discovered [18]. Pup is a 64 amino acid protein that modifies and targets mycobacterium proteins to the proteasome for degradation. Pup is similar in size to Ubiquitin but the two proteins have different sequences and lack structural homology. Pupylated proteins, which have been tagged with Pup, interact with *Mtb* Mpa [15,19].

The *Mtb* proteasome complex presents a tractable in-cell system for studying the interactions between Pup and the proteasomal ATPase, Mpa. Crystal structures of the *Mtb* proteasome CP [14,20], the Pup-Mpa coiled coil domain complex [21], as well as *in vitro* NMR solution studies of Pup-Mpa interactions [19,22,23] are available. *E. coli* is a relevant prokaryotic host that provides a proper milieu for studying the maturation of β subunit, which proceeds through autocatalytic removal of an N-terminal peptide [13].

Pup contains a di-glycine motif at the penultimate position of the C-terminus, followed by either glutamate (Pup-GGE) or pupylation of proteins and to prove that no additional factors are required to be conjugated to substrate lysines [25,26]. Here we present how Pup-GGQ interacts with the Mpa-proteasome complex inside a cell by recreating the final steps in the mycobacterium degradation pathway in *E. coli*. We used Pup-GGQ for our studies since it exists as a free molecule inside the cell, not attached to its target [18].

**Materials and Methods**

**Plasmid Construction**

**pTM-Pup.** Four separate oligonucleotides, two coding strands (5–1,5–2), corresponding to amino acids 1-31 and 32-64, respectively, and two complementary strands (3–1,3–2), were combined to form a complete Pup gene containing flanking 5' HindIII and 3' BamHI restriction sites. The oligonucleotide sequences used are (5–1): 5'-CCC AAG CTT ATG GGC CAA GAG CAG ACC AAG CGT GGC GGT GCC GGC GAT GAT GAC GAC ATC GCC GGC AGC ACC GCC GGC CAG CAG CTG CGC GAA AAG CTG ACC GAG GAG ACC GAC-3' (5–2): 5'-GAT CTG CTC GAC GAA ATC GAC GAC GTC CTC GAG GAG ACC GCC GAC GAC TTC GTC CGC GCA TAC GTC CAA AAG GGC GGA CAG TGA GGA TCC AA-3' (3–1): 5'-GCT CTC CTC GTG CAG CTT TTC GGC ACG CTC CGT GCC CGC CGG GTG GCC GCC GGC GGT GTC ATC ATC GCC GGC GCC ACC GCC ACG CTT GTG CTC CTG TTG GAC CAT AAG CTT GGG-3' and (3–2): 5'-TTG GAT CCT CAC TGT CCG CCC TTT TGG AGC TAT CGG CCG ACG AAG TCC TCG GGC TTC TCG TCC AGG ACG TCG ATG ATT TCG TCG AGC AGA TCG TC-3'. Duplexes were formed between (5–1,3–1) and between (5–2,3–2) prior to ligation. The product was restricted and ligated into pTM vector to yield pTM-Pup. The resulting plasmid, pTM-Pup, expresses Pup-GGQ from a T7 promoter/lac operator (P$_{T7}$/lacOp), which is induced by isopropyl-β-D-thiogalactoside (IPTG). This plasmid confers kanamycin resistance and contains a Bsal linker sites. The resulting plasmid, pASK-Pup, expresses Pup-GGQ from a tet promoter/operator (P$_{tet}$/tetOp), which is induced by tetracycline or anhydrotetracycline. This plasmid confers ampicillin resistance and contains an f1 origin and codes for the lacI gene, which encodes for Lac repressor.

**pASK-Pup-His7x.** DNA coding for full-length Pup-GGQ was PCR amplified from pTM-Pup using Taq polymerase and the oligonucleotides 5'-AAA AAA GGT TTC TAA TGG CCC AAG GCC ACG CCA AGC GTG G-3' and 5'-AAA AAA GGT TTC AGC GCT TCA CTG TCC GCC CTT TTG GAC G-3'. The gene was ligated into pASK3+ [IBA] using the Bsal linker sites. The resulting plasmid, pASK-Pup-His7x, expresses a N-terminal His-tagged Pup from P$_{T7}$/tetOp. This plasmid confers ampicillin resistance and contains an f1 origin and codes for the tet gene, which codes for Tet repressor.

**pRSF-Msm Mpa.** DNA coding for full-length Pup-GGQ was PCR amplified from pASK-Pup by using Taq polymerase and the oligonucleotides 5'-AAA AAA GGT TTC TAA TGG CCC AAG GCC ACG CCA AGC GTG G-3' and 5'-AAA AAA GGT TTC AGC GCT TCA CTG TCC GCC CTT TTG GAC G-3'. The gene was ligated into pASK3+ [IBA] using the Bsal linker sites. The resulting plasmid, pASK-PupHis7x, expresses a C-terminal His-tagged Pup from P$_{T7}$/tetOp. This plasmid confers ampicillin resistance and contains an f1 origin and codes for the tet gene, which codes for Tet repressor.

**pRSF-Msm Mpa.** DNA coding for full-length Pup-GGQ was PCR amplified from isolated genomic DNA extracted from *M. smegmatis* MC2 7500 [27] by using Phusion polymerase and oligonucleotides 5'-AAA AAA GGT TTC GTC AGT GCC GCT GCT TC-3' and 5'-AAA AAA GGT TTC TAA TGG CCC AAG GCC AGC GTG G-3'. The gene was ligated into pRSF-1b [Novagen] using the Kpnl and HindIII linker sites. The resulting plasmid, pRSF-Msm Mpa, expresses C-terminal His-tagged Msm Mpa from P$_{T7}$/lacOp, which is induced by IPTG. This plasmid confers kanamycin resistance, contains an RSF replication origin and codes for the lacI gene, which encodes for Lac repressor.

**pRSF-Mtb Mpa.** DNA coding for full-length Pup-GGQ was PCR amplified from isolated genomic DNA extracted from strain H37Rv by using Phusion polymerase and oligonucleotides 5'-AAA AAA GGT TTC GTC AGT GCC GCT GCT TC-3' and 5'-AAA AAA GGT TTC TAA TGG CCC AAG GCC AGC GTG G-3'. The gene was ligated into pRSF-1b [Novagen] using the Kpnl and HindIII linker sites. The resulting plasmid, pRSF-Mtb Mpa, expresses N-terminal His-tagged Mtb Mpa from P$_{T7}$/lacOp, which is induced by IPTG. This plasmid confers kanamycin resistance, contains an RSF replication origin and codes for the lacI gene, which encodes for Lac repressor.
Expression of PrcA and His-tagged PrcB, contain a PPR7 
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 expression plasmid, pRSF-Mtb Mpa, expresses N-terminal His-tagged Mtb Mpa from PPR7::lacOp, which is induced by IPTG. This plasmid confers kanamycin resistance, contains an RSF replication origin and codes for the lacI gene, which encodes for Lac repressor.

