Quercetin Targets Cysteine String Protein (CSPα) and Impairs Synaptic Transmission

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Abstract

Background: Cysteine string protein (CSPα) is a synaptic vesicle protein that displays unique anti-neurodegenerative properties. CSPα is a member of the conserved J protein family, also called the Hsp40 (heat shock protein of 40 kDa) protein family, whose importance in protein folding has been recognized for many years. Deletion of the CSPα in mice results in knockout mice that are normal for the first 2–3 weeks of life followed by an unexplained presynaptic neurodegeneration and premature death. How CSPα prevents neurodegeneration is currently not known. As a neuroprotective synaptic vesicle protein, CSPα represents a promising therapeutic target for the prevention of neurodegenerative disorders.

Methodology/Principal Findings: Here, we demonstrate that the flavonoid quercetin promotes formation of stable CSPα-CSPα dimers and that quercetin-induced dimerization is dependent on the unique cysteine string region. Furthermore, in primary cultures of Lymnaea neurons, quercetin induction of CSPα dimers correlates with an inhibition of synapse formation and synaptic transmission suggesting that quercetin interferes with CSPα function. Quercetin’s action on CSPα is concentration dependent and does not promote dimerization of other synaptic proteins or other J protein family members and reduces the assembly of CSPα:Hsc70 units (70kDa heat shock cognate protein).

Conclusions/Significance: Quercetin is a plant derived flavonoid and popular nutritional supplement proposed to prevent memory loss and altitude sickness among other ailments, although its precise mechanism(s) of action has been unclear. In view of the therapeutic promise of upregulation of CSPα and the undesired consequences of CSPα dysfunction, our data establish an essential proof of principle that pharmaceutical agents can selectively target the neuroprotective J protein CSPα.

Introduction

In neurons, there are significant demands on cellular folding events. Complex interactions between multiple cellular components underlie synaptic transmission, a process that occurs with speed, precision and plasticity for extended periods of time. Rigorous synaptic quality control mechanisms likely provide defense against the detrimental effects of functionally impaired synaptic proteins. Indeed, Huntington’s, Alzheimer’s, Parkinson’s, Amyotrophic lateral sclerosis and prion diseases are caused by defects in protein folding, underlying the biological importance of the problem of aberrant protein folding in neurons. What synaptic mechanisms mediate the balance between protecting proteins and preventing accumulation of misfolded proteins remains a current biological question.

Cysteine string protein (CSPα) is a 34 kDa synaptic vesicle protein and molecular chaperone that is critical in the defense against neurodegeneration. CSPα, so called because it contains a 25 amino acid domain comprising a string of 13–15 cysteine residues, is a member of the conserved J protein family based on the presence of a 70 amino acid signature J domain [1]. CSPα is abundant in neural tissue and displays a characteristic localization to synaptic vesicles [2] as well as clathrin coated vesicles [3]. Furthermore, CSPα null-organisms display widespread neurodegeneration [4–7]. Deletion of the CSPα gene generates mice that are normal for the first 2–3 weeks of life followed by a progressive loss of muscle strength and motor coordination, neurodegeneration, blindness and premature death [5,6]. Although the underlying molecular mechanisms of neurodegeneration in CSPα-null mice have not yet been established, electron microscopic analysis indicates that degeneration begins presynaptically [5]. In Drosophila, CSPα knockout flies that survive to adulthood show paralytic uncoordinated sluggish movements, spasnic jumping, shaking and temperature sensitive paralysis. While the precise sequence of pathogenic events remains to be identified, the reported defects include a 50% reduction of nerve-evoked neurotransmitter release at 18–22° C, a drastic reduction in evoked release above 29° C, a reduced ability to maintain normal
presynaptic Ca\textsuperscript{2+} levels and reduction of synaptic boutons at neuromuscular junctions [8–13]. Reduction in synaptic transmission, temperature sensitive paralysis and premature lethality are reversed by the expression of normal CSP\textsubscript{\alpha} [9,10,12].

The J domain of CSP\textsubscript{\alpha} interacts with and activates the ATPase activity of Hsc70 (70 kDa heat shock cognate protein) [14,15] and Hsp70 (70 kDa heat shock protein) [16]. Together with Hsc70 and SGT (small glutamine-rich tetra-tricopeptide repeat domain protein), CSP\textsubscript{\alpha} assembles into an enzymatically active chaperone complex [17,18]. Following activation of the heat shock response, another member of the J protein family, Hsp10 (heat shock protein of 40kDa) assembles with the CSP\textsubscript{\alpha} complex [19]. The presence of this chaperone complex on secretory vesicles suggests that CSP\textsubscript{\alpha} is a coordinating anchor in key conformation/activity change(s) of client protein(s) critical in synaptic transmission. A number of client proteins for the CSP\textsubscript{\alpha} system have been proposed including: G proteins, SNAREs (soluble N-ethylmaleimide-sensitive factor attachment receptor), synaptotagmin, rab3, voltage sensitive calcium channels and CFTR (cystic fibrosis transmembrane conductance regulator) [20] and it is likely that neurodegeneration in null-organisms is due to the progressive misfolding and accumulation of dysfunctional client protein(s).

