On the Mechanism of Nonspecific Inhibitors of Protein Aggregation: Dissecting the Interactions of α-Synuclein with Congo Red and Lacmoid†

Christofer Lendel,1,4 Carlos W. Bertoncini,1,4 Nunilo Cremades,4 Christopher A. Waudby,2 Michele Vendruscolo,2 Christopher M. Dobson,1,7 Dale Schenk,3 John Christodoulou,2,8 and Gergely Toth*1,8

†Department of Chemistry, University of Cambridge, Lensfield Road, Cambridge CB2 1EW, U.K., ‡Elan Pharmaceuticals, 700 Gateway Boulevard, South San Francisco, California 94080, and §Department of Structural & Molecular Biology, University College London, Gower Street, London WC1E 6BT, U.K. *C.L. and C.W.B. contributed equally to this work

Received July 26, 2009

ABSTRACT: Increasing evidence links the misfolding and aberrant self-assembly of proteins with the molecular events that underlie a range of neurodegenerative diseases, yet the mechanistical details of these processes are still poorly understood. The fact that many of these proteins are intrinsically unstructured makes it particularly challenging to develop strategies for discovering small molecule inhibitors of their aggregation. We present here a broad biophysical approach that enables us to characterize the mechanisms of interaction between α-synuclein, a protein whose aggregation is closely connected with Parkinson’s disease, and two small molecules, Congo red and Lacmoid, which inhibit its fibrilization. Both compounds are found to interact with the N-terminal and central regions of the monomeric protein although with different binding mechanisms and affinities. The differences can be attributed to the chemical nature of the compounds as well as their abilities to self-associate. We further show that α-synuclein binding and aggregation inhibition are mediated by small oligomeric species of the compounds that interact with distinct regions of the monomeric protein. These findings provide potential explanations of the nonspecific antiamyloid effect observed for these compounds as well as important mechanistical information for future drug discovery efforts targeting the misfolding and aggregation of intrinsically unstructured proteins.

A number of severe disorders, including Alzheimer’s, Parkinson’s (PD) and prion diseases and frontotemporal dementia are linked to protein misfolding and aggregation (1). The mechanisms by which proteins of wide structural diversity are transformed into morphologically similar aggregates seem to be a generic property of the peptide backbone (1). Structural transformations into fibrillar assemblies have been observed for a range of globular proteins and in particular from intrinsically unstructured polypeptides, including the amyloid β (Aβ) peptide, microtubule-associated protein tau and α-synuclein (αS) (2).

The growing class of protein misfolding diseases provides a significant opportunity and a challenge for drug discovery. Though a large number of small molecules have been shown to inhibit fibril formation in vitro, the details of their mechanisms of action are often poorly understood. Presently, no approved therapeutic agent that blocks the formation of amyloid structures exists. The fact that many protein aggregation inhibitors are reported to act on several amyloidogenic proteins (3) certainly raises questions about their specificity. Furthermore, different inhibitors seem to affect different steps along the fibril formation pathway, as recently illustrated for Aβ by Necula and co-workers (4). However, the lack of understanding of the degree of cytotoxicity of the various protein species populated along the aggregation pathways (5) suggests that an effective therapeutic strategy should maintain the proteins in their native states and thus hinder the formation of all oligomeric and prefibrillar species. In the case of globular proteins, partial unfolding has been found to trigger aggregation and, therefore, stabilization of the globular fold would provide a possible avenue for aggregation inhibition (1). In cases where the amyloidogenic precursors are intrinsically unstructured proteins, the approach for stabilizing the native state by a small molecule binder is less obvious.

Interestingly, some small molecules, for example, Congo red (CR, Figure 1a), Lacmoid (Lac, Figure 1b) and the polyphenol EGCG (6), have recently been reported to affect fibril formation of several amyloidogenic proteins, including αS and Aβ (3, 4) and to interact with the monomeric intrinsically unstructured proteins (6, 7). Such compounds thus represent exceptionally interesting model systems for characterizing the nature of protein binding and aggregation inhibition mechanisms. Lac, similarly to EGCG, contains polyphenol groups and resembles a class of compounds called phenothiazines that serves as a scaffold for several neuroleptic antipsychotic drugs such as Thorazine, Prolixin and perphenazine. CR is a histological dye frequently used to detect the presence of amyloid fibrils through a shift in absorption spectrum, induced fluorescence and the display of yellow-green birefringence under cross-polarized light (8). CR is reported to interact with a number of structured and unstructured proteins and to interfere with protein misfolding and aggregation processes (3, 9, 10). The details of the mechanism by which CR binds to proteins are not well understood, and substantial discrepancies exist in the reported effects on protein...
aggregation (8). Such discrepancies could be related to the ability of the dye to self-associate into micelle-like species (11–13) which is a behavior that appears to be characteristic of many nonspecific (promiscuous) inhibitors (14) frequently encountered among “hits” originating from high-throughput drug discovery screening. In fact, a recent report suggests that the ability of self-assembly may be a general property of small molecule inhibitors of amyloid fibril formation and that the formation of large colloidal particles is mechanistically linked to the inhibition of protein aggregation (15). Such an effect could indeed be related to the broad antiamyloid effect observed for many small molecules and increased understanding of the inhibition mechanisms of these compounds would provide important information when designing approaches for drug development targeting aggregation-related diseases.