Plasmids encoding PrcAB and PrcΔAB. Plasmids expressing PrcA and His-tagged PrcB, containing a PPR7::lacOp, a P15A replication origin, confer chloramphenicol resistance and code for the lacI gene.

Sequential over-expression and labeling

E. coli strain BL21(DE3) codon+ [Novagen] was co-transformed with pASK-Pup and pRSF-Msm Mpa (Pup-GGQ/Msm Mpa interaction); or pASK-Pup, pRSF-Mpa and pACYCDuet-PrcAB (Pup-GGQ/Msm Mpa/wild type proteasome CP interaction); or pASK-Pup, pRSF-Mpa and pACYCDuet-PrcΔAB (Pup-GGQ/Msm Mpa/Opengate proteasome CP interaction) or pASK-PupHis7x, pRSF-Mpa and pACYCDuet-PrcAB (Pup-GGQ-His7x/Msm Mpa/wild type proteasome CP interaction); or pASK-PupHis7x, pRSF-Mpa and pACYCDuet-PrcΔAB (Pup-GGQ-His7x/Msm Mpa/Opengate proteasome CP interaction). Cells were grown overnight at 37 °C to an OD600 of ~1.6 in Luria-Bertani (LB) medium supplemented with 150 mg/L of carbenicillin for cultures containing pASK-Pup or pASK-PupHis7x, and 35 mg/L of kanamycin for cultures containing pRSF-Msm Mpa or pRSF-Mtb Mpa, and 33 mg/L of chloramphenicol for cultures containing pACYCDuet-PrcAB and pACYCDuet-PrcΔAB. Four protocols were employed, as described below.

Protocol 1: Expression of [U-15N] Pup-GGQ. Transformed cells from an overnight culture grown in LB containing the appropriate antibiotics were washed once with minimal medium (M9) salts and re-suspended to an OD600 of ~0.5 in M9 medium containing the appropriate antibiotics. The cells were incubated at 37 °C for 10–15 minutes and IPTG was added to induce over-expression of Msm Mpa; induction was allowed to proceed for 16 hours. 100 mL samples were collected, centrifuged, washed twice with 10 mM potassium phosphate buffer [pH 6.5], re-suspended with 1 mL of 10 mM potassium phosphate buffer [pH 6.5] containing 10% glycerol and stored at -80 °C for subsequent NMR analysis. 50 mL culture samples were collected pre- and post-induction for SDS-PAGE analysis.

Expression of Msm Mpa. Following over-expression of labeled Pup-GGQ, the culture was centrifuged and washed once with M9 salts before re-suspending a sufficient number of cells to yield an OD600 of ~0.5 in LB medium supplemented with the appropriate antibiotics. The culture was incubated at 37 °C for 10–15 minutes and IPTG was added to a final concentration of 1 mM to induce over-expression of Msm Mpa; induction was allowed to proceed for 16 hours. 100 mL samples were collected, centrifuged, washed twice with 10 mM potassium phosphate buffer [pH 6.5], re-suspended with 1 mL of 10 mM potassium phosphate buffer [pH 6.5] containing 10% glycerol and stored at -80 °C for subsequent NMR analysis. 50 mL culture samples were collected pre- and post-induction for SDS-PAGE analysis.

Protocol 2: Expression of Msm Mpa. Transformed cells from an overnight culture grown in LB containing the appropriate antibiotics were washed once with LB medium and re-suspended to an OD600 of ~0.5 in LB medium containing the appropriate antibiotics. The cells were incubated at 37 °C for 10–15 minutes and IPTG was added to induce over-expression of Msm Mpa; induction was allowed to proceed for 16 hours. Following the first induction, a 100 mL control sample of culture was collected, the cells were centrifuged, washed twice with 50 mL of 10 mM potassium phosphate buffer [pH 6.5], re-suspended with 1 mL of 10 mM potassium phosphate buffer [pH 6.5] containing 10% glycerol and stored at -80 °C for subsequent NMR analysis. Samples were collected pre- and post-induction for SDS-PAGE analysis.

Transformed cells from the same overnight culture were washed once with M9 salts and re-suspended to an OD600 of ~0.5 in M9 medium containing the appropriate antibiotics, 15N-ammonium chloride (0.7 g/L) as the sole nitrogen source and 0.2% glucose as the sole carbon source. The cells were incubated at 37 °C for 10–15 minutes and anhydrotetracycline was added to a final concentration of 0.2 µg/mL to induce over-expression of Pup-GGQ. Induction was allowed to proceed for 4 hours at which time a 100 mL sample of culture was collected and prepared for NMR analysis.

Expression of [U-15N]-Pup. Following Msm Mpa over-expression the culture was washed once with minimal medium (M9) salts and re-suspended to an OD600 of ~0.5 in M9 medium containing the appropriate antibiotics, 15N-ammonium chloride (0.7 g/L) as the sole nitrogen source and 0.2% glucose as the sole carbon source. The cells were incubated at 37 °C for 10–15 minutes and anhydrotetracycline was added to a final concentration of 0.2 µg/mL to induce over-expression of Pup-GGQ. Induction was allowed to proceed for up to 4 hours. Following the second induction, a 100 mL sample of culture was collected and prepared for NMR analysis. Samples were collected pre- and post-induction for SDS-PAGE analysis.

Protocol 3: Expression of [U-15N]-Pup-GGQ. Samples were over-expressed, labeled, collected and stored as described in protocol 1.

Expression of Msm Mpa and wild type or Opengate proteasome CP. Following Pup-GGQ over-expression and labeling, the culture was centrifuged and washed once with M9
Protocol 4 was also used to co-express Msm Mpa and the Opengate proteasome CP. Culture samples were taken pre- and post-induction for SDS-PAGE analysis.

**Protocol 4: Expression of Msm Mpa and wild type or Opengate proteasome CP.** Cells from the overnight culture were washed once LB medium and re-suspended to an OD of ~0.5 in LB medium containing the appropriate antibiotics. For all induced cultures we substituted carbenicillin (150 mg/L) for ampicillin. The cells were incubated at 37 °C for 10–15 minutes and Msm Mpa and wild type proteasome over-expression was induced by adding 1.0 M IPTG to a final concentration of 1.0 mM. Induction was allowed to proceed for 4 hours. A 100 mL sample of culture was collected, the cells were centrifuged, washed twice with 10 mM potassium phosphate buffer [pH 6.5], re-suspended with 1 mL 10 mM potassium phosphate buffer [pH 6.5] containing 10% glycerol and stored at -80 °C for subsequent NMR analysis. Protocol 3 was also used to co-express Msm Mpa and the Opengate proteasome CP. 50 mL Culture samples were taken pre- and post-induction for SDS-PAGE analysis.

