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1 **Interactions of amyloid coaggregates with biomolecules and its relevance to**
2 **neurodegeneration**

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18

19 **Abstract**

20 The aggregation of amyloidogenic proteins is a pathological hallmark of various
21 neurodegenerative diseases, including Alzheimer's disease, Parkinson's disease, and
22 amyotrophic lateral sclerosis. In these diseases, oligomeric intermediates or toxic aggregates of
23 amyloids cause neuronal damage and degeneration. Despite the substantial effort made over
24 recent decades to implement therapeutic interventions, these neurodegenerative diseases are not
25 yet understood at the molecular level. In many cases, multiple disease-causing amyloids overlap
26 in a sole pathological feature or a sole disease-causing amyloid represents multiple pathological
27 features. Various amyloid pathologies can coexist in the same brain with or without clinical
28 presentation and may even occur in individuals without disease. From sparse data, speculation
29 has arisen regarding the coaggregation of amyloids with disparate amyloid species and other
30 biomolecules, which are the same characteristics that make diagnostics and drug development
31 challenging. However, advances in research related to biomolecular condensates and structural
32 analysis have been used to overcome some of these challenges. Considering the development of
33 these resources and techniques, herein we review the cross-seeding of amyloidosis, e.g.,
34 involving the amyloids amyloid β , tau, α -synuclein, and human islet amyloid polypeptide, and
35 their cross-inhibition by transthyretin and BRICHOS. The interplay of nucleic acid-binding
36 proteins, such as prions, TAR DNA-binding protein 43, fused in sarcoma/translated in
37 liposarcoma, and fragile X mental retardation polyglycine, with nucleic acids in the pathology
38 of neurodegeneration are also described, and we thereby highlight potential clinical applications
39 in central nervous system therapy.

40

41 **Keywords:** amyloid, coaggregation, neurodegenerative disease, oligomer, DNA, RNA, nucleic
42 acid-binding protein, G-quadruplex

43

44 Introduction

45 Neurodegenerative diseases are characterized by the progressive degeneration of the
46 neuronal system. Many of these diseases are age-related, as exemplified by Alzheimer's disease
47 (AD) and other tauopathies, Parkinson's disease and other synucleinopathies, prion diseases,
48 and other sporadic or genetic proteinopathies. Aggregation of amyloidogenic proteins, such as
49 amyloid β (A β), tau, α -synuclein (α Syn), prion protein, transactive response DNA-binding
50 protein 43 (TDP-43), and fused in sarcoma/translated in liposarcoma (FUS/TLS), is thought to
51 be a cause or a major deleterious result in most cases of these diseases. Such disease-related
52 amyloids are prone to self-assembly into matured amyloid fibrils (1). In some
53 neurodegenerations, oligomers, which have structurally metastable intermediates in a wide
54 range of molecular sizes, commonly serve as neuronal toxins rather than the structurally stable
55 fibrils. Oligomeropathy is responsible for the molecular pathogenesis of the associated diseases
56 (2-5) and causes impaired synaptic function and neuronal death through oxidative stress,
57 inflammation, apoptosis, and dysfunction of proteostasis (6). Thus, oligomers have gained
58 attention as targets for research and drug development related to diagnostics and therapeutics.

59 In relation to amyloid assembly, "oligomer" is an ambiguously defined term used to
60 describe dimers as well as hundreds of monomers when they are water-soluble, which further
61 complicates drug targeting (7-9). Despite, over recent decades, substantial research effort
62 directed at therapeutic interventions, there is no cure for oligomeric assembly as a therapy for
63 neurodegeneration, which has two possible explanations as follows. (1) Reversible equilibrium:
64 the formation of toxic oligomers resistant to degradation occurs, whereas the self-assembly
65 process redirects toward dissociation back to nontoxic monomers. These properties increase the
66 difficulty involved in targeting specific dimensions of oligomers. (2) Nonuniformity: the
67 mechanism of amyloid propagation is generally explained using a uniform stacking model, as
68 represented by a nucleation-dependent polymerization model (10) or a template-dependent
69 dock-lock model (11); however, most neurodegenerative diseases show overlapping clinical
70 symptoms, e.g., among AD, tauopathies, and synucleinopathies (12-14). Concomitantly,
71 coaggregation or co-oligomerization of amyloids occurs in the brains of patients (15, 16). In
72 addition, other biomolecules, such as nucleic acids (e.g., total RNA and noncoding RNA) (17,
73 18), interact with aggregates as cofactors and have been characterized as inducers of further
74 unsettled co-oligomerization. Due to these two characteristics, the epidemiology, diagnosis, and
75 treatment of mixed dementia remains complex and challenging.

76 Structural analysis is a powerful approach used to achieve molecular understanding and
77 drug development related to amyloidosis. The analysis of biomolecules under conditions that
78 imitate biological environments within cells or tissues has attracted the attention of researchers.
79 For example, Tycko and colleagues have pioneered research in this challenging field using
80 solid-state nuclear magnetic resonance (ssNMR) and were the first to represent *ex vivo*
81 structures of amyloid β 40 (A β 40) (19) from AD brain tissue, comprising two or three identical
82 filaments with a C-shaped fold and right-handed twist. Compared with the *in vitro* structure of
83 A β 40 fibrils (20), the monomer units resemble each other, but several differences exist at the
84 single-residue level between *ex vivo* and *in vitro* A β 40 fibril structures in the side-chain

85 orientation and the contact mode in interfilament packing. Subsequently, Tycko's group
 86 analyzed the *ex vivo* structures of amyloid β 42 (A β 42) (21), its more aggregative isoform, in
 87 variable AD clinical subtypes. Although the structure of A β 42 fibrils largely differs from that of
 88 A β 40 fibrils and has structural heterogeneity with at least two prevalent structures in most
 89 patients, detailed structural modeling of A β 42 fibrils was not presented. In prior studies, three
 90 independent groups reported the *in vitro* structures of A β 42 fibrils (22-24) based on ssNMR
 91 experiments, with results indicating an S-shaped conformation with a right-handed twist in the
 92 middle and C-terminal regions, in addition to a disordered region at the N-terminal region. The
 93 salt bridge between Lys28 and the carbonyl group of the C-terminus could be involved in
 94 stabilization of the fibril structure. Given the structural difference between fibrils from A β 40
 95 and A β 42, the coaggregation of A β 40 with the seed of A β 42 fibrils did not occur (22).

96 Structural analysis of *ex vivo* A β fibrils from post mortem human AD brains has been
 97 expanded to include cryogenic electron microscopy (cryo-EM). In 2019, Fändrich and
 98 colleagues demonstrated that an *ex vivo* A β 40 fibril fold purified from AD brain tissue was
 99 C-shaped with a right-hand twisted, in which its N- and C-terminal ends formed arches (25).
 100 These structures differed from that of the *in vitro* A β 40 fibril (26, 27), a structure that was also
 101 proposed by Fändrich's group using cryo-EM. Ultimately, Yang et al. succeeded in clarifying *ex*
 102 *vivo* A β 42 fibrils from the brains of patients with AD (28), showing two types of S-shaped
 103 filaments, including a N-terminal region around Y10 and V12 and a turn position that slightly
 104 shifted accordingly as well as a salt bridge formation between Lys 16 and Glu22 that was more
 105 contributable compared with that between Lys 28 and the carbonyl group of the C-terminus.
 106 Moreover, the participation of the N-terminal region to the S-shaped domain was not implied in
 107 the structures of *in vitro* A β 42 fibrils based on ssNMR (22-24) (Fig. 1a). These differences
 108 between *ex vivo* and *in vitro* results suggested that biomolecular cofactors were required for the
 109 formation of amyloid fibrils in brains with AD pathology. *Ex vivo* cryo-EM analyses have also
 110 been applied to other neurodegeneration-associated amyloid fibrils (e.g., tau (29) and TDP-43
 111 (30)). As observed in comparisons of A β 40 and A β 42, the structures of *in vitro* tau filaments by
 112 X-ray diffraction and Fourier transform infrared spectroscopy (31) do not reflect those of *ex*
 113 *vivo* tau filaments derived from patients with AD (29) (Fig. 1b). Moreover, *ex vivo* TDP-43
 114 filaments (30) have a double spiral-shaped fold in their low-complexity domain, including turns
 115 and β -strands composed of glycine and neutral polar residues, which shows little similarity to
 116 that of *in vitro* TDP-43 filaments (32, 33) (Fig. 1c). Despite the recognition that coaggregation
 117 is important in neurodegeneration, structural determination of coaggregates reflecting their
 118 biological environment and formation process has not been achieved.