Formulation of intracellular aggregates containing the presynaptic protein αS is a hallmark of a group of neurodegenerative disorders called α-synucleinopathies, with the most well-known variant being PD (16). αS is a 14 kDa intrinsically unstructured protein whose normal function is not yet well understood, but its overexpression and aggregation is a pathological event associated with Lewy bodies and neuritis, a hallmark of this group of neurodegenerative disorders (17). Characterization of the conformational properties of αS using NMR spectroscopy has revealed the presence of both local and long-range interactions within the fluctuating structural ensemble (18–20). Many aggregation promoting conditions, including high ionic strength, low pH or the presence of polyamines, have been shown to be accompanied by structural changes in the protein (20, 21) suggesting that the observed long-range interactions are fundamental for the definition of the native state of the protein and may provide an opportunity for the stabilization of protective native-like interactions and impair amyloid aggregation. In addition, the protein exist in a lipid-bound state in which the α-helical content is dramatically increased (18, 22, 23). Rao and co-workers recently investigated the interaction between monomeric αS and selected small molecules, including Lac and CR, using NMR and CD spectroscopy (7). However, neither the details of the protein aggregation inhibition mechanisms nor the impact of the self-association of the small molecules on the binding affinity and specificity were addressed. Such characterization is crucial to fully understand the way the investigated inhibitors modulate protein aggregation. In this study, that was conducted in parallel and independently of the work reported by Rao and collaborators, we attempt to dissect the mechanisms by which Lac and CR interact with monomeric αS.

We combine residue-specific NMR data with a wide range of complementary biophysical techniques to link the protein:ligand interactions to the inhibition of protein aggregation. Moreover, we investigate the key role that the self-association capabilities of the small molecule compounds play in their antiamyloid activities.

EXPERIMENTAL PROCEDURES

Materials. αS was expressed in Escherichia coli BL21 cells using a pT7-7 vector as described in ref 24. 15N- and 13C-15N-labeled protein was produced using M9 minimal medium supplemented with 15NH4Cl and 13C-glucose (Cambridge Isotope Labs). The protein was purified using heat treatment, ammonium sulfate precipitation, anion exchange and size exclusion chromatography as previously described (24). Purified protein samples
were dialyzed against water, flash frozen in liquid nitrogen and stored at −80 °C. Production of A24C and Q62C αS mutants has been reported elsewhere (19). Labeling with the fluorescent dye IAEDANS (Sigma) was performed following standard protocols. Briefly, 1 mg of purified cysteine-containing protein was reduced by incubation in 10 mM DTT for 30 min. The reducing agent was removed by fast desalting in PD-10 columns (GE Healthcare) and a 10:1 molar excess of the dye was immediately added. Proteins were incubated with the dye for 1 h at 4 °C, and the excess of fluorophore was removed with another fast desalting step in PD-10 columns. The yield of labeling reached 99%. Lac and CR were dissolved in buffer to prepare stock solutions of 1−50 mM. These solutions were sonicated before further dilution. Purified fraction V albumin from bovine serum (BSA, Sigma) was employed without further purification.

NMR Spectroscopy. NMR samples were prepared in 25 mM Tris buffer pH 7.4 with 100 mM NaCl and 10% 2H2O if nothing else is stated. Data were recorded on Bruker Avance 400 MHz, 500 MHz, and 700 MHz spectrometers (the latter two equipped with cryoprobes), processed using NMRPIPE (25) or TopSpin (Bruker) and analyzed in CCPNMR (26), TopSpin (Bruker) or Sparky (T. D. Goddard and D. G. Kneller, SPARKY 3, University of California, San Francisco, CA). All NMR experiments, except for the diffusion measurements, were performed at 10 °C, to avoid the enhanced line broadening at higher temperatures (27).

2D 1H−15N heteronuclear single quantum correlation (HSQC) (28) spectra were recorded with 512 × 128 complex points and spectral widths of 10 × 29.5 ppm. The HSQC spectrum of 100 μM αS was monitored during addition of 0.5 to 16 and 1 to 10 molar equiv of Lac and CR respectively. Assignment of the 1H−15N correlation spectrum of free αS has previously been reported (18, 29). The assignments of the αS spectra in the presence of Lac or CR were done by following the peaks in the correlation map during the titration.

2D 1H−15N correlation spectra of 1 μM αS with and without 4 μM CR were recorded using SOFAST-HMQC (30) with 512 × 64 complex points, 12 × 29 ppm spectral widths and 1,152 or 1,280 scans per increment.

13CO−15N correlation spectra were recorded with direct 13C detection (31, 32) using the 500 MHz spectrometer equipped with a TCI probe (Bruker). 512 × 256 complex points were recorded with spectral widths of 10 × 40 ppm (to include Pro residues) and with 16 scans per increment. Two spectra were collected and added to reduce noise. Samples contained 100 μM 15N-labeled αS with the addition of 0.9 mM CR or 0.6 mM Lac.