**SDS-PAGE Analysis**

For protocols 1 and 3, gel analyses were performed by taking 100 mL samples at time points 0 h, 2 h, 4 h, and 6 h post-Pup-GGQ over-expression followed by 12 h and 24 h post-Msm Mpa or Msm Mpa/proteasome CP over-expression. For protocols 2 and 4, 100 mL samples were taken at 12 h and 24 h post-Msm Mpa or Msm Mpa/proteasome CP over-expression followed by 0 h, 2 h, 4 h, and 6 h post-Pup-GGQ over-expression. Samples were collected, centrifuged and sonicated. Lysates were analyzed on a 18% acrylamide gel and developed by either Commassie Blue staining or Western blotting using the SuperSignal West HisProbe Kit [Pierce].

**NMR spectroscopy**

Cells containing free \([U\text{-}^{15}\text{N}]\) Pup-GGQ, \([U\text{-}^{15}\text{N}]\) Pup-GGQ/ Mpa, or \([U\text{-}^{15}\text{N}]\) Pup-GGQ/Mpa/Mtb proteasome CP complexes were re-suspended in 0.5 mL of NMR buffer, 10 mM potassium phosphate, pH 6.5, 90%/10% H2O/D2O, and transferred to an NMR tube. All NMR experiments were performed at 293K using a Bruker Avance 500 MHz NMR spectrometer equipped with a cryoprobe. We used a Water gate version of the \([H\text{-}^{15}\text{N}]\)-edited HSQC [28]. Data were recorded with 32 transients as 512x64 complex points in proton and nitrogen dimensions, respectively, apodized with a squared cosine–bell window function and zero-filled to 1k(128) points prior to Fourier transformation. The corresponding sweep widths were 12 and 35 ppm in the \(\text{H}\) and \(\text{N}\) dimensions, respectively. Chemical shifts of \([U\text{-}^{15}\text{N}]\) Pup-GGQ inside the cell are slightly different from purified Pup-GGQ. We reassigned the backbone chemical shifts of Pup-GGQ using a clarified lysate of \([U\text{-}^{15}\text{C}]\text{-}[N\text{-}^{15}\text{N}]\)-Pup-GGQ and a standard suite of triple resonance experiments. To reassign the \([U\text{-}^{15}\text{N}]\) Pup-GGQ peaks that changed their positions due to complex formation, we assumed minimum chemical shift changes [29], calculated as \(\Delta_{\text{ext}} = (\delta_{2\text{u}} + \delta_{2\text{n}}/4)^{1/2}\), where \(\delta_{2\text{u}}\) represents the change in hydrogen and nitrogen chemical shifts. After each NMR experiment, the cells were pelleted and the \(\text{H}[^{15}\text{N}]-\text{HSQC}\) spectrum of the supernatant was collected. No NMR signal was observed above the noise level implying that no leakage or cell lysis was occurring during the experimental acquisition time. Cell viability after in-cell NMR experiments was tested by plating bacteria at 1:10,000, 1:100,000, and 1:1,000,000 dilutions on plates containing the appropriate antibiotics before and after in-cell NMR experiments. After counting the colonies, it was established that cell viability was 95 +/- 3%.

The changes in chemical shifts of amide nitrogens and covalently attached amide protons, \(\Delta\), were calculated by using \(\Delta = (\delta_{\text{u}}^{2} + (\delta_{\text{u}}/4)^{2})^{1/2}\), where \(\delta_{\text{u}}\) represents the change in hydrogen and nitrogen chemical shifts. After each NMR experiment, the cells were pelleted and the \(\text{H}[^{15}\text{N}]-\text{HSQC}\) peaks of side chain amide protons and nitrogens of \([U\text{-}^{15}\text{N}]\) Pup-GGQ/Mpa/Mtb proteasome CP complexes were re-suspended in 0.5 mL of NMR buffer, 10 mM potassium phosphate, pH 6.5, 90%/10% H2O/D2O, and transferred to an NMR tube. All NMR experiments were performed at 293K using a Bruker Avance 500 MHz NMR spectrometer equipped with a cryoprobe. We used a Water gate version of the \([H\text{-}^{15}\text{N}]\)-edited HSQC [28]. Data were recorded with 32 transients as 512x64 complex points in proton and nitrogen dimensions, respectively, apodized with a squared cosine–bell window function and zero-filled to 1k(128) points prior to Fourier transformation. The corresponding sweep widths were 12 and 35 ppm in the \(\text{H}\) and \(\text{N}\) dimensions, respectively. Chemical shifts of \([U\text{-}^{15}\text{N}]\) Pup-GGQ inside the cell are slightly different from purified Pup-GGQ. We reassigned the backbone chemical shifts of Pup-GGQ using a clarified lysate of \([U\text{-}^{15}\text{C}]\text{-}[N\text{-}^{15}\text{N}]\)-Pup-GGQ and a standard suite of triple resonance experiments. To reassign the \([U\text{-}^{15}\text{N}]\) Pup-GGQ peaks that changed their positions due to complex formation, we assumed minimum chemical shift changes [29], calculated as \(\Delta_{\text{ext}} = (\delta_{2\text{u}} + \delta_{2\text{n}}/4)^{1/2}\), where \(\delta_{2\text{u}}\) represents the change in hydrogen and nitrogen chemical shifts. After each NMR experiment, the cells were pelleted and the \(\text{H}[^{15}\text{N}]-\text{HSQC}\) spectrum of the supernatant was collected. No NMR signal was observed above the noise level implying that no leakage or cell lysis was occurring during the experimental acquisition time. Cell viability after in-cell NMR experiments was tested by plating bacteria at 1:10,000, 1:100,000, and 1:1,000,000 dilutions on plates containing the appropriate antibiotics before and after in-cell NMR experiments. After counting the colonies, it was established that cell viability was 95 +/- 3%.

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GGQ/ Msm Mpa/Mtb proteasome CP, or Pup-GGQ/ Msm Mpa/Mtb Opengate proteasome CP complex. Positive changes in relative intensities denote peak broadening due to binding interactions. Negative changes in intensities are due to overlapping peaks in the bound state. To resolve these overlaps, we assumed that two overlapped peaks are of equal intensity and, consequently, the intensity can be scaled down in relative intensities denote peak broadening due to binding overlaps, we assumed that two overlapped peaks are of equal intensity and, consequently, the intensity can be scaled down by two. Due to the qualitative nature of this procedure, all overlapped peaks were marked on the bar diagrams by crosses. To distinguish between residues directly and indirectly affected by complex formation, chemical shift changes above 0.1 ppm and differential peak broadening above 60% were considered to be significant.

