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Clioquinol promotes the degradation of metal-dependent amyloid-β (Aβ) oligomers to restore endocytosis and ameliorate Aβ toxicity

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Alzheimer’s disease (AD) is a common, progressive neurodegenerative disorder without effective disease-modifying therapies. The accumulation of amyloid-β peptide (Aβ) is associated with AD. However, identifying new compounds that antagonize the underlying cellular pathologies caused by Aβ has been hindered by a lack of cellular models amenable to high-throughput chemical screening. To address this gap, we use a robust and scalable yeast model of Aβ toxicity where the Aβ peptide transits through the secretory and endocytic compartments as it does in neurons. The pathogenic Aβ1–42 peptide forms more oligomers and is more toxic than Aβ1–40 and genome-wide genetic screens identified genes that are known risk factors for AD. Here, we report an unbiased screen of ~140,000 compounds for rescue of Aβ toxicity. Of ~30 hits, several were 8-hydroxyquinolines (8-OHQs). Clioquinol (CQ), an 8-OHQ previously reported to reduce Aβ burden, restore metal homeostasis, and improve cognition in mouse AD models, was also effective and rescued the toxicity of Aβ-secreting yeast. This model allowed us to take advantage of yeast genetics to identify compounds that may eventually help treat AD. We have taken advantage of the great conservation of the secretory and endocytic pathways between yeast and neurons to study Aβ in a simple, highly tractable model organism—budding yeast. By expressing Aβ as a fusion to an endoplasmic reticulum (ER) targeting signal, we have mimicked in yeast the multicompartmental distribution of Aβ (8). This approach bypasses the need to recapitulate the entire APP processing pathway and generates an Aβ peptide with exactly the same sequence as is found in the human brain. The ER targeting signal directs cotranslational transport of the primary translation product into the ER, where the signal sequence is removed by signal peptidase. The peptide then transits through the secretory pathway and is secreted from the cell. In yeast, the cell wall prevents secreted Aβ from diffusing away, allowing it to interact with the plasma membrane and undergo endocytosis. As in the human nervous system (2), the aggregation-prone 42-aa peptide is more prone to forming oligomeric species than the 40-aa peptide (9) and is more toxic (8).

This model allowed us to take advantage of yeast genetics to perform a completely unbiased screen of the yeast genome for suppressors or enhancers of Aβ toxicity. Of the ~6,000 genes we tested, we recovered only a handful of modifiers. There are phenotypic small-molecule screen | metal chelation

Significance

Identifying disease-modifying therapies for Alzheimer’s disease (AD) has been an insurmountable challenge. To provide a new discovery tool for high-throughput compound screening, we used a simple yeast model that makes toxic amounts of β-amyloid (Aβ), a peptide central to AD pathology. Previous genetic analysis established that Aβ compromises yeast biology in a manner relevant to human AD. We screened 140,000 compounds for reversal of toxicity and identified a class of protective metal-binding compounds related to clioquinol (CQ), a compound that alleviates Aβ toxicity in mouse AD models. Treating yeast with CQ promoted rapid degradation of Aβ oligomers, rescuing cellular processes perturbed by this insidious peptide and restoring viability. Our approach provides a method for identifying compounds that may eventually help treat AD.


The authors declare no conflict of interest.

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~25,000 genes in the human genome and less than 20 (10) have been shown to confer risk for AD. However, several of the yeast genes that alter Aβ toxicity are either direct homologs or interacting partners of human risk factors (8, 10). For example, YAP1P02, the yeast homolog of PICALM, one of the most highly validated risk factors for AD (10, 11); NIP52, is homologous to SYNJ1, which interacts with the risk factor BIN1 (12, 13); and SLAI is homologous to SH3KBP1, which interacts with the risk factor CD2AP (14–16). All of these proteins are involved in clathrin-mediated endocytosis in yeast and humans. In addition to ameliorating the toxicity of Aβ in yeast, these proteins reduced Aβ toxicity in both Caenorhabditis elegans and rat cortical neuronal models (8). The recovery of genes that promote clathrin-mediated endocytosis in unbiased genome-wide screens suggested that Aβ poisons this process (8). Indeed, the peptide that provided sufficient endocytosis of a transmembrane receptor (Ste3), an activity crucial for neuronal function. Importantly, the mechanism of action of these risk factors had not previously been linked to Aβ. Thus, the yeast model has already provided key insights on the nature of Aβ's cellular toxicity in the human brain.