Pulsed field gradient (PFG) NMR experiments were acquired at 15 °C on the 700 MHz spectrometer using a 100 μM unlabeled protein sample in 50 mM phosphate buffer (pH 7.6) (uncorrected), 100 mM NaCl, in 99.9% 2H2O and containing 10 mM dioxane as an internal radius standard and viscosity probe (33). 24 1D 1H spectra were collected as a function of gradient strengths from 1.60 G cm−1 to 32.0 G cm−1, in a linear manner. Each 1H spectrum comprised 32 scans (or 128 scans in the case of CR). 8,192 complex points were acquired with a spectral width of 12 ppm. The dioxane peak and selected signals in the aromatic and aliphatic regions of the 1H protein spectrum were integrated and the decay of the signals as a function of the gradient strength was fitted to a Gaussian function using Sigma plot 7.0 to determine the hydrodynamic radii (33). The radius of hydration for the protein (R_H,prot) was calculated from the decay rates of the protein and the dioxane peaks, using the formula R_H,prot = (d_diox/d_prot) R_H,diox, whereas R_H,diox is 2.12 Å, and d_diox and d_prot are the dioxane and protein decay rates, respectively.

1D NMR spectrum of Lac were recorded at 400 MHz 1H frequency at 25 °C. The sample contained 0.5 mM Lacmoid in 25 mM Tris buffer pH 7.4 with 100 mM NaCl. 512 scans were added in the final spectrum.

Fluorescence Spectroscopy. The binding affinities of the αS:ligand complexes were determined from fluorescent quenching of 5 μM IAEDANS conjugated αS (αS-62C-AEDANS for CR and αS-24C-AEDANS for Lac) when titrated with 1 μM to 0.5 mM CR or Lac. The experiments were performed in 25 mM Tris buffer pH 7.4, 100 mM NaCl, using a Cary-Eclipse spectrophluorimeter (Varian). An excitation wavelength of 337 nm was used and the emission was measured from 400 to 600 nm. The decrease in fluorescence intensity at 480−520 nm, after subtracting contributions from the free ligand, was fitted to a one-site ligand binding model. Control experiments of free IAEDANS titrated with Lac or CR also showed ligand concentration-dependent quenching of the fluorophore. The apparent K_d values for these interactions are an order of magnitude higher than for the protein binding (19 ± 2 μM and 154 ± 6 μM for CR and Lac respectively). Fitting the protein titration data to two independent binding events using the K_d derived from the control experiments were not successful indicating that specific binding of the compounds to the attached AEDANS is not likely to occur.

Circular Dichroism Spectroscopy. CD was measured on a Chirascan spectrometer (Applied photophysics) equipped with a Peltier temperature control system. The samples contained 5 μM αS in 25 mM Tris buffer pH 7.4, 100 mM NaCl with or without 10 μM to 0.5 mM of the compounds. Samples with the same compound concentrations but without any protein were used as references. The measurements were performed at 10 °C, 25 °C and 37 °C using a cell with 1 mm path length.

Isothermal Titration Calorimetry. ITC experiments were carried out using VP-ITC titration microcalorimeters (MicroCal) at 25 °C. αS, Lac and CR were dissolved in PBS buffer pH 7.4 and the samples were degassed before the measurements. Each experiment involved a preliminary 2 μL injection followed by 25−27 injections of 10 μL using a 300 μL syringe. The cell volumes were 1.416 or 1.4242 mL. ITC raw data was analyzed using Origin 7 (OriginLab Corporation). In the CR experiments 2 μM αS (in the cell) was titrated with 0.1 mM or 0.2 mM CR. In the Lac experiments, 5 μM, and 100 μM αS (in the cell) was titrated by 0.1 mM and 1 mM Lac respectively. For all the experiments, relevant reference experiments were performed by titrating the solution in the syringe into pure buffer.

Dynamic Light Scattering. DLS was measured using an ALV/CGS-3 compact geometer system operating at 632.8 nm wavelength (ALV-GmbH). The samples were filtered through 0.2 μm filter before the measurements and the scattered light was detected at 150° angle. The samples contained 1 mM or 0.1 mM CR or 0.5−0.6 mM Lac in 25 mM Tris pH 7.4, 100 mM NaCl, and the data was recorded at 25 °C. The acquired data was analyzed by the regularization algorithm in the ALV correlator 3.0 software (ALV-GmbH). Additional data for Lac was acquired using a Zetasizer Nano ZS instrument (Malvern Instruments) operating at 532 nm and a detection angle of 173°. The correlation data was exported and analyzed using the ALV software.