Over-expression and purification of proteins for in vitro assays

E. coli strain BL21(DE3) codon+ [Novagen] was individually transformed with pASK-Pup His7x, pRSF-Msm Mpa or pRSF-Mtp Mpa. Cells expressing Pup-GGQ-His7x were grown in LB medium containing 150 µg/mL of carbenicillin at 37 °C to an OD600 of ~0.7 and induced with 1 µg/mL of anhydrotetracycline in DMF to a final concentration of 0.2 µg/mL for 6 h. Cells expressing Msm Mpa or Mtp Mpa were grown in LB broth containing 35 mg/L of kanamycin at 37 °C to an OD600 of ~0.7 and induced with 1.0 mM IPTG for 12 h. The induced cells were harvested by centrifugation at 4000g for 10 min at 4 °C. Cells were suspended in 30 mL of ice-cold lysis buffer, 50 mM NaH2PO4 [pH 8.0], 300 mM NaCl, 10 mM imidazole, and were lysed by sonication using a Branson sonicator at 300 W for 0.3 s intervals followed by a 1 s rest for a total of 5 min. The lysate was centrifuged at 20000g for 20 min at 4 °C. The supernatant was loaded onto a Ni-NTA resin-filled column (Qiagen). The column was washed with 20 mL of lysis buffer and 30 mL of wash buffer, 50 mM NaH2PO4 [pH 8.0], 300 mM NaCl, 20 mM imidazole. Fusion proteins were eluted with elution buffer, 50 mM NaH2PO4 [pH 8.0]. 300 mM NaCl, 250 mM imidazole. The eluted proteins were buffer exchanged into 50 mM Tris-HCl, pH 7.5, 0.1 M NaCl. eluted proteins were buffer exchanged into 100 mL of reaction buffer R.

In vitro degradation assay

In vitro degradation assay of Pup-GGQ by the Mpa/proteasome CP complex was performed as described by Striebel et al. (2010) (15). In summary, the reaction was carried out in buffer R containing 25 mM phosphocreatine (Sigma), 1 U/mL creatine phosphokinase (Sigma), 0.3 µM Msm Mpa, 0.4 µM Opengate proteasome CP, 3 µM Pup-GGQ and 5 mM ATP. Assays in which WT proteasome CP was used in place of the Opengate proteasome CP or where Mtb Mpa was used in place of Msm Mpa were carried out under identical conditions. Reactions were performed at 25 °C. Samples were collected at 0, 30, 60, 120, 240 min and quenched by adding SDS-sample buffer and analyzed by SDS–PAGE and/or Western blot using the SuperSignal West HisProbe Kit [Pierce].

Proteasome activity assay

The WT and Opengate proteasome CPs were purified as described and exchanged into buffer PA, 20 mM Heps [pH 7.5], 0.5 mM EDTA. Proteasome CPs (10 ng of each subunit) were incubated at 37 °C with 20 µM Suc-LLVY-AMC in a 96-well black plate (Corning) inside a fluorescence spectrophotometer with continuous stirring. The release of AMC was monitored at λex =440 nm using λem = 360 nm; experiments were performed in triplicate. Inhibition of proteasome CPs was accomplished by adding (0.8 nmol) of Bortezomib to each reaction to ensure that degradation occurred via the proteasome.

Results

Pup-GGQ is disordered inside E. coli

Pup is a small intrinsically unstructured protein with NMR relaxation properties favorable for in-cell NMR analysis [22,23,30,31]. To determine if intracellular Pup contains any regions of induced secondary structure due to macromolecular crowding, [U-15N] Pup-GGQ was over-expressed in E. coli and an in-cell 1H(15N)-HSQC NMR spectrum was collected (Figure 1A). The spectrum of Pup exhibited a signal to noise ratio of better than 5:1 and contained well-resolved backbone amide peaks. To determine that the signals were due solely to intracellular protein, after collecting the spectrum, the sample was centrifuged and the supernatant was examined. No NMR signal was observed above the noise level implying that no leakage or cell lysis was occurring during the experimental acquisition time (Figure S1A).

After obtaining the in-cell spectrum, cells containing [U-15N] Pup-GGQ were lysed and the spectrum of the lysate was collected (Figure 1B). The peaks from the in vitro spectrum are sharpened, relative to those of the in-cell spectrum, because of
the increased rate of tumbling associated with the decreased viscosity of the lysate. The chemical shift of all but three peaks, D16, V46, and D53, from both the in-cell and lysate spectra are found to be within 0.1 ppm of each other (Figures S1B and S1C). Changes in protein conformation typically result in significant/larger changes (>0.5 ppm) in the chemical shifts. Since only minor changes in chemical shifts and peak intensities were observed, we conclude that Pup remains an intrinsically disordered protein in vivo.

Both the N- and C-termini of Pup engage the proteasome ATPase, Mpa

Pup residues 21 through 51 exhibit a propensity to form a transient α-helical structure [22,23]: 13C chemical shifts of α- and β-carbons in this region of the protein are consistent with partial ordering of the structure [32], while the lack of dispersion in the amide region of the NMR spectrum suggests a disordered state. The crystal structure of the Pup-Mpa complex shows a helix conformation when Pup-GGE binds to the coiled coil domain of Mpa [21]. The binding induces a stable helical conformation encompassing amino acids 21-51 of Pup-, while the N- (aa 1-21) and C-termini (aa 52-64) remain unstructured in the Pup-Mpa complex [21]. To investigate the structural role of the interaction between the α-helix and Mpa under in-cell conditions, STINT-NMR [6] was used to characterize the interaction surface of Pup-GGQ when bound to Mycobacterium smegmatis Mpa, Msm Mpa.

Mpa was first overexpressed in non-labeling medium followed by over-expression of Pup-GGQ in [U-15N] labeling medium. In-cell 1H{15N}-HSQC NMR spectra were collected as the concentration of Pup-GGQ increased (Figure 2A). Because Mpa is in excess and the binding affinity of Pup for full length Mpa is sub-micromolar [21], as Pup-GGQ is over-expressed, a complex is expected to form. Depending on the chemical exchange rate between the free and bound states of the target, affected NMR peaks, corresponding to backbone amides, can shift, broaden their line shape or disappear completely. In the case of peak broadening, changes are monitored relative to the spectrum of the free target since only this species gives rise to visible peaks.

Indeed, peak broadening was observed when Pup-GGQ was over-expressed in the presence of Mpa. The most pronounced effects were seen for residues K7 in the N-terminal region, L39, E42, I43, D44, D45, L47, E49, and A51 of the α-helix region, and D53, F54, V55, A57, V59, K61, G63 and Q64 in the C-terminus of the protein (Figure 2A, S2A, and S2B). The largest changes in chemical shifts are primarily localized to the C-terminal half of the α-helix (Figure S2A). These results suggest that several regions of Pup-GGQ, including most of the C-terminal region (residues 52-64), part of the α-helix (residues 39-51), as well as a portion of the N-terminal region of Pup-

Figure 1. Pup is disordered inside E. coli. A. In-cell 1H{15N}-HSQC spectrum of [U-15N] Pup-GGQ. B. In vitro 1H{15N}-HSQC spectrum of [U-15N] Pup-GGQ. The comparatively high viscosity of the cytosol, relative to that of the cell lysate, results in a lower rate of tumbling and consequent peak broadening in the in-cell spectrum. Conversely, the lower viscosity of the cell lysate allows faster tumbling and generates sharper resonances in the in vitro spectrum. For quantitation of changes in chemical shifts and peak intensities see Figure S1. Asterisks denote NMR peaks originating from small metabolites.