It has been suggested that interference with CSP\textsubscript{\alpha} function (eg toxic proteins, environmental toxins) may be an underlying mechanism leading to neurodegenerative diseases [5]. It follows that relatively small changes to CSP\textsubscript{\alpha}’s activity would be expected to significantly affect neural survival, however there is currently no direct support for this notion. In this study, we begin to address the hypothesis that inhibition of CSP\textsubscript{\alpha} activity may be common to the pathological sequence of events that underlies neurodegenerative disease and that the neuroprotective synaptic vesicle protein CSP\textsubscript{\alpha} represents a promising therapeutic target for the treatment or prevention of neurodegeneration. Here we identify CSP\textsubscript{\alpha} as a target for quercetin, a naturally occurring flavonoid. The Western diet contains ~25 mg/day of mixed flavonoids (quercetin ~70%). Quercetin is particularly high in apple skins, green tea and red grapes. It is also a major component of the nutrient supplements Ginkgo Biloba and St. John’s Wort. Ginkgo Biloba is widely heralded as a memory enhancer but is also commonly taken for altitude sickness and cancer. St. John’s Wort is commonly taken for depression. Ginkgo biloba leaves have been used for many centuries in traditional Chinese medicine and justification for quercetin supplements is historical rather than mechanistic. In this study we show that quercetin promotes CSP\textsubscript{\alpha} dimerization and inhibits synaptic transmission as well as synapse development. The identification of quercetin as an agent that targets CSP\textsubscript{\alpha} is a first step in the identification of pharmaceuticals that target members of the large J protein family and as such serves as proof of principle that pharmaceutical tools can selectively target J proteins. Given CSP\textsubscript{\alpha}’s anti-neurodegenerative properties, upregulation of CSP\textsubscript{\alpha} may hold therapeutic promise in protecting nerve terminals from misfolded or toxic proteins [21].

Results

Quercetin stimulates formation of CSP\textsubscript{\alpha}-CSP\textsubscript{\alpha} dimers

70 kDa detergent-resistant CSP\textsubscript{\alpha} dimers have been extensively reported in rat brain [16,22,23], various cell lines [24,25] and purified preparations [26] however, the role this dimer plays in CSP\textsubscript{\alpha}-mediated conformational work is not known. In order to investigate the possibility that the neuroprotective synaptic vesicle protein CSP\textsubscript{\alpha} can be targeted by pharmaceutical agents, we screened for drugs which initiate changes in CSP\textsubscript{\alpha} dimerization. Figure 1A shows that under control conditions, a small fraction of total CSP\textsubscript{\alpha} is detectable as a dimer in CAD cells (CNS-derived catecholaminergic neuronal cells) in addition to monomeric CSP\textsubscript{\alpha} (both unmodified and palmitoylated). Post translational modification of CSP\textsubscript{\alpha}, involving extensive fatty acylation (*) results in its retarded migration upon SDS-PAGE [22,27]. Interestingly,
Figure 2. Quercetin exerts a concentration-dependent effect on the formation of the CSPα dimer in rat cortical neurons and CAD cells. (A) Western blot of cultured rat cortical neurons were treated with indicated concentrations of quercetin for 24 hours prior to lysis. Equal amounts of cellular protein were resolved by SDS-PAGE as confirmed by ponceau S staining. (B) Native CSPα was detected in adult rat brain by Western analysis with a monoclonal anti-CSPα antibody. Twenty-five micrograms of unfractionated tissue homogenate isolated from the indicated regions of rat brain were separated by SDS-PAGE, transferred to PVDF and probed. Arrows indicate the CSPα dimer at ~72 kDa; * indicates a palmitoylated CSPα monomer at ~34 kDa. Actin is shown as a loading control. doi:10.1371/journal.pone.0011045.g002

Quercetin Targets CSP

We next examined the effect of quercetin on the dimerization of CSPα in rat cortical neuronal cultures. Figure 2A shows that quercetin promoted CSPα dimerization in a concentration (20 μM-100 μM) dependent manner in cortical neurons expressing endogenous levels of CSPα. Increases in dimer were detectable at 20 μM quercetin. The expression levels of Hsc70 and Hsp40 chaperones which are known to associate with CSPα [14,15,16,19], were not altered by quercetin in treated cortical cultures. Actin is shown as a loading control. No CSPα dimerization was initiated by the DMSO vehicle control. The concentration-dependent induction of the CSPα dimer by quercetin in CAD cells is shown in Figure 2B. No changes in either the expression or formation of dimers were observed for the chaperones Hsc70, Hsp40, Rdj2 or the putative CSPα client proteins Gαs and syntaxin, consistent with findings in cortical neurons. Figure 2C shows that relative differences in CSPα monomer and dimer expression are further observed in different rat brain regions. The CSPα dimer was highest in the thalamus, midbrain, entorhinal cortex and pons and low in the spinal cord and medulla. Taken together, these results show that a) in intact brain, the expression of CSPα-CSPα dimers is region specific, b) in cortical neuronal CSPα dimerization is induced in the presence of quercetin in a concentration dependent manner and c) quercetin does not cause a generalized oligomerization of J proteins.