Aggregation Assays. Aggregation of αS was assayed in 100 μM protein samples in 20 mM Tris buffer pH 7.4, 100 mM
NaCl with the addition of 0.01% NaN₃. 500 μL of protein sample were incubated at 37 °C under constant shaking at 300 rpm, and 50 μL aliquots were withdrawn on a daily basis, assayed for thioflavine T (ThioT) binding (24, 34) and stored at 4 °C until the end of the assay for further determinations (SDS–PAGE and electron microscopy). Fibril formation was monitored by the ThioT assay (24, 34). Briefly, 10 μL aliquots were diluted in 1 mL of 20 μM ThioT and the fluorescence was measured in a FlashScan spectrophotometer (Jena Analytik), with an excitation wavelength of 446 nm. Emission wavelengths from 460 to 600 nm were collected and the integrated fluorescence between 470 and 490 nm was employed for determination of the relative content of αS fibrils in the sample. Quenching of ThioT fluorescence by Lac and CR was initially investigated by heteronuclear 1H–15N correlation NMR spectrum of αS, observed during titrations of the protein with increasing amounts of each compound, indicate that Lac and CR bind to the protein in a concentration dependent manner (Figure 1 and Figure S1 of the Supporting Information). The observed changes are more pronounced for CR than for Lac, which suggests different affinities for the ligands. The chemical shift perturbations report on changes in the chemical environment and/or in the relative populations of different conformations in the αS structural ensemble. Both ligands affect the 100 most N-terminal residues of the protein significantly more than the C-terminus (Figure 1). Within this region, two broadly distinct binding domains can be identified with the first being approximately residues 1–40 and the second, residues 50–100. The latter domain includes the NAC region, which is believed to be the most aggregation-promoting segment of the protein (35, 36). The most N-terminal binding domain experiences a higher degree of peak intensity loss for both compounds while the region showing the largest chemical shift changes differs between the two binders; for CR, the largest shift changes are observed in the second binding domain (the region 50–77) although it cannot be excluded that even larger perturbations occur for residues in the N-terminal part that are too broad to be detected. Lac, on the other hand, mainly affects the chemical shifts of residues 1 to 22 in the N-terminal region.

Ligand Binding Causes Exchange Broadening of αS NMR Signals. The changes in peak intensity observed in the 1H–15N correlation spectrum of αS bound to CR and Lac could be due to several processes: (i) increase in the rotational and translational correlation times, e.g. by binding to large ligand aggregates or by ligand-induced oligomer formation of the protein itself; (ii) chemical exchange on the μs–ms time scale due to changes in the conformational properties of the αS polypeptide ensemble and/or interaction with the ligands; (iii) changes in amide hydrogen exchange rates.

Alterations in peak intensities resulting from hydrogen exchange can be expected to result from interactions that alter the exposure of the peptide backbone to solvent. In order to exclude this effect as a cause for the changes in peak intensity, 13C direct detected NMR experiments were performed on 13C–15N-labeled αS with and without CR and Lac (31). 13CO–15N correlation spectra showed the same profiles for intensity changes as the 1H–15N correlation (Figure 2a,b and Figure S2 of the Supporting Information) suggesting that hydrogen exchange effects are not a major cause of line broadening under these conditions. Furthermore (as described in detail below), PFG NMR experiments suggest that the line broadening observed upon ligand binding is not due to significant increases in the molecular size of the complexes, and points toward exchange broadening on a slow to intermediate time scale as the major cause for the decreased NMR peak intensity.

In order to probe whether complex formation perturbs the content of secondary structure in the ensemble of αS conformations, we employed CD spectroscopy. The far-UV CD spectrum of αS at 10 °C is in agreement with a largely unstructured protein (Figure 2c,d) as previously reported for αS (7, 21, 37). The addition of increasing amounts of CR results in ellipticity changes at 196 and 220 nm (Figure 2c), similar to those reported for other structure-promoting conditions (21, 38, 39). Addition of Lac causes changes in the 196–200 nm region of the CD spectrum, similar to those seen with CR but with lower magnitude (Figure 2d). Similar, but less significant, changes in the αS CD spectrum are observed for both compounds at higher temperatures (Figure S3 of the Supporting Information). Taken together, the NMR and CD data suggest that the ensemble of conformations populated by αS is strongly perturbed upon binding to CR but is less affected by Lac.

Lac and CR Form Supramolecular Assemblies in Solution. The ability of CR itself to form aggregates or micelle-like species in solution is well-known (11–14), and self-assembly has as well been reported for some phenothiazines similar to Lac (40). The propensities of the compounds to self-associate were therefore investigated using DLS.

In 1 mM solution, the DLS data show that CR forms two types of species (Figure 3a). The smaller of these (representing the major population) has a R₄₄ distribution centered around 1.9 nm, which is in agreement with other reports of CR self-assembly (11). Due to the elongated shape of the CR molecule this apparent R₄₄ could correspond to the monomeric compound as well as a small oligomer. The larger particles have an R₄₄ of approximately 40 nm (Figure 3a). At lower CR concentrations (0.1 mM), a similar size distribution is observed but with a slightly shifted average (~1.4 nm) of the smaller species (Figure 3a). Such concentration dependence strongly suggests an oligomeric nature of this molecular species.

The use of DLS to investigate Lac self-assembly is limited by the overlap of its absorbance spectrum with the laser wavelengths of the DLS instruments. However, measurements from two instruments, operating at different wavelengths, indicate that Lac, at 0.5 mM concentration, does indeed form supramolecular...
aggregates. The average $R_{H}$ of these species are approximately 15 and 90 nm for the two populations respectively (Figure 3b).