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Figure 2. Pup-GGQ interacts with Msm Mpa through its N-terminus, C-terminus and α-helix. A. Overlay of E. coli $^1$H-$^{15}$N-HSQC spectra showing $[^{15}$N] Pup-GGQ after 4 h of over-expression (red), and after 4 h of over-expression following 16 h of Msm Mpa over-expression (black). B. Overlay of E. coli $^1$H-$^{15}$N-HSQC spectra showing $[^{15}$N] Pup-GGQ after 4 h of over-expression (red) followed by 16 h of Msm Mpa over-expression (black). C. Ribbon model of Pup docked with Msm Mpa. The sequence of Pup-GGQ is below the image. Residues that broadened the most upon complex formation with Msm Mpa are shown in red. The model is created by Swiss-PDB Viewer [43] based on the Pup-GGE-Mtb Mpa complex [21]. For quantitation of changes in chemical shifts and peak intensities see Figure S2.

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GGQ are involved in the interaction with Msm Mpa (Figure 2C). The engagement of N-terminal residues of Pup by Mpa is a novel finding that likely results from interactions that can only be observed in vivo or in-cell.

To deduce whether the presence of Pup-GGQ affects the assembly of Msm Mpa and therefore the formation of the Pup-GGQ/Mpa complex, Pup-GGQ was first over-expressed in [U-15N] labeling medium followed by over-expression of Mpa in non-labeling medium. In-cell 1H(15N)-HSQC NMR spectra were collected as the concentration of Mpa increased (Figure 2B). The differential broadening observed in the resulting spectra are characteristic of intermediate exchange and reflect an equilibrium between free and bound Pup-GGQ since Pup-GGQ is in excess and Mpa is not over-expressed to a sufficiently high level to form a large population of a complex. In general, the same regions of Pup-GGQ are most strongly perturbed as in the previous experiment, and while the magnitudes of the chemical shift changes are comparable (Figure S2C), the magnitudes of the intensity changes are reduced, consistent with sub-stoichiometric populations of Pup-GGQ and Mpa (Figure S2D). The order of expression appears to have no significant effect on the regions of Pup-GGQ affected by Mpa binding. We conclude that Msm Mpa assembles into the same conformational state regardless of the absence or presence of a high concentration of its physiological ligand.

Pup-GGQ is not degraded when bound to Mpa/proteasome complex in-cell

To examine the interaction of Pup-GGQ with Mpa in the context of the intact proteasome, STINT-NMR experiments were performed in which Pup-GGQ and the interactor complex, consisting of Mpa and the Mtb wild type (WT) proteasome CP were over-expressed. Simultaneous over-expression of Mpa and the Mtb WT proteasome CP allows the assembly of a functional mycobacterial proteasome complex in E. coli. However, high levels of over-expression of Mtb Mpa and the Mtb proteasome CP could not be achieved in the same cell. Because of 91% sequence identity between Msm Mpa and Mtb Mpa, and the ability to form the same hexameric structure, Msm Mpa is often used in functional studies of Mtb proteasome activity [25,33]. To ensure the highest levels of protein over-expression, we used Msm Mpa in all experiments.

To create the Pup-GGQ/proteasome complex, simultaneous over-expression of Msm Mpa and the α and β subunits of the Mtb proteasome CP were induced off compatible plasmids in non-labeling medium followed by over-expression of [U-15N]-Pup-GGQ in labeling medium. In-cell 1H(15N)-HSQC NMR spectra were collected as the concentration of Pup-GGQ increased (Figure 3A). Under these conditions, a complex between Pup-GGQ and the Msm Mpa/Mtb proteasome CP is formed. Signal intensities are uniformly broadened for all but T22, R56, V59, and Q60 of Pup-GGQ reflecting the interaction of Pup-GGQ with the much larger proteasome complex (Figure S3). As in the interaction of Pup-GGQ with Msm Mpa, the largest changes in chemical shifts are primarily localized to the C-terminal region of the α-helix (Figures 3A and S3A). Thus, Pup-GGQ appears to be more extensively engaged with Msm Mpa in the Msm Mpa/Mtb proteasome CP complex than with Msm Mpa alone, and involves residues from the N-terminus, α-helix and C-terminus of the protein (Figure 3C). We believe that the complete disappearance of the NMR spectrum is due to the N-terminal tail of Pup falling into the central cavity of the Mpa hexamer.

The order of over-expression was reversed, i.e. [U-15N] Pup-GGQ followed by Msm Mpa and the Mtb proteasome CP, and in-cell 1H(15N)-HSQC spectra were acquired. As observed for the Pup-GGQ/Msm Mpa interaction, the resulting spectra are characteristic of intermediate/slow exchange and the inability to form a large population of complex implying that the end point to the titration was not reached (Figure 3B). A smaller set of residues are broadened, L32, L40, and E42, in the α-helix (Figure 3B and S3). As seen with previous interactions, the largest chemical shifts affect primarily residues in the C-terminus of the α-helix and the C-terminal region of the protein (Figure S3C).

Pup is conjugated to proteins targeted for degradation through its C-terminus; an interaction confirmed by examining N-terminal deletion mutants of Pup [15,34]. Furthermore, the N-terminus is necessary for pupylated substrates to be directed into the Mpa central pore [15]. It was expected that in the presence of the Msm Mpa/Mtb proteasome CP complex, the N-terminus of Pup-GGQ would be engaged, thereby initiating degradation by the proteasome. As the concentration of the Msm Mpa/Mtb proteasome CP complex increased, no degradation of Pup-GGQ was observed, as monitored by Western blots and/or in-cell NMR. We rationalized that the Msm Mpa/Mtb proteasome CP complex may require additional factors not present in E. coli to remove inhibition for proteolysis of Pup-GGQ.

We substituted the OpenGate CP for the WT proteasome CP to circumvent this requirement [12,15]. The OpenGate CP lacks the eight N-terminal amino acids of the α-chains that regulate gated entry into the proteasome complex, and shows increased proteolytic activity for its substrates [14,15]. STINT-NMR experiments were carried out as described above to form the Msm Mpa/OpenGate proteasome complex and in-cell 1H(15N)-HSQC NMR spectra were collected. As the concentration of [U-15N]-Pup-GGQ increased in the presence of the Msm Mpa/OpenGate proteasome, a complex is formed (Figure 4). The signal intensities observed are uniformly broadened for the majority of Pup-GGQ residues, albeit to a lesser extent than seen with the Mtb proteasome CP, implying that most of the protein is still engaged with the proteasome complex (Figure 4A, 4C, and S4). A small number of Pup-GGQ residues exhibit significant changes in chemical shifts, D38, V46 and N50 of the α-helix (Figure S4A). The interaction between Pup-GGQ and the Msm Mpa/OpenGate proteasome complex is very similar to but less extensive than that of Pup-GGQ and the Msm Mpa/Mtb proteasome CP complex, affecting primarily the C-terminus of the α-helix, the C-terminal region and parts of the N-terminus of Pup-GGQ. These differences suggest that the Pup-GGQ-Mpa complex may contact the 8-amino acid terminal tail of the WT α-subunit when bound to the proteasome CP.