119 When attempting to overcome the chaotic state in metastable protein assemblies in the
 120 brain, approaches in which molecular dynamic analysis is focused on biomolecular condensates
 121 and technology advancements in NMR and cryo-EM could be useful if the coaggregating
 122 molecules are localized and condensed in small organelles. In this review, we provide an update
 123 on the mechanistic insights into coaggregation (or cross-seeding) of neurodegeneration-related
 124 proteins with a focus on A β , tau, α Syn, prions, TDP-43, and FUS/TLS, the molecular sizes of
 125 which range across an order of magnitude 40–520-mer residues. The interplay with RNA, which

126 functions as an amyloid trapping molecule in the misplacement of encoding messages and
 127 thereby induces cellular deterioration, is an emerging “hot topic” in the proteinopathy research
 128 field, as exemplified by nucleic acid-binding proteins (e.g., prions, TDP-43, and FUS/TLS) and
 129 fragile X mental retardation polyglycine (FMRpolyG). The application of each amyloid
 130 coaggregation with different amyloid species or biomolecular cofactors has been separately
 131 reviewed in the past; however, to our knowledge, there has been no comprehensive review of
 132 amyloid coaggregates from the perspective of their interaction with cytosolic or nuclear
 133 biomolecules and potential overlap among neurodegenerative diseases. In this review, we also
 134 discuss how these coaggregates could play supramolecular roles in potential strategies for
 135 discovery of central nervous system (CNS)-targeting drugs and the development of therapeutics
 136 for neurodegeneration.

137

138 **Disparate amyloid cross-seeding and cross-inhibition in neurodegeneration**

139 Neurodegenerative diseases are characterized by aggregates of proteins such as A β , tau,
 140 and α Syn, the pathological forms of which appear to spread through the brain in characteristic
 141 patterns (34, 35). Although each disease exhibits the accumulation of specific characteristic
 142 protein aggregates, many cases exist in which aggregation of multiple pathological proteins is
 143 exhibited. Studies in *in vitro*, cellular, and *in vivo* systems have revealed several potential types
 144 of interactions between the different pathological proteins involved in neurodegeneration,
 145 including cross-seeding of aggregates in one protein initiating misaggregation of another. To
 146 explain the mechanisms of fibril formation of amyloidogenic proteins *in vitro*, a
 147 nucleation-dependent polymerization model has been used (10). This model consists of two
 148 phases: nucleation and seeding extension. Nucleus formation requires a series of association
 149 steps of monomeric proteins that are thermodynamically unfavorable, representing the
 150 rate-limiting step in fibril formation. Once the nucleus (seed) has been formed, the further
 151 addition of monomers becomes thermodynamically favorable, resulting in the seeding extension
 152 of fibrils. This model was originally advocated as a model for a single amyloid (A β) (10), but it
 153 has also been applied with a combination of different amyloids; accordingly, it is thought that
 154 various types of seeding aggregation can occur depending on the number of amyloids involved
 155 (Fig. 2). Below, we attempt to shed light on the amyloid proteins and their cross-seeding effects
 156 (Table 1) and occasionally cross-inhibition effects (Table 2).

157

158 *A β cross-seeding with tau aggregation*

159 AD is characterized by the accumulation of extracellular A β plaques and intracellular tau
 160 neurofibrillary tangles (NFTs) pathologically. It was found that A β binds to multiple tau
 161 peptides, especially those in exons 7 and 9, whereas tau binds to multiple A β peptides in the
 162 middle portion to C-terminal regions of A β (36). Such binding affinity between A β and tau was
 163 almost 1,000-fold higher than that of tau for itself. In P301L mutant tau transgenic mice,
 164 injection of A β 42 fibrils can significantly accelerate NFT formation, which further induces the
 165 phosphorylation of tau (37), indicating that the cross-seeding interaction of A β 42 with P301L
 166 tau generates many more NFTs than are generated by either A β 42 or P301L tau alone. Similarly,

167 the introduction of tau in Tg2576 transgenic mice was found to enhance the expression of
 168 mutant A β precursor protein (APP) and subsequent aggregation of A β (38).

169

170 *A β cross-seeding with α Syn aggregation*

171 Up to 50% of AD cases exhibit significant Lewy bodies (LBs) pathology in addition to
 172 plaques and tangles (39, 40). Compared with pure AD, AD with LBs pathology as a secondary
 173 lesion has been reported with lower mini mental state examination scores and more advanced
 174 dementia, suggesting that the severity of the disease increases due to complications related to
 175 LBs pathology (41). Likewise, patients with dementia with LB (DLB) frequently exhibit AD
 176 pathology, particularly senile plaques (42). Autopsy studies of 213 patients with LBs disorder in
 177 which the burden of tau NFTs and neuritic plaques was assessed revealed 26% with low-level
 178 AD neuropathology, 21% with intermediate-level AD neuropathology, and 30% with high-level
 179 AD neuropathology (43). As levels of AD neuropathology increased, cerebral α Syn scores also
 180 increased, and the interval between onset of motor and dementia symptoms and disease duration
 181 was shorter. In the same study, multivariate regression revealed independent negative
 182 associations between the cerebral tau NFT score and the interval between onset of motor and
 183 dementia symptoms (43).

184 Using transgenic mice with neuronal expression of A β and α Syn, it was shown that A β
 185 enhances α Syn accumulation and neuronal deficits (44). An NMR study showed that A β and
 186 α Syn might interact directly at a few sites (45). Although various studies have identified A β and
 187 α Syn oligomers as central toxic events during AD and LBs disease, leading to cell death and
 188 synaptic dysfunction (3, 46), a specific *in vitro* study found that A β and α Syn might interact
 189 directly to form hybrid pore-like oligomers that contribute to neurodegeneration (47).
 190 Previously, Ono and colleagues showed that fibrils and oligomers of A β 40, A β 42, and α Syn
 191 acted as seeds and affected the aggregation pathways within and among species *in vitro* (48).
 192 The seeding effects of α Syn fibrils were increased relative to those of A β 40 and A β 42 fibrils in
 193 the A β 40 and A β 42 aggregation pathways, respectively. It was also shown that A β and α Syn
 194 acted as seeds and each affected the aggregation pathway of the other *in vitro* (48).

195

196 *Tau cross-seeding with α Syn aggregation*

197 Lee's group found that one strain of preformed α Syn fibrils can be directly cross-seeded
 198 for tau aggregation, both in neuron cultures and an *in vivo* model of tau (49). This group
 199 injected α Syn preformed fibrils into mice with abundant A β plaques, and the A β deposits
 200 dramatically accelerated α Syn pathogenesis and spread throughout the brain. Remarkably,
 201 phosphorylated tau was induced in α Syn fibril-injected 5 \times FAD mice, and these mice showed
 202 neuron loss that was correlated with the progressive decline of cognitive and motor performance.
 203 These findings suggest the existence of a feed-forward mechanism in which A β aggregates
 204 enhance endogenous α Syn aggregation and spreading, which exacerbates the pathogenesis of
 205 A β and tau temporally postinjection with preformed fibrillar seeds of α Syn (50).

