# Identification of Fibril-Like Tertiary Contacts in Soluble Monomeric $\alpha$ -Synuclein

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ABSTRACT Structural conversion of the presynaptic, intrinsically disordered protein  $\alpha$ -synuclein into amyloid fibrils underlies neurotoxicity in Parkinson's disease. The detailed mechanism by which this conversion occurs is largely unknown. Here, we identify a discrete pattern of transient tertiary interactions in monomeric  $\alpha$ -synuclein involving amino acid residues that are, in the fibrillar state, part of  $\beta$ -strands. Importantly, this pattern of pairwise interactions does not correspond to that found in the amyloid state. A redistribution of this network of fibril-like contacts must precede aggregation into the amyloid structure.

# INTRODUCTION

In many neurodegenerative amyloid-linked human conditions, such as Alzheimer's and Parkinson's diseases, fibril formation proceeds via crucial steps that involve monomer to oligomer transition, oligomer maturation, and amyloid formation (1). Recent studies suggest that a complex structural reorganization of the implicated proteins along this process governs neurotoxicity (2-4). However, the precise nature of this conformational rearrangement remains elusive. In this work, we sought to shed light on the early stages of the amyloid pathway of the intrinsically disordered protein (IDP)  $\alpha$ -synuclein ( $\alpha$ S), a presynaptic protein involved in Parkinson's disease (5). Recent evidence indicates that in vivo  $\alpha$ S may coexist between an aggregationprone monomeric disordered state, an aggregation-resistant tetrameric helix-rich conformation, a helical vesicle-bound state, and different  $\beta$ -sheet-rich species of prefibrillar and fibrillar nature (6–9). Although the structure of  $\alpha S$  fibrils has not been unambiguously determined at atomic resolution due to fibril polymorphism (10), the most accepted model suggests that in its fibrillar state  $\alpha$ S monomers adopt an antiparallel in-register five  $\beta$ -strand fold ( $\beta$ 1 to  $\beta$ 5) with monomer units stacked in a parallel fashion forming the fibril protofilament (11). Such structural plasticity has stimulated research toward the structural characterization of  $\alpha S$ , which has now become the prototypical example of an IDP.

Despite being largely disordered in solution, monomeric  $\alpha$ S is known to populate collapsed conformations (12–14), which are associated with an increase in the rate of protein aggregation (12,14). Several authors have identified the presence of long-range interactions in the protein that put in contact distant regions of the protein (15–20). Studies of the protein in aggregation-prone conditions suggest that these contacts may play a key role in the control of self-oligomerization

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(15,20,21). Tertiary contacts are thus a fundamental structural feature of this protein that underlies conversion of  $\alpha$ S into a neurotoxic, likely  $\beta$ -sheet rich, conformation. NMR studies of soluble  $\alpha$ S in supercooling conditions have demonstrated that the monomeric protein transiently populates  $\beta$ -strandrich conformations within its fibril-forming domain, as judged by negative secondary C $\alpha$  and C' deviations as well as positive secondary 3J(HNH $\alpha$ ) couplings (22). No transient tertiary interactions between residues forming  $\beta$ -strands in the amyloid state have yet been described in the monomeric state of the protein; however, close examination of NMR (15–17), single-molecule Förster Resonance Energy Transfer (FRET) (18,19), and electron transfer kinetics data (20) suggest a more complex pattern of transient tertiary interactions in solution than the one identified so far (Fig. 1, *a* and *b*).

Here, we mapped at high resolution this network of transient tertiary interactions present in the soluble monomeric state of  $\alpha$ S and found a fingerprint of its amyloid fibrillar structure. To calculate a high-resolution interresidue contact map of  $\alpha$ S, we employed an extensive set of intramolecular paramagnetic relaxation enhancement (PRE) data for the monomeric state of the protein (15, 16, 23, 24). PREs are an extremely powerful NMR parameter for detecting longrange transiently formed contacts in IDPs due to the highly nonlinear distance-dependent enhancement of relaxation rates in nuclei induced by the presence of paramagnetic spin probes (25). Our contact map of  $\alpha$ S provides novel, to our knowledge, insights into the relationship between the structural properties of amyloid fibrils and those of their monomeric precursors as well as on the conformational changes that occur during amyloid formation.

