Chaperone Proteins Select and Maintain [PIN⁺] Prion Conformations in Saccharomyces cerevisiae

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Background: Prion proteins adopt different conformations, known as variants, each with a distinct phenotype. Mechanisms by which prion variants are determined remain unclear. Here we use the Saccharomyces cerevisiae prion Rnq1p/[PIN⁺] as a model to investigate the effects of chaperone proteins upon prion variant determination. We show that deletion of specific chaperone genes alters [PIN⁺] variant phenotypes, including [PSI⁺] induction efficiency, Rnq1p aggregate morphology/size and variant dominance. Mating assays demonstrate that gene deletion-induced phenotypic changes are stably inherited in a non-Mendelian manner even after restoration of the deleted gene, confirming that they are due to a bona fide change in the [PIN⁺] variant. Together, our results demonstrate a role for chaperones in regulating the prion variant complement of a cell.

The mammalian prion protein, PrP, was originally identified as the causative agent of a group of neurodegenerative disorders collectively known as the transmissible spongiform encephalopathies (1). When the prion-determining domain (PrD) of PrP misfolds, PrP can switch from a non-infectious conformation (PrP°C) to a conformation prone to forming self-propagating, β-sheet-rich, amyloid polymers (PrPSc) (1, 2). These amyloids spread by catalyzing the transformation of PrPSc into PrPSc, eventually forming large aggregates (for review, see Ref. 3).

Although genetic polymorphisms of the PrP gene do affect its disease pathology (4, 5), distinct sets of symptoms arising from genetically identical PrPSc, called strains, have been described (6, 7). Protease treatment of different strains of PrPSc aggregates revealed that they differed by the size and composition of their amyloid core region, suggesting that it is specific conformations of the amyloid that give rise to PrPSc strains (8).

Prions are proteins that can adopt different infectious conformations known as “strains” or “variants,” each with a distinct, epigenetically inheritable phenotype. Mechanisms by which prion variants are determined remain unclear. Here we use the Saccharomyces cerevisiae prion Rnq1p/[PIN⁺] as a model to investigate the effects of chaperone proteins upon prion variant determination. We show that deletion of specific chaperone genes alters [PIN⁺] variant phenotypes, including [PSI⁺] induction efficiency, Rnq1p aggregate morphology/size and variant dominance. Mating assays demonstrate that gene deletion-induced phenotypic changes are stably inherited in a non-Mendelian manner even after restoration of the deleted gene, confirming that they are due to a bona fide change in the [PIN⁺] variant. Together, our results demonstrate a role for chaperones in regulating the prion variant complement of a cell.

Prions have also been characterized in non-mammalian model organisms, including bakers’ yeast, Saccharomyces cerevisiae, where they have been demonstrated to occur frequently in the wild (9–13). Yeast prions can also form different strains, called variants (14–16). Studies have shown that different prion conformations distinguish each prion variant (17–19).

The prion protein Rnq1p can adopt different variants, each with a distinct phenotype. Although the function of its non-prion conformation remains unknown, the prion conformation of Rnq1p, [PIN⁺], acts to help another yeast prion, Sup35p/[PSI⁺], adopt its own prion state (20, 21). Rnq1p has a prion-determining domain between amino acids 153 and 405, with a non-prion domain N terminus (11). Some [PIN⁺] variants include [PIN⁺]low, [PIN⁺]medium, and [PIN⁺]high, each named for its respective [PSI⁺] induction efficiency (16). Another [PIN⁺] variant-linked phenotype is observed when GFP-tagged Rnq1p is overproduced in vivo (22). [PIN⁺]high strains form multiple Rnq1-GFP foci per cell, whereas [PIN⁺]medium and [PIN⁺]low strains generally form only a single Rnq1-GFP focus. Another phenotypic difference between [PIN⁺] variants is the size and stability of their amyloid aggregates. [PIN⁺]high contains less stable aggregates that break down into smaller subparticles when heated, as opposed to [PIN⁺]low or [PIN⁺]medium aggregates, which remain stable when heated (16, 23, 24). When two [PIN⁺] variants are introduced into the same cell, either through mating or cytoduction, the diploid and all haploid progeny adopt the phenotype of the dominant variant. [PIN⁺]high is dominant over [PIN⁺]medium, which is in turn dominant over [PIN⁺]low (16). These multiple, distinct phenotypes make [PIN⁺] an ideal model system with which to study the etiology of prion variants.

Mutations in a prion protein can affect its amyloid structure and, through that, the type of variant it adopts (25, 26). Still, variants can arise from prion proteins with identical sequences (14–16, 27), suggesting that other cellular factors may influence the variant conformation that a given prion will adopt. Chaperone proteins are strong candidates for such variant regulating factors. Chaperones are known to affect the conformation of a wide array of client proteins (for review, see Ref. 28) and have been implicated in other aspects of prion biology, including the de novo formation, propagation, and curing of...
prions (for review, see Ref. 29–31). Additionally, changes to chaperone activity and levels have been shown to affect prion variants. Yeast strains expressing N- and C-terminal truncations of the primary stress response transcriptional regulator, Hsf1p, which were shown to increase Hsp104p and decrease Hsp90 levels, respectively, preferentially, preferentially formed specific [PSI+] variants upon de novo induction (32). Likewise, strains over- or underexpressing SSE1, which is important to Hsp70 activity, gave rise to specific [PSI+] variants when induced (33). To date, only one genetic mutation has been reported to lead to a change in a pre-existing variant. Sondheimer et al. (27) demonstrated that deletion of the Sis1p G/F domain altered Rnq1-GFP aggregation pattern in a manner stably propagated even after reintroduction of wild-type Sis1p.

Here, we report the findings of our investigations into the actions of chaperone proteins upon already established [PIN+] variants. We found that disruption of several chaperone genes gives rise to shifts in [PIN+] variant-linked phenotypes. Genetic analysis showed that the phenotypic shifts are inherited in a non-Mendelian manner, confirming that a bona fide change in [PIN+] variant was achieved. Our findings provide evidence that chaperones can affect established prion variants and highlight a potential role for chaperones in regulating prion-linked phenotypes through their modulation of prion variants.