206

207 *A β or α Syn cross-seeding with IAPP aggregation*

208 Islet amyloid polypeptide (IAPP) is an amyloidogenic protein secreted as a randomly
 209 unstructured peptide. It plays a vital role in the progression of type 2 diabetes (T2D) mellitus;
 210 indeed, autopsies of bodies with this disease displayed IAPP aggregates in the pancreatic islets
 211 (51). The conformation of IAPP is assumed to be changed from a random structure to β -sheets
 212 before aggregation (52). Several studies have shown that individuals with AD develop signs and
 213 symptoms of T2D or other glucose-related disorders, whereas individuals with T2D are at a
 214 higher risk than healthy individuals of developing AD (53, 54). A study on the interaction of $A\beta$
 215 and IAPP showed that IAPP promotes $A\beta_{42}$ oligomerization and the formation of larger
 216 heteroaggregates with enhanced toxicity in neuronal cells (55). In the same study, $A\beta_{42}$ and
 217 IAPP interacted to form heterocomplex aggregates, which induced cell death in neuroblastoma
 218 cells (55). In transgenic mice, an intravenous injection of preformed $A\beta$ fibrils triggered IAPP
 219 aggregation in the pancreas, suggesting that $A\beta$ could enhance IAPP aggregation through
 220 cross-seeding (56). Several studies have reported the presence of α Syn in pancreatic β cells (57,
 221 58). One study showed that the octapeptide TKEQVTNV from α Syn can cross-seed with IAPP
 222 monomers and facilitate IAPP fibrillization (59). Contrary to expectations, this cross-seeding
 223 increased cell viability and reduced IAPP-induced cytotoxicity by shifting into a different
 224 seeding pathway of IAPP (59).

225

226 *TTR cross-inhibition with $A\beta$ or IAPP aggregation*

227

228 Along with cross-seeding between discrete amyloidogenic proteins, there are smaller but
 229 respectable literatures on the cross-inhibition against fibrillogenesis in $A\beta$ or IAPP by two other
 230 amyloids, i.e. TTR and BRICHOS. These amyloids are known as a paradox of amyloidogenic
 231 proteins with anti-amyloid aggregation properties, but the structural analysis using NMR and
 232 cryo-EM has not yet carried out. TTR is a 127-mer homotetrameric protein, which can be
 233 expressed mostly in the liver and be secreted into the plasma (60, 61). TTR molecules can
 234 misfold and form amyloid fibrils in the heart and peripheral nerves in the patients with TTR
 235 amyloidosis. The initial step in TTR aggregation is rate-limiting, and is involved in the
 236 dissociation of the native tetramer into monomers that subsequently undergo conformational
 237 changes forming aggregation-prone intermediates (60, 61).

237

238 Johnson and colleagues demonstrated that neutralization of TTR by chronic infusion of an
 239 anti-TTR antibody into the hippocampus of Tg2576 mice as a $A\beta$ -overexpressing AD model
 240 exacerbates $A\beta$ accumulation, tau phosphorylation, and neuronal loss (62). The same group
 241 reported that hemizygous deletion of TTR in APP^{sw}/PS1 Δ E9 mice resulted in earlier $A\beta$
 242 deposition in the cortex and hippocampus compared to control mice (63). These results suggest
 243 that TTR plays a critical role in the prevention of several AD pathologies. To explore the effect
 244 of TTR on $A\beta$ aggregation, thioflavin-T (Th-T) fluorescence assay and TEM were carried out
 245 by Olofsson and colleagues (64). They demonstrated that TTR inhibited fibril formation
 246 primarily by interfering the nucleation stage, resulting in the formation of Th-T-negative
 247 non-amyloid aggregates. It is noteworthy that TTR did not affect the seeding extension process
 248 in $A\beta$ aggregation (64). Further studies by Knowles and Chiti using atomic force microscopy
 (AFM) and dynamic light scattering (DLS) revealed that TTR inhibited both the primary and

249 secondary nucleation phases, but not fibril elongation, and then A β oligomers-induced
 250 cytotoxicity was reduced by TTR treatment (65).

251 TTR is expressed within the IAPP producing β -cells. Although there are no in vivo reports
 252 on cross-inhibition of IAPP aggregation by TTR, it was shown that TTR not only delayed the
 253 lag-phase but also impaired the elongation phase during the process of IAPP aggregation by
 254 Th-T assay (66). In addition, the interfering potential of TTR could be correlated inversely to
 255 thermodynamic stability, but no such correlation was observed in the dissociation rate of the
 256 tetramer (66). In AD model mice (*App*^{NL-F/NL-F}), high fat diet (HFD) treatment caused obesity
 257 and impaired glucose tolerance (i.e., T2D-like phenotypes), and an impaired cognitive function
 258 accompanied by marked increases in both A β deposition and microgliosis in the hippocampus
 259 were observed (67). Further to investigate, HFD treatment decreased TTR expression in
 260 *App*^{NL-F/NL-F} mice, indicating that the depletion of TTR could underly the increased A β
 261 deposition in AD pathology (67). These results imply TTR as a potential target of disease
 262 treatment for AD and T2D.

263

264 *BRICHOS cross-inhibition with A β or IAPP aggregation*

265 BRICHOS is a 100-mer protein domain found in 12 protein families including over 300
 266 proteins with a chaperon function (68). Especially, integral membrane protein 2B (ITM2B or
 267 Bri2) is a protein that in humans is encoded by the ITM2B gene, which is related to familial
 268 Danish dementia and familial British dementia, and Bri3 is a mutant of Bri2. BRICHOS domain
 269 from both Bri2 and Bri3 interacted with A β in neurons of AD patients (69). Studies on
 270 transgenic *Drosophila melanogaster* showed that co-expression of A β 42 and BRICHOS domain
 271 in the brain delayed A β 42 aggregation and significantly improved both lifespan and locomotor
 272 function compared with only A β 42-expressing flies (70). Moreover, BRICHOS increased the
 273 ratio of soluble to insoluble A β 42, and bound to A β aggregates (70), but the effects of each Bri2
 274 or Bri3 were not studied in this study. In further studies using *Drosophila melanogaster*
 275 expressing Bri2 by the same group, the neurotoxic effects of A β 42 were downregulated in the
 276 fly brains (71).

277 There have been several in vitro reports that prosurfactant protein C (proSP-C) BRICHOS
 278 and Bri2 BRICHOS significantly reduced the aggregation speed at substoichiometric levels by
 279 directly interacting with A β 42. Bri2 BRICHOS also suppressed the formation of toxic A β 42
 280 oligomers by specifically preventing the secondary nucleation pathway to remove the dominant
 281 source of A β 42 oligomers (72). 3D reconstruction of Bri2 BRICHOS analysis using TEM
 282 revealed that the monomers of Bri2 potently prevented A β 42-induced cytotoxicity. In particular,
 283 the dimers strongly suppressed A β 42 fibril formation by assembling into high molecular weight
 284 oligomers with a two-fold symmetry and the oligomers inhibited non-fibrillar aggregation.
 285 These data imply that Bri2 BRICHOS could harbor the molecular chaperone diversity by
 286 forming quaternary structures (73). As a comparison study, Bri3 BRICHOS also inhibited A β
 287 fibrillization and non-fibrillar protein aggregation in vitro by forming high molecular weight
 288 oligomers although the inhibitory effect of BRICHOS from Bri3 was weaker compared to that
 289 of Bri2 (74), raising a possibility of different roles for Bri2 and Bri3 BRICHOS against A β

290 pathology.

291 Regarding the cross-inhibition with IAPP, the effects of BRICHOS on IAPP aggregation
 292 and toxicity have been explored using in vitro studies, fly studies, and T2D patient materials
 293 (75). The BRICHOS domain of Bri2 intracellularly colocalized with IAPP in amyloid deposits
 294 of T2D patients. Bri2 BRICHOS showed a strong inhibitory activity against IAPP aggregation
 295 through targeting the secondary nucleation and redirecting the reaction towards formation of
 296 amorphous aggregates. Moreover, IAPP-induced toxicity was exacerbated in the human β -cell
 297 line EndoC- β H1 whose endogenous expression of Bri2 was downregulated by siRNA, whereas
 298 a concomitant overexpression of Bri2 BRICHOS recovered the cell viability. Similarly, the
 299 coexpression of IAPP and Bri2 BRICHOS in lateral ventral neurons of a *Drosophila* model
 300 increased the survival rate (75). These findings suggest that BRICHOS can be a potential
 301 endogenous inhibitor of IAPP pathologies, and then can be important therapeutic target T2D as
 302 well as AD.