Opposite to CR, the larger assemblies represent the major population. The DLS data recorded for 0.1 mM Lac were
inconclusive (data not shown), most likely because of sample absorbance, and neither confirms nor excludes the presence of aggregated species.

The tendency of Lac to self-associate was further investigated by NMR spectroscopy. The intensities/integrals of the Lac signals in the 1D NMR spectrum at 0.5 mM total Lac concentration were found to be significantly lower than what would be expected for a monomeric compound of low molecular weight (Figure S4 of the Supporting Information), which, in line with the DLS data, strongly suggests the presence of a substantial population of large species. Using the high intensity Tris buffer peak (25 mM peak at 3.7 ppm) as an integration reference, the
The thermodynamic characteristics of the binding process is entropically driven with a small favorable enthalpy contribution, which is in agreement with CR binding to similar molecules binding with similar affinities in a cooperative process. The presence of small oligomeric CR species or several individual CR molecules binding with similar affinities in a cooperative process. The small magnitude of the measured heat output offers an explanation for why no interaction was detected in the low concentration experiments and suggests an entropically driven binding process, as observed for CR. The absence of signal baseline recovery at the end of the titration indicates that the binding process does not reach saturation and prohibits the

To explore further a potential 1:1 binding of CR to αS, additional NMR experiments were carried out at low micromolar concentrations. The $^1$H–$^{15}$N correlation spectrum of 1 μM αS with 4 μM CR shows substantial line broadening of the 100 most N-terminal residues, most of them beyond detection (Figure 4c). This result is closely similar to the observations of the high concentration experiments (see above) and suggests that CR interacts with monomeric αS at a low μM concentration which is below its reported critical concentration for self-association (13).

Affinities and Thermodynamics of the αS:Ligand Interactions. The thermodynamic characteristics of the binding events of CR and Lac to αS were investigated using ITC. Control titration of CR into buffer shows substantially larger endothermic heats for the first few injections compared to later ones (Figure 5a), suggesting that CR is in an aggregated state in the syringe. Such calorimetric behavior is typically observed for disaggregation of micellar surfactants (43, 44) and has previously been reported for CR (45). Similar results were observed for 1.0 mM Lac titrated into buffer (Figure 5b), which confirms the presence of oligomeric species observed by DLS. Titration of 0.1 mM Lac into buffer, however, showed exothermic heat effects as expected for a process dominated by the heat of dilution (Figure 5b). The self-association properties of Lac thus seem to change precisely in this concentration interval.

Titrations of 0.2 mM (Figure 5a) and 0.1 mM (data not shown) CR into 2 μM αS solution indicated a one-site interaction mechanism where several ligand molecules (~8) of CR bind to the protein with a $K_d$ of 0.4–1 μM. This could reflect the binding of small oligomeric CR species or several individual CR molecules binding with similar affinities in a cooperative process. The binding process is entropically driven with a small favorable enthalpy contribution, which is in agreement with CR binding to hydrophobic patches of αS.

ITC experiments with the Lac:αS system confirm the lower affinity of this compound compared to CR. Titration of 0.1 mM Lac into 5 μM αS does not result in significant heat changes (data not shown); however, small exothermic heat effects were generated when 1.0 mM Lac was titrated into 100 μM αS (Figure 5b). The small magnitude of the measured heat output offers an explanation for why no interaction was detected in the low concentration experiments and suggests an entropically driven binding process, as observed for CR. The absence of signal baseline recovery at the end of the titration indicates that the binding process does not reach saturation and prohibits the
The fluorophore was specifically attached to a cysteine residue introduced by mutagenesis in the vicinity of the NMR-derived binding regions (position 24 for Lac and 62 for CR). Fluorescence, they seem to affect the majority of the N-terminal peaks due to compound addition almost fully disappeared, and peaks along the whole sequence became visible again. This clearly shows that BSA competes with αS for the binding of both compounds in a similar way as has been reported for promiscuous inhibitors (14, 15). The results also show that the binding of the compounds to αS are reversible events.

**CR and Lac Modulate Differently the αS Aggregation Pathway.** The effect of CR and Lac on αS aggregation and fibril formation was investigated by incubating the protein with various concentrations of the ligands. Both compounds have previously been reported to inhibit the formation of amyloid fibrils by αS (3), and, as determined by reduced ThioT fluorescence, they seem to affect the αS fibril formation significantly, even at very low stoichiometric ratios (Figure 7a,b). Interestingly, however, protein aggregates were visible in most of the incubated samples. This observation could be due to the presence of nonamyloid aggregates that do not bind ThioT or to specific reduction of ThioT fluorescence by the compounds as a result of quenching or competition of binding sites in the fibrillar species. The potential influence on ThioT fluorescence by the compounds was investigated by adding dye to mature compound-free αS fibrils, preincubated with Lac or CR, and comparing the observed fluorescence with a negative (compound-free) control. The results clearly show that high concentrations of

determination of any dissociation constant or binding stoichiometry.