EXPERIMENTAL PROCEDURES

Yeast Strains, Culture, and Genetic Manipulation—S. cerevisiae strains used in this study are listed in supplemental Table S1. All yeast strains were cultured at 30 °C with the exception of diploids undergoing sporulation, which were cultured at 25 °C. Media were as follows: YEPD (1% yeast extract, 2% peptone, 2% glucose); CSM (0.67% yeast nitrogen base without amino acids, 2% glucose, 1 × Complete supplement mixture (Bio 101, Vista, CA)); CSM auxotrophic marker growth medium (same as CSM but containing 1 × CSM minus the appropriate auxotrophic selection (ADE, −HIS, −LEU, −URA, −LYS-URA, −TRP-URA)); CSM auxotrophic marker induction medium (same as CSM but with 2% galactose in place of 2% glucose). For solid media, the same recipes were used with 2% agar added. Deletions were made using a HIS3 cassette as described (34) and confirmed by PCR. Isogenic MATα strains were made using YCpGAL::HO and mating-type switching (35). Mating, diploid sporulation, and dissection of haploids were performed as previously described with modified sporulation medium (0.3% potassium acetate, 0.02% raffinose) (36). Haploid progeny of previously described with modified sporulation medium (0.3% potassium acetate, 0.02% raffinose) (36). Haploid progeny of Hsf1p, which were shown to increase Hsp104p and decrease Hsp90 levels, respectively, preferentially, preferentially formed specific [PSI+] variants upon de novo induction (32). Likewise, strains over- or underexpressing SSE1, which is important to Hsp70 activity, gave rise to specific [PSI+] variants when induced (33). To date, only one genetic mutation has been reported to lead to a change in a pre-existing variant. Sondheimer et al. (27) demonstrated that deletion of the Sis1p G/F domain altered Rnq1-GFP aggregation pattern in a manner stably propagated even after reintroduction of wild-type Sis1p.

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Chaperone-mediated Determination of [PIN\textsuperscript{+}] Variants

**FIGURE 1. Effect of chaperone gene deletion on [PIN\textsuperscript{+}] induction efficiency in relation to [PIN\textsuperscript{+}] variants.** A, shown is a nonsense suppression assay. Chaperone gene deletions were made in [pin\textsuperscript{−}] [PIN\textsuperscript{+}\textsubscript{low/med/high}] strains that were induced to become [PIN\textsuperscript{+}] by overproduction of Sup35NM-GFP. Normal growth is shown on CSM medium, and [PIN\textsuperscript{+}]-dependent growth is shown on ADE medium. B, [PIN\textsuperscript{+}] induction efficiency is shown. The efficiency of [PIN\textsuperscript{+}] induction was determined by quantification of the nonsense suppression assay, expressing average ADE cfu as a percentage of average CSM cfu. Error bars represent S.E. Student’s t tests were done to compare the [PIN\textsuperscript{+}] induction efficiency of deletion strains with that of their parental wild-type strain. *p < 0.05; **p < 0.01; ***p < 0.001.

Yeast Two-hybrid Analysis—Yeast two-hybrid analysis was performed using the yeast strain HF7c and the plasmids pGAD424 (prey) and pGBT9 (bait) (Clontech, Mountain View, CA) following standard protocols (41). Genes encoding Rnq1p or chaperone proteins were ligated in-frame into pGAD424 and pGBT9 for expression. Interactions were scored by growth of cells on CSM-HIS medium after 4 days of incubation relative to growth on YEPD. At least three independent experiments were done for each tested pair of proteins.

**RESULTS**

Deletion of Chaperone Genes Affects [PIN\textsuperscript{+}] Induction Efficiency in a [PIN\textsuperscript{+}] Variant-dependent Manner—To investigate the role of chaperones in [PIN\textsuperscript{+}] variant determination, we made deletions of several chaperone-encoding genes in strains carrying characterized [PIN\textsuperscript{+}] variants and measured changes in [PIN\textsuperscript{+}] induction efficiency. Each strain was [pin\textsuperscript{−}] and carried a nonsense-suppression reporter gene (ADE1–14 UGA) that allowed growth on CSM – ADE medium only when [PIN\textsuperscript{+}] was induced (Fig. 1A). We measured the relative [PIN\textsuperscript{+}] induction efficiencies of deletion strains using a nonsense suppression assay (Fig. 1B) as described under “Experimental Procedures.” In brief, we overproduced Sup35NM-GFP using the galactose-driven expression vector pYES2.0-SUP35NM-GFP and quantified cfu on CSM – ADE medium plates relative to cfu on CSM medium plates. The strength of the induced [PIN\textsuperscript{+}] variants was not factored into our calculations of [PIN\textsuperscript{+}] induction efficiency; only the number of colonies and not their size or pigmentation was considered.

Deletion of HSC82, CPR6, CPR7, or TAH1 in the [PIN\textsuperscript{+}\textsubscript{low}] strain as well as HSC82, AHA1, CPR6, CPR7, or SSE1 in the [PIN\textsuperscript{+}\textsubscript{medium}] strain increased strain growth on CSM – ADE medium to levels significantly higher than those of wild-type [PIN\textsuperscript{+}\textsubscript{low}] and wild-type [PIN\textsuperscript{+}\textsubscript{medium}] strains and similar to those of the wild-type [PIN\textsuperscript{+}\textsubscript{high}] strain. Deletion of these same genes in the [PIN\textsuperscript{+}\textsubscript{high}] strain did not affect [PIN\textsuperscript{+}] induction efficiency. Conversely, deletion of SBA1 in the [PIN\textsuperscript{+}\textsubscript{medium}] and [PIN\textsuperscript{+}\textsubscript{high}] backgrounds decreased the efficiency of [PIN\textsuperscript{+}] induction to levels matching those of the wild-type [PIN\textsuperscript{+}\textsubscript{low}] strain but did not significantly decrease the efficiency of [PIN\textsuperscript{+}] induction when deleted in the [PIN\textsuperscript{+}\textsubscript{low}] strain. Nearly all strains gave rise to colonies with variable levels of pigment ranging from pink to almost white (Fig. 1A). The only exception was the [PIN\textsuperscript{+}\textsubscript{low}] strain deleted for TAH1 in which colonies were almost entirely white after induction, suggesting that the induced [PIN\textsuperscript{+}] colonies all carry a strong [PIN\textsuperscript{+}] variant.