In this study, we used our yeast Aβ model to identify small molecules that ameliorate toxicity. In an extremely stringent and unbiased screen of 140,000 compounds, we identified a small number of cytoprotective compounds, including 8-hydroxyquinolines (8-OHQu). Members of this family bind metals and are among the few compounds that have been shown to alleviate Aβ toxicity in mouse models of AD (17, 18), and to show early potential as an AD therapeutic (19). Here, we investigate the mechanism of action for the most efficacious member of this family, clioquinol (CQ).

**Results**

**Screen for Compounds That Rescue Aβ Toxicity in Yeast.** To screen for compounds that ameliorate Aβ toxicity in yeast, we used a strain with a compromised drug efflux system that expresses Aβ under the control of a galactose-inducible promoter (8). This strain is a model of Aβ toxicity that provided sufficient dynamic range and high signal-to-background signal to confidently identify rescuing compounds. Approximately 140,000 compounds were tested in duplicate in 384-well plates using increased growth (optical density) as an end point. The high reproducibility enabled robust detection of “hits” (Fig. L4). Only ~30 compounds passed our rigorous scoring criteria (Z score > 3), establishing the high stringency of the screen. Strikingly, one-half of all recovered hits were 8-OHQu (Fig. 1B and Fig. S1).

Notably, the effectiveness of this class of compounds against Aβ toxicity has been established in neurons and mouse models. Two closely related 8-OHQu—CQ and PBT2—decrease Aβ accumulation and alleviate cognitive and behavioral symptoms in mouse models of AD (17, 18). 8-OHQu are moderate-strength metal chelators that can extract metals weakly bound to proteins and both redistribute metals within the cell and act as ionophores (17, 18). PBT2 is one of only a few compounds effective in phase II clinical trials with human AD patients (19). CQ rescued our yeast Aβ model in a highly dose-dependent manner (Fig. 1 C and D), but at higher concentrations it inhibited the growth of both Aβ-expressing cells and control strains (Fig. 1C). The increase in optical density of Aβ-expressing cells with CQ was a direct consequence of increased viable colony-forming units and not an artifact of increased cell size (Fig. S2).

Due to our previous findings that different 8-OHQu have distinct activities in different models of proteotoxicity (20), we compared the efficacy of CQ in all of our yeast proteotoxicity models. The Aβ, α-synuclein (i.e., Parkinson disease), TDP-43 (i.e., frontotemporal dementia and ALS), and htt2Q (i.e., Huntington disease) yeast models have comparable levels of toxicity, but in each case the nature of the cellular toxicity is distinct as are the genetic hits obtained from unbiased genome-wide modifier screens (8, 21–23). CQ had significant, yet modest efficacy against α-syn and htt2Q and no effect on TDP-43 (Fig. 1E). However, it rescued the Aβ model more potently than it did any other model, indicating a considerable degree of selectivity.

Although CQ showed some protection in other models, we previously showed that CQ does not rescue the toxicity caused by diverse yeast genes when overexpressed from the same galactose-regulated promoter (20). In addition, CQ was unable to rescue growth defects caused by five different gene deletion mutations and seven toxic compounds with diverse modes of action (Table S1). Given the selectivity of CQ, we focused on investigating its activity in the yeast Aβ toxicity model.