We next established the time course of quercetin’s induction of CSPα dimers. Transfected CAD cells were treated with 100 μM quercetin, lysed at 5 min, 8 hrs, 24 hrs, 48 hrs and the formation of CSPα dimers evaluated by Western analysis with an anti-myc monoclonal or anti-CSPα polyclonal antibodies. The structure of quercetin (3,3’,4’,5’,7-pentahydroxyflavone), a common dietary flavonoid, is shown in Figure 1B. In contrast to quercetin, the Hsp90 inhibitor geldanamycin, the neurotoxin MPP+, the oxidizing agent H2O2 and the proteasome inhibitor lactacystin, did not stimulate CSPα dimerization, indicating that quercetin’s effect on CSPα is selective. The quercetin-induced CSPα dimer is resistant to disruption by incubation in SDS-containing sample buffer at either 37°C or 80°C (Figure 1C). SDS separates the vast majority of cellular protein complexes to monomers by treatment 37°C and more stable protein complexes at 80°C. The SDS and temperature resistance of the CSPα-CSPα complex demonstrates the great stability of the dimer. In contrast, no oligomerization or changes in expression of Hsc70 were observed. Actin detection is shown as a loading control. Our data demonstrate that the extremely stable CSPα-CSPα complex is selectively increased by quercetin.
Quercetin Targets CSP

Quercetin inhibits synapse formation and synaptic transmission

Deletion of CSPα is ultimately linked to a state where the integrity of synaptic terminals is compromised. Therefore, the ability of quercetin to target CSPα prompts the question: does quercetin promote or inhibit synaptic function? To discern between these possible scenarios we evaluated synapse formation in the fresh water snail Lymnaea stagnalis primary neuronal culture model that is uniquely suitable for precise measurements of synaptic transmission at a resolution not achievable elsewhere. In these experiments, the functionally defined respiratory neurons VD4 (visceral dorsal 4; presynaptic, cholinergic) and LPeD1 (left pedal dorsal 1; postsynaptic) were plated with their somata juxtaposed onto poly-L-lysine-coated dishes and the excitatory synapses allowed to develop for 12–18 hrs either in the presence or absence of 25 or 100 μM quercetin (Figure 4A). Prior to intracellular recordings, quercetin was washed off for 1 to 2 hrs and VD4 presynaptic action potentials and LPED1 excitatory postsynaptic potentials (EPSP) were recorded. As expected, in control synapses, current injection-induced action potentials in VD4 generated 1:1 EPSPs with averaged amplitudes of 9.8 ± 1.7 mV in LPED1 (n = 8) (Figure 4A and B) similar to that seen in vivo [28–30]. However, in the presence of 25 μM quercetin, the mean amplitude of evoked EPSPs was significantly reduced to 2.5 ± 0.7 mV (n = 5, p < 0.05) (Figure 4A and 4B, insert). Five VD4/LPED1 cultures failed to form synapses in the presence of 100 μM of quercetin, while synaptic transmission was detected in 5 neuron pairs but with greatly reduced amplitude (0.6±0.4 mV).

Figure 3. Quercetin increases the stability of the CSPα dimer in CAD cells. (A) CAD cells were transfected with 1.0 μg of c-myc CSPα DNA and treated with 100 μM quercetin as shown. Cells were lysed at indicated times following quercetin treatment. 30 μg of protein was resolved. Upper panel: CSPα was detected by Western analysis with the c-myc antibody. Lower panel: Quantification of the CSPα dimer at 5 minutes and 24 hours in control and quercetin-treated cells. Results are expressed as mean +/- SE for a total of 4 separate experiments. (B) Equal volumes of purified recombinant rat CSPα were treated with indicated concentrations of quercetin for 24 hrs (C) CAD cells were transfected with 1.0 μg CSPα A-CSPα D, CSPα A-CSPα D, and Rdj2 DNA and treated with 100 μM quercetin for 24 hours. Arrows indicate CSPα dimer at ~72 kDa. Data are representative of three separate experiments.

doi:10.1371/journal.pone.0011045.g003
Furthermore, quercetin caused clamping of tetanic action potential firing in 90% of presynaptic VD4 cells examined (n = 21) (Figure 4A). We next measured the amplitude of potentiated EPSP (pEPSP) following a tetanus compared to the amplitude of the EPSPs before tetanus (Figure 4B, insert). The ratio of pEPSP and EPSP was significantly reduced by 100 μM (n = 8) but not by 25 μM (n = 5) of quercetin. Overall, our data establish that quercetin impaired synapse formation and reduced synaptic plasticity in *Lymnaea* and thus support the hypothesis that quercetin targets and inhibits CSPα function leading to a loss of synaptic integrity.

As expected, Western blot analysis revealed both CSPα monomers and quercetin-induced CSPα-CSPα dimers in *Lymnaea* (Figure 5A). *Lymnaea* were maintained in pond water containing 100 μM quercetin for 12–18 hrs and the VD4 ganglia were then harvested and neuronal proteins were resolved by SDS-PAGE. Native CSPα was detected by Western analysis with the CSPα polyclonal antibody. The panels shown are from the same experiment and are representative of three independent experiments. Arrow indicates CSPα dimer at ~72 kDa. Hsc70 is shown as a loading control. (B) Paired *Lymnaea* soma were cultured overnight and subjected to immunostaining with CSPα polyclonal antibody. Stacks of 0.28 μm slices were collected and collapsed into Z projections in maximum intensity using ImageJ. Images are representative of five experiments.

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**Figure 4.** Quercetin inhibits synapse formation in *Lymnaea stagnalis* primary cultured neurons. The presynaptic, cholinergic neuron, visceral dorsal 4 (VD4) and the postsynaptic neuron, left pedal dorsal 1 (LPeD1) were juxtaposed and cultured in the absence or presence of quercetin (25 or 100 μM) for 12–18 hours. Prior to intracellular recordings, quercetin was washed off. (A) Sample traces of presynaptic action potentials on VD4 cells and excitatory postsynaptic potentials (EPSPs) on LPeD1 cells. (B) The mean amplitude of EPSP and the ratio of potentiated EPSP (pEPSP) over EPSP was reduced in the presence of both 25 μM quercetin (n = 5) and 100 μM quercetin (n = 7) (inserts). Statistical significance was determined using Student’s t-test. * indicates significant difference at the level of p = 0.05. Error bars indicate S.E.