ADE-competent colonies were confirmed to be prion-linked because they lost ADE competence after treatment with guanidine HCl (supplemental Fig. S1). Also, the extent of [PIN\textsuperscript{+}] induction in a strain being dependent on the levels of plasmid retention by that strain was excluded as a possibility, as both wild-type and deletion strains retained plasmid at levels between 90 and 95% whether or not the cells were ADE-competent (data not shown).

**Deletion of Chaperone Genes Affects the Aggregation of Rnq1p in a [PIN\textsuperscript{+}] Variant-dependent Manner**—Our results showing that the effects of gene deletion were [PIN\textsuperscript{+}] variant-specific

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Table A

|          | Low deletions CSM | ADE CSM | Med. deletions CSM | ADE CSM | High deletions CSM | ADE CSM |
|----------|------------------|--------|-------------------|--------|-------------------|--------|
| w1       |                  |        |                   |        |                   |        |
| hsp82Δ   |                  |        |                   |        |                   |        |
| hsc82Δ   |                  |        |                   |        |                   |        |
| ahα1Δ    |                  |        |                   |        |                   |        |
| hch1Δ    |                  |        |                   |        |                   |        |
| cpr6Δ    |                  |        |                   |        |                   |        |
| cpr7Δ    |                  |        |                   |        |                   |        |
| sba1Δ    |                  |        |                   |        |                   |        |
| sfl1Δ    |                  |        |                   |        |                   |        |
| tαh1Δ    |                  |        |                   |        |                   |        |
| sse1Δ    |                  |        |                   |        |                   |        |
| sse2Δ    |                  |        |                   |        |                   |        |
| sso2Δ    |                  |        |                   |        |                   |        |
| sso3Δ    |                  |        |                   |        |                   |        |
| yqf1Δ    |                  |        |                   |        |                   |        |

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**FIGURE 1.** Effect of chaperone gene deletion on [PIN\textsuperscript{+}] induction efficiency in relation to [PIN\textsuperscript{+}] variants. A, shown is a nonsense suppression assay. Chaperone gene deletions were made in [pin\textsuperscript{−}] [PIN\textsuperscript{+}\textsubscript{low/med/high}] strains that were induced to become [PIN\textsuperscript{+}] by overproduction of Sup35NM-GFP. Normal growth is shown on CSM medium, and [PIN\textsuperscript{+}]-dependent growth is shown on ADE medium. B, [PIN\textsuperscript{+}] induction efficiency is shown. The efficiency of [PIN\textsuperscript{+}] induction was determined by quantification of the nonsense suppression assay, expressing average ADE cfu as a percentage of average CSM cfu. Error bars represent S.E. Student’s t tests were done to compare the [PIN\textsuperscript{+}] induction efficiency of deletion strains with that of their parental wild-type strain. *p < 0.05; **p < 0.01; ***p < 0.001.
and resulted in shifts in $[PSI^+]$ induction efficiency from levels found in one wild-type variant strain, e.g. $[PIN^+]_{low}$, to a different wild-type variant strain, e.g. $[PIN^+]_{high}$, suggesting that the chaperone gene deletions may have given rise to a change in the $[PIN^+]$ variant carried by the cell. If the $[PIN^+]$ variant of a strain had been altered by deletion of a chaperone gene, then other variant-specific phenotypes should also have been affected. Accordingly, we characterized the effects of chaperone gene deletion upon the phenotype of Rnq1p aggregates.

We first quantified the localization of Rnq1-GFP in all the deletion strains affected in $[PSI^+]$ induction efficiency (Fig. 2A). When GFP-tagged Rnq1p was overproduced from the galactose-driven plasmid pYES2.0-Rnq1-GFP, $\approx 75\%$ of focus-containing wild-type $[PIN^+]_{low}$ and wild-type $[PIN^+]_{medium}$ cells contained a single Rnq1-GFP focus, whereas $\approx 25\%$ contained more than one focus. $\approx 50\%$ of wild-type $[PIN^+]_{high}$ focus-containing cells contained multiple foci, in agreement with previous findings (22). $[PIN^+]_{low}$ and $[PIN^+]_{medium}$ strains deleted for the genes HSC82, CPR6, CPR7, TAH1, or SSE1 that gave rise to increased $[PSI^+]$ induction also displayed multiple Rnq1-GFP foci at the frequency displayed by the wild-type $[PIN^+]_{high}$ strain. Deletion of AHA1 in the $[PIN^+]_{medium}$ strain but not in the $[PIN^+]_{low}$ strain led to increased $[PSI^+]$ induction (Fig. 1B). Likewise, deletion of AHA1 in the $[PIN^+]_{medium}$ strain but not in the $[PIN^+]_{low}$ strain led to an increase in multiple Rnq1-GFP foci to the frequency exhibited by the wild-type $[PIN^+]_{high}$ strain. In contrast, deletion of the SBA1 gene from the $[PIN^+]_{low}$ strain led to fewer cells with multiple Rnq1-GFP foci, consistent with the frequencies exhibited by wild-type $[PIN^+]_{low}$ and wild-type $[PIN^+]_{medium}$ strains. Interestingly, increases in the number of multiple Rnq1-GFP foci were also observed when TAH1 and SSE1 were deleted in $[PIN^+]_{medium}$ and $[PIN^+]_{low}$ strains, respectively, even though there was no observed increase in $[PSI^+]$ induction efficiency in these deletion strains.

**FIGURE 2. Effect of chaperone gene deletion on $[PIN^+]$ variant-dependent phenotypes.** A, Rnq1-GFP was overproduced in deletion strains of interest, and the frequency of multiple Rnq1-GFP foci in foci-containing cells was calculated as a percentage of total cells. Error bars represent S.E. Student’s $t$ tests were performed to compare the frequency of multiple Rnq1-GFP foci in cells containing foci with that of their parental wild-type strain. **, $p < 0.01$; ***, $p < 0.001$. B, the Rnq1p aggregate size in deletion strains of interest was compared by SDD-AGE. Samples were incubated at room temperature (top) or 55°C (bottom) before loading.
We next characterized the sizes of Rnq1p amyloid subparticles in the deletion strains. The sizes of Rnq1p amyloid subparticles have been shown to vary depending on whether the \([PIN^+]\) variant displays a single Rnq1p-GFP focus per cell, as with the wild-type \([PIN^+]_{\text{low}}\) and wild-type \([PIN^+]_{\text{medium}}\) strains, or multiple foci per cell as observed in the wild-type \([PIN^+]_{\text{high}}\) strain (23). Strains with a single focus display a relatively narrow range of large subparticles, whereas strains with multiple foci display subparticles ranging in size from monomer to as large as, or larger than, those found in strains with a single focus. Accordingly, we produced HA-tagged Rnq1p from the plasmid pGRE535-Rnq1 in the wild-type \([PIN^+]_{\text{low}}\) strain. The wild-type strain was then mated with the wild-type \([PIN^+]_{\text{low}}\) strain Eliminated small HA-Rnq1p aggregates, also proved to be sensitive to temperature, degradation in part to a monomer at 55 °C (Fig. 2B). It is important to note that the pattern of protein migration displayed by these partly degraded protein aggregates is easily distinguished from that produced by protein aggregates isolated from a \([pin^-]\) strain. \([pin^-]\) aggregates were monomeric and of low molecular weight, whereas partially degraded aggregates from \([PIN^+]\) strains still displayed species of high molecular weight. Our results suggest that deletion of specific chaperone genes leads to changes in the prion amyloid physical properties, thereby affecting its heat sensitivity.