As the concentration of Msm Mpa/OpenGate proteasome complex was increased in the presence of [U-15N]-Pup the resulting in-cell 1H(15N)-HSQC NMR spectra show that the most
Figure 3. Pup-GGQ is extensively engaged with Msm Mpa in the wild type proteasome complex. A. Overlay of E. coli $^1$H($^{15}$N)-HSQC spectra showing [$U^{-15}$N] Pup-GGQ after 4 h of over-expression (red), and after 4 h of over-expression following 16 h of Msm Mpa/WT proteasome complex over-expression (black). B. Overlay of E. coli $^1$H($^{15}$N)-HSQC spectra showing [$U^{-15}$N] Pup-GGQ after 4 h of over-expression (red) followed by 16 h of Msm Mpa/WT proteasome complex over-expression (black). C. Ribbon model of Pup docked with Msm Mpa in the context of the Msm Mpa/WT proteasome complex. The sequence of Pup-GGQ is below the image. Residues that broadened the most upon complex formation with the proteasome complex are shown in red. The model is created by Swiss-PDB Viewer [43] based on the Pup-GGE-Mtb Mpa complex [21]. For quantitation of changes in chemical shifts and peak intensities see Figure S3.

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pronounced chemical shift changes and intensity changes are limited to the C-terminal half of the α-helix and the C-terminal region of Pup-GGQ (Figure 4B and S4). As before, the relatively small number of perturbed residues is due to the fact...
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that the titration endpoint was not reached and the resulting spectra represent an equilibrium between free and bound Pup-GGQ. Western blots indicate that the β subunit of the CP is processed and there is no degradation of Pup-GGQ resulting from its interaction with the proteasome complex (Figure 4D and 4E). The STINT-NMR results coupled with Western blots provide compelling evidence that the N-terminus of Pup is engaged by the Msm Mpa/proteasome complex, but is not degraded.

Pup-GGE has been shown to be a poor substrate for proteasomal degradation [15]. We used the in-cell E. coli system to determine whether or not Pup-GGE is degraded under these conditions. Pup-GGE cloned into the pASK-3+ expression vector failed to produce a visible band on Western blots. However, Pup-GGE cloned into the pCDF-1b expression vector, which utilizes the strong T7 phage promoter for protein over-expression, produced a weak band on Western blots (Figure S5). Over-expression of Pup-GGE was suppressed following simultaneous over-expression of Msm Mpa and the Mtb proteasome CP (Figure S5). The over-expression peaked at 12 h and decreased over time suggesting that Pup-GGE may be degraded by an active proteasome in E. coli, is subject to proteolysis by endogenous E. coli proteases, or may be a result of cell death. Proteasomal degradation of Pup-GGE [15] and pupylated substrates [15,34] was reconstituted in vitro. In both cases, degradation was extremely slow, requiring more than 6 hours to detect. Our results are consistent with the conclusion that Pup-GGE is a very poor substrate for proteasomal degradation both in-cell and in vitro.

Re-creation of the Pup proteasome pathway in vitro reveals that Pup-GGQ is not degraded

Proteins that are targeted for degradation in eukaryotes are tagged with Ubiquitin, a small (76 aa), highly conserved regulatory protein [17]. One of the key motifs of this pathway is the highly conserved mechanism of how Ubiquitin is recycled. Although Pup may be recycled [40], it has also been reported to be degraded when bound to target proteins and when free in the cytosol [15]. To determine whether Pup-GGQ is degraded by either the Msm Mpa/WT proteasome or Msm Mpa/Opengate proteasome complex, the degradation assay of Striebel et al. [15] was employed. The reaction was initiated by adding ATP to a final concentration of 5 mM, samples were collected at 30 minute intervals, quenched with loading buffer and analyzed by SDS-PAGE. No degradation of Pup-GGQ was observed after 4 h when using either the WT or Opengate CP (Figure 5A). This in vitro result is consistent with in-cell observations of the proteasome complex reconstituted in E. coli.

To determine if the absence of Pup-GGQ degradation is a result of using Msm Mpa, purified Mtb Mpa was substituted for Msm Mpa in the degradation reaction [15]. No degradation of Pup-GGQ was observed when using either the Opengate (Figure S6A) or WT CP in combination with Mtb Mpa (Figure S6B). We concluded that Pup-GGQ is not a substrate for the mycobacterial proteasome CP.

To verify that the purified Mtb proteasome CPs are active, a known substrate, N-Succinyl-Leu-Leu-Val-Tyr-7-aminomethylcoumarin, Suc-LLVY-7-AMC, was used in the activity assay; when cleaved the AMC moiety fluoresces [13]. Suc-LLVY-7-AMC was added to purified WT or Opengate CP, and, over time, an increase in fluorescence was observed, consistent with proteasome-induced degradation of the substrate (Figure 5B). To confirm that the increase in fluorescence is due to cleavage and not from contaminating proteolytic activity, a specific proteasome inhibitor, Bortezomib [35], which forms a tetrahedral adduct with the catalytic T10 in the PrCB subunit [14], was added to increasing concentrations of WT proteasome CP. Over time, in the presence of Bortezomib, fluorescence emission was reduced verifying that substrate degradation is due specifically to Mtb proteasome activity (Figure 5C).

Discussion

Assembly of macromolecular machinery in the presence of native ligands in the crowded cytosol presents a complicated system for study by amino-acid residue resolution techniques. We used in-cell STINT NMR to map the interactions of the prokaryotic ubiquitin-like protein, Pup, with the mycobacterial proteasome in E. coli. The intracellular medium provides a prokaryotic environment for structural study of Mtb proteasome function without the complications of additional factors that may specifically interact with this system [31]. Reconstructing the interactions between the mycobacterial Msm Mpa/Mtb proteasome CP complex and Pup-GGQ inside a cell at amino-acid residue resolution has allowed us to examine intracellular processes that are not accessible by in vitro investigations.

In vitro studies showed that Pup is a disordered protein possessing a transient helical structure in its C terminal region [22,23]. As in the case of α-synuclein, physiological conditions result in a seemingly disordered protein [30] may acquire stable secondary and even tertiary structure [36,37]. Only minor changes in the in-cell NMR spectrum of Pup-GGQ occur when compared to the cell lysate spectrum. This suggests that Pup-GGQ does not possess a stable secondary structure in the cytosol. Since Pup-GGQ acts as an anchor for the proteasome system, with the N-terminus assuming an extended structure [21,38], the disorder may be important for its function.