303

304 **Coaggregation of amyloidogenic proteins with nucleic acids in neurodegeneration**

305 In 1998, a study on the detection of cytoplasmic RNAs in the pathological lesions of
 306 diverse neurodegenerative diseases was reported (76). Two pathological characteristics of AD,
 307 senile plaques and neurofibrillary tangles, contain RNA (77, 78). Mammalian nucleic acids have
 308 also been studied as cofactors for aggregation of several amyloidogenic proteins in
 309 proteinopathies. RNA and DNA molecules are postulated to interact with amyloids either
 310 directly or indirectly, resulting in conformational conversion, misfolding, aggregation, and
 311 infection. Inherently, most amyloids bind to polyanions, such as nucleic acids,
 312 glycosaminoglycans, and lipids (79-81). It was assumed that amyloid aggregates and nucleic
 313 acids would act as polyelectrolytes based on electrostatic forces (82). Each DNA or RNA has
 314 specific advantages and limitations in terms of chemical properties and structure, which are
 315 determined by Watson-Crick-type and Hoogsteen-type base pairings. Faced with enzymatic
 316 degradation, DNA oligonucleotides are more stable than their RNA counterparts. In contrast, the
 317 presence of the 2'-OH in ribose, as opposed to deoxyribose, potentially enables the higher
 318 conformational stability and diversity of RNA (83) (Fig. 3a). The absence of the 5'-methyl
 319 group in uracil, in contrast to its presence in thymine, has a similar impact on RNA properties.
 320 Indeed, nucleic acid (i.e., RNA and DNA) aptamers acting as synthetic oligonucleotides
 321 targeting A β , tau, α Syn, and prions have been extensively investigated as a means of disturbing
 322 the interaction of amyloids with nucleic acids (reviewed by Murakami et al. (9)). In comparison,
 323 studies of coaggregation of amyloids with endogenous nucleic acids causing pathologies have
 324 concentrated primarily on four examples of nucleic acid-binding proteins, i.e., prions, TDP-43,
 325 FUS/TLS, and FMRpolyG; therefore, these proteins are the focus of the subsections that follow.
 326 The binding characteristics of amyloidogenic proteins and nucleic acids and the subsequent
 327 nucleic acid-binding amyloids included in this review are summarized in Table 3.

328

329 *Prion coaggregation with DNA or RNA*

330 Human and animal prion diseases, including Creutzfeldt-Jakob disease, Kuru, Gerstmann-

331 Strüssler–Scheinker disease, and fatal familial insomnia in humans, bovine spongiform
 332 encephalopathy in cattle, scrapie in sheep and goats, and chronic wasting disease in deer and elk
 333 (84-86), are characterized by aberrant accumulation of misfolded prion protein (PrP). PrP
 334 consists of 253 amino acids and contains RNA recognition motif (RRM) and glycine rich
 335 domain (GXXXG) (87) (Fig. 4a). PrP exists physiologically as PrP^C (cellular form), which
 336 functions in neuroprotection and trophic signaling, whereas PrP^C can misfold into a toxic
 337 conformation as PrP^{Sc} (scrapie form) due to genetic and environmental causes (88, 89). PrP^{Sc}
 338 can form various conformational strains that are self-propagating and transmissible from cell to
 339 cell. The infectivity of PrP^{Sc} within the same species and sometimes across species is contingent
 340 on the specific strain and strain barrier. PrP^C is normally rich in α -helices, yet PrP^{Sc} forms a
 341 cross- β structure of amyloid fibrils upon aggregation due to some cofactors and acquires
 342 resistance to proteinases and denaturing, which leads to neurotoxicity (90). Thus, conversion of
 343 PrP^C to PrP^{Sc} or aggregation of PrP^{Sc} is a potential therapeutic target for the development of
 344 drug modalities.

345 Nucleic acids have attracted attention as key physiological factors required for the
 346 transformation of PrP^C to PrP^{Sc}. In 1997, Nandi first identified a bovine papilloma virus-derived
 347 plasmid DNA (16 kb) as a nucleic acid binder of PrP, showing that it bound to human
 348 PrP106-126 to possibly induce its structural change (91). A subsequent study by the same group
 349 revealed that human PrP106-126 generated amyloid fibrils with the addition of plasmid DNA
 350 but not without such DNA (92), suggesting that DNA plays a role as a cofactor in prion
 351 aggregation. For murine PrP23-231, a longer isoform, this was also the case in terms of its
 352 coaggregation with DNA (93), implying that nucleic acid metabolism is modulated by PrP.

353 Cordeiro et al. demonstrated the relevance of DNA to the pathology of prion-related
 354 diseases (94), finding, via circular dichroism (CD) spectrometry analysis, that the
 355 double-stranded DNA (18–34 bp; e.g., *recA1/2*, *Lexcons24*, *Lexcons 28*, and *E2DBS*) in molar
 356 excess (>2:1) over prions transformed murine PrP23-231 (i.e., PrP^C) to PrP^{Sc} and triggered the
 357 aggregation of PrP^{Sc} to induce fibril formation in a light-scattering assay. In contrast, the
 358 aggregation of Syrian hamster PrP109-141 and PrP109-149 was prevented by the presence of an
 359 equal or lower molar level of DNA oligonucleotides. A further study by this group showed that
 360 artificial single-stranded DNA oligonucleotides (18 or 21 nt in length) with different GC
 361 contents enhanced the aggregation of murine PrP23-231 in a light-scattering assay and
 362 transmission electron microscopy (TEM) as well as inducing the neurotoxicity of murine
 363 PrP23-231 in murine neuroblastoma cells (Neuro-2a cells) in a MTT test and caspase release
 364 assay (95). These findings indicate that the abundance of cellular DNA can contribute to PrP
 365 misfolding and neuronal death by modulating the equilibrium between PrP^{Sc} and PrP^C in
 366 neurodegeneration, and the possible interaction between DNA and PrP might originate from GC
 367 sequences. The dependency of prion–DNA binding on GC content is consistent with the
 368 preferable binding of small-length DNA aptamers (12-mer) that form the G-quadruplex, a
 369 noncanonical structure of nucleic acids induced via Hoogsteen-type base pairing, to ovine
 370 PrP23-231 (96). The G-quadruplex structure includes a stable planar core comprising four
 371 guanine bases in the same plane that form G-tetrads with Hoogsteen-type base pairing (Fig. 3b)

372 (97). G-quadruplex formation is involved in the protein–nucleic acid association through π – π
 373 interactions. The studies conducted to date suggest that endogenous DNA may facilitate prion
 374 propagation and aggregation by acting as a scaffold or molecular glue through the interaction
 375 between PrP^C and PrP^{Sc}. We propose that excess nucleic acids might modulate the balance
 376 between physiological PrP and misfolded PrP by making protein–protein interactions more
 377 likely.

378 Structural insights into the recognition of DNA oligonucleotides by mouse PrP23-231 have
 379 been provided using NMR and small-angle X-ray spectroscopy (SAXS) (98). SAXS is a
 380 small-angle scattering technique that can be used to determine the dynamics and structural
 381 information of molecules via analysis of the elastic scattering mode of X-rays at small angles.
 382 The SAXS data confirmed that mouse PrP23-231 forms a complex with 18-bp DNA in which
 383 the globular domain of C-terminal PrP, rather than the disordered region in the N-terminal
 384 portion, might contribute to complex formation (98). Perturbation experiments of the chemical
 385 shift in ¹⁵N-¹H HSQC NMR using ¹⁵N-labeled Syrian hamster PrP90-231 suggested that α -helix
 386 structures in the C-terminal region of PrP could be involved in the association with DNA (98).