**CQ Restores Endosomal Trafficking.** Our recent unbiased screen for suppressors of Aβ toxicity uncovered several proteins involved in clathrin-mediated endocytosis whose homologs are AD risk factors in humans (8). We directly tested the effect of Aβ on clathrin-mediated endocytosis using Ste3, a G-protein–coupled receptor for the mating pheromone. Indeed, steady-state Ste3 endocytosis was severely perturbed by Aβ and rescued by the genetic suppressors involved in this process (8). Here, we extended the analysis with another widely used reporter of endocytic function, a putative 13-pass transmembrane transporter for methionine, Mup1. This reporter provides a facile means of investigating stimulus-dependent trafficking: the endocytosis of Mup1 is rapidly induced by the addition of methionine.

As expected, Mup1-GFP localized primarily to the plasma membrane of WT cells in the absence of methionine (Fig. 2, Upper Left). After 30 min of methionine treatment, Mup1-GFP was endocytosed and exclusively detected in endosomes and the vacuole (Fig. 2, Upper Right). In the absence of methionine, Aβ expression perturbed steady-state trafficking of Mup1, detected by its localization to both the plasma membrane and to intracellular foci (Fig. 2, Lower Left). The addition of methionine promoted...
Mup1-GFP internalization, but delivery to the vacuole was greatly impeded (Fig. 2, Lower Center). CQ completely restored endosomal transport of Mup1-GFP to the vacuole in the strain expressing Aβ (Fig. 2, Lower Right). Thus, Aβ causes defects in endosomal transport that are manifested downstream of initial transporter internalization and these defects are rescued with CQ treatment.

CQ Rescues a C. elegans Model of Aβ Toxicity. To establish that the ability of CQ to rescue the toxicity of Aβ in the secretory compartment was conserved in neurons, we tested a nematode model of Aβ toxicity. In this model, Aβ is expressed in the secretory pathway of glutamatergic neurons (a neuronal subtype particularly relevant to AD) using the eat-4 promoter and a C. elegans ER signal sequence (8). An advantage of C. elegans is that the transparency of the cuticle allows surviving neurons to be visualized in living animals expressing GFP in the same neurons. Aβ expression produced an age-dependent loss of glutamatergic neurons. Importantly, the endocytosis-related genes we originally identified in our yeast screen, which are also known AD risk factors in humans, suppress Aβ toxicity in this nematode model (8). Expressing Aβ in glutamatergic neurons enables a quantitative measure of neurodegeneration and differs from previous approaches in the nematode where Aβ expressed within the body wall muscle cells caused a motor phenotype (24, 25).

As previously described, the percentage of worms with the WT number of neurons decreased from 3 to 7 d post larval hatching [Fig. 3A] (8)]. Because the cuticle of the worm is relatively impenetrable, we used an established protocol to administer CQ acutely at the earliest larval stage (26). After 24 h, the worms were transferred to normal growth media and allowed to develop and age in the absence of CQ. Even with this single early dosing, CQ increased the percentage of worms with the WT number of neurons. Rescue was apparent at both 3 and 7 d post CQ treatment (Fig. 3A and B). Thus, as in yeast, the toxicity of secreted forms of Aβ is rescued by CQ in glutamatergic neurons. This encouraged us to take advantage of the yeast model to further investigate CQ’s mechanism of action.

CQ Rescue of Aβ Toxicity Requires Metal Binding. Because 8-OHQs are known metal binding compounds, we first asked whether this functionality was required for the rescue of Aβ toxicity. We tested CQ analogs lacking either the hydroxyl group or the aromatic nitrogen. Together, these groups coordinate the bidentate binding of transition metals ions. Because direct analogs of CQ lacking these groups were not available, we tested analogs of chloroxine (CQ₂Cl) (Fig. 4A). This 8-OHQ is identical to CQ except that it has a chlorine atom in place of the iodine atom. It was as effective at alleviating Aβ toxicity in yeast as was CQ (Fig. 4B, “Max”). We then tested CQ₂Cl analogs lacking either the hydroxyl group (CQ₂Cl₂OH) or the nitrogen atom (CQ₂Cl₂N). Both were inactive against Aβ, indicating CQ efficacy requires bidentate metal binding (Fig. 4B).