The relative affinities of the αS:ligand complexes were then estimated from the quenching of the fluorescence emission of AEDANS-conjugated αS upon titration with the compounds. The fluorophore was specifically attached to a cysteine residue introduced by mutagenesis in the vicinity of the NMR-derived binding regions (position 24 for Lac and 62 for CR). Fluorescently labeled proteins were titrated with the compounds and a dose-dependent reduction of fluorescence intensity was observed (Figure 6). Fitting the data to a single site binding model resulted in apparent $K_d$ for Lac. The observed lower binding affinity of Lac compared to CR is in good agreement with NMR experiments described above.

**Specificity of CR and Lac Interactions with αS.** In order to evaluate the specificity of the compounds for αS, we studied the binding behavior of β-synuclein (βS), a closely related protein which does not aggregate under pathological conditions. βS also lacks a defined secondary structure in solution and populates a highly dynamic ensemble of conformations (42), however it does not readily fibrillate due to the absence of a stretch of highly hydrophobic amino acids in the central region of the protein (46). Employing $^1H-^15N$ correlation NMR experiments we found that βS is able to bind both CR and Lac in a similar fashion to αS (Figure S6 of the Supporting Information), suggesting that the compounds do not discriminate between amyloidogenic and nonamyloidogenic conformers.

As an additional control for the specificity of the binding mechanisms of the compounds, the heteronuclear NMR experiments were repeated in presence of bovine serum albumin (BSA). This test is commonly used for determining the promiscuity of binding by small molecule ligand (14, 15). Addition of 25 mg/mL (8:1 BSA:αS molar ratio) BSA to αS samples containing CR or Lac resulted in a dramatic change of the NMR resonances (Figure S6 of the Supporting Information). The line broadening observed in the majority of the N-terminal peaks due to compound addition almost fully disappeared, and peaks along the whole sequence became visible again. This clearly shows that BSA competes with αS for the binding of both compounds in a similar way as has been reported for promiscuous inhibitors (14, 15). The results also show that the binding of the compounds to αS are reversible events.

**Figure 5:** Calorimetric characterization of the interactions between the small molecules and αS. Baseline corrected ITC raw data (top) and integrated heat data (bottom) of CR and Lac titrated into αS solution and pure PBS buffer at 25 °C. (a) 0.2 mM CR titrated into buffer (red) and 2 μM αS (black). The best fit of the integrated data to a one-site model is shown as a black line and corresponds to 8 CR molecules binding with an $K_d$ of 0.4 μM. (b) 1 mM Lac titrated into buffer (red) or 100 μM αS (black). The baseline corrected raw data of 0.1 mM Lac titrated into buffer is also included (blue line at −0.25 μcal/sec for better comparison) to show the concentration dependence of the heat of dilution.
While Lac addition does not show differences with gates but not mature amyloid fibrils, shows an increased amount of Sarkosyl, a detergent capable of solubilizing amorphous aggregates (Figure 7e,f). Only in samples containing 5:1 molar excess of insoluble protein aggregates in all the conditions studied, the amount of soluble protein in the samples was evaluated at different time points of the aggregation assays by centrifuging an aliquot at 16000 g. Quenching data fitted to a single-site binding model yields apparent $K_d$ values of $1.05 \pm 0.05 \mu M$ for CR (a) and $12 \pm 2 \mu M$ for Lac (b). The fluorescence raw data in the absence (top line) and presence of 2.5 $\mu M$ CR (a) and 1 $\mu M$ to 0.5 mM Lac (b) are shown as insets.

The compounds alter the ThioT fluorescence of the fibrils (Figure 7c,d). However, the influence on ThioT fluorescence of low concentrations of the compounds is not significant. (Figure 7c,d). However, the influence on ThioT fluorescence of low concentrations of the compounds is not significant. To verify further the presence of aggregated protein material, the amount of soluble protein in the samples was evaluated at different time points of the aggregation assays by centrifuging an aliquot at 16000 g and analyzing the protein remaining in the supernatant using SDS–PAGE. The results confirm the presence of insoluble protein aggregates in all the conditions studied (Figure 7e,f). Only in samples containing 5:1 molar excess of Lac does $\alpha$S remain in a monomeric soluble form until the end of the assay. Indeed, treatment of aggregated protein with 1% Sarkosyl, a detergent capable of solubilizing amorphous aggregates but not mature amyloid fibrils, shows an increased amount of soluble protein both in control experiments and with the compounds. While Lac addition does not show differences with the control at concentrations of 50 $\mu M$ or lower, CR increases the extent of Sarkosyl soluble protein even at low concentrations (Figure 7e,f).

Such observations are corroborated by TEM of the aggregated samples at the end of the assays. In the control experiment (i.e., no added ligands) $\alpha$S formed fibrils as expected (Figure 7h) (21, 36). Incubation with low concentrations of Lac does not, under the applied experimental conditions, distinctively change the morphology of the fibrillar material (Figure 7h). However, incubation with 10 $\mu M$ CR produces amorphous aggregates as well as fibrillar species (Figure 7g). At higher CR concentrations, some fibrils are visible but amorphous aggregates are still the dominating morphology (Figure 7g). Interestingly, in the samples incubated with 0.5 mM Lac, no aggregates are visible (Figure 7h), confirming its ability to reduce the overall protein insolubilization.