387 In contrast to smaller length single-stranded DNA (i.e., <50 bp), longer length
 388 single-stranded RNA (several hundreds of nucleotides) plays a role in binding to PrP, which has
 389 RNA binding and chaperoning activities in relation to nucleocapsid retroviral proteins, such as
 390 NCp7 of human immunodeficiency virus (HIV) type 1. HIV-derived RNA has some resistance
 391 to proteinase K digestion through the formation of a complex with PrP, leading to PrP
 392 aggregation (99). According to NMR measurements using the N-terminal truncated peptide of
 393 PrP, the N-terminal region could participate in RNA binding in a similar manner to DNA.
 394 Additionally, RNA sources from mammals, yeast, or bacteria induced the aggregation of mouse
 395 PrP. Furthermore, PrP23-231 aggregated by incubation with total RNA from mouse
 396 neuroblastoma cells (Neuro-2a cells) was cytotoxic to such cells, and this cytotoxicity was
 397 consistent with the conformational change from an α -helix to β -sheet according to CD
 398 spectrometry analysis (99). The conversion of PrP^C to PrP^{Sc} was also stimulated by total RNA
 399 isolated from the hamster brain (100). Based on an *in vitro* conversion assay following the
 400 protein-misfolding cyclic amplification method and using prion-infected brain homogenate as a
 401 propagation seed, Saborio et al. demonstrated that RNA can be a requisite for the conversion
 402 and accumulation of pathogenic PrP^{Sc} (101). In this method, amplification is based on multiple
 403 cycles of PrP^{Sc} incubation in the presence of excess PrP^C followed by sonication. During the
 404 incubation periods, further PrP^{Sc} aggregation occurs through the incorporation of PrP^C, whereas
 405 the aggregates dispersed by sonication expand the population of converting units. Recently, the
 406 G-quadruplex formation of PrP^C mRNA was implied as the missing link in the initial conversion
 407 of PrP^C to PrP^{Sc} (102), suggesting that G-quadruplex binders or inhibitors could be therapeutics
 408 for prionoid diseases. Pseudoknot, a functional nucleic acid structure that contain stem-loop
 409 structures through Watson-Crick interaction and Hoogsteen interaction (103) (Fig. 3c), was
 410 reported to function as a recognition motif with human PrP^C similar to that of tRNA (104).
 411 These RNA structures can form stable nucleoprotein complex with human prion proteins.

412 Whether the PrP–RNA interaction occurs as a pathological trigger has not been shown

413 conclusively *in vivo*. PrP^C is localized physiologically at the plasmatic membrane, whereas
 414 misfolded PrP^C is believed to translocate to the nucleus of neuronal and endocrine cells where it
 415 interacts with chromatin (105, 106). It has been speculated that abnormal nuclear
 416 compartmentalization of PrP^C causes prion pathogenesis through encounters with RNA
 417 counterparts. Alternatively, crosstalk between PrP^C and RNA might be possible in the endocytic
 418 pathway because exogenous PrP binds endocytotically to nucleic acid. The endosomal recycling
 419 compartment was also identified as the likely site of the structural conversion of PrP^C (107).
 420 Indeed, cytosolic PrP^C was shown to form various RNA granule forms derived from nuclear
 421 RNA, 5S ribosomal RNA, or total RNA in Neuro-2a cells (108). Other studies reported that the
 422 PrP^C to PrP^{Sc} conversion occurred on the plasma membrane following the infection of the host
 423 with scrapie from external sources (109, 110). Collectively, the plasmatic membrane, nuclear
 424 compartment, cytosol, and plasma membrane can be considered crosstalk locations.

425 As mentioned above, experimental evidence has been accumulated on PrP–nucleic acids
 426 interactions, PrP^C to PrP^{Sc} conversion catalysts, and the induction of cytotoxicity. Sometimes,
 427 the coaggregation of prions can be accelerated by additional cofactors, such as copper. Indeed,
 428 PrP^C is a copper-binding protein with superoxide dismutase activity, whereas PrP^{Sc} is dependent
 429 on its copper-binding capacity (111, 112). One study found that the association of CuCl₂ in the
 430 interaction between ovine PrP^C and total RNA was fundamental for structural conversion to
 431 β-sheet-rich PrP^{Sc} and the acquisition of resistance to proteinase K (113). More studies on this
 432 relationship will be required to facilitate the development of antiprion drugs.

433

434 *TDP-43 coaggregation with RNA*

435 The DNA/RNA-binding protein TDP-43 (114, 115) plays a critical role in RNA processing,
 436 such as in alternative splicing, RNA stability, and transcriptional regulation in the CNS (116,
 437 117). Hyperphosphorylated and ubiquitinated TDP-43 were accumulated in inclusion bodies in
 438 the brain and spinal cord of patients with frontotemporal lobar degeneration (FTLD) and
 439 amyotrophic lateral sclerosis (ALS) (118). Almost cases (90%) of ALS are sporadic, whereas
 440 familial ALS cases (10%) include the inheritance of mutations. The mutation (10%) in *TARDBP*
 441 that encodes TDP-43 and the remaining 90% are due to mutations in other genes (e.g.,
 442 *C9ORF72*, *SOD1*, and *FUS*). From the unique gene *C9ORF72*, the transcribed RNA forms foci
 443 in neurons and glial cells and sequesters RNA-binding proteins, such as hnRNP43, in a
 444 mechanism that includes loss of *C9ORF72* function (119). Assemblies of SOD1 (120) and FUS
 445 (114, 115) were also found in the inclusion bodies. Although SOD1, a metalloprotein that binds
 446 to copper and zinc ions, is not related to RNA binding, FUS is a known RNA-binding protein;
 447 thus, FUS is described in the next subsection.

448 TDP-43 consists of 414 amino acids and contains two RNA recognition motifs (also
 449 known as ribonucleoproteins: RRM1 at aa 101–176 and RRM2 at aa 191–262) (121, 122), and a
 450 C-terminal low-complexity domain (LCD: aa 274–414) (118, 123) (Fig. 4a). The LCD includes
 451 glutamine/asparagine rich and glycine rich domains and is unstructured and flexible with
 452 functions that differ from those in normal regions involved in structured regions. Molliex et al.
 453 reported that the phase separation by the LCD induced from TDP-43 promoted stress granule

454 assembly and TDP-43 aggregation (124). An accumulation of studies also suggest that the LCD
 455 is responsible for the propensity to form amyloid fibrils and stress granules (125). Notably,
 456 based on the moderate similarities in the sequence between the LCD and prion proteins from
 457 *Homo sapiens* and *Pan troglodytes*, TDP-43 is postulated to have prion-like properties (126);
 458 indeed, the LCD was found to be propagated intercellularly (cell to cell) as a trigger of disease
 459 progression (127). In particular, TDP-43 downregulated splicing of the exon 9 of cystic fibrosis
 460 transmembrane conductance regulator by binding to a UG repeat site in RRM1 based on
 461 electrophoretic mobility shift assay (EMSA) ($K_D = 27$ nM) or isothermal titration calorimetry
 462 (ITC) ($K_D = 32$ nM) (128, 129), and the fragment of TDP-43 showed stronger affinity ($K_D = 5.3$
 463 nM) to RNA including (UG)₆ using EMSA (130). NMR studies based on ¹H-¹⁵N
 464 SOFAST-HMQC revealed the binding site in RRM1 loop3 and RRM2 pocket around V220 of
 465 TDP-43 (131). As observed in the case of RNA, TDP-43 associated with ssDNA containing
 466 (TG)₁₂ with potent binding affinity using ITC (132).

467 NMR studies by Conicella et al. showed that TDP-43 generated dimers through helix–helix
 468 contact (aa 321–343) in the LCD (133). Liquid-to-liquid phase separation (LLPS) is known to
 469 occur when two liquid phases coexist in nonmembrane organelles, and it drives the formation of
 470 biomolecular compartments, such as lipid droplets, for local biological reactions and signaling
 471 systems (134, 135). The experiments of these authors also revealed that the intermolecular
 472 helix–helix contact in the LCD induced the formation of liquid droplets of TDP-43, which
 473 generated amyloid fibrils upon their incubation. As validation, the nonpathological mutation
 474 (A321G) in the LCD disturbed the helical structure as well as liquid droplet formation, whereas
 475 an ALS-related mutation (G335D) in the LCD increased LLPS through stabilization of the helix
 476 structure (133). Fonda et al. conducted ssNMR analysis that showed that the liquid droplets
 477 from TDP-43 were transformed into β -strand-rich fibrils through stabilization of a region (aa
 478 365–400) within the LCD (136). Cryo-EM analysis by Li et al. indicated that amyloid fibrils of
 479 TDP-43 in the LCD harbored a core architecture, including several β -strands that were linked
 480 by loop structures (33). These findings demonstrate that the LCD is indispensable to the
 481 aggregation of TDP-43.