#### MATERIALS AND METHODS

## Intramolecular PRE data

PRE-derived interresidue distance restraints for  $\alpha$ S at pH 7.4 were collected from works published by leading laboratories in the study of the structural

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FIGURE 1 High-resolution mapping of tertiary contacts in monomeric  $\alpha$ S. (a) Spin labeling of single Cys-containing  $\alpha$ S variants with MTSL leads to mapping of tertiary contacts by NMR. Nuclei within a distance of up to 25 Å from the PRE label suffer attenuation of its NMR intensity, which is proportional to r<sup>-6</sup>. Positions of PRE labels used in the structure calculations are shown in red. (b) Left panel: Experimental PRE data (red) and fitted PRE data (black) are shown for a representative position of the MTSL label (76-MTSL). The expected profile for a random coil is shown in gray. Tertiary contacts are highlighted by arrowed dashed lines. Right panel: Results of the structure calculation procedure that best fit the data. Forty independent calculations were performed and the structures were pooled before contact map analysis. (c) Interresidue  $C\alpha$ - $C\alpha$  contact map resulting from the structure calculations. Contact fraction denotes the number of conformers from the pool that contain a given contact. Contacts previously reported in the literature are highlighted as Known contacts (15,16,20). Contacts discovered in this work are highlighted as Novel contacts.

properties of this protein (see Table S1 in the Supporting Material) (15– 17,23). In total, we analyzed 1104 intensity ratios arising from 11 spin probes (MTSL: (S-(2,2,5,5-tetramethyl-2,5-dihydro-1H-pyrrol-3-yl)methylmethanesulfonothioate) located at various positions along the polypeptide chain (amino acids 18, 20, 24, 42, 62, 76, 85, 87, 90, 103, and 140). Under these experimental conditions of low temperature and low concentration,  $\alpha$ S remains monomeric and no oligomerization is detectable by any ensemble spectroscopic method or by control intermolecular PREs experiments employing mixtures of PRE-tagged <sup>14</sup>N protein and untagged <sup>15</sup>N protein (16). Because of this experimental setup, we can be confident that PRE effects arise predominantly from intramolecular interactions, however we cannot completely rule out a marginal contribution of intermolecular nature arising from transient low order oligomeric species (26) (see discussion in the Supporting Material). A scaling factor had to be introduced in the case of some PRE labels to compensate for slight variations in experimental conditions between data sets, temperature, and reduction of the MTSL label. Scaling factors were determined by normalization of each data set to their respective average values. Scaling factors applied were 1.17 for labels at positions at 18, 90, and 140 (15); 1.25 for labels at positions 20 and 85 (17); 1.0 for positions 24, 42, 62, 87, and 103 (16); and 1.0 for the label at position 76 (23).

### Determination of contacts maps

Structure calculations were performed using full-atom Monte Carlo (MC) simulations method guided by PRE experimental restraints (27) (see extended details in the Supporting Material, Supplementary data). The calculation method was benchmarked with synthetic PRE data generated for the protein  $\alpha$ S, as well as other intrinsically disordered and chemically denatured proteins (27). PRE intensity ratios on HN atoms (intensity ratio between <sup>1</sup>H-<sup>15</sup>N amide crosspeaks in oxidized and reduced states of the spin label) were calculated at each MC move. The MTSL spin label was explicitly built and rotated at each step to sample the accessible conformational space. Calculations were performed for ensembles sizes of 1, 2, 5, 10, 20, and 50 conformers (see Fig. S4). The calculation procedure was repeated 40 times for each ensemble size and structures were pooled before contact map analysis. For ensemble size 1 (7.8 restrains/amino acid), data were fitted within experimental error ( $R^2 = 0.96$ , root mean-square deviation (rmsd) = 0.09). Only a marginal increment in the goodness of the fit was obtained for ensemble sizes larger than 1 ( $\Delta rmsd = 0.01$ ). As shown before, due to the  $r^{-6}$  weighted averaging of the PRE effect, employing a limited number of conformers to satisfy the PRE distance restraints avoids the presence of unrestrained conformers (28). Energy minimization of the structures determined using the AMBER force field in GROMACS (29) left the calculated structures virtually unchanged. C $\alpha$ -C $\alpha$  contact maps were generated by pooling 40 structures obtained from independent runs, which formed a well-defined cluster of conformations. A contact was defined as a C $\alpha$ -C $\alpha$  interresidue distance up to 20 Å (range of PRE action). The contacts reported for the fibril correspond to ssNMR data previously reported (11). This data considers contacts between residues within 6 Å, which have been extracted from <sup>13</sup>C-<sup>13</sup>C proton-driven spin diffusion spectra (experiment with a mixing time of 250 ms).