Next, we tested the ability of exogenous metals to alter CQ activity. We used copper, iron, and zinc, all of which are known to bind CQ (27). On their own, none of these metals affected Aβ toxicity (Fig. 4C). When coadministered at a concentration that was equimolar with CQ, each metal had completely different effects (Fig. 4C). Copper abolished rescue of Aβ by low concentrations of CQ and also abolished the growth inhibition caused by higher CQ concentrations. Iron shifted the dose–response, increasing the required dose for rescue and reducing the growth inhibitory effects of CQ. Zinc had no effect on CQ efficacy or growth inhibition at higher concentrations. These data suggest that CQ binds copper, and to a lesser extent iron, to alleviate Aβ toxicity.

Effects of Metals and CQ on in Vitro Aβ Assembly. Metal ions, including copper, iron, and zinc, bind to Aβ both in vitro and within AD plaques from the brains of AD patients (28, 29). In in vitro assembly reactions with Aβ, the addition of copper favors the formation of nonfibrillar, amorphous aggregates (29) and oligomers (30, 31) that are more toxic to cells than amyloid fibers of Aβ (18, 30–32). Metal binding compounds, such as CQ and PBT2, can remove metals from Aβ in vitro (17, 18, 33, 34). These compounds also increase the amount of soluble Aβ in the brains of AD mouse models in vivo and stimulate the extraction of Aβ from plaques in brain samples from AD patients (17). Because the predominant form of Aβ 1–42 in yeast is small oligomers (8) and copper appears central to rescue of Aβ toxicity (Fig. 4C), we wondered whether exogenous copper would similarly rescue Aβ toxicity in C. elegans.
The decrease in oligomeric Aβ could be due to either its reduced accumulation or accelerated degradation. To distinguish between these two possibilities, we used two methods to assess degradation rates. First, we treated cells with the protein synthesis inhibitor, cycloheximide (CHX), and monitored Aβ levels by immunoblotting. After expressing Aβ for 24 h with either DMSO or CQ, cells were treated with CHX for 15 or 45 min. In CQ-treated cells, Aβ was rapidly degraded within 15 min of inhibiting translation (Fig. 6D). This increased degradation was not observed with a control protein (Pgk1). In contrast, the Aβ peptide that accumulated in cells treated with DMSO did not significantly degrade during this same time frame (Fig. 6D).

Inhibiting translation in this global manner might alter the cellular degradation pathway and confound interpretations. We therefore used \[^{35}S\]methionine pulse labeling and immunoprecipitation. Cells expressing Aβ were transiently labeled with \[^{35}S\]methionine to allow incorporation of radiolabeled methionine into newly synthesized proteins. After chasing cultures with nonlabeled methionine, we monitored the remaining Aβ by immunoprecipitating the peptide from cell lysates. In agreement with the CHX experiment, CQ specifically increased degradation of Aβ (Fig. 6E and Fig. S3). Together, the two experiments indicate that the stability of Aβ oligomers formed within the secretory pathway is dependent on metals and that, by affecting these oligomers, CQ can directly antagonize the cellular pathologies caused by Aβ.

**Discussion**

We have used an unbiased phenotypic small-molecule screen of ~140,000 compounds to identify a clinically relevant class of compounds that prevent the toxic effects of Aβ expressed within the secretory pathway. Although multiple metal-related Aβ pathologies and modes of action for CQ—and PBT2—have been described and postulated (17, 18, 28), our data suggest a mechanism of action in the yeast model that is not ionophore based (18). Rather, CQ promoted the degradation of metal-dependent Aβ oligomers within the secretory and endosomal compartments.

![Figure 4](https://www.pnas.org/legacy/pnas1402228111/supplemental/fig4.jpg)

**Fig. 4.** CQ rescues Aβ through a metal-dependent mechanism. (A) Structures of CQ derivative, chloroxine (CQ,3), and analogs lacking either the hydroxyl group (CQ\(_{-}\)-OH) or aromatic nitrogen (CQ\(_{-}\)-N). (B) Dose–response curves of CQ\(_{-}\) analogs. Data are expressed as the difference in OD\(_{500}\) between drug-treated and untreated cultures. Values above graph indicate toxicity. Only the maximum effective concentration (1 μM, “Max”) is shown for CQ and CQ\(_{-}\). (C) Dose–response curves of Aβ cultures treated with CQ and equimolar copper, iron, or zinc indicate interactions between CQ and copper and iron. Data reflect differences in OD\(_{500}\) between treated and untreated samples normalized to maximal rescue by CQ (micromolar) in the absence of supplemented metals.