**DISCUSSION**

A variety of compounds have been reported to interact with $\alpha$S and inhibit its aggregation and/or fibril formation (3, 6, 47–52). However, the molecular mechanisms for their action are, in most cases, only vaguely understood. We have conducted an extensive characterization of the protein:ligand interactions at a residue-specific resolution, together with a careful description of the effects on $\alpha$S aggregation, in an attempt to provide a picture of the inhibition mechanisms for CR and Lac.

The heteronuclear NMR data show that CR and Lac mainly interact with the N-terminal and central regions of $\alpha$S. Of the two compounds, CR binds more strongly and has a more pronounced affinity for the aggregation-prone NAC region, features that might be related to the presence of negative charges (7). Moreover, CR and Lac interactions differently perturb the ensemble of structures populated by $\alpha$S, as shown by NMR and CD data. In both cases NMR resonance intensities are lost due to line broadening, mainly caused by conformational exchange on a slow-intermediate time scale. For CR, the observed changes in secondary structure content are accompanied by a reduction in the hydrodynamic radius of the $\alpha$S complex, while Lac has less effect on the secondary structure content and the hydrodynamic radius. These observations could, in principle, be interpreted in terms of the accumulation of specific intermediate states, as previously suggested for other effectors of $\alpha$S aggregation (21), but it is more likely that they represent changes in the population distribution across the conformational energy landscape of the protein.

The self-association characteristic of the compounds indeed complicates the analysis of the binding mechanisms. Based on the diffusion measurements, interactions between monomeric $\alpha$S and large colloidal species can be excluded, which leaves the options that the interaction is mediated by monomeric or small oligomeric compound species. Our findings point to small oligomeric compound species as the main effectors; however, weak 1:1 interactions cannot be excluded. The presence of small oligomeric CR species is supported by DLS, and the ITC data imply a mechanism where several CR molecules bind to each $\alpha$S molecule. The apparent 1:1 binding between the compounds and $\alpha$S observed with fluorescence quenching experiments could actually be a 1:n binding if the compounds bind as an oligomer to a single site on the protein. Previous studies reported the binding of several CR molecules to different types of proteins (11, 53, 54) indicating that multivalent binding is characteristic of CR regardless of the nature of the specific protein involved. The critical concentration of CR needed to form aggregates has been reported to be approximately 5 $\mu M$ (13), suggesting that the binding observed by NMR at low micromolar CR concentrations might represent a 1:1 interaction. However, the self-association ability is most likely modulated by factors such as pH, ionic strength and temperature which could lower the critical concentration for CR to form soluble aggregates. In addition, the presence of $\alpha$S itself could affect this concentration, as shown for certain aggregation promoting detergents (55).

In the case of Lac, the absence of binding site saturation in the ITC prevents the determination of the stoichiometry. Such behavior could indicate a mechanism of binding involving several...
Lac molecules per αS molecule, or a competition between ligand:ligand and ligand:protein interactions, reducing the effective concentration of binding competent species for the latter. The reference ITC experiments indicate a change of the self-association properties of Lac in the range 0.1–0.5 mM, and the critical aggregation concentrations have indeed been reported to be in the mM range for other phenothiazine compounds (40). Hence, besides the intrinsic chemical properties of the compounds, the observed difference in binding affinity could be related to the difference in critical concentrations of aggregation as well as the propensities to form oligomeric species of appropriate size and structure.

The apparent $K_d$ values for the CR and Lac complexes with αS are in the low micromolar region, which represent binding of

![Figure 7](image-url)
moderate strength. Such interactions are expected to be easily competed out, which indeed is observed by addition of BSA. Albumin is well-known to bind a variety of different small molecules, such as negatively charged compounds (56), and has been shown to interact with small oligomers of CR (53). Our results strongly suggest that BSA interacts with both CR and Lac by binding to either or both monomeric and small oligomeric species. Previous studies of CR binding to globular proteins, such as immunoglobulin light chain variable domain (45) and human growth hormone (9), have also reported $K_d$ values in the micromolar range.

The differences in the interaction mechanisms for the two compounds are also reflected in their anti-amyloid behavior. For CR we observe an increased proportion of amorphous aggregates at very low concentrations, accompanied with a more rapid loss of soluble material compared to the reference experiment (Figure 7). On the basis of these observations, we hypothesize that binding of CR to $\alphaS$ induces conformational changes that weaken the (protective) interactions of the native state allowing the formation of new intermolecular interactions between the N-terminal and C-terminal part of the protein. These events direct the protein into an alternative pathway leading to amorphous aggregates that might be kinetically more favorable compared to fibrils. The interaction with the NAC region might also impair with the formation of amyloid-like structures. Moreover, the fact that CR has high affinity for amyloid fibrils could render early amyloid nuclei incompatible for elongation, stabilizing oligomers and amorphous aggregates. Lac, on the other hand, seems to stabilize a native-like state with the concomitant protection of the NAC region of $\alphaS$ and thereby inhibit self-assembly. The effect is, however, only observed at high concentrations of the compound, which could be related to the lower self-association ability compared to CR. Another source for the different inhibitory effects could be the charge differences between the compounds. Negatively charged surfaces have been suggested to promote protein aggregation (57), and anionic detergents have indeed been shown to accelerate $\alphaS$ self-association while uncharged detergent did not have this effect (53). The effect on $\alphaS$ aggregation observed for CR might thus be a rather complex process resulting from a combination of aggregation promoting and inhibitory effects.