doi:10.1371/journal.pone.0011045.g004

**Figure 5.** Quercetin induces the CSPα dimer in *Lymnaea stagnalis*. (A) *Lymnaea* were maintained in pond water containing 100 μM quercetin over night as indicated prior to harvesting of the VD4 ganglia from several snails. Equal numbers of ganglia were combined and resolved on a gel. Native CSPα was detected by Western analysis with the CSPα polyclonal antibody. The panels shown are from the same experiment and are representative of three independent experiments. Arrow indicates CSPα dimer at ~72 kDa. Hsc70 is shown as a loading control. (B) Paired *Lymnaea* soma were cultured overnight and subjected to immunostaining with CSPα polyclonal antibody. Stacks of 0.28 μm slices were collected and collapsed into Z projections in maximum intensity using ImageJ. Images are representative of five experiments.

doi:10.1371/journal.pone.0011045.g005
polyclonal antibody generated against the C terminus of rat CSPz. Quercetin-induced CSPz dimer formation in Lymnaea respiratory neurons (Figure 5A) was similar to that found for transfected CAD cells and cortical neurons (Figures 1 and 2). Confocal microscopy confirmed that CSPz is abundant in paired Lymnaea neurons, with highest localization observed at the plasma membrane (Figure 5B). No signal was obtained with either quercetin alone or secondary antibody alone (data not shown). In addition to Lymnaea, CSPz homologues have been reported in Torpedo, Xenopus, Drosophila and numerous mammals; this high degree of evolutionary conservation of the CSPz system makes a strong case for an important cellular function. Of course, J proteins are even more extensively conserved and have been reported in bacteria, plants and viruses [20] where they most certainly are involved in conformational work. Our data provide unequivocal evidence that the the sensitivity of the CSPz chaperone system to quercetin is conserved from mammals to Lymnaea.

To ask whether the functional defects observed following quercetin treatment of intact neurons were time dependent, we next tested the effects of acute application of quercetin on synaptic transmission. Strikingly, Figure 6 shows that bath application of 25 μM quercetin to functional VD4/LPeD1 cultured neurons rapidly reduced synaptic transmission within 10 minutes. Neurons were paired overnight and on day two, simultaneous pre- and postsynaptic intracellular recordings under control conditions clearly showed that induced action potentials in VD4 produced 1:1 EPSPs in LPeD1 and these were potentiated significantly after a brief presynaptic tetanus. The amplitude of EPSPs in LPeD1 was greatly diminished within minutes of perfusion with 25 μM quercetin (Figure 6A) indicating that quercetin rapidly inhibited synaptic transmission. A typical action potential is shown before (red) and after (blue) exposure to quercetin. Quercetin prolonged the presynaptic repolarization phase of the action potential, thereby lengthening action potential duration and rendering the neuron incapable of rapid and repetitive firing. Washout of quercetin (10-20 min) did not reverse the reduction in EPSP (Figure 6B) suggesting that quercetin-dependent alteration of CSPs is irreversible in the short term. In the absence of an action potential quercetin did not stimulate exocytosis and spontaneous EPSPs were not observed following exposure to quercetin (data not shown). Larger reductions in EPSP amplitude were observed with 100 μM compared to 25 μM quercetin. The ratio of the pre-tetanus EPSPs to that of post-tetanus was reduced following quercetin treatment (Figure 6C) demonstrating a reduction in the ability of neurons to exhibit short-term synaptic plasticity.

Even more telling, quercetin did not alter the postsynaptic response to acetylcholine (Figure 7A). Postsynaptic LPeD1 cells were cultured overnight (12-18hrs) and 1 μM Acetylcholine (ACh), the transmitter released from presynaptic VD4, was exogenously applied before and after exposure to quercetin (25 or 100 μM) for 30 mins while the postsynaptic potential was monitored. ACh elicited excitatory postsynaptic potentials which triggered firing of action potentials both before and after the exposure to either 25 or 100 μM quercetin for 30 mins (n = 4). Higher concentrations of quercetin LPeD1 action potentials showed a slower rate of repolarization.

To further test for quercetin-induced changes in presynaptic function, action potentials were recorded simultaneously with intracellular Ca2+ concentration [Ca2+]i measurements before and after perfusion of quercetin (25 μM) (Figure 7B). VD4 cells were cultured overnight and then loaded with the fluorescent Ca2+ indicator, Fura-2 AM. Elevations in intracellular free [Ca2+]i in response to single action potential and action potential bursts were significantly reduced after exposure to quercetin for 20 mins (n = 4) compared to control neurons (Figure 7B insert). Because 100 μM quercetin induced action potential clamping (Figure 4A and 6A), we opted to directly measure presynaptic Ca2+ currents (ICa) by whole cell voltage clamp with direct depolarizing steps from holding potential of −80 mV to +70 mV in 10 mV increments. Exposure to 100 μM quercetin...
for 15 min reduced depolarization-induced Ca\(^{2+}\) entry (Figure 7c). Figure 7C left panel shows raw traces of ICa induced by a depolarization step from $-80 \text{ mV}$ to $+20 \text{ mV}$ before and after exposure to quercetin 100 \(\mu\)M for 15 min. Normalized current-voltage relations of ICa, (Figure 7C right panel) shows that the peak current occurred at $+20 \text{ mV}$ and the normalized peak current was significantly reduced from a control level of 0.90±0.10 to 0.49±0.10 (\(n = 4\), \(P < 0.05\)) after 15 mins of 100 \(\mu\)M quercetin. These observations indeed indicate that quercetin indeed rapidly inhibits action potential-induced Ca\(^{2+}\) entry and reduces synaptic transmission in Lymnaea neurons, consistent with the hypothesis that quercetin binds to and inactivates CSP\(\alpha\), leading to a sequence of events that involves disabling ion channels (eg voltage sensitive Ca\(^{2+}\) channels). We have previously shown that CSP\(\alpha\) promotes G protein-mediated inhibition of N-type Ca\(^{2+}\) channels [23,31] thereby directly regulating Ca\(^{2+}\) channel activity. Of note, CSP\(\alpha\) has been suggested to be a Ca\(^{2+}\) channel chaperone, but this notion has been controversial [32–38]. In addition, previous studies have concluded that quercetin is both an L type Ca\(^{2+}\) channel activator [39] and inhibitor [40,41] as well as a BKCa channel activator [42], Kir channel inhibitor [43] and Ca\(^{2+}\) ATPase inhibitor [44,45]. Although our data in Lymnaea VD4/ LPeD1 respiratory neurons support the idea that quercetin’s inhibition of CSP\(\alpha\) activity leads to downstream inhibition of synaptic transmission, these findings together with the above stated reports could potentially also be explained by a direct block of presynaptic Ca\(^{2+}\) channels by quercetin. Previous studies have demonstrated that lowering extracellular Ca\(^{2+}\) also blocks synapse formation between cultured Lymnaea neurons [46] showing the importance of Ca\(^{2+}\) in the activation of synaptogenesis.