In-cell, Pup-GGQ interacts with the hexameric proteasomal ATPase, Msm Mpa, and its interaction surface does not change with the order of expression. This result was expected since intramolecular binding of Mpa subunits is very tight; the complex fails to dissociate on a chromatographic sizing column [10], consistent with an association constant greater than ~10^8 M^{-1} [39]. Pup-GGQ binds to Msm Mpa primarily via the C-terminal half of the helical region. In addition, a short region of the N-terminus, T6-R8, and C-terminus, E52-Q64, are affected by Msm Mpa binding. Titrating Msm Mpa into [U-15N] Pup-GGQ in vitro results in a gradual uniform broadening of the peaks from residues S21 to Q60 [23], excluding the C-terminal residues, K61, G62, G63 and Q64. In-cell, peak intensities do not decrease for all the residues in the Pup-GGQ helix, S21-A51. We suggest that this difference is due to the interactions of Pup-GGQ with components of cytosol that block part of the interaction surface between Pup and Mpa observed in vitro. The in-cell observations are also in general agreement with
specific contacts in the helical region, identified in the Pup-Mpa co-crystal [21], although in this instance, truncated Mpa was used. The use of a truncated Mpa may not accurately represent the Pup-Mpa interaction due to the possibility of altered conformations resulting from the truncation and from in vitro conditions that fail to duplicate the cellular environment in which this interaction normally occurs.

In cells expressing Pup-GGQ, Msm Mpa, and WT or Open gate proteasome CPs, the intracellular concentration of CP is significantly less than that of Msm Mpa and Pup-GGQ (Figure 4D and 4E). In this case, the Pup-GGQ/Msm Mpa complex will be the predominant species (Figures 3 and 4). Nevertheless, the in-cell NMR spectrum of the Pup-GGQ/Msm Mpa complex in cells expressing non-stoichiometric amounts of proteasome CPs is different from that of cells expressing only Pup-GGQ and Msm Mpa: the presence of proteasome CPs results in the complete broadening of peaks associated with the N-terminal tail of Pup-GGQ. The Pup-GGQ/Msm Mpa complex appears to be stabilized by non-stoichiometric amounts of proteasome CPs. Since the proteasome CP binds to Msm Mpa with low affinity [14], we postulate that transient binding of the proteasome CP to the Pup-GGQ/Msm Mpa complex results in the N-terminal tail of Pup-GGQ being occluded by the Msm Mpa central cavity, which leads to complete broadening of the Pup-GGQ spectrum (Figure 3C). Different in-cell spectra result when the order of over-expression of [U-15N] Pup-GGQ and the Msm Mpa/proteasome CP complex are reversed (Figures 3 and 4). The Msm Mpa/proteasome CP complex can take up to 12 hours to fully assemble in the cell following induction of over-expression. When [U-15N] Pup-GGQ is over-expressed first, the result is a mixed population of cells containing free Pup-GGQ, Pup-GGQ bound to Msm Mpa and Pup-GGQ bound to the Msm Mpa/proteasome complex. The corresponding in-cell NMR spectrum will represent an average of these three spectra. In this case, the spectrum is very similar to that of [U-15N] Pup-GGQ in complex with Msm Mpa, reflecting the interaction between Msm Mpa and the helical region of Pup-GGQ (Figure 2). When the Msm Mpa/proteasome CP complex is over-expressed first, the in-cell spectrum is completely broadened. Western analysis demonstrated that this is not due to degradation of Pup-GGQ (Figure 4).
Unlike Ubiquitin, which is recycled in the eukaryotic proteasome [17], both free and target-bound Pup-GGE are degraded in the mycobacterium proteasome [15], but may be recycled by depupyrase/deamidase Dop [34,40]. Pup-GGQ is a precursor molecule that is converted to Pup-GGE before being ligated to a substrate targeted for degradation [26,41]. Unlike Pup-GGE, which is a very poor substrate [15], Pup-GGQ is not a substrate for proteasomal degradation, consistent with its function as a precursor molecule. Indeed, free Pup-GGQ can be detected in Dop mutants of both *M. smegmatis* and *M. tuberculosis* strains [33,42], albeit in low concentration. STINT-NMR experiments present a dynamic picture of the fate of Pup-GGQ inside a cell containing Mpa and proteasome CPs (Figure 3 and 6): In the absence of the proteasome CP, the C- and N-terminal regions of Pup-GGQ interact with *Msm* Mpa only weakly; non-specific interactions are blocked by the cytosol. In the presence of non-stoichiometric amounts of proteasome CP, Pup-GGQ is completely bound to *Msm* Mpa with the C terminus binding to the mouth of the hexamer and the N-terminus falling into the central cavity (Figures 3 and 6). The charge difference between glutamine and glutamate in Pup may affect proteasomal degradation.

In conclusion, we assembled a functional 1.2 megadalton mycobacterium proteasome, consisting of Mpa and the proteasome CP, inside *E. coli* to study its interactions with the prokaryotic ubiquitin-like protein (DOP) in the presence of ATP. Pup-GGE is irreversibly bound to the Mpa-proteasome CP complex and possibly degraded.

![Figure 6. The precursor Pup-GGQ must be deamidated to Pup-GGE to activate proteasome-mediated degradation. Pup-GGQ is reversibly bound to the Mpa-proteasome CP complex, with the C terminus (blue) binding to the mouth of the hexamer and the N-terminus (yellow) falling into the central cavity, and is not degraded. Upon deamidation by Deamidase of Prokaryotic Ubiquitin like Protein (DOP) in the presence of ATP, Pup-GGE is irreversibly bound to the Mpa-proteasome CP complex and possibly degraded. doi: 10.1371/journal.pone.0074576.g006](https://www.plosone.org/fig6)

Unlike Ubiquitin, which is recycled in the eukaryotic proteasome [17], both free and target-bound Pup-GGE are degraded in the mycobacterium proteasome [15], but may be recycled by depupyrase/deamidase Dop [34,40]. Pup-GGQ is a precursor molecule that is converted to Pup-GGE before being ligated to a substrate targeted for degradation [26,41]. Unlike Pup-GGE, which is a very poor substrate [15], Pup-GGQ is not a substrate for proteasomal degradation, consistent with its function as a precursor molecule. Indeed, free Pup-GGQ can be detected in Dop mutants of both *M. smegmatis* and *M. tuberculosis* strains [33,42], albeit in low concentration. STINT-NMR experiments present a dynamic picture of the fate of Pup-GGQ inside a cell containing Mpa and proteasome CPs (Figure 6): In the absence of the proteasome CP, the C- and N-terminal regions of Pup-GGQ interact with *Msm* Mpa only weakly; non-specific interactions are blocked by the cytosol. In the presence of non-stoichiometric amounts of proteasome CP, Pup-GGQ is completely bound to *Msm* Mpa with the C terminus binding to the mouth of the hexamer and the N-terminus falling into the central cavity (Figures 3 and 6). The charge difference between glutamine and glutamate in Pup may affect proteasomal degradation.

