Are stress granules the RNA analogs of misfolded protein aggregates?

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ABSTRACT
Ribonucleoprotein (RNP) granules are ubiquitous features of eukaryotic cells. Several observations argue that the formation of at least some RNP granules can be considered analogous to the formation of unfolded protein aggregates. First, unfolded protein aggregates form from the exposure of promiscuous protein interaction surfaces, while some mRNP granules form, at least in part, by promiscuous intermolecular RNA–RNA interactions due to exposed RNA surfaces when mRNAs are not engaged with ribosomes. Second, analogous to the role of protein chaperones in preventing misfolded protein aggregation, cells contain abundant “RNA chaperones” to limit inappropriate RNA–RNA interactions and prevent mRNP granule formation. Third, analogous to the role of protein aggregates in diseases, situations where RNA aggregation exceeds the capacity of RNA chaperones to disaggregate RNAs may contribute to human disease. Understanding that RNP granules can be considered as promiscuous, reversible RNA aggregation events allow insight into their composition and how cells have evolved functions for RNP granules.

Keywords: RNA aggregates; RNA chaperones; eIF4A; stress granules

RNP GRANULES ARE UBIQUITOUS AND DIVERSE
Ribonucleoprotein (RNP) granules are a ubiquitous feature of eukaryotic cells and exist in both the cytosol, such as stress granules and P-bodies, and the nucleus, such as the nucleolus, paraspeckles, speckles, and Cajal bodies. There are also cell-type specific RNA granules such as maternal mRNP granules in a variety of oocytes and embryos (Buchan 2014), neuronal mRNA transport granules (Dalla Costa et al. 2021), and myogranules in developing muscle (Vogler et al. 2018).

RNP granules contain different subsets of RNAs with paraspeckles primarily consisting of NEAT1 RNA (Clemson et al. 2009); the nucleolus consisting of nascent rRNA transcripts and snoRNAs (Berry et al. 2015); Cajal bodies being enriched in scaRNAs, snRNAs, and snoRNAs (Machyna et al. 2014); and P-bodies and stress granules being enriched in untranslating mRNAs (Hubstenberger et al. 2017; Khong et al. 2017). For each type of RNP granule, there are also characteristic RNA-binding proteins (RBPs) and other constituents that complete their distinct composition (Jain et al. 2016; Markmiller et al. 2018; Youn et al. 2018; Matheny et al. 2019). Stress granules have been particularly useful to study the composition and assembly parameters of an RNP granule. Stress granules form under conditions where most mRNAs exit translation. For example, during the integrated stress response, phosphorylation of the eukaryotic translation initiation factor-2α (eIF2α) inhibits translation initiation, leading to polysome run-off and condensation of a subset of untranslated mRNPs into stress granules (Kedersha et al. 1999). RNA sequencing of purified stress granules indicates that they contain a highly diverse transcriptome enriched in long, poorly translated mRNAs (Khong et al. 2017; Namkoong et al. 2018; Matheny et al. 2019).

Several observations now argue that stress granules form from the summation of protein–protein, protein–RNA, and RNA–RNA interactions between individual mRNPs (Van Treneck et al. 2018; Van Treneck and Parker 2018; Roden and Gladfelter 2021). Evidence for protein interactions comes from the observation that cell lines lacking the Ras GTPase-activating protein SH3-domain-binding protein 1 and 2 (G3BP1 and G3BP2) paralogs are deficient at stress granule formation under a variety of stresses (Kedersha et al. 2016). Evidence that intermolecular RNA–RNA interactions are important in stress granule formation is that RNA self-assembly in vitro can largely recapitulate the
stress granule transcriptome (Van Treeck et al. 2018), and proteins that inhibit RNA self-assembly in vitro limit stress granule formation in cells (Tauber et al. 2020a; Budkina et al. 2021). Thus, stress granule assembly can be understood as ribosome run-off leading to newly exposed RNA sequences, which can allow both additional protein–RNA interactions (Yang et al. 2020) and promiscuous intermolecular RNA–RNA interactions (Van Treeck et al. 2018). The protein–protein or RNA–RNA interactions between individual mRNPs can be thought of as a highly cooperative equilibrium binding reaction leading to a larger assembly.

Multiple observations suggest that the ability of RNA to form assemblies based on promiscuous intermolecular RNA–RNA interactions is robust. First, all RNA homopolymers are able to condense into droplets, tangles, or aggregates in vitro (Van Treeck et al. 2018; Boeynaems et al. 2019). Second, isolated total RNA from yeast forms RNA condensates under physiological conditions (Van Treeck et al. 2018). Third, the estimated concentration of exposed coding regions after ribosome run-off in yeast (170–800 µg/mL) or in mammalian U-2 OS cells (~180 µg/mL) is in the range that can trigger RNAs to condense (Bounedjah et al. 2012; Van Treeck et al. 2018). Granule-enriched RNAs are on average very long and therefore contain multiple sites for potential interactions (Khong et al. 2017). This allows RNA molecules to have stable interactions within an mRNP granule through the summation of multiple weak interactions (Banani et al. 2016; Matheny et al. 2021).

The specific types of intermolecular RNA–RNA interactions that promote RNA condensation remain to be established in biological conditions. However, one anticipates that any type of interaction between two RNA molecules can contribute to RNA condensation including base-pairing, non-Watson–Crick base interactions, the formation of triple helices, ribose zippers, and co-axial stacking between helices in different RNA molecules (Bevilacqua et al. 2022).

The formation of mRNP granules is a general phenomenon that occurs in multiple biological contexts where there is an increased pool of untranslating mRNPs. For example, P-bodies are constitutive RNP granules that assemble from untranslating mRNPs associated with the RNA decay machinery (Sheth and Parker 2003) and that increase in size when the pool of untranslating mRNPs is increased (Teixeira et al. 2005). Similarly, P-granules are present in Caenorhabditis elegans oocytes and embryos that contain pools of untranslating mRNPs and are enhanced upon stress when the pool of untranslating mRNPs is increased (Lee et al. 2020; Parker et al. 2020). Other examples are mRNP granules found in neurons, where untranslating mRNAs condense into granules that are transported to the synapse for their disassembly and local translation (Tsang et al. 2019).

The tendency of untranslating mRNPs to assemble into mRNP granules can be generally understood as the loss of ribosomes causing three alterations to mRNP organization that promotes their condensation (Fig. 1). First, when mRNPs are no longer associated with ribosomes, they expose additional RNA sequences that could be capable of forming intermolecular RNA–RNA interactions, leading to mRNP condensation. Second, ribosomes are highly effective helicases and elongating ribosomes would remove any transient intermolecular RNA–RNA interactions that occur between the coding regions of different mRNPs. Third, we anticipate that the 80S ribosome has evolved to be relatively inert in forming promiscuous intermolecular RNA–RNA interactions and as such would be excluded from the RNP granule, although this final possibility remains to be examined experimentally.

**ARE STRESS GRANULES ANALOGOUS TO MISFOLDED PROTEIN AGGREGATES?**

We suggest that promiscuous assembly of untranslating mRNPs into RNP granules can be considered analogous to the formation of misfolded protein aggregates and occurs by the same three critical steps (Fig. 2). First, regions of proteins or RNAs capable of forming promiscuous interactions need to be exposed. For proteins, this happens with misfolding exposing hydrophobic surfaces that can then form aggregates through promiscuous interactions. For mRNAs, we suggest that this occurs when mRNAs are released from translation, exposing the coding region for promiscuous intermolecular RNA–RNA base-pairing and non-Watson–Crick interactions. Second, the exposed promiscuous surfaces need to overwhelm the existing mechanisms that limit nonspecific aggregation. For proteins, this would involve overwhelming protein chaperones, while for RNA aggregation, this would involve overcoming the anti-RNA aggregation effects of abundant RBPs and/or DEAD-box RNA helicases. Third, the initial assembly of proteins or mRNPs into promiscuous assemblies creates a high local concentration that would promote additional intermolecular interactions thereby stabilizing the “aggregate” (Tauber et al. 2020b).

Another similarity is that both misfolded proteins in aggregates and RNAs in stress granules show slow exchange rates, while protein chaperones in protein aggregates and many RBPs in stress granules show fast exchange rates (Kim et al. 2002; Winkler et al. 2010; Moon et al. 2019; Tauber et al. 2020b).

**THE “RNA CHAPERONE” NETWORK**

We suggest that there is an “RNA chaperone” network, which we define as a set of proteins that limits promiscuous intermolecular RNA–RNA interactions. This RNA chaperone network would be analogous to the protein chaperone network that is known to limit misfolded protein
aggregation and consists of Heat shock proteins (HSPs) and other components.

One key component of the RNA chaperone network is the eIF4A protein (Fig. 3). The key observations are that by binding RNA in an ATP-dependent manner, eIF4A can limit the self-assembly of RNA in vitro and limit stress granule formation in vivo (Tauber et al. 2020a). The biochemical properties of eIF4A are ideal for an RNA chaperone. eIF4A is a DEAD-box protein and can bind to, and destabilize, short duplexes (Rogers et al. 1999, 2001), leading to duplex disassembly before ATP hydrolysis and eIF4A release. eIF4A is not efficient at resolving stable duplexes (Rogers et al. 2001) because it does not directly use the energy of ATP hydrolysis to unwind RNA duplexes (Liu et al. 2008).

FIGURE 1. Possible mechanisms by which elongating ribosomes limit stress granule formation. The figure shows three possible mechanisms by which untranslating mRNAs assemble into mRNP granules due to loss of ribosomes: (i) Ribosome runoff leads to exposure of multiple RNA sequences (highlighted in orange) that can form promiscuous interactions. (ii) Ribosome acts as a helicase and removes transient intra- and intermolecular RNA–RNA interactions. (iii) Translating ribosomes do not engage in unspecific interactions and are excluded from RNP granules.

FIGURE 2. Analogous formation of stress granules and misfolded protein aggregates. Upon ribosome runoff, the newly exposed coding regions are free to form promiscuous intermolecular RNA–RNA interactions. Similarly, upon protein misfolding, hydrophobic surfaces get exposed that can then form aggregates through promiscuous interactions.
However, the ability to bind and destabilize short RNA duplexes that could form promiscuously is an ideal property for a general RNA chaperone. This suggests that additional DEAD-box and related DEVH-box proteins will play roles in limiting inappropriate RNA–RNA interactions.

A role for eIF4A, and other general RNA helicases, in limiting RNA condensation can be considered analogous to protein chaperones, such as HSP70, limiting the aggregation of misfolded proteins (Fig. 3). Multiple protein chaperones, including HSP70 proteins, bind to protein aggregates to disassemble aberrant interactions, thereby allowing for aggregate solubilization and protein refolding, using ATP hydrolysis as a switch for binding (Kampinga and Craig 2010; Mogk et al. 2018). We suggest that RNA condensation and inappropriate aggregation occur when the amount of exposed RNA in the cell exceeds the capacity of the cellular machinery limiting RNA condensation. Thus, the intrinsic aggregation properties of both proteins and RNAs are countered by abundant cellular machinery to keep these macromolecules correctly folded and dispersed for proper function.

Other members of the RNA chaperone network include abundant RBPs that limit RNA aggregation by binding to single-stranded regions in RNAs and limiting intermolecular RNA–RNA interactions. For example, the abundant RBP YB-1 limits RNA assembly in vitro and, when overexpressed, blocks stress granule formation in cells (Budkina et al. 2021). Such abundant RBPs can be considered analogous to small heat shock proteins (sHsps), a class of protein chaperones promoting the formation of reversible protein aggregates by limiting irreversible protein aggregation, presumably because sHSPs bind, and limit, interactions of aggregation-prone regions in proteins (Fig. 4; Zwirowski et al. 2017; Mogk et al. 2018). By analogy, RBPs could limit irreversible RNA aggregation by limiting the degree of intermolecular RNA–RNA interactions. A prediction of this model is that stress granules formed in the absence of YB-1, or similar RBPs, would show decreased RNA dynamics and prolonged persistence. In this light, G3BP1 has been suggested to prevent stable RNA entanglements by promoting the formation of a less stable, dynamic RNA assembly, although this has only been observed with poly(G) homopolymers (Guillén-Boixet et al. 2020). Since poly(G) homopolymers typically form hyper stable G-quadruplex structures (Williamson et al. 1989), it remains to be seen how G3BP1 affects the condensation of more typical mRNAs.

Other components of the RNA chaperone network not only remain to be identified but may also include RNA modification enzymes that destabilize RNA–RNA interactions, as well as nucleases that limit the intracellular concentration of RNA.

The importance of the RNA chaperone network will be amplified in some biological contexts. For example, during transcription in eukaryotic cells, a high local concentration of RNA can be produced, which would be prone to intermolecular RNA–RNA interactions potentially creating aberrant RNA-based assemblies. We hypothesize that cells counter this tendency by the higher concentration of RBPs and DEAD-box proteins per RNA in the nucleus as compared to the cytosol (Khong and Parker 2020). This elevated concentration of RBPs may effectively bind nascent RNAs and prevent intermolecular RNA–RNA interactions from forming.

We anticipate that the RNA chaperone network will be important at limiting intermolecular RNA–RNA interactions during cold shock. Moreover, the biophysical properties of proteins and RNAs allow an understanding of the fundamental differences between the heat and cold shock responses (Fig. 5). Specifically, proteins can experience
heat-induced misfolding due to temperature increase. This causes cells to induce the protein chaperone network that counters protein misfolding and restores homeostasis. Conversely, compared to proteins, RNA structures are more stable and therefore less likely to be severely perturbed by heat shock. However, RNA structures, as well as inappropriate RNA interactions, will have increased stability in the cold (e.g., Noble and Guthrie 1996; Zhang et al. 2018) increasing the demands for the RNA chaperone network. Notably, the cold shock response, most carefully documented in eubacteria, consists of the induction of RBPs, DEAD-box proteins, including an ortholog of eIF4A, and RNA nucleases (Zhang et al. 2018). This network has been proposed to function in alleviating cis

FIGURE 4. Do some RNA-binding proteins function analogously to small heat shock proteins (sHSPs)? RNA-binding proteins interact with RNAs, form dynamic assemblies, and facilitate disassembly of promiscuous interactions to limit RNA self-assembly into aggregates. Similarly, sHSPs bind to hydrophobic protein regions and form reversible assemblies to prevent their aggregation into insoluble aggregates that are more difficult to disassemble by the protein chaperone network.

FIGURE 5. Different properties of RNA and protein explain fundamental difference in heat and cold shock response: During heat shock, temperature increase dramatically induces protein unfolding and misfolding, while the reduction in temperature during cold shock on the contrary stabilizes preliminary RNA–RNA contacts in cis or trans. Both lead to different stress signaling cascades and induction of proteins to resolve those promiscuous interactions.
folding of RNAs, but we anticipate an additional contribution in limiting inappropriate trans interactions that would otherwise inhibit RNA function.

The RNA chaperone network might also be very important in plants under drought conditions, where the intracellular concentration of salt is expected to increase. This is relevant since high intracellular salt would be expected to stabilize RNA duplexes by charge neutralization and higher intracellular salt is known to promote stress granule formation, while low intracellular salt inhibits stress granule formation (Boundedjah et al. 2012). Strikingly, work in a number of plant species has shown that overexpression of eIF4A increases plants’ tolerance to salt stress (e.g., Rao et al. 2017), which might be due to eIF4A limiting RNA aggregation under these conditions. Moreover, plants may limit RNA aggregation by the production of solutes destabilizing RNA–RNA interactions (Bevilacqua et al. 2022).

**IMPLICATIONS OF “RNA AGGREGATION” AND THE RNA CHAPERONE NETWORK**

The possibility that promiscuous intermolecular RNA–RNA interactions are a prevalent biophysical force within cells, that are generally countered by an “RNA chaperone network,” provides insight into the formation and composition of RNP granules.

There are four basic features of mRNAs that dominate their enrichment in mRNP granules. First, the probability of an mRNA being enriched in an mRNP granule will be enhanced by being translationally repressed, thereby allowing the possibility of more productive intermolecular RNA–RNA interactions (Khong et al. 2017; Matheny et al. 2019; Lee et al. 2020). Second, longer RNAs will in general be preferentially enriched in granules, as increased length allows for multiple weak interactions distributed along an mRNA to act in summation (Matheny et al. 2021). Third, mRNAs with increased binding sites for protein components of mRNP granules will have increased partitioning into the assembly (Matheny et al. 2021). Finally, one anticipates that the mRNAs that fold into structures with more exposed single-stranded RNA (ssRNA) will partition more efficiently than structured mRNAs with less exposed ssRNA regions into mRNP granules due to their presumed ability to form additional intermolecular interactions. This possibility is suggested by the observation that mRNAs predicted to fold into structures with more exposed ssRNA form RNA-based assemblies with more stable structures in vitro than mRNAs with less exposed ssRNA (Ma et al. 2021), although whether these principles will occur in cells has not been demonstrated.

A second important implication is that RNP assembly into RNP granules will generally be dominated by protein–protein interactions under most normal physiological conditions, where RNA aggregation can be effectively countered by the RNA chaperone network. We anticipate that promiscuous intermolecular RNA–RNA interactions will play major roles in two contexts. First, intermolecular RNA–RNA interactions will likely dominate in RNP granule formation when the capacity of the RNA chaperone network capacity is exceeded, such as during the stress response, where the large pools of untranslated mRNAs condense into stress granules (Van Treeck et al. 2018; Tauber et al. 2020a). A similar phenomenon may occur in specialized cell types, such as neurons and oocytes, with larger pools of untranslated mRNPs. We anticipate that promiscuous intermolecular RNA–RNA interactions will also occur once an RNP granule is assembled by protein–protein interactions, due to the formation of a high local concentration of RNA, and thereby reinforce the formation of the RNP granule (Tauber et al. 2020b).

A third point is that promiscuous interactions between untranslated mRNPs can be utilized by cells to build functional assemblies, and this biophysical driving force can explain the otherwise perplexing composition of some RNP granules. One example that can now be understood is the diverse composition of mRNAs in P-granules in C. elegans, which are mRNP granules that segregate specific mRNAs to the developing germline during embryogenesis. Surprisingly, these mRNAs are a very diverse set of mRNAs including mRNA-encoding housekeeping proteins such as ribosomal proteins, and only a few specific mRNAs that are critical to germline development when properly localized (Lee et al. 2020). This composition can be understood as a case where evolution has used the stochastic formation of untranslated mRNAs in bulk to create an RNP granule, which then recruits the specific mRNAs needed for germline development and thereby properly segregates those mRNAs. To ensure that those critical mRNAs are present in P-granules, they are strongly translationally repressed (Lee et al. 2020). Similarly, a wide diversity of mRNAs is localized to dendrites and include mRNAs for housekeeping genes such as ribosomal proteins that have no apparent need to be locally translated in dendrites (Ohashi and Shiina 2020). Such a diversity of mRNAs at dendrites could be explained by these mRNAs being components of an RNP transport granule built by promiscuous interactions between untranslated mRNPs.

One anticipates that evolution will alter the surfaces of RNAs to enhance specific and limit promiscuous intermolecular RNA–RNA interactions. For example, we hypothesize that there will be selective pressure on functional RNAs (such as snRNA, tRNAs, snoRNAs, and rRNAs) to evolve surfaces limited in their ability to form inappropriate intermolecular RNA interactions, through compact RNA secondary and tertiary structure and the binding of proteins. Moreover, evolution can create intermolecular RNA–RNA interactions that target specific mRNAs to
mRNP granules. For example, during Drosophila oogenesis, specific trans interactions between the Oskar mRNA (Jambor et al. 2011) or Bicoid mRNA (Ferrandon et al. 1997) 3’ untranslated regions (3’ UTRs) allow specific dimerization and target them to distinct RNP granules located at opposite poles of the oocyte. In contrast, mRNAs may also evolve structures that limit promiscuous interactions and therefore influence their assembly into mRNP granules. Such a possibility is suggested by the observations that altering the folding of the Cln3 mRNA in Ashiyba can lead to increased interactions with some distally localized mRNAs in vitro and a corresponding mislocalization of the Cln3 mRNA from RNP granules near the nucleus to distally localized mRNP granules (Langdon et al. 2018).

**DOES “RNA AGGREGATION” CONTRIBUTE TO HUMAN DISEASE?**

We suggest that in some conditions, RNA aggregation will contribute to human disease progression. For example, stress granule formation, which is driven in part by RNA aggregation, contributes to tumor progression and some degenerative diseases (Anderson et al. 2015; Shukla and Parker 2016; Taylor et al. 2016). In addition, some diseases thought to be based on protein aggregation may in fact be driven by a combination of both protein and RNA aggregation. This is suggested by the observation that tau aggregates in both cell and mouse models of disease are enriched in specific RNAs that may contribute to disease progression (Lester et al. 2021). Moreover, since many of the proteins that aggregate in neurodegenerative diseases are RBPs (e.g., FUS, TDP-43, Annexin-11, Ataxin-2), a role for RNA aggregation contributing to “protein aggregation” may be more prevalent. RNA could contribute to the pathology of protein aggregates both by enhancing the aggregation process and reducing the functional level of key RBPs whose loss of function contributes to disease (Lester et al. 2021).

**CONCLUSION**

In this review, we hypothesize that stress granules are the RNA equivalent of misfolded protein aggregates and that they form when increased levels of mRNP surfaces are exposed due to widespread mRNA release from ribosomes. The understanding that promiscuous intermolecular RNA–RNA interactions can be a driving force leading to RNA aggregation highlights the need for an “RNA chaperone network” of proteins that act to limit promiscuous RNA aggregation. A fuller understanding of the RNA chaperone network and its role in regulating RNA structure and function remains to be determined.

It is also clear that cells have taken advantage of the propensity of untranslated mRNAs to form RNP granules for function. Untranslating mRNP granules with important biological roles include neuronal granules, which can be important for synaptic plasticity (Bakhthavachalu et al. 2018) and maternal mRNP granules, which can be important for early development (Buchan 2014; Lee et al. 2020). Similarly, stress granules have functions in promoting survival during stress (Kedersha et al. 2013) and limiting viral infections (Eiermann et al. 2020), although the specific mechanisms by which stress granules exert these functions remain to be determined. Further understanding of the underlying mechanisms of RNA aggregation, and how cells regulate and utilize that process, should both illuminate new principles of cellular physiology and might provide new insights to develop therapeutic strategies in a number of disease states.

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