To establish more definitely if channel blockers generally altered CSP\(\alpha\) dimer levels, we again utilized the transiently transfected CAD cell model. No significant differences in CSP\(\alpha\) dimer formation was observed following treatment of transiently transfected CAD cells with CdCl\(_2\), nifedipine, D-2-amino-5-phosphonovalerate (D-APV) or tetrodotoxin (TTX) (Figure 7C). Furthermore, lowering cytosolic free [Ca\(^{2+}\)] by incubation with the membrane permeable Ca\(^{2+}\) chelator BAPTA-AM (20 \(\mu\)M), a Ca\(^{2+}\) chelator, for 1 hour did not stimulate CSP\(\alpha\) dimer formation or influence quercetin’s induction of the dimer. Taken together, these data indicate that none of the channel blockers examined promoted CSP\(\alpha\)-CSP\(\alpha\) dimer formation, suggesting that ion channel block per se or reduction in cytosolic free Ca\(^{2+}\) does not stimulate CSP\(\alpha\) dimerization, while quercetin impairs presynaptic Ca\(^{2+}\) influx and reduced neuronal ability to maintain normal Ca\(^{2+}\) levels. Independent of whether quercetin directly targets CSP\(\alpha\) and secondarily disables channels or directly targets channels as well as CSP\(\alpha\), the results shown in Figures 4, 5 and 6 demonstrate that quercetin promoted dimerization of CSP\(\alpha\), reduced the amplitude of EPSPs in LPeD1 Lymnaea neurons and rapidly triggered a prolongation of presynaptic repolarization associated with inhibition of synaptic transmission and reduced synapse formation.

Increases in the CSP\(\alpha\):CSP\(\alpha\) dimer correlate with a decrease in active CSP\(\alpha\):Hsc70 units

CSP\(\alpha\)-mediated conformational work relies on the assembly of a CSP\(\alpha\) with the ATPase Hsc70 [14–17]. To analyze the influence of quercetin on the assembly of the CSP\(\alpha\) active chaperone complex, CSP\(\alpha\) was immunoprecipitated from CAD cells treated with quercetin and the co-association of Hsc70 was evaluated. The CSP\(\alpha\) dimer, monomeric palmitoylated CSP\(\alpha\), and monomeric unpalmitoylated CSP\(\alpha\) immunoprecipitated from CAD cell lysates. Following treatment with quercetin, higher levels of CSP\(\alpha\) dimer immunoprecipitated as expected. Immunoprecipitation of the monomeric palmitoylated CSP\(\alpha\) (but not the unpalmitoylated) species was relatively lower following quercetin. Figure 8 shows
that quercetin treatment reduced Hsc70 association with CSPα, indicating that quercetin inhibits assembly of the active chaperone complex. These results suggest that the quercetin-induced changes in CSPα dimerization and synaptic transmission involve inhibition of CSPα chaperone activity.

Finally, we asked whether other flavonoids also elicit an increase in the abundance of CSPα-CSPα dimers. Exposure to epigallocatechin gallate (EGCG) increased CSPα dimerization in CAD cells like that observed for quercetin (Figure 9). The CSPα dimer was detected by Western analysis with anti-myc monoclonal antibody. Figure 9B shows that the ratio of CSPα dimer to monomeric palmitoylated CSPα was higher following quercetin treatment than EGCG treatment. Actin is shown as a loading control. We conclude that quercetin and EGCG are potent activators of CSPα dimerization and predict that the common polyphenol structure is a central structure involved in increasing CSPα dimerization.