482 Wang et al. used NMR to show that the interaction between N-terminal domains was
 483 involved in the dimerization of TDP-43 (137). Their study also showed that the S48E mutation
 484 inhibited the aggregation of TDP-43 and that liquid droplet formation was prevented by the
 485 failure to phosphorylate Ser48 (137). In TDP-43, RRM binding to RNA (e.g., long noncoding
 486 RNA) drove liquid-like granule formation within cells (138) together with the self-association
 487 of N-terminal domains as well as helix–helix contact within the LCD (137, 139). TDP-43 is also
 488 known as a shuttling protein that travels between the nucleus and cytoplasm to facilitate cellular
 489 functions. Indeed, it is normally localized in the nucleus (140), but it moves to the cytoplasm
 490 and aggregates to form insoluble inclusions in disease states (141). Under transient stress
 491 conditions, TDP-43 forms nuclear bodies for sheltering by binding long noncoding RNA in the
 492 nucleus. Subsequently, the transferred TDP-43 generates stress granules by bundling
 493 heterogeneous nuclear ribonucleoproteins (hnRNPs) and RNAs in the cytosol. Moreover,
 494 long-term physiological stress and senescence induce the formation of stress granules, leading

495 to maturation and transformation into inclusion bodies.

496

497 *FUS/TLS coaggregation with RNA*

498 The 526-mer RNA-binding protein FUS/TLS is a heterogeneous nuclear ribonucleoprotein
 499 P2 encoded by *FUS* ([142](#), [143](#)). In 2009, mutations in *FUS* were found to be causative in
 500 subtypes of ALS cases ([144](#), [145](#)). FUS play a pivotal role in RNA processing, splicing, and
 501 transport ([146](#)). It is mainly composed of a serine–tyrosine–glycine–glutamine rich LCD (aa 1–
 502 165) in the N-terminal region, three arginine–glycine–glycine (RGG) rich domains (aa 166–267,
 503 aa 372–422, and aa 453–501, respectively), a RRM (aa 285–371), a zinc finger motif (aa 422–
 504 453), and a highly conserved nuclear localization signal domain (aa 501–526) in the C-terminal
 505 region ([147](#), [148](#)). Using a bioinformatics approach, the presence of two prion-like domains was
 506 also discovered (aa 1–239 and 391–407) ([149](#)) (**Fig. 4a**). Tycko and colleagues first determined
 507 the structure of *in vitro* fibrils obtained from the LCD of FUS by ssNMR, showing that a
 508 segment of 61 residues formed the fibril core with a S-shaped fold and right-handed twist,
 509 which plays a role in LLPS ([150](#)). The following study using a different segment (91 residues)
 510 in LCD domain based on cryo-EM supports for this structural feature by ssNMR ([151](#)) (**Fig. 4b**).
 511 However, the *ex vivo* structure of FUS fibrils has not yet been determined.

512 FUS is normally localized in the nucleus, but it can shuttle between the nucleus and
 513 cytoplasm in a similar manner to TDP-43 ([152](#)). The FUS mutation in familial ALS hampers the
 514 signal for nuclear localization, leading to the accumulation of FUS in the cytoplasm and
 515 acquisition of gain-of-function toxicity via sequestration of RNA ([153](#), [154](#)). In one study, the
 516 accumulation of RNA-binding proteins in the pathogenesis of ALS was found to be seeded by
 517 granules of ribonucleoprotein ([155](#)). The granules composed of membraneless organelles were
 518 stabilized by LLPS, in which self-association was induced in the liquid droplets ([156](#)). These
 519 observations are in agreement with the immunoreactivity of FUS in the cytoplasmic inclusions
 520 of the brain of patients with ALS and FTLD. Moreover, these findings indicate that the
 521 mislocalization of FUS to the cytoplasm is a pathogenic trigger ([157](#)).

522 Lerga et al. found that RNA oligoribonucleotides bind to the FUS protein at the GGUG
 523 motif with a 250 nM affinity using EMSA test ([158](#)). Another study showed that the recruitment
 524 of FUS to DNA damage sites was modulated by the RGG domain ([159](#)). In addition to RGG
 525 domains, the zinc finder motif in FUS is related to its RNA binding ([160](#), [161](#)), and then the
 526 binding constant was determined to be 56 nM by Wang et al ([162](#)). Indeed, RGG domain in
 527 FUS extensively recognized G4RNA from r(UUAGGG)₄ ($K_D = 6.2$ nM) ([163](#)). In the following
 528 two studies, RGG domain was also reported to bind G4RNA deduced from post-synaptic
 529 density protein 95 (PSD-95) using a steady-state fluorescence spectroscopy ($K_D = 28$ nM) ([164](#))
 530 and surface plasmon resonance (SPR) ($K_D = 3.2$ nM) ([165](#)). On the other hand, the binding
 531 affinity of stem-loop RNA structure in hnRNPA2/B1 to RGG was weaker based on ITC analysis
 532 ($K_D = 9.2$ μ M) ([166](#)).

533 The C-terminal nuclear localization signal domain in FUS is known to bind to the nuclear
 534 input receptor to enable transportation of FUS from the cytoplasm to the nucleus. Most of the
 535 mutations found in familial ALS are concentrated in this C-terminal signaling domain.

536 Loss-of-function mutations for such signaling prevent FUS from being transported into the
 537 nucleus, resulting in its accumulation in the cytoplasm (167, 168). Evidence suggests the LCD
 538 is localized at the N-terminus of FUS and contributes to the phase transition of the protein,
 539 facilitating the self-assembly of the LCD (169). Taken together, these findings indicate that FUS
 540 is transported from the nucleus to the cytoplasm before phase separation forms granules, after
 541 which FUS reversibly aggregates. However, it is conceivable that self-assembly of the LCD in
 542 the disease state originates from mutations in familial ALS; in this case, FUS may irreversibly
 543 aggregate, possibly through the LLPS pathway.

544

545 *FMRpolyG coaggregation with RNA*

546 Fragile X-related tremor/ataxia syndrome (FXTAS) is a neurodegenerative disease that is
 547 characterized by CGG triplet repeat expansions in *FMRI* (170), a gene that encodes
 548 repeat-associated non-AUG (RAN) translation and produces the 153-mer RNA-binding protein
 549 FMRpolyG. FXTAS is characterized by neuronal death and ubiquitin-positive inclusions within
 550 neurons, which are composed of FMRpolyG aggregates (Fig. 4a). According to confocal
 551 microscopy observations, FMRpolyG forms dot-like aggregates in the mitochondria as well as
 552 in the nuclear inclusion for subcellular localization of FMRpolyG. Using neuronal cell culture
 553 models and a FXTAS transgenic mouse model, these assemblies were related to the disturbance
 554 of mRNA splicing and impairment of the respiratory chain (171). Kumar and colleagues
 555 screened a small molecule library (>250,000 molecules) and identified three candidates that
 556 prevent impaired mRNA splicing and bind CGG repeat RNAs [r(CGG × 20–60)] using a
 557 fluorescence binding test and isothermal calorimetry titration binding assay (172). These RNA
 558 binders also inhibited the aggregation of FMRpolyG and RAN translation, indicating that they
 559 are lead molecules for anti-FXTAS drug development.

560 G-quadruplex of RNA (G4RNA) plays an important role in mRNA translocation and
 561 translation in the axons, dendrites, and dendritic spine of neuronal networks (Fig. 3b). Transport
 562 of mRNA to the synapse contributes to synaptic plasticity and learning memory (173, 174). In
 563 particular, dendritic mRNA generates a complex with RNA-binding proteins in RNA granules.
 564 Bioinformatics analysis showed that the function of ~30% of known dendritic mRNA was
 565 related to G-quadruplex consensus in the 3'-untranslated region (175). Given that FMRpolyG
 566 recognizes G4RNA in target *FMRI* mRNA (176, 177), Brown et al. performed a microarray of
 567 immunoprecipitation of the mouse brain with FMRpolyG, identifying 432 mRNAs, ~70% of
 568 which included a G-quadruplex-forming sequence (178). Crystal structure analysis also showed
 569 that the RNA-binding motif in FMRpolyG possesses three K-homology domains (also known as
 570 KH domains), which were identified in human hnRNP, and one RGG box (179). Notably, based
 571 on X-ray crystallography analysis, RGG peptides could stabilize the G-tetrad unit for
 572 facilitation of the G-quadruplex (180). The recognition of FMRpolyG to the G-quadruplex
 573 probably disturbs protein translation and RNA localization in the pathology of FXTAS.
 574 FMRpolyG is also known to suppress the mRNA translation of other genes, including *APP*
 575 (181), *PP2Ac* (182), and *MAP1B* (183), in a similar manner to its effects on *FMRI*. These
 576 results provide a potential modulator of FMRpolyG for FXTAS therapy.