# Physicochemical characteristics of the primary sequence of $\alpha$ S

Data were generated using the Expasy protscale server (http://web.expasy. org/cgi-bin/protscale/protscale.pl). Hydrophobicity at pH 7.5 obeys the scales reported by Cowan et al (30). A sliding average window of seven residues was employed. The profile of predicted amyloid aggregation follows the commonly used algorithms WALTZ (high sensitivity filter) and Zyggregator (Zagg) (31,32).

# RESULTS

As a starting point in our calculations, we collected a large set of intramolecular PRE data from 11 paramagnetic labels previously measured for monomeric  $\alpha$ S under conditions that minimize the presence of oligomeric species (pH 7.4, 100  $\mu$ M protein concentration, temperature of 10 to 15°C). To ensure consistency between the data sets we implemented a scaling factor (Table S1). Thus, our calculation strategy employs a total of 1104 distance restraints, two to three times the number of PRE restraints used in previous analysis of the protein (15,16,23,24). The interresidue contact map was derived by fitting the PRE data to structural models by using a full-atom MC structure calculation method that we recently developed (27) (Fig. 1 *c*, Fig. S1, and Supplementary data). It is important to clarify that  $\alpha$ S populates an ensemble of rapidly interconverting conformations in solution and, therefore, the interresidue contact map is a projection of all tertiary interactions, populated up to a few percent by the disordered state of the protein, which may not be simultaneous.

The interresidue contact map we determined for  $\alpha$ S highlights with unprecedented detail previously known longrange interactions between the C-terminus of the protein, with both the N-terminal and central parts of the protein, termed NAC (Fig. 1 *c*, *Known contacts* and Fig. S3) (15,16,20). We find that the C-terminal region possesses a strong avidity for establishing interactions: the region <sub>124</sub> AYEMPSEEGYQDYEP<sub>138</sub> contacts residues <sub>1</sub>MVDFM KGLMK<sub>10</sub>, whereas the region comprising residues <sub>104</sub> EEGAPQEGILEDMPVDPNEAYEMPSE<sub>130</sub> interacts with three segments involving residues <sub>38</sub>LYVGSKTKEGVV HGVATVA<sub>56</sub>, <sub>74</sub>VTAVAQK<sub>80</sub>, and <sub>90</sub>AATGFVKKDQL<sub>100</sub>. The charged nature of these regions underlies their contact tendency.

Due to the high resolution achieved in our calculations, a number of short-range contacts could also be resolved indicating a strong degree of collapse present at both the N- and C-terminal regions of the protein (Fig. 1 c). At the N-terminus, the collapsed state spans up to residue 30, which could reflect the population of conformations with nascent helical content because this region folds into an  $\alpha$ -helix upon oligomerization or in the presence of phospholipids (33). Indeed, studies of <sup>15</sup>N relaxation and secondary chemical shifts indicate partial population of helical conformations by this domain in the momomeric state of the protein (34,35). The collapse at the C-terminus is centered on residue 120, the region with the highest degree of residual structure in the protein, as judged by unusually high NMR residual dipolar couplings (15), and being also a major metal binding site (36).