Discussion

Our work fulfills the proof of principle that pharmacological compounds like quercetin can target select members of the J protein family to modulate their cellular behavior. Specifically, our data show that the flavonoid, quercetin, interferes with CSPα function to maintain synaptic integrity. A current working model of the CSPα complex is illustrated in Figure 10. In this model, CSPα is anchored to the secretory vesicle and on its own is inactive. At least two other CSPα complexes exist: the active Hsc70/SGT1/CSPα complex, and the CSPα dimer. CSPα displays unique anti-neurodegenerative properties and impairments in CSPα activity lead to impairment in synaptic transmission. In our model, quercetin targets CSPα, increases the abundance of stable CSPα-CSPα dimers and reduces assembly of CSPα with Hsc70, thereby reducing the folding capacity of the CSPα complex. CSPα dimerization may be a cause for the inhibition of neurotransmission, synaptic plasticity and synapse formation, essential features of normal brain function but other possibilities exist. Mechanistic insight into neurodegeneration following CSPα-inoperation remains limited but ultimately a reduction in synaptic folding capacity and a progressive buildup of unfolded presynaptic client proteins most likely creates a situation where the integrity of synaptic terminals is compromised.

Quercetin does not initiate widespread oligomerization of J proteins (Figures 2 and 3). The human genome encodes for over 40 J proteins with specific cellular and subcellular distributions [1]. The importance of the J protein family in protein folding has been recognized for many years [Reviewed: [47,48]]. Members of the J protein family have a modular architecture in which a signature J domain is grafted to other sequences that impose specific cellular functions. The J domain is a 70 amino acid signature region comprised of four helices with a highly conserved tripeptide of histidine, proline and aspartic acid (HPD motif) located between helices II and III that is critical for chaperone activity. Hsc70 is targeted to a particular substrate through binding to the “J domain” of its partner and regulates the conformation and activity of the target protein via mechanisms that appear to involve cycles of substrate binding and release, which are governed by ATP binding and hydrolysis. Since J proteins are thought to provide the basis for selective chaperone action in the cell they are also promising therapeutic targets for the manipulation of specific protein folding processes [21].

Many questions remain about the biochemical pathway(s) responsible for CSPα-mediated neuroprotection. It is obvious that the CSPα chaperone unit is highly conserved machinery built upon the universal J domain/Hsc70 association and contributes to presynaptic protection. Furthermore, it is known that other J proteins do not compensate for the absence of CSPα, consistent with the idea that CSPα is designed to facilitate specific synaptic folding events and that the unfolded CSPα client is toxic. That said, significant efforts towards establishing the identity of CSPα client proteins and the underlying molecular details of CSPα’s essential synaptic quality control are ongoing. Regulators of the degeneration seen in CSPα knockout models have been identified. Chandra and colleagues have shown that α-synuclein, a small neural protein whose biological function is unclear, selectively modulates the CSPα neurodegeneration pathway [7]. CSPα deficient mice are rescued from presynaptic degeneration and lethality by the overexpression of α-synuclein. Moreover, ablation of endogenous α-synuclein accelerates the degeneration of presynaptic terminals observed in mice lacking CSPα. Furthermore, α-synuclein specifically rescues CSPα deletion, but transgenic α-synuclein does not rescue the spinal cord degeneration in mice that express mutant superoxide dismutase. Precisely how α-synuclein abolishes neurodegeneration triggered by the absence of CSPα is unclear. Results reported here show that the cellular

Figure 9. Epigallocatechin gallate (EGCG) stimulates formation of CSPα-CSPα dimers. (A) CAD cells were transiently transfected with 1.0 µg c-myc-CSPα DNA and treated with indicated concentrations of EGCG for 24 hours prior to lysis. Following separation of cellular protein (30 µg) by SDS-PAGE, CSPα was detected by Western analysis. β-actin is shown as a loading control. Arrow indicates CSPα dimer at ~72 kDa. (B) Quantification of the CSPα dimer to monomeric palmitoylated CSPα ratio under control, 100 µM quercetin and 200 µM EGCG conditions. Numbers in parentheses indicate the numbers of experiments; error bars denote standard errors.

doi:10.1371/journal.pone.0011045.g009
events following exposure to quercetin include i) a reduction in presynaptic Ca\(^{2+}\) and ii) an increase in the presynaptic repolarization phase rendering neurons incapable of firing continuously, diminishing the amplitude of EPSPs and preventing additional synapse formation. One possibility is that following reduction of CSP\(\alpha\) folding activity, a downstream progressive misfolding of protein(s) central to Ca\(^{2+}\) homeostasis leads to inhibition of synaptic transmission. Another possibility is that quercetin has cellular targets in addition to CSP\(\alpha\) perhaps directly blocking voltage dependent Ca\(^{2+}\) channels. Although the precise mechanism by which quercetin alters Ca\(^{2+}\) homeostasis remains to be established in detail, it is notable that lowering of cytosolic free [Ca\(^{2+}\)] with either the Ca\(^{2+}\) chelator BAPTA-AM or the Ca\(^{2+}\) channel blocker CdCl\(_2\) does not stimulate CSP\(\alpha\) dimerization, demonstrating that CSP\(\alpha\) dimerization is not a general cellular response to Ca\(^{2+}\) channel blockade. Independent of which explanation is correct, our results illustrate that quercetin targets CSP\(\alpha\) and impairs synaptic function.

If quercetin is so powerful in inhibiting synaptic transmission, why do so many choose to supplement their diet with flavonoid mixtures (eg Ginkgo biloba)? Flavonoids are a class of compounds with polyphenolic structures and in all likelihood a spectrum of physiological functions can be expected depending on the position of hydroxyls and side chains. Our results indicate that the common structural features of quercetin and EGCG are important for triggering CSP\(\alpha\) dimerization ([Figures 1 and 9]). Such compromised CSP\(\alpha\) activity could contribute to the rate of progression of neurodegenerative (misfolding) diseases (eg. Alzheimer’s disease, Huntington’s disease) however there is no direct evidence for this notion. Neurotoxicity would depend on the flavonoid mixture utilized, metabolites and final concentration in the CSF (cerebrospinal fluid) after oral intake. Not all related compounds would necessarily inhibit CSP\(\alpha\) activity, and it remains to be established if select flavonoids would enhance rather than inhibit CSP\(\alpha\) function. Direct evidence supporting a role for flavonoids in memory enhancement is currently not available.