In conclusion, we assembled a functional 1.2 megadalton mycobacterium proteasome, consisting of Mpa and the proteasome CP, inside *E. coli* to study its interactions with the prokaryotic ubiquitin-like protein, Pup-GGQ at amino acid residue resolution. By using STINT-NMR, we show that in-cell proteasomal degradation is dynamically regulated by transient interactions between Mpa and the proteasome CP. Differences between the binding of Pup and Mpa in vivo and *in vitro* underscore the importance of studying interactions under close to physiological conditions.
Supporting Information

Figure S1. Pup-GGQ is disordered inside the cell. A. Cell leakage test. An in-cell NMR sample of overexpressed [U-15N] Pup-GGQ was re-suspended in 500 μL of NMR buffer, 10 mM KPO4, pH 7.0, and incubated for 1 hour at RT. The cells were pelleted and the 1H(15N)-HSQC spectrum of the supernatant was collected. No NMR signal was observed above the noise level implying that no leakage or cell lysis was occurring during the experimental acquisition time. B. Differences in the chemical shifts of the 15N-HSQC spectra between in-cell and in vitro Pup-GGQ. The changes in chemical shifts of amide nitrogens and covalently attached amide protons, Δ, were calculated by using Δ = (δ_i - δ_i0)2/4, where δ_i represents the change in hydrogen and nitrogen chemical shifts. Based on the comparison of chemical shifts, we conclude that in-cell Pup-GGQ does not contain any structure induced by macromolecular crowding. C. Differences in peak intensities of the 15N-HSQC spectra between in-cell and in vitro Pup-GGQ. Since the 1H(15N)-HSQC peaks of side chain amide protons and nitrogens of [U-15N] Pup-GGQ glutamines do not change their positions in in-cell and in vitro Pup-GGQ, we used the intensities of these peaks (Iint) to scale the intensities of backbone amide protons and nitrogens. Changes in intensity were calculated by using ΔI = (Iref/Iint)_in_vitro - (Iref/Iint)_in_cell, where (Iref/Iint)in_vitro is the scaled intensity of an individual peak in the in-cell spectrum of Pup-GGQ and (Iref/Iint)in_vitro is the scaled intensity of individual peaks in the in vitro spectrum of Pup-GGQ. Positive changes in intensity reflect peak broadening due to the decrease in viscosity of the lysate relative to that of the cytosol. Changes in the chemical shifts or intensities above the continuous lines are considered to be significant. (PDF)

Figure S2. Pup-GGQ forms multiple contacts with Msm Mpa. In A. and B. Msm Mpa was over-expressed first followed by over-expression of [U-15N] Pup-GGQ. A. Differences in the chemical shifts of the 15N-HSQC spectra between free Pup-GGQ and the Pup-GGQ/ Msm Mpa complex. B. Relative changes in peak intensities of the 15N-HSQC spectra between free Pup-GGQ and the Pup-GGQ/ Msm Mpa complex. In C. and D. [U-15N] Pup-GGQ was over-expressed first followed by over-expression of Msm Mpa. C. Differences in the chemical shifts of the 15N-HSQC spectra between free Pup-GGQ and Pup-GGQ in complex with Msm Mpa. D. Relative changes in peak intensities of the 15N-HSQC spectra between free Pup-GGQ and Pup-GGQ in complex with Msm Mpa. The order of over-expression of Msm Mpa and Pup-GGQ does not change the Pup-GGQ-Msm Mpa interaction. The changes in chemical shifts of amide nitrogens and covalently attached amide protons Δ(ppm) and changes in peaks intensities ΔI were calculated as described in Materials and Methods. Overlapped peaks in the Pup-GGQ-Mpa complex are indicated by crosses. Changes in the chemical shifts or intensities above the continuous lines are considered to be significant. (PDF)

Figure S3. Pup-GGQ is extensively engaged by the Msm Mpa/WT proteasome CP complex in-cell. In A. and B. Msm Mpa and the WT proteasome CP were simultaneously over-expressed first followed by over-expression of [U-15N] Pup-GGQ. B. Differences in the chemical shifts of the 15N-HSQC spectra between free Pup-GGQ and the Pup-GGQ-Msm Mpa/WT proteasome CP complex. C. Relative changes in peak intensities of the 15N-HSQC spectra between free Pup-GGQ and the Pup-GGQ/ Msm Mpa/WT proteasome CP complex. In D. and E. [U-15N] Pup-GGQ was over-expressed first followed by simultaneous over-expression of Msm Mpa and the WT proteasome CP. In this case, the Msm Mpa/Opengate proteasome CP complex interacts with the C-terminus of Pup-GGQ. D. Differences in the chemical shifts of the 15N-HSQC spectra between free Pup-GGQ and Pup-GGQ in complex with the Msm Mpa/WT proteasome CP complex. The changes in chemical shifts of amide nitrogens and covalently attached amide protons Δ(ppm) and changes in peaks intensities ΔI were calculated as described in Materials and Methods. Overlapped peaks in the Pup-GGQ-Msm Mpa/WT proteasome CP complex are indicated by crosses. Filled dots above the bars indicate that the peaks are at the noise level. Changes in the chemical shifts or intensities above the continuous lines are considered to be significant. (PDF)

Figure S4. Pup-GGQ is extensively engaged by the Msm Mpa/Opengate proteasome CP complex in-cell. In A. and B. Msm Mpa and the Opengate proteasome CP were over-expressed first followed by over-expression of [U-15N] Pup-GGQ. A. Differences in the chemical shifts of the 15N-HSQC spectra between free Pup-GGQ and Pup-GGQ in complex with the Msm Mpa/Opengate proteasome CP complex. B. Relative changes in peak intensities of the 15N-HSQC spectra between free Pup-GGQ and the Pup-GGQ/Msm Mpa/Opengate proteasome CP complex. In C. and D. [U-15N] Pup-GGQ was over-expressed first followed by over-expression of Msm Mpa and the Opengate proteasome CP. In this case, the Msm Mpa/Opengate proteasome CP complex interacts with the C-terminus of Pup-GGQ. C. Differences in the chemical shifts of the 15N-HSQC spectra between free Pup-GGQ and Pup-GGQ in complex with the Msm Mpa/Opengate proteasome CP complex. D. Relative changes in peak intensities of the 15N-HSQC spectra between free Pup-GGQ and Pup-GGQ in complex with the Msm Mpa/Opengate proteasome CP complex. The changes in chemical shifts of amide nitrogens and covalently attached amide protons Δ(ppm) and changes in peaks intensities ΔI were calculated as described in Materials and Methods. Overlapped peaks in the Msm Mpa/Opengate proteasome CP complex are indicated by crosses. Filled dots above the bars indicate that the peaks are at the noise level. Changes in the chemical shifts or intensities above the continuous lines are considered to be significant. (PDF)
Figure S5. Over-expression of Msm Mpa and the Opengate proteasome CP suppresses Pup-GGE over-expression. (top panel) Western blot showing over-expression of Pup-GGE alone or following simultaneous over-expression of Msm Mpa and the Opengate proteasome CP. Samples were collected 6, 12 and 24 h post-Pup-GGE induction. Material loaded into each lane was normalized by the OD of the cell culture (lower panel). Intracellular Pup-GGE decreases over time in the presence of Msm Mpa and the Opengate proteasome CP. Pup-GGE bands were quantified using a Bio-Rad ChemiDoc XRS imager. The integrated density per mm² (Density INT/mm²) is shown. The experiments were repeated in triplicate. (TIF)

Figure S6. Pup-GGQ is not degraded in the presence of the Mtb Mpa/WT or Opengate proteasome complex in vitro.

References