**CQ Dramatically Reduced Aβ Peptide Levels by Increasing Degradation.**

To determine whether CQ has a similar ability to modulate Aβ within the context of the secretory compartment of a living cell, we returned to yeast. First, we examined the effects of CQ on the accumulation of Aβ using denaturing SDS gels. CQ dramatically reduced the accumulation of Aβ, with a sharp dose dependence that mirrored the sharp dose dependence of CQ's ability to rescue Aβ toxicity (Fig. 6A, Upper). CQ had no effect on total protein levels (Fig. 6A, Lower).

We next tested whether exogenous metals antagonized the ability of CQ to reduce Aβ levels. Indeed, they did so in a manner similar to their effects on the rescue of Aβ toxicity (Fig. 4C). Copper completely eliminated CQ's effects on Aβ accumulation; iron affected it modestly; and zinc had no effect (Fig. 6B). We next asked what species of Aβ were most affected by CQ. Nondenaturing gels, which preserved oligomer structure, demonstrated a decrease in most forms of Aβ when cultures were treated with CQ (Fig. 6C, Right). The levels of a control protein (YFP) expressed from the same promoter did not change (Fig. 6C, Left). Thus, the ability of CQ to rescue Aβ toxicity directly correlated with its ability to prevent the accumulation of oligomeric Aβ in a manner dependent on metal binding.

![Figure 5](https://www.pnas.org/legacy/pnas1402228111/supplemental/fig5.jpg)

**Fig. 5.** Effects of copper and CQ on Aβ oligomers. (A) ThT fluorescence of Aβ assembly reactions in the presence of copper (micromolar) at indicate molar ratios compared with Aβ. EM shows representative fields of Aβ with no or 50 μM Cu\(^{2+}\). (B) Dot blot analysis of Aβ assembly (10 μM) in the presence of both copper and CQ at 1:1 and 5:1 molar ratios. A11 monitors oligomeric Aβ conformations and 6E10 monitors total Aβ. (C) Dot blot (Left) and quantitation (Right) of Aβ assembly reactions in the presence of indicated concentrations of copper and CQ.
Thus, our yeast Aβ model has allowed us to focus on an early aspect of Aβ misfolding and the identification of CQ indicates that these early events are directly modulated by metals. Moreover, the results from this chemical screen indicate that modulating Aβ:metal complexes directly ameliorates the defects caused by Aβ oligomers within the secretory/endoosomal pathways, which we previously linked to AD risk alleles through genetic screens (8). This connection further establishes the potential for this class of small molecules to mitigate both early and late aspects of Aβ cellular pathologies.

Endogenous transition metals play an important role in Aβ pathology and their modulation has potential promise in treating AD patients (19, 36, 37). We suggest that the metal-dependent enhanced degradation of Aβ we observe here for CQ rescue of Aβ toxicity plays a role in neuronal systems as well. However, it is important to note that multiple mechanisms likely operate on the longer timescales of Aβ accumulation in mouse models and in human disease. For example, accumulation of transition metals in extracellular Aβ plaques formed late in disease can limit intracellular metal availability (28). CQ can dissolve plaques that have formed in late stages of disease (in both mice and humans) (17) and liberate trapped metals. This then presumably allows the ionophoric capability of CQ and PBT2 to simultaneously restore metal homeostasis and up-regulate proteases that degrade extracellular Aβ (18, 28, 36). Deciphering both early and late aspects of Aβ pathology will be important in understanding cellular pathologies and developing new therapeutic approaches in the treatment of AD.