Interestingly, CR and Lac also differ in the mechanisms by which they inhibit aggregation of the $\alphaB$ peptide (4). In agreement with our results, Lac inhibits the formation of all types of $\alphaB$ aggregates, CR, on the other hand, appeared to inhibit formation of oligomers but not fibrils. During the preparation of this manuscript, Rao and co-workers (7) presented an NMR and CD study of the CR and Lac interactions with $\alphaS$. Our results agree well with theirs regarding that the compounds interact with monomeric $\alphaS$ and the observed binding sites on $\alphaS$. However, a broad multidisciplinary experimental strategy allows us to characterize the differences between the compounds with respect to the binding mechanisms, affinities and effects on amyloid formation. Furthermore, our detailed investigation establishes the critical role of the self-association of the small molecules for their interaction with $\alphaS$.

Recently it was suggested that self-association might be a common property among aggregation inhibitors and that the inhibition would be a result of sequestering the protein molecules to large colloidal particles (14). Our data confirm that both CR and Lac are able to self-associate into such particles. However, the binding of the compounds to monomeric $\alphaS$ proceeds via interactions with small oligomeric species. Such species could be pre-existing or formed as a consequence of binding to the protein. In both cases, the self-association abilities of the compounds are of critical importance. This type of compounds, especially CR, has been reported to interact with a wide range of proteins, which is indeed confirmed by the observed BSA competition. However, their mechanisms of action might be of particular importance when binding to intrinsically unstructured proteins because of their ability to provide a large interaction surface that could compensate for the entropic cost of binding. This might be an explanation for their frequent occurrence as aggregation inhibitors. The full impact of these findings remains to be investigated, but it is indeed notable that self-associating small molecules have shown anti-amyloid effects in vivo (8, 15, 58, 39), and it has been suggested that the self-association properties could also be of relevance for oral drug delivery (60).

CONCLUSIONS

Our investigation illustrates well the complexity associated with the discovery and in vitro characterization of aggregation inhibition mechanisms of small molecules. We show that a broad biophysical approach is needed to obtain a complete characterization of such systems and to avoid false conclusions based on experimental artifacts such as low specificity or quenching of ThioT fluorescence by the tested compounds. This approach allows us to describe the mechanisms by which CR and Lac affect the amyloid fibril formation of $\alphaS$. Our experiments show that CR and Lac bind to monomeric $\alphaS$ as small oligomeric species, although to alternate regions and with different affinities, and that these binding events are directly responsible for affecting the fibril formation of $\alphaS$ in distinct ways. These differences can be attributed to the chemical nature of the compounds as well as their abilities to self-associate. Our results differ from reported cases in which such compounds inhibit the biochemical function of enzymes and the aggregation of other misfolding proteins in a promiscuous fashion by forming macromassemblies (14). These findings provide potential explanations for the nonspecific anti-amyloid effect observed for these compounds as well as important mechanistical information for future drug discovery efforts targeting the misfolding and aggregation of intrinsically unstructured proteins.

ACKNOWLEDGMENT

We thank Prof. Astrid Gräslund (Stockholm University) for the use of DLS, CD, and NMR equipment; Dr. Verna Frasca (MicroCal), Louise Creagh (University of British Columbia) and Dr. Terence Hui (Elan Pharmaceuticals) for their assistance with the ITC experiments; Dr. Don Walker (Elan Pharmaceuticals) for LC/MS analyzes; Dr. Shang-Te D. Hsu (University of Cambridge) for the setup of $^{13}$C detected NMR experiments; Dr. Rick Arthis (Elan Pharmaceuticals) for critical review of the manuscript and for his suggestions; and the staff of the Biomolecular NMR facility at the Department of Chemistry, University of Cambridge.

SUPPORTING INFORMATION AVAILABLE

Comparison of $^1$H–$^{15}$N correlation NMR spectra of $\alphaS$ in presence and absence of the investigated compounds (Figure S1), comparison of $^{13}$CO–$^{15}$N correlation NMR spectra of $\alphaS$ in presence and absence of Lac (Figure S2), CD spectra of $\alphaS$ in...
presence and absence of the investigated compounds at three different temperatures (Figure S3), 1D NMR spectrum of Lac (Figure S4), detailed analysis of pulsed field gradient NMR data measured for $\alpha S$ (Figure S5), and binding of the compounds to $\beta S$ and the effect of BSA on the binding of the compounds to $\alpha S$ investigated using $^{1}H-^{15}N$ correlation NMR (Figure S6). This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES


