Principles of self-organization in biological pathways: a hypothesis on the autogenous association of alpha-synuclein

Andreas Zanzoni1,2, Domenica Marchese1,2, Federico Agostini1,2, Benedetta Bolognesi1,2, Davide Cirillo1,2, Maria Botta-Orfila1,2, Carmen Maria Livi1,2, Silvia Rodriguez-Mulero1,2 and Gian Gaetano Tartaglia1,2,*

1Gene Function and Evolution, Bioinformatics and Genomics, Centre for Genomic Regulation (CRG), 08003 Barcelona, Spain and 2Universitat Pompeu Fabra (UPF), 08003 Barcelona, Spain

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ABSTRACT

Previous evidence indicates that a number of proteins are able to interact with cognate mRNAs. These autogenous associations represent important regulatory mechanisms that control gene expression at the translational level. Using the catRAPID approach to predict the propensity of proteins to bind to RNA, we investigated the occurrence of autogenous associations in the human proteome. Our algorithm correctly identified binding sites in well-known cases such as thymidylate synthase, tumor suppressor p53, synaptotagmin-1, serine/arginine-rich splicing factor 2, heat shock 70 kDa, ribonucleic particle-specific U1A and ribosomal protein S13. In addition, we found that several other proteins are able to bind to their own mRNAs. A large-scale analysis of biological pathways revealed that aggregation-prone and structurally disordered proteins have the highest propensity to interact with cognate RNAs. These findings are substantiated by experimental evidence on amyloidogenic proteins such as TAR DNA-binding protein 43 and fragile X mental retardation protein. Among the amyloidogenic proteins, we predicted that Parkinson’s disease-related a-synuclein is highly prone to interact with cognate transcripts, which suggests the existence of RNA-dependent factors in its function and dysfunction. Indeed, as aggregation is intrinsically concentration dependent, it is possible that autogenous interactions play a crucial role in controlling protein homeostasis.

INTRODUCTION

Although proteins are involved in almost every cellular process, increasing evidence indicates that coding and non-coding RNAs play fundamental roles in gene regulation (1,2) and disease (3,4). Recent studies showed that establishment of aberrant associations or disruption of functional protein–RNA interactions occurs in neurological disorders (5,6). For instance, interaction with RNA favors conversion of alpha-helix rich prion protein PrPC into the pathogenic beta-structure-rich insoluble conformer PrPSc that propagates in Creutzfeldt–Jakob disease (7). In Alzheimer’s disease, the association between Amyloid Precursor Protein mRNA and iron regulatory protein 1 is disrupted, resulting in compromised translation efficiency and elevated cytotoxicity (8).

Protein–RNA associations regulate several processes such as synthesis, folding, translocation, assembly and clearance of molecules. Previous studies suggested that ribonucleoprotein interactions might be able to facilitate protein and RNA folding (9,10). As a matter of fact, it has been observed that there is strong affinity between amino acids and their corresponding codons (11,12), which could imply a direct interaction between proteins and their own mRNAs (13,14). Indeed, TAR DNA-binding protein 43 (TDP-43) and Fragile X Mental Retardation protein (FMRP) have been found to interact with their own mRNAs (15,16). In these cases, expression is regulated by a negative feedback loop involving the 3’ untranslated region (UTR). Other autogenous associations have been observed in proteins associated with cell proliferation and gene expression (17,18). Also structurally disordered proteins such as Serine/Arginine-rich splicing factor 2 (SRSF2) (19) as well as heterogeneous ribonucleoprotein
members (20,21) are able to inhibit their translation by associating with their own mRNAs.

How often do autogenous associations occur in the human proteome? Recent technological advances revealed that a large number of proteins have RNA-binding abilities (22), which suggests that interaction with cognate mRNAs could be more frequent than previously thought. Are autogenous associations linked to specific functions? It is possible that autoregulatory mechanisms are involved in controlling protein production. For instance, in the case of TDP-43 and FMRP, inhibition of expression via autogenous interaction is a way to preserve protein functionality (15,16). Overexpression leads to high protein production and enhanced amyloidogenicity, resulting in harmful gain- or loss-of-function effects on cellular metabolism (23).

In this work, we focused on the ability of proteins to establish autogenous associations. Using our computational approach catRAPID (24), we studied the occurrence of these interactions in the human proteome. A large-scale analysis was performed to identify the role of autogenous associations in biological pathways and characterize their properties.

**MATERIALS AND METHODS**

**Biological pathway annotations**

We downloaded (September 2012) pathway data from two manually curated and high-quality resources: Reactome (25) and the NCI Pathway Interaction Database (NCI-PID) (26). The Reactome annotations (version 41) were gathered via the BioMart query interface returning a list of 167 canonical pathways containing 5375 unique protein coding genes, whereas the NCI-PID pathways were fetched directly from the database website (241 pathways, 2053 unique protein coding genes). In both cases, UniprotKB (27) accession numbers were converted to Ensembl (version 68) gene identifiers using the UniprotKB id-mapping file (version 2012.08). Subsequently, the gene pathway annotations were transferred to the corresponding polypeptides and coding/non-coding transcripts.

**Protein–RNA interaction prediction**

We used the catRAPID algorithm (24) to predict interaction propensity among all peptides and transcripts belonging to Reactome and NCI-PID pathways. catRAPID was trained on a large set of protein–RNA pairs available in the Protein Data Bank to discriminate interacting and non-interacting molecules using secondary structure propensities, hydrogen bonding and van der Waals contributions (28). The method was tested on the non-nucleic acid-binding database (NNBP; area under the receiver operating characteristic curve of 0.92), the NPIter database (area under the receiver operating characteristic curve of 0.88) and a number of individual interactions (e.g. RNAse mitochondrial RNA MRP and X-inactive specific transcript XIST networks; average accuracy of 78%). Owing to CPU limitations in the calculation (29), we restricted the predictions to RNA sequences with a length between 50 and 1500 nt as well as to polypeptides with a length between 50 and 750 amino acids. The ‘fragment’ and ‘strength’ algorithms were used to identify regions involved in the binding and compute the specificity with respect to random protein–RNA associations (5,29). For each protein–RNA pair under investigation, a reference set of \(10^5\) protein and \(10^2\) RNA molecules (total of \(10^4\) interactions) was used as a control (29). Reference sequences have the same lengths as the pair of interest to guarantee that the measure is independent of protein and RNA lengths (29).

**Gene partition**

The gene partition function \(gp(\pi,P)\) depends on the interaction propensity \(\pi\) and type of protein–RNA association, which is defined as autogenous \((P = a)\), intra- \((P = i)\) or inter-pathway \((P = l)\):

\[
gp(\pi,P) = \frac{c(\pi,P)}{c(\pi,a) + c(\pi,i) + c(\pi,l)}
\]

The number of counts \(c(\pi,P)\) is the fraction \(n(p,r,\pi)\) of protein \((p)\) and RNA \((r)\) molecules with interaction propensity higher than \(\pi\):

\[
c(\pi,P) = \sum_{p \in A(p)} n(p,r,\pi) / N(p,r)
\]

The function \(N(p,r)\) is the total number of interactions in the autogenous, intra- or inter-pathways \(A(P)\).

**Disorder propensity**

We predicted disorder propensities using the IUPred algorithm (30) with the ‘long disorder’ prediction option. We defined a residue as disorder prone if its IUPred score was above 0.5, as in a previous experimental study (31).

**Comparison of the propensity distributions**

We analyzed the disorder propensity of proteins involved in autogenous interactions by comparing them with the distributions of all the proteins annotated in the respective pathway data set. We evaluated the statistical significance using the Kolmogorov–Smirnov test (two-sided, alpha = 0.05).

**Pathway enrichment analysis**

We assessed the enrichment of autogenous interactions in biological pathways using the Gene Set Enrichment Analysis (GSEA) method (32). For each pathway data set, we used as background the whole list of autogenous interactions predicted by the catRAPID algorithm. We tested only those pathways annotated with autogenous interactions and containing at least five and not >500 genes (Supplementary Tables S1–S4). We ran GSEA with default parameters and performing 1000 permutations.

**Protein and RNA abundances**

Protein abundances were retrieved from the integrated whole organism Human PeptideAtlas (33,34), as assembled in http://pax-db.org (versions 2009, 2010,
RESULTS

Protein–RNA interactions in biological pathways

Using the catRAPID method (5,29), we systematically investigated the role of autogenous interactions in biological networks. For this purpose, we collected protein and RNA sequences annotated in two pathway resources: Reactome (25) and NCI-PID (26).

We first computed the interaction potential of 295 \times 10^6 protein–RNA pairs (10 376 protein sequences against 28 493 RNA sequences) in Reactome and 65 \times 10^6 protein–RNA pairs (4754 protein sequences against 13 608 RNA sequences) in NCI-PID (Supplementary Table S1; ‘Materials and Methods’ section). We then classified interactions as follows (Figure 1a): (i) intra-pathway or between proteins and RNAs coded by different genes belonging to the same pathway; (ii) inter-pathways or between proteins and RNAs coded by different genes belonging to different non-overlapping pathways; and (iii) autogenous or between proteins and RNAs coded by the same genes. To quantify the proportion of genes that are preferentially involved in intra-, inter-pathways or autogenous interactions, we introduced the ‘gene partition’ function, which is the fraction of associations predicted at a certain interaction score (Figure 1b; ‘Materials and Methods’ section).

In both Reactome and NCI-PID (Figure 1b; Supplementary Figure S1), we found that intra-pathway and inter-pathway interactions are strongly depleted at high interaction propensities, whereas autogenous associations are enriched (Figure 1b; ‘Materials and Methods’ section). We observed the same trend for both coding and non-coding transcripts (Supplementary Figure S1).

Biological pathways enriched in autogenous interactions

To uncover biological processes in which autogenous interactions play a functional role, we performed a GSEA (32) on both Reactome and NCI-PID pathways. In Reactome, we found 10 pathways enriched and 4 pathways depleted in autogenous interactions with a false discovery rate \(< 5\%\) (Supplementary Table S2). The top enriched pathway is ‘Amyloids’ (q-value = 3.2 \times 10^{-3}) followed by ‘Base Excision Repair’ (q-value = 3.8 \times 10^{-3}) and ‘Amine compound SLC transporters’ (q-value = 4.4 \times 10^{-3}) (Supplementary Table S2). Similarly, we identified 13 NCI-PID enriched pathways in autogenous interactions (Supplementary Table S3). We did not find any significantly depleted pathway at false discovery rate \(< 5\%\). The top enriched NCI-PID pathways are ‘Signaling events mediated by HDAC Class III’ (q-value = 2.9 \times 10^{-2}), ‘C-MYC pathway’ (q-value = 3.0 \times 10^{-2}) and ‘Hypoxic and oxygen homeostasis regulation of HIF-1-alpha’ (q-value = 3.1 \times 10^{-2}) (Supplementary Table S3). We identified the ‘Botulinum neurotoxicity’/Effect of Botulinum toxin’ pathway enriched in both databases (Reactome, q-value = 1.1 \times 10^{-2}; NCI-PID, q-value = 3.5 \times 10^{-2}) as well as the ‘\alpha\text{-synuclein signaling’ pathway enriched in

![Figure 1](image_url)

Figure 1. Autogenous versus intra- and inter-pathways interactions. (a) Sketch of biological pathways (pink and gray boxes connected by arrows; Supplementary Table S1). For each pathway, we studied three types of protein–RNA interactions: autogenous (green line), intra- (orange line) and inter-pathway (blue line); (b) From low to high interaction propensities (24), we found that the autogenous associations dominate over intra- and inter-pathway interactions present in Reactome (statistics for coding genes is shown; mean and s.e.m. of bins are shown; Supplementary Figure S1) (25). The gene partition is defined as the total fraction of genes showing preferential enrichment for autogenous, intra- or inter-pathway interactions (propensity \(> 50\%\): number \(n\) of genes enriched in autogenous interactions = 1238 of 1704; propensity \(> 100\%\): \(n = 211\) of 242; propensity \(> 150\%\): \(n = 20\) of 20; Supplementary Figure S2; ‘Materials and Methods’ section).
NCI-PID (q-value = 4.2 × 10⁻²) and related to the Reactome ‘Amyloids’ pathway (Supplementary Table S3).

**Known autogenous interactions in biological pathways**

To assess whether known cases of autogenous interactions were linked to the pathways identified in our analysis, we carried out a literature search. Indeed, amyloidogenic proteins such as TDP-43 and FMRP (Reactome pathway ‘Amyloids’) have strong propensities to bind to their mRNAs and have been discussed in our previous work (5).

We found that tumor suppressor p53, involved in the two top-enriched pathways ‘Signaling events mediated by HDAC Class III’ and ‘Hypoxic and oxygen homeostasis regulation of HIF-1-alpha’ (both in NCI-PID), is able to bind to its own mRNA (17,39,40). In *Mus musculus*, the RNA-binding site of p53 is a stable stem-loop structure that involves the 5′ UTR plus a region of 280 nucleotides in the coding sequence (5′ terminal region) (17). A similar mechanism has been observed in *Homo Sapiens*, but no conclusive evidence has been reported on the interaction (39). In agreement with experimental evidence on murine p53, our predictions indicated that the 5′ terminal region has strong propensity to bind (Figure 2a), and the interaction is specific (interaction strength = 89%) with respect to a control set of molecules of same size (Figure 2b; ‘Materials and Methods’ section) (17). We found similar results for human p53, although the region involved in the binding is not known (Supplementary Figure S2a and b). It is worth mentioning that p53 has strong propensity to form amyloid fibrils (42), and interaction with nucleic acids represents a way to control its aggregation potential by limiting the amount of protein product (17,43). Moreover, we note that p53 can be associated with ‘Base Excision Repair’ pathway (Reactome) (44,45). Indeed, as ‘Base Excision Repair’ deficiency affects genome stability and is implicated in many human diseases, including...
premature aging neurodegeneration and cancer (46,47), we expect that a self-regulatory mechanism in its components could greatly contribute to system’s efficiency.

In both NCI-PID and Reactome, we found that the pathways ‘Botulinum neurotoxicity’ and ‘Effect of Botulinum toxin’ are significantly enriched in autogenous interactions. One of the key-players in Botulinum toxicity is synaptotagmin-1, which is essential in Ca(2+) dependent neurotransmitter release (48). Synaptotagmin-1 interacts with the 3’ UTR of its own mRNA (49). In agreement with in vitro experiments, our predictions indicated that the 3’ UTR of synaptotagmin-1 RNA is involved in the autogenous interaction (Supplementary Figure S3a and b). Importantly, translation of synaptotagmin-1 is downregulated by the 3’ UTR, which is compatible with a negative feedback mechanism (49). As reported by light-scattering assays and electron microscopy, the protein forms large aggregates in a calcium-dependent manner (48), suggesting that the autogenous interaction protects against production of toxic oligomers (49).

Thymidylate synthase catalyzes the reaction generating thymidine monophosphate, which is phosphorylated to thymidine triphosphate for use in DNA synthesis and repair. Thymidylate synthase forms a ribonucleoprotein complex with C-MYC mRNA (50) and interacts with its own mRNA (51). The protein is not reported in the ‘C-MYC pathway’ (NCI-PID), but solid evidence exists on its interaction with C-MYC network (52). The RNA binding site for thymidylate synthase is within the 5’ UTR of the transcript (52). catRAPID correctly located the interaction within the first 188 nt of the 5’ UTR (Supplementary Figure S4a) and predicted high specificity for the binding (interaction strength = 99%; Supplementary Figure S4b). We note that the bacterial homologue of thymidylate synthase is able to associate with its cognate mRNA (53), which highlights the crucial role of autogenous interactions across different species.

Autogenous interactions and structural disorder

A recent study (31) showed that many RNA-binding proteins contain intrinsically disordered regions. Using the IUPred algorithm (30), we investigated the role of structural disorder in biological pathways. We found that proteins involved in autogenous interactions have a significant higher fraction of disorder prone residues compared with all proteins annotated in Reactome and NCI-PID (Figure 3 and Supplementary Figure S5; same results were observed for both coding and non-coding RNAs). To investigate whether known cases of autogenous interactions are enriched in unstructured regions, we performed a literature search.

We found that the human SRSF2, which has a long disordered C-terminus spanning amino acids 117–221, interacts with its own transcript (19) (Figure 4a). Notably, catRAPID correctly identified the binding site between the RRM domain (amino acids 14–92) and region 1/II of the terminal exon (55) (Figure 4b; interaction strength = 84%). The disordered region was predicted to be not interacting with the terminal exon. As a matter of fact, the C-terminal region participates in processes that only indirectly relate to the RNA-binding activity of the protein: facilitation of Ser/Arg phosphorylation to allow entrance in the nucleus (56) and establishment of low-affinity interactions to enhance the splicing activity (57).

Human Heat Shock 70 kDa (HSP70) interacts with its own mRNA (58) and has a disordered C-terminal region of ~10 kDa [highly conserved across species (59)] containing the Glu-Glu-Val-Asp regulatory motif. Using catRAPID, we predicted that the binding occurs at the 3’ UTR, which is in agreement with previous observations (Supplementary Figure S6a) (60). Indeed, HSP70 has a strong tendency to bind to AU-rich sequences that are located at the 3’ UTR (Supplementary Figure S6b) (61). Importantly, we predicted that both the N-terminal ATPase domain and the disordered C-terminus are involved in the interaction, which is consistent with the observation that HSP70 RNA-binding affinity depends on the ATPase domain but is considerably reduced when the disordered region is removed (62) (Supplementary Figure S6c). A high concentration of HSP70 is toxic, as the protein has a strong tendency to aggregate (58). Hence, interaction with mRNA represents...
a mechanism to control protein production and formation of toxic aggregates.

The small nuclear RiboNucleic Particle-specific U1A protein also binds to its own transcript with high affinity and specificity (18). Human U1A protein comprises two RNP domains separated by a disordered linker containing a nuclear localization signal. The determinants of protein–RNA specificity reside within amino acids 1–102 of the first RNP domain and a region in the linker (63). Using catRAPID, we predicted interactions between the 3' UTR and the first RNP domain as well as a region of the disordered linker (Supplementary Figure S7a and b; protein domains take from UniProtKB entry P09012) (18). We also predicted interaction within the second RNP domain, which has not been reported in literature but is involved in pre-mRNAs recognition (64). We note that unstructured regions of the protein participate in base recognition by forming direct and water-mediated hydrogen bonds with RNA (65).

Also ribosomal protein S13 shows moderate presence of secondary structure (66) and is able to bind to its own mRNA (67). Indeed, even though no information is available on its native state, it is possible that S13 contains disordered regions as most ribosomal proteins (68,69). In agreement with experimental evidence, catRAPID identified S13 binding site within the first and second exon (Supplementary Figure S8a and b) (67).

We note that also p53 and synaptotagmin-1 contain disordered regions, as shown by previous studies (41,70).

In the calculations of autogenous interactions involving disordered proteins, we used protein and RNA sequences as reported in the original papers. As the length of the RNA sequences exceeded catRAPID size limitation, the ‘fragmentation’ algorithm (5,29) was used to identify regions involved in the binding (‘Materials and Methods’ section).

**Autogenous interactions in control of protein translation**

Autogenous interactions regulate gene expression at the translational level by controlling protein concentration. If protein concentration is high, binding to mRNA is expected to have a major effect on translation efficiency.

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**Figure 4.** The autogenous interaction of SRSF2. (a) The catRAPID algorithm (5,29) was used to reproduce experimental evidence on the autogenous interaction of SRSF2 (19). The binding site is located in the final exon of SRSF2 mRNA (the blue box marks the region I/II that was determined experimentally) (19). (b) The interaction strength between SRSF2 and region I/II is significantly specific (84%) with respect to a control set of protein–RNA associations (‘Materials and Methods’ section). The RNA-binding domain RRM and disorder regions were reported as indicated in a previous study (54) and in agreement with the Uniprot (27) annotation (entry Q01130).
We investigated the relationship between protein abundance and propensity for autogenous interactions. From low to high expression levels, we found that reduction in translation efficiency (protein to RNA abundance ratio < 1) is accompanied by an increase in autogenous interaction propensity (Figure 5), which is fully compatible with a negative feedback loop mechanism. If the negative feedback loop is active, translation slows down, and the ratio between protein and RNA abundance decreases. By contrast, in case of efficient translation (protein to RNA abundance ratio > 1), the propensity for autogenous interactions is low (Figure 5), suggesting that the feedback loop does not occur.

Hence, our findings indicate that autogenous interactions might be involved in the modulation of protein expression by inducing regulatory feedback loops at the translational level. According to our analysis and in agreement with previous reports (15,49,58), negative feedback loops are a common type of mechanism involving autogenous interactions (for a list of predictions, see Supplementary Table S4).

**A hypothesis on alpha-synuclein association**

Among the amyloid proteins, we predicted that alpha-synuclein (SNCA gene) is significantly prone to autogenous interactions. The average interaction propensity with SNCA transcripts = 68 ± 4, and $P = 1.2 \times 10^{-12}$ (the average interaction propensity with transcripts coding for the major protein isoform = 72 ± 5 and $P = 1.3 \times 10^{-8}$; $P$-values were calculated with Mann–Whitney U test). The most abundant isoforms ENST00000394986, ENST00000336904 and ENST00000394991 (36,37) have interaction propensities of 81, 71 and 61, respectively. Intriguingly, the transcripts coding for the major protein isoform have a protein to RNA abundance ratio of 0.58, which is compatible with a negative feedback mechanism. The protein has been found to be present in both the cytoplasm and nucleus (71,72).

Alpha-synuclein is a 14 kDa protein composed of an amphipathic, positively charged 100-residue N-terminal domain with a lysine-rich N-terminus that binds reversibly to anionic membranes (73) and a 40-residue acidic C-terminal domain (74). The protein is predominantly monomeric in solution with a smaller fraction of multimeric species and is intrinsically unstructured (75,76). Importantly, interactions with double- or single-stranded DNA are able to convert alpha-synuclein into a highly structured protein (77). Circular dichroism shows that the alpha-helical content increases from 5 to 64% upon binding to DNA, whereas the random coil decreases from 95 to 33% (77).

The fact that alpha-synuclein interacts with DNA suggests that nucleic acid interactions might be relevant for its regulation. Our calculations indicate that ENST00000394986, ENST00000336904 and ENST00000394991 bind to alpha-synuclein at the 5’ UTRs (Figure 6a and Supplementary Figure S9). We predicted that the 5’ UTR interaction is specific (the interaction strengths of ENST00000394986, ENST00000336904 and ENST00000394991 are 100, 95 and 99%, respectively; see Figure 6b) and within GC-rich regions (Supplementary Figure S10), in agreement with previous evidence showing alpha-synuclein preference for G and C nucleotides (77). Moreover, a lysine-rich region spanning residues 40–60 was predicted by catRAPID to be involved in RNA recognition, which is consistent with previous results indicating an anion binding ability of the N-terminus (72).

At present, it is unknown whether RNA associations protect alpha-synuclein against formation of toxic aggregates. As a matter of fact, interaction with DNA sensibly increases alpha-synuclein amyloidogenicity (78) and a study on the Hofmeister series showed that anion binding promotes fibrilization (79). As GC-rich DNA aptamers have been found to associate with both monomeric and oligomeric alpha-synuclein (80), it is likely that stable RNA secondary structures, enriched in GC content, facilitate the disorder-to-order transition of alpha-synuclein, which could result in production of partially folded and highly amyloidogenic intermediates.

**DISCUSSION**

In this work, we used the catRAPID method (24) to compute the propensity of proteins to interact with coding and non-coding transcripts. In a number of biological pathways annotated in Reactome and NCI-PID, we found enrichment for autogenous interactions.

Our results are in agreement with available experimental evidence on the amyloidogenic TDP-43 and FMRP (the ‘Amyloids’ pathway has the highest enrichment in autogenous interactions) (15,16), tumor suppressor p53 (‘Signaling events mediated by HDAC Class III’,
‘Hypoxic and oxygen homeostasis regulation of HIF-1-alpha’ and ‘Base Excision Repair’ (39), synaptotagmin-1 (‘Botulinum neurotoxicity’ and ‘Effect of Botulinum toxin’) (49) and thymidylate synthase (connected to ‘C-MYC pathway’) (51). We expect that other autogenous interactions will occur in these pathways, although few cases have been reported in literature. For instance, the RNA-binding chaperone HSP90 (81,82) is involved in ‘Hypoxic and oxygen homeostasis regulation of HIF-1-alpha’ pathway, and it might have the ability to associate with its own transcript as other heat shock proteins (58,83). Moreover, we found enrichment in pathways such as ‘amine compound SLC transporters’, ‘metabolism of nitric oxide’, ‘iron uptake and transport’, ‘ABC-family proteins mediated transport’, ‘metabolism of vitamins and cofactors’, ‘amino acid transport’, ‘energy metabolism’, where autogenous interactions could be playing a role in metabolic regulation (84).