In view of the crucial importance of CSP\(\alpha\) in synaptic integrity, our data identify a key role for compounds that interfere with its specialized presynaptic function. We speculate that neuropathological abnormalities could be due to or exacerbated by toxins with inhibitory actions towards CSP\(\alpha\) similar to that of quercetin. Conversely, compounds related to quercetin may be found to enhance rather than inhibit CSP\(\alpha\) activity. Thus, the identification of quercetin as tool that selectively modulates CSP\(\alpha\)'s neuroprotective function is a promising lead towards the identification of agents that enhance CSP\(\alpha\)'s neuroprotective function and thereby have a high potential in therapy development for neurodegenerative diseases. Our study further predicts that compounds that selectively target J proteins may have considerable potential as novel therapeutic agents.

**Materials and Methods**

**Reagents and chemicals**

Anti-CSP\(\alpha\) rabbit polyclonal was prepared as described previously [22]. Anti-Hsp40 rabbit polyclonal were from Stressgen. Anti-Hsc70 mouse monoclonal, anti-\(\beta\)-actin mouse monoclonal, anti-syntaxin mouse monoclonal, quercetin, forskolin, MPP\(^{+}\) and lactacystin was from Sigma. Anti-c-myc mouse monoclonal was from Clontech. Anti-CSP\(\alpha\) mouse monoclonal was from BD Biosciences. Anti-G\(\alpha\), rabbit polyclonal and geldanamycin were from Calbiochem. Anti-Rdj2 mouse monoclonal was from Abnova. H\(_2\)O\(_2\) was from VWR. ACh was obtained from Research Biochemicals (Natick, MA; product A-112).

**CAD mouse neuroblastoma cells** [49,50] were seeded into 6 well plates and grown in DMEM/F12 medium supplemented with 10% fetal bovine serum and 1% penicillin/streptomycin as previously described. For differentiation cells were grown in Opti-MEM for 3 days. Cells were lysed in 40 mM Tris (pH 7.4), 150 mM NaCl, 2 mM EDTA, 1 mM EGTA, 1 mM Na\(_3\)VO\(_4\), 0.1% SDS, 1% T-X100, 0.5 mM PMSF and protease inhibitor (Sigma) at 4°C for 1 hour. Lysates were centrifuged at 15"000xg.
for 5 minutes at 4°C and the supernatant was collected. Protein concentration was determined using a Bradford style assay kit (BioRad). For transient transfection, CAD cells were washed in PBS and transiently transfected with e-myc tagged rat CSPα1-198 DNA using Lipofectamine-2000 (Invitrogen) in Opti-MEM and maintained in culture for 24 hours prior to treatments.

**Rat cortical neurons**

Rat primary cortical neurons (Cryopreserved) were purchased from QBM Cell Science (Ottawa, Canada) and stored in liquid nitrogen. Prior to culture, cells were thawed and gently transferred into pre-warmed neurobasal medium (Invitrogen, No. 21103-049) supplemented with 2% B27 (Invitrogen, 17504-044). Cells were then plated onto poly-D-lysine and laminin coated cover slips and maintained in neurobasal medium for 7 days at 37°C in 5% CO₂ prior to treatment with quercetin for 24 hours. Cells were harvested with lysis buffer and equal volumes of protein lysate were resolved by SDS-PAGE.

**Lymnaea ganglia**

Equal numbers of whole ganglia were harvested and lysed at 4°C for 40 minutes and resolved by SDS-PAGE.

Fresh water snails, *Lymnaea stagnalis*, were maintained at room temperature (22-23°C) in a well-aerated aquarium containing filtered pond water. Neurons were isolated from 1-2 month old snails with a shell length of 20-22 mm and the *Lymnaea* brain-conditioned medium was prepared using 3-6 month old animals with a shell length of 25-30 mm. The cell isolation and cell culture procedures have been described in detail elsewhere [51]. Briefly, the *Lymnaea* were dissected and central ring ganglia were removed. Following treatment with Trypsin (2 mg/ml) for 23 mins, the central ring ganglia were then treated with trypsin inhibitor (2 mg/ml) for another 15 mins. Identified presynaptic neuron visceral dorsal 4 (VD4, cholinergic) and postsynaptic neuron left pedal dorsal 1 (LPcD1) (for synaptogenesis and synaptic transmission experiments) were isolated by applying gentle suction through a fire-polished and Sigma-coat (Sigma, St. Louis, MO)-treated pipette. Isolated cells were then plated onto poly-L-lysine coated glass dishes in the presence of either medium (L-15; Life Technologies, Gaithersburg, MD; special order) or conditioned medium which contains trophic factors. Soma-soma synapses were prepared by juxtaposing VD4 and LPcD1 cell bodies against each other. The synapses developed overnight as described previously and were tested through direct intracellular recordings [20].

**Immunoblotting**

Proteins were electrotransferred from polyacrylamide gels to 0.45 μm nitrocellulose membrane in 20 mM Tris, 150 mM glycine and 12% methanol. Membranes were blocked with 4% glycine and 12% methanol. Membranes were blocked with 4% and QuantityOne 4.2.1 software. Antiserum were quantitated by Biorad Fluor-S MultiImager Max (Pierce Biotechnology Inc.). Immunoreactive bands were visualized secondary antibody. Antigen was detected using West Pico reagent solution and incubated with horseradish peroxidase-coupled secondary antibody. Protein lysate with a shell length of 25–30 mm. The cell isolation and cell culture procedures have been described in detail elsewhere [51]. Briefly, the *Lymnaea* were dissected and central ring ganglia were removed. Following treatment with Trypsin (2 mg/ml) for 23 mins, the central ring ganglia were then treated with trypsin inhibitor (2 mg/ml) for another 15 mins. Identified presynaptic neuron visceral dorsal 4 (VD4, cholinergic) and postsynaptic neuron left pedal dorsal 1 (LPcD1) (for synaptogenesis and synaptic transmission experiments) were isolated by applying gentle suction through a fire-polished and Sigma-coat (Sigma, St. Louis, MO)-treated pipette. Isolated cells were then plated onto poly-L-lysine coated glass dishes in the presence of either medium (L-15; Life Technologies, Gaithersburg, MD; special order) or conditioned medium which contains trophic factors. Soma-soma synapses were prepared by juxtaposing VD4 and LPcD1 cell bodies against each other. The synapses developed overnight as described previously and were tested through direct intracellular recordings [20].

**Electrophysiology**

Neuronal activity was recorded using conventional intracellular recording techniques as described previously [30]. Glass microelectrodes (1.5 mm internal diameter; World Precision Instruments, Sarasota, FL) with tip resistances of 20–50 MΩ were filled with a saturated solution of K₂SO₄. Neurons were impaled using Narashige (Tokyo, Japan) micromanipulators (MM202 and MM 204) (Axiovert 135; Zeiss, Thornwood, NY) on an inverted microscope. To test for synaptic connections, current was injected into the presynaptic neuron VD4 via an intracellular microelectrode which induced action potentials in the VD4 cell, and postsynaptic responses in LPcD1. The recorded electrical signals were displayed on a digital oscilloscope (PM 3394; Philips, Eindhoven, Netherlands) and relayed through a digitizer (Digidata 1322A, MDS Inc, Toronto, Canada) and recorded on a computer using Axoscope 9.0 software (MDS Inc, Toronto, Canada). Acetylcholine (ACh, 1 μM) was pressure applied (10–20 psi, 0.5–1 s duration) directly to the somata through a glass pipette (2–4 μm tip diameter) connected to a PV800 pneumatic picopump (World Precision Instruments).

Whole-cell recordings of voltage-gated Ca²⁺ currents (I_{Ca}) were performed using a Multiclamp 700B amplifier (Axon Instruments) connected to an analog-to-digital interface Digidata 1322 (Axon Instruments). Signals were acquired and stored on a personal computer equipped with pClamp 9.2 software (Axon Instruments). Borosilicate pipettes (A-M Systems, Inc, Sequim, WA) were pulled using a Sutter P-97 microelectrode puller (Sutter) and the pipette resistance was 3–7 MΩ after being filled with pipette solution containing: 35 mM CsCl; 1 mM CaCl₂; 2 mM MgATP; 10 mM EGTA; 10 mM HEPES; adjusted to pH 7.4 with CsOH. The external bath solutions used to isolate Ca²⁺ currents contains: 10 mM CaCl₂; 1 mM MgCl₂; 45.7 mM TEA-Cl; 10 mM HEPES; 5 mM 4-AP; adjusted to pH 7.9 with TEA-OH. I_{Ca} in the presynaptic VD4 cells were elicited by depolarizing the cells from a holding potential of −80 mV to +70 mV in 10 mV steps. The Ca²⁺ current data were analyzed using Clampfit 9.0 software (Axon Instruments) and traces were plotted using OriginPro 8.0 SRO (Northampton, MA, USA).

**Ca²⁺ imaging**

Fura-2 AM (Molecular Probes, Carlsbad, CA), a membrane permeable and ratiometric Ca²⁺ sensor, was used to determine changes in the intracellular Ca²⁺ levels. A detailed Ca²⁺ measurement procedure has been described elsewhere [52]. In brief, neurons were loaded with Fura-2 AM (10 μM) at room temperature (21–22°C) for 45 min. and were then exposed alternately to excitation wavelengths 340 and 380 nm using a high-speed wavelength switcher LAMBDA DG4 (Sutter Instruments, Novato, CA). The emitted fluorescence signal was collected.
at 510 nm by a Regiga Exi camera. Images were acquired with Northern Eclipse software running ionwave program (Empix Imaging, Canada). The free intracellular Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\(_i\)) was estimated based on values obtained with a fura-2 Ca\(^{2+}\) imaging calibration kit (F-6774, Molecular Probes) according to [53].

Immunoprecipitation
Immunoprecipitation was achieved by incubating detergent solubilized cells with the myc- monoclonal overnight at 4\(^\circ\)C, followed by protein A/G agarose for 2 hrs at 4\(^\circ\)C. Samples were washed, resuspended in 30ul of sample buffer, separated by SDS-PAGE, transferred to nitrocellulose and probed with antibodies for Western blot analysis.

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Acknowledgments
We thank Drs. Andy Braun and Gerald Zamponi for their critical review of this manuscript. The technical assistance of Wali Zaidi and Svetlana Farkas is gratefully acknowledged.

Author Contributions
Conceived and designed the experiments: NS, JEAB. Performed the experiments: FX, JN, SJG. Analyzed the data: FX, NS, JEAB. Contributed reagents/materials/analysis tools: RW, NJ, JEAB. Wrote the paper: FX, NS, JEAB. Principal Investigator: JEAB.
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