Most notably, our high-resolution contact map reveals an up to now unreported pattern of transient tertiary interactions in  $\alpha$ S (Fig. 1 *c*, *Novel contacts*, Fig. S3). A comparative analysis with ssNMR data measured for  $\alpha$ S fibrils indicates that these contacts involve amino acids that form the  $\beta$ -strands within the amyloid fibril core of  $\alpha$ S ( $\beta$ 1 to  $\beta$ 5), therefore we termed these fibril-like contacts (Fig. 2, *a* and *b*). In Fig. 2 we compare the tertiary contacts found in solution with contacts derived from the ssNMR model of the fibril. According to ssNMR, H/D exchange mass spectrometry and electron paramagnetic resonance data,



FIGURE 2 Fibril-like contacts are present in soluble monomeric  $\alpha S$ . (*a*) Overlay of contacts determined by ssNMR in the amyloid core of  $\alpha S$  fibrils (*black dots*) with fibril-like contacts described for soluble  $\alpha S$  in this work.  $\beta$ -strands found in the fibrillar core are depicted as grayscale arrows numbered 1 to 5. (*b*) Frequency of contacts established by residues within the amyloid core of  $\alpha S$  in its soluble state (*red line*) and in the fibril (*black line*). Contacts established by residues of the region 30–100 are considered in this analysis.

the amyloid core of the fibril comprises, approximately, residues 30 to 100 (11,37,38). Therefore, we decided to focus our analysis on this stretch of residues. From this figure it is noteworthy that, although some of these fibril-like interactions correlate directly with contacts established in the amyloid fold of  $\alpha$ S, referred to here as *fibrillar contacts*, the remaining establish a mismatched pattern of pairwise interactions with respect to the arrangement found in its amyloid state, referred to herein as nonfibrillar contacts (Fig. 2 a, Fig. 3 a). Among the fibrillar contacts, we were able to resolve tertiary contacts between  $\beta 1 - \beta 2$  and  $\beta 4 - \beta 5$ , two fibril-like interactions that could seed amyloid conversion. We cannot exclude contacts between  $\beta 2 - \beta 3 - \beta 4$ , which are likely to occur, but could not be resolved due to the shortrange nature of these contacts and the presence of overlapping interactions.

As mentioned, we also detected in monomeric  $\alpha$ S a pattern of nonfibrillar contacts between residues that are, in the fibrillar state, part of  $\beta$ -strands. The most remarkable of such interactions involves the stretch of residues  $_{35}$ EGVLYV<sub>40</sub> contacting residues  $_{70}$ VVTGVTAVA<sub>78</sub>. These two regions correspond to  $\beta$ 1 and  $\beta$ 4 strands of the amyloid fibril (Fig. 2 *a*, Fig. 3 *a*) and contain Tyr<sub>39</sub> and the stretch of hydrophobic residues 71–82, both known to be critical for fibril formation (39,40). We also observed



FIGURE 3 Reorganization of contacts between  $\beta$ -strands during aggregation of  $\alpha$ S. (a) Overview of contacts described for soluble  $\alpha$ S in this work. The regions involved in the establishment of each particular contact are identified with green labels if they are off the aggregation pathway (Nterminus, NAC, C-terminus) or with red if they are on the aggregation pathway (strands  $\beta$ 1 to  $\beta$ 5). (b) Fibril-like contacts are established between hydrophobic regions predicted as highly amyloid prone. Predictions for  $\alpha S$ aggregation computed by WALTZ (amyloid-forming regions, blue line) and Zyggregator (Zagg, amyloid aggregation propensity, red line). Hydrophobicity is shown as a black dashed line. (c) Our high-resolution contact map for monomeric  $\alpha S$  suggests that residues within the fibril-forming domain of the protein adopt a different spatial arrangement in solution than in the fibrillar state, demanding structural reorganization of contacts between residues in  $\beta$ -strands before amyloid formation. Such reorganization would probably take place at the level of oligomeric and prefibrillar species.

nonfibrillar contacts between regions involving residues that correspond to strands  $\beta 1-\beta 3$ ,  $\beta 2-\beta 4$ , and  $\beta 2-\beta 5$  in the amyloid fibril. Nonfibrillar interactions established by these residues are mismatched contacts that would require reorganization before fibril formation, similar to nonnative interactions that modulate the folding of certain globular proteins (41–45).