The nearly complete elimination of Aβ accumulation that is mediated by CQ in yeast far exceeds any effect on protein stability that we have observed for any other compound or genetic suppressor in any of our yeast models of protein-folding pathologies (20, 38). CQ modestly rescues α-syn and htt2Q toxicity [Fig. 1E (20)]. However, it has little effect on the accumulation of these proteins (20). Presumably, this difference in the activities and efficacy of CQ is due to some, as yet unknown, difference in the cellular context and biophysical properties of different metal-binding proteins [of which α-syn is one (39) ]. We found that the degradation of Aβ was not affected by the ablation of any single pathway (i.e., autophagy; Fig. S4). This, together with the extreme rate of CQ-mediated Aβ degradation, suggests that multiple mechanisms are at play.

![Fig. 6. CQ promotes Aβ degradation. (A) Denaturing SDS-PAGE and Aβ immunoblot show reduction of Aβ levels in response to CQ treatment compared with total protein (Coomassie). (B) Aβ immunoblots of cells treated with CQ and equimolar Cu²⁺, Fe³⁺, and Zn²⁺. Pgk1 is a loading control. (C) Aβ immunoblot of nondenaturing gel shows a decrease in all forms of Aβ. Aβ oligomeric states are indicated to the right of the monomer control lane. A YFP control strain is shown on the left. (D) Immunoblot analysis of Aβ-expressing yeast with DMSO control or CQ after a short CHX time course. Short and long exposures are shown for comparison. (E) The percentage of Aβ remaining after β2Methionine pulse-labeling:Aβ-expressing yeast in presence or absence of CQ. Aβ was immunoprecipitated and quantified at 2-h intervals after pulse labeling.](image-url)

Empowered by significant improvements in DNA sequencing, a considerable effort is now underway to catalog polymorphisms in the human genome and identify those that contribute to pathologies related to protein homeostasis. It is now clear that the total number of human polymorphisms is much greater than previously realized (40), creating an intense need for rapid and tractable model systems to explore genotype–phenotype causality. Our yeast Aβ system meets this need. A previous unbiased genetic screen for modifiers of Aβ toxicity allowed us to link toxicity to recently discovered human AD risk alleles (8) with no previously known relationship to Aβ. Here, the facile yeast system enabled high-throughput compound screening that allowed us to rapidly identify relevant compounds and interrogate their mechanism of action. This ultimately validates our approach as we pursue new compounds with unknown protective mechanisms. Although this initial screen primarily revealed the central importance of metals in cellular Aβ pathology, we anticipate that other compounds with distinct modes of action will be identified. In general, for AD and other protein-misfolding diseases, yeast chemical genetic approaches will help elucidate new targets and “druggable” pathways and aid discovery and validation (38). It is imperative to integrate several orthogonal model systems and many other recent advances (10, 22, 38, 41, 42) to further our understanding of, and eventually our ability to treat, devastating diseases such as AD.

**Methods**

**Constructs and Yeast Strains.** The signal sequence: Aβ (ssAβ) construct has been described (8). The chemical screening strain used here was generated by integrating either GAL1-regulated Aβ or YFP constructs into the trp1 and ura3 loci of a WT strain and deleting the drug efflux pump (pdr5::KanMX) using standard methods (Table S1). The MUP1-GFP Aβ strain was generated by recombining a PCR product of GFP and a downstream HIS3 auxotrophic marker amplified from a chromosomal MUP1-GFP WT yeast strain (38).

**Small-Molecule Screen and Compounds.** Small-molecule screening was carried out as described (20). Briefly, late log phase Aβ cultures grown in 2% (wt/vol) raffinose-containing media were diluted (OD₆₀₀ = 0.03) into galactose-containing media. Compounds (100 nL) were pipetted to 384-well plates containing diluted culture and incubated for 40 h at 30 °C. OD₆₀₀ values were then used to calculate Z scores (OD₆₀₀ well – OD₆₀₀ plate average)/OD₆₀₀ plate average. Compounds were purchased as follows: CQ (Sigma-Aldrich); CQ₉-OD (Sigma-Aldrich); CQ₉-OD (Santa Cruz Biotechnology); CQ₉-N (Frontier Scientific Services).