Our results show a correlation between chaperone gene deletion strains displaying altered \([PIN^+]\) induction efficiency and those displaying changes in other \([PIN^+]\)-linked phenotypes. The \(\text{tah}1\Delta\) strains were exceptions. They were unaffected in the sizes of their HA-Rnq1p aggregates vis à vis the sizes of the aggregates in their corresponding wild-type \([PIN^+]\) variant background. Also of note is that although the \(\text{sse}1\Delta\) strain in the \([PIN^+]_{\text{low}}\) background showed multiple Rnq1p-GFP foci per cell, its aggregates were limited to a narrow range of large sizes.

Deletion-induced Phenotypes Are Inherited in a Non-Mendelian Manner—The \([PIN^+]\) variant-linked phenotypic changes we observed in strains deleted for chaperone genes could be due directly to the effects of gene deletion or could be due simply to changes in the cell chaperone complement and to the overproduction of tagged protein resulting from our methodology. For example, loss of a chaperone could impair the ability of the cell to deal with the overexpression of tagged Rnq1p, leading to formation of denatured, non-prion aggregates that are detectable in our assays as multiple Rnq1p-GFP foci and/or low molecular weight aggregates in SDD-AGE. To eliminate this trivial explanation and to confirm that it is indeed deletion of the chaperone genes that gives rise to the observed \([PIN^+]\) variant shifts, we reintroduced a wild-type copy of a deleted chaperone gene by crossing a deletion strain with an isogenic wild-type strain. This wild-type strain was \([pin^-]\) so as to avoid any convolution related to introducing dominant \([PIN^+]\) variants (16). If the effects of deletions were specific to the deletion of the gene, then restoring the chaperone gene would be expected to restore the original \([PIN^+]\) variant phenotype. If on the other hand a permanent change in the \([PIN^+]\) variant had occurred in the deletion strain, then the phenotypic changes should persist after reintroduction of the chaperone gene. Fig. 3 shows that, for the most part, \([PSI^+]\) induction efficiencies, the frequency of
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Rnq1-GFP foci, and the sizes of Rnq1p aggregates of diploid strains did not revert to the phenotypes of the original wild-type strains from which the deletion strains were derived. Instead, these diploid strains maintained the phenotypes of the deletion strains. The only exceptions were diploids arising from matings with \(\text{sse}1\Delta\) in the \([\text{PIN}^+]\)\text{low} background and with \(\text{tah}1\Delta\) in both the \([\text{PIN}^+]\)\text{low} and \([\text{PIN}^+]\)\text{medium} backgrounds. The increased frequency of Rnq1-GFP foci in the original haploid strains deleted for \(\text{sse}1\) or \(\text{tah}1\) was eliminated in the diploid strains, suggesting that these changes were \([\text{PIN}^+]\)\text{-independent}. It should be noted that under our experimental conditions we were unable to statistically distinguish \([\text{PIN}^+]\)\text{low} from \([\text{PIN}^+]\)\text{medium} based upon \([\text{PIN}^+]\) induction efficiencies (supplementary Fig. S2). As such, we could categorize our results only as being consistent with either \([\text{PIN}^+]\)\text{low/medium} or \([\text{PIN}^+]\)\text{high} levels.

There was the remote possibility that spurious mutations had been introduced into our deletion strains and had produced dominant effects mimicking an apparent variant switch. If this were the case, it would be expected that this trait would be co-inherited in a Mendelian manner as opposed to the non-Mendelian manner of a true change in variant. To eliminate the possibility of introduced spurious mutations, we sporulated and dissected our diploid strains, yielding wild-type and deletion-carrying haploid progeny. The presence of the chaperone gene deletion cassette (\(\text{HIS}3\)) was detected by growth on CSM-HIS medium and showed a 2:2 ratio in the progeny as expected (data not shown). We then measured \([\text{PSI}^+]\) induction efficiency of the dissected tetrads by nonsense suppression assay (Fig. 4). The specificity of this assay was demonstrated by mating wild-type strains carrying different \([\text{PIN}^+]\) variants with a strain deleted for \(\text{HSP104}\), which is required for both \([\text{PSI}^+]\) and \([\text{PIN}^+]\) maintenance (20, 42). The wild-type haploid progeny were inducible at levels similar to their parental strains, whereas the \(\text{hsp}104\Delta\) haploids were unable to be induced. When the chaperone gene deletion strains were mated with the wild-type \([\text{PIN}^-]\) strain, we found that the haploid progeny of 24 tetrads all maintained the same \([\text{PSI}^+]\) induction efficiency as the parental deletion strain regardless of the presence or absence of the chaperone gene of interest. These levels were consistent with either \([\text{PIN}^+]\)\text{low/medium} or \([\text{PIN}^+]\)\text{high} levels.

Haploid progeny were found to retain pYES2.0-Sup35NM-GFP at a rate consistent with the parental strains, with no strain losing or retaining the plasmid at a markedly higher level than any other strain. Also, putative \([\text{PIN}^+]\) colonies that arose after \([\text{PSI}^+]\) induction were shown to be curable by growth on medium containing guanidine HCl (supplemental Fig. S3).