A close analysis of the nature of the regions involved in establishing fibril-like contacts in soluble monomeric  $\alpha$ S indicates that these interactions are of hydrophobic nature and engage residues commonly found in intermolecular complexes involving IDPs (46). Note that these hydrophobic regions form a discrete pattern of contacts instead of an all-to-all interacting pattern, suggesting a complex hierarchy in the establishment of tertiary interactions. Fibrillike contacts appear to be modulated by amino acid regions with a high propensity to aggregate into amyloid fibrils, according to the predictions made by two commonly used algorithms (31,32) (Fig. 3 *b*), and further validated by experimental data (39,40).

# DISCUSSION

The investigation of aggregation-prone conformations populated in the monomeric ensemble of amyloidogenic IDPs such as  $\alpha$ S, Tau, or A $\beta$  aims to identify structural elements that could initiate protein misfolding and aggregation in pathologies such as Alzheimer's and Parkinson's. During the last years a large number of research groups have implemented NMR and computational methodologies to characterize such heterogeneous conformations with atomic detail (47). In the absence of persistent secondary structure, one of the primary elements that restrict the conformations populated by IDPs is tertiary interactions, which are detected by NMR methodologies such as PREs (48). With the aim of uncovering low populated contacts that were previously not identified in  $\alpha$ S, in this work we characterized tertiary contacts present in the monomeric state of the protein that are consistent with a large set of intramolecular PRE data. Indeed, our contact map allowed us to describe for the first time, to our knowledge, tertiary interactions present in the soluble state of  $\alpha S$  that arise from regions of sequence involved in the cross- $\beta$  core of the fibrils, but that do not correspond to those found in the amyloid state (Fig. 3). Although we positively identified fibrillar contacts between residues forming  $\beta 1 - \beta 2$  and  $\beta 4 - \beta 5$  strands, we also found nonfibrillar contacts between residues embedded in  $\beta_{1-\beta_{4}}$ ,  $\beta_{1-\beta_{3}}$ ,  $\beta_{2-\beta_{4}}$ , and  $\beta_{2-\beta_{5}}$  strands of the amyloid fibril. Importantly, close examination of the PRE profiles shows that all of these interactions are indeed present in the experimental data and are not a consequence of the structure calculation protocol (Fig. S3).

These findings have mechanistic implications for the current knowledge of the fibrillation pathway of  $\alpha$ S. Intramolecular long-range N/C-terminal tertiary interactions in the soluble state of  $\alpha$ S have been proposed to maintain the monomeric protein in an autoinhibited, less aggregation-prone conformation (15,16,20). These contacts thus lie off the pathway of amyloid aggregation (Fig. 3 *a*). The transient tertiary contacts between hydrophobic regions participating in the fibril core of the protein would, by contrast, direct the formation of early aggregation-prone species (Fig. 3 *a*). Conceptually related to a folding nucleus, these contacts would lead to the buildup of metastable species rich in cross- $\beta$  structure but with a network of interresidue interactions different to that of mature fibrils (3,11).

According to our model, initial oligomerization would then be followed by a step where such network reorganizes to lead to the thermodynamically stable 5  $\beta$ -strand fold of the amyloid fibril. Spectroscopy studies indicate that initial steps of  $\alpha$ S aggregation give rise to prefibrillar oligometic species that have different  $\beta$ -sheet signatures than mature amyloid fibrils, as judged by ssNMR and FTIR (49,50). In addition, single-molecule FRET studies showed that the structural conversion between oligometric forms of  $\alpha$ S into  $\beta$ -sheet-rich oligomers is remarkably slow and progresses through a rough energy landscape as a result of multiple possibilities of interstrand interactions (3). All this experimental evidence is consistent with our finding that structural signatures of soluble and fibrillar  $\alpha S$  species would have a different  $\beta$ -strand arrangement (Fig. 3 c). In the broad context of protein misfolding and aggregation, our data suggests that on-pathway oligomeric intermediates may be stabilized by nonfibrillar contacts akin to the nonnative interactions that stabilize certain folding intermediates in globular proteins (41–45). Such nonfibrillar contacts sustain the idea of a rugged energy landscape in the misfolding and aggregation of proteins (51). It could be of particular interest to unravel this proposed mechanism experimentally by, for example, enforcing some of these contacts by covalent linkage of Cys-containing mutants of  $\alpha$ S. Such a procedure was recently applied to the  $A\beta$  peptide and yielded novel insights into the earliest stages of aggregation (52). In an analogous fashion it should be possible to isolate and characterize the transient oligomeric intermediates formed by  $\alpha$ S by stabilizing the nonfibrillar contacts that we discovered in this work.