Our analysis showed that disordered proteins have significant propensity for autogenous interactions. The results are in agreement with experimental evidence on SRSF2 (19), HSP70 (58), U1A (18), p53 (17) and synaptotagmin-1 (49). The fact that proteins containing disordered regions have a high potential for autogenous interactions suggests that RNA interactions could protect unstructured domains from aberrant interactions or aggregation (85,86). Indeed, it has been observed that polyanionic molecules increase the solubility of nascent polypeptides, and that RNA molecules can act as molecular chaperones helping proteins to fold into their

Figure 6. A hypothesis on the autogenous association of α-synuclein. Among the amyloid proteins, we found that α-synuclein (SNCA gene) is significantly prone to autogenous interactions (average interaction propensity with SNCA transcripts = 68 ± 4). (a) Our calculations indicated that transcript ENST00000394986, which is abundant in brain, binds to α-synuclein at the 5’ UTR (gray box). (b) The 5’ UTR association was predicted to be specific (interaction strength = 100%) and involving a GC-rich region (Supplementary Figure S9), in agreement with previous experimental evidence showing that α-synuclein binds to G and C nucleotides (77).
native structures (87). As a matter of fact, interaction with cognate nucleic acids directly influences the aggregation propensity of TDP-43 (15,88), FMRP (89,90), p53 (17,43), synaptotagmin-1 (48,49) and HSP70 (58,83).

In the ‘Amyloids’ pathway, we focused on α-synuclein, a highly disordered and amyloidogenic protein linked to pathogenetic processes such as Parkinson’s disease and Lewy body dementia. Previous evidence indicated that α-synuclein forms partially folded multimers and aggregates impairing neuronal viability (75). We predicted that α-synuclein is able to establish autogenous interactions, which suggests the existence of RNA-dependent factors in the etiopathogenesis of Parkinson’s disease and other synucleinopathies. Indeed, our findings indicate that α-synuclein solubility might be modulated in vivo by associations with molecules such as nucleic acids (91,92). At present, it is has been shown that interaction with GC-rich DNA increases α-synuclein amyloidogenicity (77).

Although α-synuclein is present at the presynaptic terminals of neurons (93), several studies show that it can localize in the ‘nucleus’ (94,95). It should be considered that some mRNAs are shuttled to the axonal periphery where local synthesis takes place (96). Hence, the α-synuclein localization pattern and the presence of mRNAs at the neuronal terminals suggest that the predicted interaction of α-synuclein with RNA molecules may occur in the cellular context.

As in the case of other ribonucleoprotein interactions (28,97), we expect that ancillary elements present in vivo could increase α-synuclein affinity for nucleic acid associations. We do not exclude that the ribosomal components themselves could contribute to the formation of autogenous associations (10,98). As a matter of fact, the ribosome is the cellular component where autogenous interactions and translational control could take place simultaneously. Although our predictions on α-synuclein are compatible with a feedback loop mechanism (15,16), experimental evidence is required to determine the binding affinity of α-synuclein for RNA molecules and evaluate whether the interactions are mediated by other factors present in the cellular context.

Our findings suggest that autogenous interactions are able to reduce protein expression by inducing a negative feedback loop at the translational level. We previously observed that a tight anti-correlation (97%) exists between mRNA expression levels and aggregation rates of proteins (99,100). This relationship suggests that an evolutionary pressure acts against formation of toxic aggregates (101), and a molecular mechanism is in place to control expression of amyloidogenic proteins (102). In the light of our new findings, it is possible to speculate that autogenous interactions directly reduce the aggregation potential of proteins by controlling expression via feedback loops. As a number of genes have been reported to be dosage-sensitive (86,103), it is tempting to hypothesize that autogenous interactions play an important role in regulating their expression.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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