The Rnq1p aggregates of wild-type and mutant haploid progeny derived from the mating of deletion strains that gave rise to \([\text{PIN}^+]\) variant-related phenotypic changes were characterized using SDD-AGE (Fig. 5). We found that the deletion-induced changes in aggregate size were stable even after tetrad dissection. Deletion strains that exhibited no change in aggregate size were found to be consistent with the wild-type strain in which they were made (data not shown). Taken altogether, our results show that deletion of chaperone genes gives rise to \textit{bona fide} changes in \([\text{PIN}^+]\) variant.

The Hierarchy of Inheritance of Induced Variants Is Consistent with Characterized \([\text{PIN}^+]\) Variants—Another characteristic of prion variants is that one is often dominant over another. In the case of \([\text{PIN}^+]\) variants, \([\text{PIN}^+]\)\text{high} is dominant over \([\text{PIN}^+]\)\text{medium}, which in turn is dominant over \([\text{PIN}^+]\)\text{low} (16). In most cases, deletion-induced variants display phenotypes comparable with \([\text{PIN}^+]\)\text{high}, with the exception of \(\text{sha}1\Delta\), which gives rise to a variant similar to \([\text{PIN}^+]\)\text{low/medium}. Still, it remained unclear if these variants were actually \([\text{PIN}^+]\)\text{high}, \([\text{PIN}^+]\)\text{low/medium}, or novel, previously uncharacterized types of variants. If these variants were not novel, then they would be expected to follow the same hierarchy of variant dominance in relation to other \([\text{PIN}^+]\) variants. To determine the hierarchy of the induced variants, we crossed deletion strains to isogenic wild-type strains carrying \([\text{PIN}^+]\)\text{low}, \([\text{PIN}^+]\)\text{medium}, or \([\text{PIN}^+]\)\text{high}. We then documented the \([\text{PIN}^+]\) variant-related phenotypes of the resulting diploid strains (Fig. 6). As before, we were unable to distinguish statistically \([\text{PIN}^+]\)\text{low} from \([\text{PIN}^+]\)\text{medium} based on their \([\text{PSI}^+]\) induction efficiencies (supplemental Fig. S4).

We found that diploids generated by mating putative \([\text{PIN}^+]\)\text{high} strains with any wild-type strain exhibited phenotypes consistent with wild-type \([\text{PIN}^+]\)\text{high}. Also, the presumed \([\text{PIN}^+]\)\text{low/medium} variant induced by the deletion of \(\text{SBA}1\) was shown to be eliminated by the introduction of the more dominant \([\text{PIN}^+]\)\text{high}. In the case of \(\text{tah}1\Delta\) generated in \([\text{PIN}^+]\)\text{low}, its \([\text{PIN}^+]\)\text{high}-like \([\text{PSI}^+]\) induction efficiency was dominant over \([\text{PIN}^+]\)\text{low/medium} wild-type phenotypes, whereas wild-type \([\text{PIN}^+]\)\text{high} was dominant over its other \([\text{PIN}^+]\)\text{-linked} phenotypes. Finally, the frequency of \([\text{PIN}^+]\)\text{high}-like Rnq1-GFP foci observed when \(\text{sse}1\) was deleted in a \([\text{PIN}^+]\)\text{low} strain reverted to a \([\text{PIN}^+]\)\text{low/medium}-like phenotype when mated with a wild-type \([\text{PIN}^+]\)\text{low} strain or a wild-type \([\text{PIN}^+]\)\text{medium} strain. All other traits followed wild-type patterns of inheritance.

\textit{Rnq1p Interacts Physically with Chaperone Proteins}—We performed a yeast two-hybrid analysis to investigate possible physical interactions between our chaperones of interest and Rnq1p (Fig. 7). With growth on CSM-HIS medium as a reporter

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**Table 1: Chaperone-mediated Determination of [PIN\(^+]\) Variants**

| Chaperone gene | Low deletions x WT [PIN\(^-\)] | Med. deletions x WT [PIN\(^-\)] | High deletions x WT [PIN\(^-\)] |
|----------------|---------------------------------|---------------------------------|---------------------------------|
| WT             | +                               | +                               | +                               |
| hsc82\Delta    | +                               | +                               | +                               |
| shah1\Delta    | +                               | +                               | +                               |
| cpr6\Delta     | +                               | +                               | +                               |
| cpr7\Delta     | +                               | +                               | +                               |
| sb\Delta       | +                               | +                               | +                               |
| tah1\Delta     | +                               | +                               | +                               |
| sse1\Delta     | +                               | +                               | +                               |

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\[ x_{\text{hsp104A}} [\text{pin}] \quad x_{\text{WT}} [\text{pin}] \]

\[
\begin{array}{cccccccc}
\text{WT low} & \text{WT high} & \text{WT high} & \text{WT low} & \text{WT high} & \text{WT high} & \text{WT low} & \text{WT high} \\
+ & + & + & + & + & + & + & + \\
\end{array}
\]

Chaperone-mediated Determination of [PIN\(^+\)] Variants

![Image](129x553 to 482x666)

**FIGURE 5. Non-Mendelian inheritance of chaperone gene deletion-induced changes in HA-Rnq1p aggregate size.** Diploids generated by mating chaperone deletion strains with a wild-type [PSI\(^-\)/pin\(^-\)] strain were sporulated and dissected. The resulting tetrads were analyzed for HA-Rnq1p aggregate size by SDD-AGE. Wild-type strains were mated with hsp104A strains to demonstrate genetic specificity for this assay. Wild-type (+) and deletion-carrying (−) haploid progeny are presented. The hsp104A samples and wild-type controls were run on a continuous gel and exposed for a longer period. Twenty-four tetrads were analyzed per diploid.

of interaction, we detected a previously documented interaction between Rnq1p and Tah1p (43). We also identified two novel interactions: Cpr7p and Sba1p with Rnq1p. No interaction was detected between the remaining chaperones and Rnq1p. When we characterized the [PIN\(^+\)] variant of our yeast two-hybrid strain (HF7c) by SDD-AGE and localization of Rnq1-GFP (supplemental Fig. S5), we found HF7c to be [pin\(^-\)], suggesting that the interactions we detected occur when Rnq1p is in its non-prion conformation.