We note that antiamyloidogenic agents that target monomeric  $\alpha$ S have been recently shown to interact with Tyr<sub>39</sub>  $(\beta 1)$  or key hydrophobic residues in the central part of the protein ( $\beta$ 4 and  $\beta$ 5) (39,53–55). Mostly of polyphenolic nature, these agents establish  $\pi$ - $\pi$  stacking interactions with the side chain of Tyr<sub>39</sub> and interact in a rather unspecific manner with the stretch of hydrophobic residues comprising amino acids 70-90. In light of our data, it is then possible that such small molecules prevent crucial structural rearrangements to take place at the level of the monomeric protein by inhibiting some of the long-range fibrillar contacts established between residues that compose the  $\beta$ -strand regions of the fibril. Identification of such regions that are necessarily brought into spatial proximity to set off protein aggregation should facilitate the design of novel, to our knowledge, small molecules that could modulate fibril formation.

The intra- versus intermolecular nature of tertiary N/C interactions provides an increased layer of complexity in the aggregation pathway of  $\alpha$ S. Here, we mapped with high resolution the intramolecular N/C-terminal interactions occurring at the level of the monomeric protein. Notably, intermolecular PRE studies of partially oligomeric  $\alpha$ S (1.1 mM protein concentration) showed transient interchain

N/C-terminal interactions involving residues 1–15/120–140 and 35-45/120-140, precisely two of the main interactions detected in our contact map of the monomeric protein (26). In this regard, it was previously shown that ligands that bind the C-terminus trigger protein aggregation by release of the intramolecular N/C interactions (15), and more recently fluorescence spectroscopy evidenced local structural reorganization at both the N- and C-termini preceding oligomerization (56). Moreover, measurements of intrachain diffusion rates in  $\alpha$ S monomers showed that intramolecular reconfiguration of tertiary contacts determines the kinetics of bimolecular association (57). It is then evident that residual structure at both the N- and C-terminus, and its interplay via long-range interactions, appears to govern the early steps in  $\alpha$ S oligomerization. The possibility of swapping between intra- and intermolecular contacts while remaining disordered offers a rationale for the initial population of oligomeric species in the amyloid pathway of  $\alpha$ S with minimal energetic costs. Even when these regions are not directly involved in the fibrillar core of the protein, they seem to communicate via intrachain contacts to dictate the conformations populated by the protein. For example, nitration of C-terminal Tyr residues perturbs binding of the N-terminus to lipid membranes via a yet unknown mechanism that could, very likely, involve reorganization at the level of N/C interactions (19).

The unexpected complexity in the nature and functionality of the network of short- and long-range tertiary interactions of monomeric  $\alpha S$  could have implications for our future understanding of the biological relevance of such transient structural features in IDPs. From this work it is noticeable that a high-resolution description of tertiary contacts using small ensembles is a very valuable complementary study to the use of large ensembles (47,48,58,59). Although the latter approaches allow a more general description of the vast conformational space populated by the disordered protein, our methodology allows identifying with higher precision biologically relevant contacting regions of the polypeptide chain. In particular, the description of fibril-like interactions in monomeric  $\alpha$ S could allow better identification and targeting of aggregation-prone species within the ensemble of  $\alpha$ S conformations.

## SUPPORTING MATERIAL

Six figures, two tables, references (60–64), and supplementary data are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(13)00866-7.

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