**Growth Assays.** Growth assays with compounds were performed by either Bioscreen C or 384-well plates analysis. In both cases, log phase raffinose cultures were diluted (OD₆₀₀ ~ 0.02) into galactose-containing media, compounds or DMSO added, and cultures grown at 30 °C with either intermittent shaking (Bioscreen C) or endpoint (384-well) OD₆₀₀ readings. Data were typically expressed as the difference in OD₆₀₀ between compound- and DMSO-treated cultures. For all metal experiments, metals were added at the same time as CQ. Specificity experiments were performed in conditions with at least 50% reduction in growth in response to gene deletion or toxic compounds (Table S2). Toxic compound concentrations were selected to inhibit growth of WT yeast by at least 50%.

**MUP1-GFP Assay.** The effects of Aβ expression on Mup1-GFP endocytosis were performed as described (38). Cultures induced for 12 h with SGA1Met in a Bioscreen C with or without CQ (0.8 μM), methionine (50 μg/mL) added for 30 min, and GFP localization then imaged by fluorescence microscopy.

**C. elegans QTP Treatment.** The C. elegans strain (UA166) expressing an ssAβ construct in glutamatergic neurons has been described (8). Compound treatment (20) and analysis of glutamatergic neurons (8) have been described.
containing 10 μM Aβ with varying concentrations of Cu and CuO were loaded onto a 0.1-μm nitrocellulose membrane using the Bio-Rad vacuum filter trap device. The membrane was then blocked in 5% (v/v) nonfat dry milk in PBS for 1-2 h followed by an overnight incubation in A11 antibody (Millipore; 1:250 in 5% (v/v) milk/PBS) and a horseradish peroxidase-conjugated anti-rabbit secondary antibody (Sigma). Quantification of dot blots was performed using ImageJ (NIH). Transmission electron microscopy was performed by incubating Aβ solutions aggregated for ~12-15 h on a Ni-Formvar grid followed by negative staining by 2% (w/v) uranyl acetate.

Effect of CQ on Aβ Levels. The sAβ and YFP strains were grown to log phase in media containing galactose with either 0.6 μM CQ or DMSO and grown in a Bioscreen C plate at 30 °C to an OD600 of ~0.6 (~16 h). Lysetes were generated and blots probed as previously described (8).

Analysis of Aβ Degradation. For CHX experiments, Aβ yeast strains were cultured and induced as above with or without CQ. After Aβ induction, CHX was added (35 μg/ml) and aliquots removed at indicated time points. Samples were brought to 40 mM sodium azide, washed with ice-cold water, and Aβ levels analyzed (8). For [35S]methionine pulse-labeling experiments, cells expressing sAβ were grown as described above, harvested, washed twice in media lacking methionine, resuspended in the same media (OD600 = 1.0), and incubated at 30 °C for 15 min. L-[35S]Methionine (Perkin-Elmer; specific activity >1,000 Ci/mmol) was then added (50 μCi/μL) for 20 min. Cultures were diluted into media containing excess cold methionine (450 μg/mL) to an OD600 of ~0.5. Aliquots were removed, brought to 20 mM Na azide and 0.1 mg/ml BSA and placed on ice. Aβ was immunoprecipitated with the 6E10 antibody (Covance) and Protein A/G Plus Agarose ( Pierce). After washing, beads were boiled in sample buffer and supernant subjected to SDS/PAGE. For total label, Aβ immunoprecipitation total extract was resolved and the entire lane quantified. Imperial blue (Pierce)-stained gels were washed with water, equilibrated with 1 M sodium salicylate in 10% (v/v) glycerol, and dried. The gel was exposed to a FujiFilm imaging plate at ~20 °C, the plate scanned with a FujiFilm BAS-2500 Bio-Imaging Analyzer, and the data quantitated with Multi Gauge, version 2.2, software.

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