**DISCUSSION**

Although recombinant *S. cerevisiae* prion proteins do not require cofactors to misfold into multiple infectious variants *in vitro*, they need the activity of different chaperone proteins to propagate stably *in vivo*. Most prominent among these chaperones are Hsp104p, Sis1p, and members of the Ssa subfamily (for review, see Refs. 29–31). Together these chaperones facilitate the fragmentation of growing amyloid fibrils, thereby exposing more fibril growing ends and generating more infectious prion seeds. It has been proposed that conformational differences between variants alter their susceptibility to fragmentation and their rate of fibril growth and that equilibrium between these processes determines the stability of prion propagation and the strength of prion phenotype (18, 44, 45). For example, compared with [PSI\(^+\)]\(^{\text{weak}}\), [PSI\(^+\)]\(^{\text{strong}}\) has a smaller amyloid core that fragments more easily, allowing for the generation of more growing ends and a higher rate of Sup35p incorporation into a greater number of prion seeds (18). These seeds in turn lead to increased nonsense suppression and more stable propagation of [PSI\(^+\)]. Chaperones have also been implicated in variant determination, with alteration of Hsf1p or Sse1p activity affecting the *de novo* induction of the [PSI\(^+\)] variant and truncation of Sis1p leading to stable changes in the established [PIN\(^+\)] variant (27, 32, 33).

Our study provides further evidence that chaperones are important for the selection of prion variants. Deletion of *AHA1* in a [PIN\(^+\)]\(^{\text{medium}}\) background or deletion of *HSC82, CPR6*, or *CPR7* in either a [PIN\(^+\)]\(^{\text{low}}\) or a [PIN\(^+\)]\(^{\text{medium}}\) background increases the efficiency of [PSI\(^+\)] induction to levels comparable to those seen for the wild-type [PIN\(^+\)]\(^{\text{high}}\) strain. Conversely, deletion of *SBA1* in the [PIN\(^+\)]\(^{\text{high}}\) background decreases the efficiency of [PSI\(^+\)] induction to wild-type [PIN\(^+\)]\(^{\text{low/medium}}\) background levels.

Could the effects of deleting these chaperone-encoding genes on [PSI\(^+\)] induction be independent of [PIN\(^+\)]? For example, the [PIN\(^+\)] variant-specific effects on [PSI\(^+\)] induction could be explained by different chaperone requirements for the *de novo* formation of [PSI\(^+\)] in the presence of different [PIN\(^+\)] variants. This scenario is not without precedent, as a [PSI\(^+\)] variant with exceptionally large aggregates was shown to require increased Hsp104p levels to propagate stably (46). However, such a scenario is unlikely in our case, because the changes we observed in [PSI\(^+\)] induction in our chaperone gene deletion strains were also accompanied by changes in the size and localization pattern of Rnq1p, consistent with a shift in [PIN\(^+\)] variant. Additionally, these phenotypic shifts persist after sporulation of diploids into wild-type and deletion-carrying haploid progeny upon reintroduction of the deleted chaperone gene. Together, our results demonstrate that the chaperone gene deletion-induced phenotypes we observed are due to stable shifts in the [PIN\(^+\)] variant.

Like other chaperone gene deletions, deletion of *TAH1* or *SSE1* in the [PIN\(^+\)]\(^{\text{low}}\) or [PIN\(^+\)]\(^{\text{medium}}\) backgrounds gave rise to changes in [PIN\(^+\)] variant-dependent phenotypes. However, in contrast to what was observed for the other chaperone gene deletion strains, not all of the tested phenotypes were altered in *TAH1* or *SSE1* gene deletion strains, and of those phenotypes that were altered, not all were maintained after the wild-type gene was restored. This suggests that at least some of the changes observed in the *tah1Δ* and *sse1Δ* strains are not the result of shifts in the [PIN\(^+\)] variant and that any putative var-
iant shift that may have occurred in these deletion strains does not correspond to a previously characterized variant (16, 22). In contrast, variants arising from deletion of HSC82, AHA1, CPR6, or CPR7 are consistent with [PIN⁺] high, whereas variants in sba1Δ strains match [PIN⁺] low or [PIN⁺] medium. These deletion-induced variants may eventually be shown to be distinct from classical [PIN⁺] variants but for the purpose of discussion are referred to here as [PIN⁺] high and [PIN⁺] low/medium, respectively.

[PIN⁺] high phenotypes were readily identified, but [PIN⁺] low and [PIN⁺] medium were difficult to distinguish from each other. Their [PSI⁺] induction efficiencies did not vary greatly, and the properties of their aggregates were indistinguishable. As such, it is interesting that deletion of AHA1, TAH1, or SSE1 had spe-

FIGURE 6. Dominance of chaperone gene deletion-induced variants. Diploids were generated by mating chaperone deletion stains with a wild-type [psi⁻] strain carrying the [PIN⁺] low or [PIN⁺] medium or [PIN⁺] high variant. These diploids were analyzed for [PSI⁺] induction efficiency, frequency of multiple Rnq1-GFP foci in foci-containing cells, and Rnq1p aggregate size. These phenotypes were then categorized as either [PIN⁺] low/medium or [PIN⁺] high. Quantification of [PSI⁺] induction efficiency and frequency of multiple Rnq1-GFP foci in foci-containing cells can be found in supplemental Fig. S4.

FIGURE 7. Yeast two-hybrid analysis. Cpr7p, Sba1p, and Tah1p were found to interact with Rnq1p based on growth of yeast on selective CSM-HIS medium. Total growth (top) and growth arising from protein interaction (bottom) are shown. The pattern presented is representative of four independent experiments. AD, activation domain; BD, binding domain.
specific effects depending upon whether they were deleted in \([PIN^+]_{low}\) or \([PIN^+]_{medium}\) backgrounds. This finding reinforces that \([PIN^+]_{low}\) and \([PIN^+]_{medium}\) are distinct variants and that these chaperone genes could serve as a genetic fingerprint for experimentally differentiating between the two variants in future studies.

By what mechanism might the variant changes we observed occur? Hsc82p, Aha1p, Cpr6p, Cpr7p, and Sba1p are known to act in the Hsp90 cycle (for review, see Ref. 47); Hsc82p is the constitutively expressed isomer of Hsp90; Aha1p, Cpr6p, and Cpr7p have been shown to increase Hsp90 ATP hydrolysis; Sba1p, the yeast homologue of p23, stabilizes the Hsp90-client complex, thereby decreasing Hsp90 ATPase activity (48–51). Our finding that deleting genes linked to increased Hsp90 ATPase activity led to a \([PIN^+]_{medium}\) phenotype whereas deleting \(SBA1\) led to a \([PIN^+]_{low/medium}\) phenotype suggests that Hsp90 activity is important for variant determination, with Rnq1p a possible Hsp90 client. This possibility is supported by the finding that a strain expressing a C-terminal truncation of Hsf1p and predisposed to form unstable \([PSI^+]\) also had markedly decreased Hsp90 levels (32).

We detected physical interactions of Rnq1p with Cpr7p, Sba1p, and Tah1p but not with the other chaperones. This suggests that some of these chaperones may act upon Rnq1p independently of Hsp90. Cpr7p, for example, has intrinsic isomerase activity (52, 53). Rnq1p contains three proline residues: one in the N-terminal region, a region shown to affect prion propagation (54), and two in a putative loop region between the \(\beta\)-sheets of the amyloid core (55). Isomerization of these prolines could potentially affect Rnq1p conformation.

Additionally, as numerous physical interactions between these chaperones and other chaperone complexes have been documented, changes in the levels of these chaperones may affect how Hsp40s, Hsp70s, and/or Hsp104p interact with Rnq1p, resulting in variant change. Cpr7p, for example, has been shown to interact with Hsp104p in a manner that is not essential for its thermotolerance activity (56, 57). Additionally, inhibition of Hsp90 ATPase activity has been shown to increase the levels of both Hsp104p and Hsp70 (58). The effect of \(SSE1\) deletion that we report here also implicates Hsp70s in variant change, as \(SSE1\) encodes an important Hsp70 nucleotide exchange factor (59). Fan et al. found that manipulation of Sse1p levels affected the \([PSI^+]\) variant (33), although our sse1\(\Delta\) strain did not display the same disposition toward an unstable weak \([PSI^+]\) variant. The difference between our and their results could be due to \([PIN^+]\) variant-specific effects, as Fan et al. (33) did not characterize the \([PIN^+]\) variant of their strain. It is also interesting that it has been reported that mutations in Sis1p give rise to an increase in Rnq1-GFP foci (27), similar to what we observed. It may be that our gene deletions indirectly impaired the activity of Sis1p and/or its association with Rnq1p.

How the loss of specific chaperones alters the levels and activities of other chaperones, in addition to their interaction with Rnq1p, will be an important avenue of future investigation to clarify the mechanisms underlying the variant changes that we observed. Chaperones could mediate \([PIN^+]\) variant changes by altering the conformation of Rnq1p. For example, chaperones could regulate the folding of monomeric Rnq1p in such a way as to predispose it to adopt a specific variant upon \(de novo\) formation or upon encountering prion seeds. More likely, because we observed changes to established variants, chaperones could work together to affect the conformation of existing amyloid polymers. For this to be effective, any growing ends of polymers need be remodeled. Alternatively, if multiple or unstable variants are present at the same time in the cell, as has been shown to occur for \([PSI^+]\) (60), changes in the chaperone environment could alter the rates of amyloid polymer fragmentation and/or prion seed generation in a variant-specific manner. In this way, one variant could be selected over others.

In closing, we have demonstrated that altering the chaperone complement of a cell can alter existing prion variants without the introduction of exogenous prion material. By modulating existing prion variants, the cell can maintain prion seeds within the cell while mitigating potential negative effects of stronger prion phenotypes. Also, when environmental pressures demand, the existing prion variant could be quickly altered to provide a more advantageous phenotype to the cell. In light of recent findings reporting the prevalence of prions in wild strains of yeast as well as the apparent survival advantages that they bestow (13), prion variant regulation represents a powerful mechanism for modulating a cell response and adaptability to changing environmental conditions.

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REFERENCES

1. Prusiner, S. (1982) Novel proteinaceous infectious particles cause scrapie. Science 216, 136–144
2. Pan, K. M., Baldwin, M., Nguyen, J., Gasset, M., Serban, A., Groth, D., Mehlhorn, I., Huang, Z., Fletterick, R. J., and Cohen, F. E. (1993) Conversion of α-helices into β-sheets features in the formation of the scrapie prion proteins. Proc. Natl. Acad. Sci. U.S.A. 90, 10962–10966
3. Colby, D. W., and Prusiner, S. B. (2011) Prions. Cold Spring Harb. Perspect. Biol. 3, a006833
4. Brown, K., and Mastroianni, J. A. (2010) The prion diseases. J. Geriatr. Psychiatry Neurol. 23, 277–298
5. Ohhashi, Y., Ito, K., Toyama, B. H., Weissman, J. S., and Tanaka, M. (2010) Differences in prion strain conformations result from non-native interactions in a nucleus. Nat. Chem. Biol. 6, 225–230
6. Fraser, H., and Dickinson, A. G. (1968) The sequential development of the brain lesion of scrapie in three strains of mice. J. Comp. Pathol. 78, 301–311
7. Bruce, M. E. (1993) Scrapie strain variation and mutation. Br. Med. Bull. 49, 822–838
8. Bessen, R. A., and Marsh, R. F. (1994) Distinct PrP properties suggest the molecular basis of strain variation in transmissible mink encephalopathy. J. Virol. 68, 7859–7868
9. Wickner, R. B. (1994) \([URE3]\) as an altered URE2 protein. Evidence for a prion analog in Saccharomyces cerevisiae. Science 264, 566–569
10. Paushkin, S. V., Kushnirov, V. V., Smirnov, V. N., and Ter-Avanesyan, M. D. (1996) Propagation of the yeast prion-like \([psi^+]\) determinant is...
mediated by oligomerization of the SLP35-encoded polypeptide chain release factor. *EMBO J.* **15**, 3127–3134

11. Sondheimer, N., and Lindquist, S. (2000) Rnq1. An epigenetic modifier of protein function in yeast. *Mol. Cell* **5**, 163–172

12. Alberti, S., Halfmann, R., King, O., Kapila, A., and Lindquist, S. (2009) A systematic survey identifies prions and illuminates sequence features of prionogenic proteins. *Cell* **137**, 146–158

13. Halfmann, R., Jarosz, D. F., Jones, S. K., Chang, A., Lancaster, A. K., and Lindquist, S. (2012) Prions are a common mechanism for phenotypic inheritance in wild yeasts. *Nature* **482**, 363–368

14. Derkatch, I. L., Chernoff, Y. O., Kushnirov, V. V., Inge-Vechtomov, S. G., and Liebman, S. W. (1996) Genesis and variability of [PSI+] prion factors in *Saccharomyces cerevisiae*. *Genetics* **144**, 1375–1386

15. Schlumberger, M., Prusiner, S. B., and Herskowitz, I. (2001) Induction of distinct [URE3] yeast prion strains. *Mol. Cell. Biol.* **21**, 7035–7046

16. Bradley, M. E., Edskes, H. K., Hong, J. Y., Wickner, R. B., and Liebman, S. W. (2002) Interactions among prions and prion “strains” in yeast. *Proc. Natl. Acad. Sci. U.S.A.* **99**, 16392–16399

17. Tanaka, M., Chien, P., Naber, N., Cooke, R., and Weissman, J. S. (2004) Conformational variations in an infectious protein determine prion strain differences. *Nature* **428**, 323–328

18. Tanaka, M., Collins, S. R., Toyama, B. H., and Weissman, J. S. (2006) The physical basis of how prion conformations determine strain phenotypes. *Nature* **442**, 585–589

19. Toyama, B. H., Kelly, M. J., Gross, J. D., and Weissman, J. S. (2007) The structural basis of yeast prion strain variants. *Nature* **449**, 233–237

20. Derkatch, I. L., Bradley, M. E., Zhou, P., Chernoff, Y. O., and Liebman, S. W. (1997) Genetic and environmental factors affecting the de novo appearance of the [PSI+] prion in *Saccharomyces cerevisiae*. *Genetics* **147**, 507–519

21. Derkatch, I. L., Bradley, M. E., Hong, J. Y., and Liebman, S. W. (2001) Prions affect the appearance of other prions. The story of [PIN+]. *Cell* **106**, 171–182

22. Bradley, M. E., and Liebman, S. W. (2003) Destabilizing interactions among [PSI+] and [PIN+] yeast prion variants. *Genetics* **165**, 1675–1685

23. Bagriantsev, S., and Liebman, S. W. (2004) Specificity of prion assembly in vivo. [PSI+] and [PIN+] form separate structures in yeast. *J. Biol. Chem.* **279**, 51042–51048

24. Liebman, S. W., Bagriantsev, S. N., and Derkatch, I. L. (2006) Biochemical and genetic methods for characterization of [PIN+] prions in yeast. *Methods* **39**, 23–34

25. Chien, P., and Weissman, J. S. (2001) Conformational diversity in a yeast prion dictates its seeding specificity. *Nature* **410**, 223–227

26. Kushnerov, V. V., Kryndushkin, D. S., Boguta, M., Smirnov, V. N., and Ter-Avanesyan, M. D. (2000) Chaperones that cure yeast artificial [PSI+] prion aggregates are formed by small Sup35 polymers fragmented by Hsp104. *J. Biol. Chem.* **278**, 94636–94643

27. Bagriantsev, S., and Liebman, S. W. (2004) Specificity of prion assembly in vivo. [PSI+] and [PIN+] form separate structures in yeast. *Mol. Cell* **17**, 565–570

28. Bagriantsev, S. N., and Derkatch, I. L. (2006) Biochemical and genetic methods for characterization of [PIN+] prions in yeast. *Methods* **39**, 23–34

29. Sondheimer, N., Lopez, N., Craig, E. A., and Lindquist, S. (2001) The role of Sis1 in the maintenance of the [RNY+] prion. *EMBO J.* **20**, 2435–2442

30. Morano, K. A., Grant, C. M., and Moye-Rowley, W. S. (2012) The response to heat shock and oxidative stress in *Saccharomyces cerevisiae*. *Genetics* **190**, 1157–1195

31. Maison, D. C., Kirkland, P. A., and Sharma, D. (2009) Influence of Hsp70s and their regulators on yeast prion propagation. *Prion* **3**, 65–73

32. Romanova, N. V., and Chernoff, Y. O. (2009) Hsp104 and prion propagation. *Protein Pept. Lett.* **16**, 598–605

33. Summers, D. W., Douglas, P. M., and Cyr, D. M. (2009) Prion propagation by Hsp40 molecular chaperones. *Prion* **3**, 59–64

34. Park, K.-W., Hahn, J.-S., Fan, Q., Thiele, D. J., and Li, L. (2006) *De novo* appearance and “strain” formation of yeast prion [PSI+] are regulated by the heat-shock transcription factor. *Genetics* **173**, 35–47

35. Fan, Q., Park, K.-W., Du, Z., Morano, K. A., and Li, L. (2007) The role of Sse1 in the *de novo* formation and variant determination of the [PSI+] prion. *Genetics* **177**, 1583–1593

36. Gietz, R. D., and Woods, R. A. (2002) Transformation of yeast by lithium acetate/single-stranded carrier DNA/polycethylene glycol method. *Methods Enzymol.* **350**, 87–96

37. Herskowitz, I., and Jensen, R. E. (1991) Putting the HO gene to work.
glutamine aggregates. *Mol. Cell. Biol.* 28, 3313–3323
55. Wickner, R. B., Dyda, F., and Tycko, R. (2008) Amyloid of Rnq1p, the basis of the [PIN⁺] prion, has a parallel in-register β-sheet structure. *Proc. Natl. Acad. Sci. U.S.A.* 105, 2403–2408
56. Abbas-Terki, T., Donzé, O., Briand, P. A., and Picard, D. (2001) Hsp104 interacts with Hsp90 cochaperones in respiring yeast. *Mol. Cell. Biol.* 21, 7569–7575
57. Mackay, R. G., Helsen, C. W., Tkach, J. M., and Glover, J. R. (2008) The C-terminal extension of *Saccharomyces cerevisiae* Hsp104 plays a role in oligomer assembly. *Biochemistry* 47, 1918–1927
58. Reidy, M., and Massion, D. C. (2010) Sti1 regulation of Hsp70 and Hsp90 is critical for curing of *Saccharomyces cerevisiae* [PSI⁺] prions by Hsp104. *Mol. Cell. Biol.* 30, 3542–3552
59. Shaner, L., Sousa, R., and Morano, K. A. (2006) Characterization of Hsp70 binding and nucleotide exchange by the yeast Hsp110 chaperone Sse1. *Biochemistry* 45, 15075–15084
60. Sharma, J., and Liebman, S. W. (2012) [PSI⁺] prion variant establishment in yeast. *Mol. Microbiol.* 4, 866–881