577 Shioda and colleagues described the direct interaction of CGG repeat-derived G4RNA
578 with the polyglycine region of FMRpolyG and the generation of FMRpolyG–G4RNA
579 coaggregates (184). Prior to liquid-to-solid transition, LLPS induced the coaggregation of
580 FMRpolyG–G4RNA, which primarily interacted with exosomal proteins (e.g., PPIA, eEF1A1,
581 PKM, and hnRNP A2B1) in the exosomes, resulting in prion-like propagation of cell to cell and
582 neuronal dysfunction. The same group previously identified a G4-binding ligand,
583 5-aminolevulinic acid, which was metabolized to porphyrins protoporphyrin IX and hemin in
584 cells (185, 186). Notably, oral administration of 5-aminolevulinic acid prevented not only RAN
585 translation of FMRpolyG but also coaggregation of FMRpolyG with G4RNA, resulting in the
586 rescue of impaired synaptic plasticity and learning behavior in a mouse model of FXTAS. Their
587 findings suggest that 5-aminolevulinic acid is a promising drug lead for G4RNA prionoids (185,
588 186).

589 Generally, RNA loss of function and RAN translation (gain of function) are considered the
590 two underlying mechanisms of proteinopathies associated with repeat expansion disorders (187).
591 These two mechanisms are not independent but synergistically induce the formation of
592 LLPS-derived FMRpolyG coaggregates with CGG repeat-derived G4RNA. To date, FMRpolyG
593 has not been structurally determined. Thus, further investigation to clarify the underlying
594 mechanism responsible for FXTAS etiology will be required to facilitate the development of
595 target-specific medicines with few adverse effects.

596

597 **Conclusions and future perspectives**

598 It is hypothesized that cross-seeding between amyloids is dependent on conformations that
599 lower the energy barrier for seeding. The mechanism of amyloid aggregate formation has yet to
600 be clarified for most proteins, and a more profound understanding is required to facilitate
601 anti-amyloid drug design and discovery. The protein components involved in cross-seeding
602 should be further investigated to clarify the role and underlying mechanism of cross-seeding
603 aggregation. The inhibitors that dually target amyloidogenic proteins participating in the
604 cross-seeding event can inhibit heterologous aggregation but also cause disassembly of the
605 aggregates. Understanding the molecular mechanism underlying the interactions in
606 cross-seeding will help researchers develop effective disease-modifying therapies for
607 protein-misfolding diseases. Furthermore, by designing site-specific inhibitors, researchers
608 could also develop new approaches to inhibit and disassemble both homologous and
609 heterologous aggregates.

610 In 2022, using proteomics data of A β plaques from AD brains, Konstantoulea et al. found
611 that heterotypic A β interacts with peptide fragments from human proteins to facilitate
612 cross-seeding (188). These proteins shared local sequence homology with aggregation-prone
613 regions within A β , and transient expression of three of these proteins (WD repeat-containing
614 protein 81, chondroitin sulfate proteoglycan 5, and interferon regulatory factor 7) accelerated
615 A β aggregation in a cellular reporter model. Although the amyloid core is believed to share a
616 common interface with other amyloid species through a cross- β unit (189), the coaggregation
617 hypothesis has been expanded to unrelated human proteins. Indeed, ectopic DNA, such as

618 bacterial extracellular DNA, has been shown to enhance aggregation of A β (190) and tau (191).
619 However, β -sheet-triggered protein–nucleic acid interactions may also play a pivotal role in the
620 stability, compartmentalization, and degradation-resistance of vital amyloid-related proteins
621 (192). Although physiological and pathological differences may be subtle, the role of nucleic
622 acids is supposedly dependent on the cellular environment for modulation of the balance
623 between the two. Although an accumulation of evidence suggests that LLPS in the nucleus and
624 cytoplasm is important in maintaining cellular homeostasis and advances have been made in the
625 structural determination of amyloid assemblies, drugs targeting amyloid coaggregation have not
626 yet advanced to clinical trials in the neurodegeneration disease field. To advance the
627 development of anticoaggregation medicines for biomedical applications, several issues should
628 be addressed as follows.

629 First, the conformational metastability and heterogeneity of coaggregates of amyloid
630 oligomers with other amyloids or nucleic acids are major impediments for structural elucidation
631 of complexes. This challenge can be addressed using cryo-EM for *ex vivo* amyloid fibrils, which
632 could partially reflect the coaggregates with biomolecules, as demonstrated in the cases of A β 40
633 (19, 25), A β 42 (21, 28), tau (29), and TDP-43 (30). Although cryo-EM analyses combined with
634 computational advancement have been applied to RNA structures (193) and structures of
635 nucleoprotein–RNA complexes (194), there have been no such studies (to the best of our
636 knowledge) on amyloid–RNA complexes.

637 Second, identification of the inhibitors harboring dual inhibition activities against different
638 pairs of coaggregates is a promising approach that could increase both the speed and success of
639 the process. Considering the existing common amyloid core in coaggregates, natural products
640 with inhibitory activities against multiple (dual or triple) amyloid aggregates have been reported
641 (as discussed above): benzylamino-2-hydroxyalkyl derivatives (195), a curcumin derivative
642 (PE859) (196), notopterol (197), and epigallocatechin-3-gallate (198) against A β and tau
643 aggregation; curcumin (199) against A β and α Syn aggregation; and a synthetic compound
644 (MG-2119) (200) against tau and α Syn aggregation. Drug repositioning could also be successful
645 in the case of entacapone and tolcapone, inhibitors of catechol-*O*-methyltransferase and
646 anti-Parkinsonian drugs, which are also available as aggregation inhibitors against A β and α Syn
647 aggregation. Development of nucleic acid binders will help obtain inhibitors of amyloid
648 coaggregation with nucleic acids, as demonstrated by CGG repeat RNA binders in FXTAS
649 therapeutics (172), and will also help clarify the molecular basis of amyloid recognition by
650 nucleic acids. The increasing use of next-generation sequencing to sequence mRNA transcripts
651 exhaustively suggests that additional data will be forthcoming.

652 Finally, the antagonistic function of nucleic acid medicine targeting nucleic acids, which
653 can participate in the formation of amyloid coaggregates, is a promising lead in
654 neurodegeneration therapeutics (9). Nucleic acid aptamers are potential candidates, especially in
655 the anti-amyloid field, and various aptamers against amyloidogenic proteins, including
656 oligomeric assembly, have been developed. We expect that these aptamers will suppress the
657 interactions of amyloid coaggregates and further aggregation by shifting equilibrium to the
658 disaggregated state. There are several delivery systems to the CNS, such as exosomes and

659 nanoliposomes, and each system has advantages and limitations; future study should address the
660 current shortcomings in the target specificity of aptamers, which could improve the delivery of
661 aptamers into the CNS, as was demonstrated successfully for a DNA aptamer against α Syn
662 (F5R1) using a model mouse with a synucleinopathy (201). Recent advances in both
663 computational and experimental approaches suggest that antinucleic acid therapeutics will be
664 realized in the near future. Overall, we have highlighted new mechanistic insights into amyloid
665 coaggregation, which could pave the way for future studies on the underlying mechanisms and
666 causes of neurodegenerative diseases.

667

668 **Competing interests**

669 The authors declare that they have no competing interests.

670

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673 **Author contributions**

674 K.M. and K.O. completed the literature search; planned, wrote, and revised the manuscript; and
675 prepared the figures and tables.

676 **DATA AVAILABILITY STATEMENT**

677 This is not applicable for this review.

678

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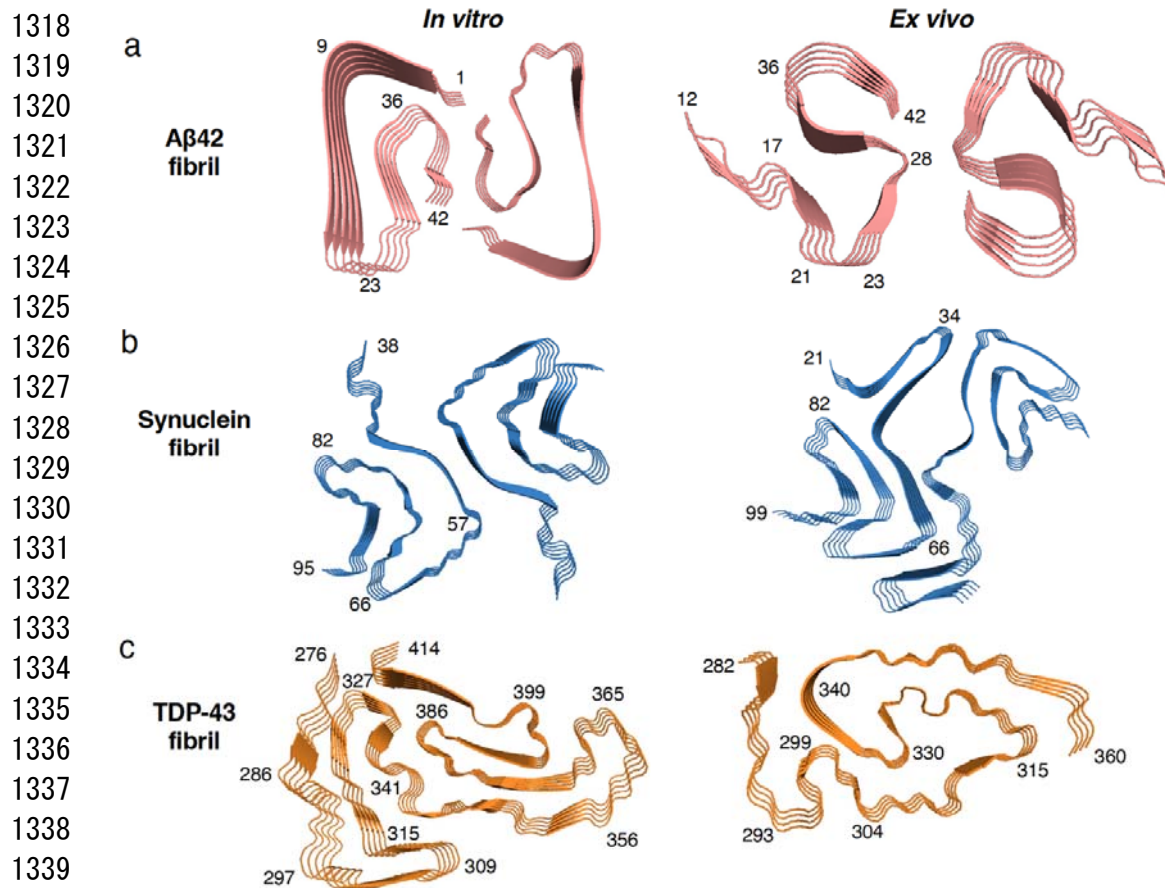


Fig. 1 Comparison of the *in vitro* and *ex vivo* structures of amyloid fibrils via cryo-EM analysis.
(a) A β 42 (PDB ID: 5OQV for *in vitro*, 7Q4M for *ex vivo*), (b) α Syn (PDB ID: 6H6B for *in vitro*, 6XYO for *ex vivo*), (c) TDP-43 (PDB ID: 7KWZ for *in vitro*, 7PY2 for *ex vivo*). PDB, Protein Data Bank. Because *ex vivo* fibrils formed from A β 42 and α Syn contained multiple conformers, the representative one is shown.

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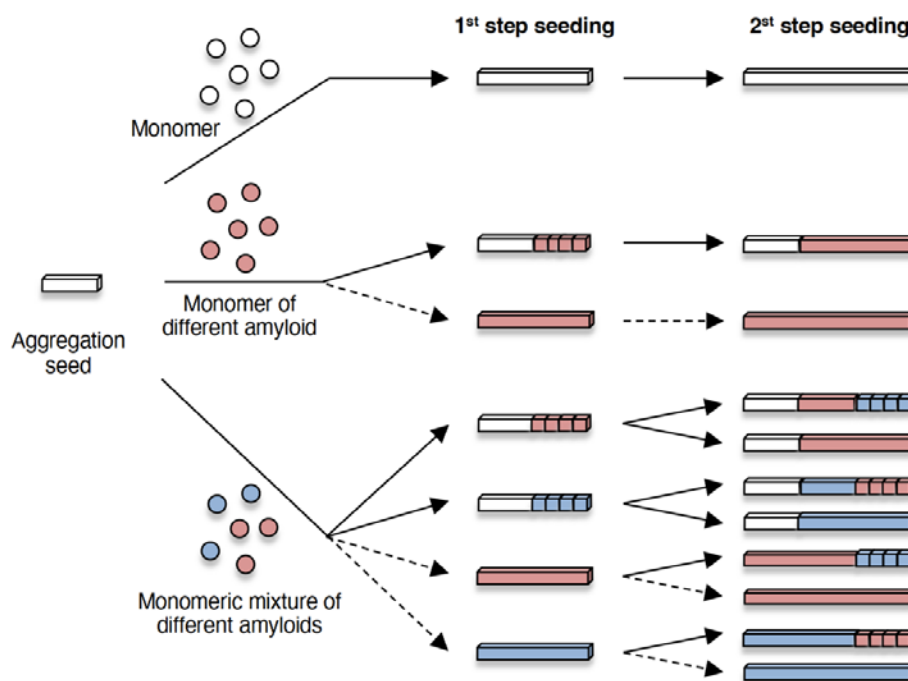


Fig. 2 Schema of the cross-seeding aggregation model. The original model proposed by Jarrett and Lansbury (10) for a single amyloidogenic proteins was applied into a combination of multiple amyloids. Various patterns of seeding aggregation depending on the number of amyloids can occur in the presence of seeds as templates of propagation.

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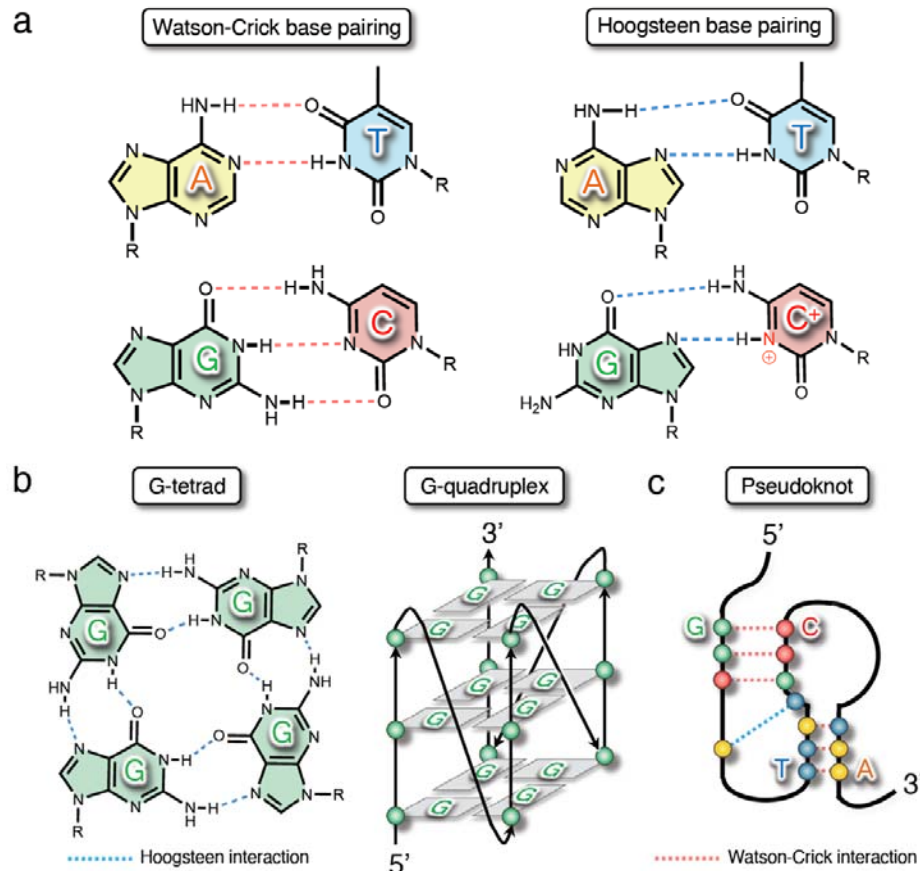
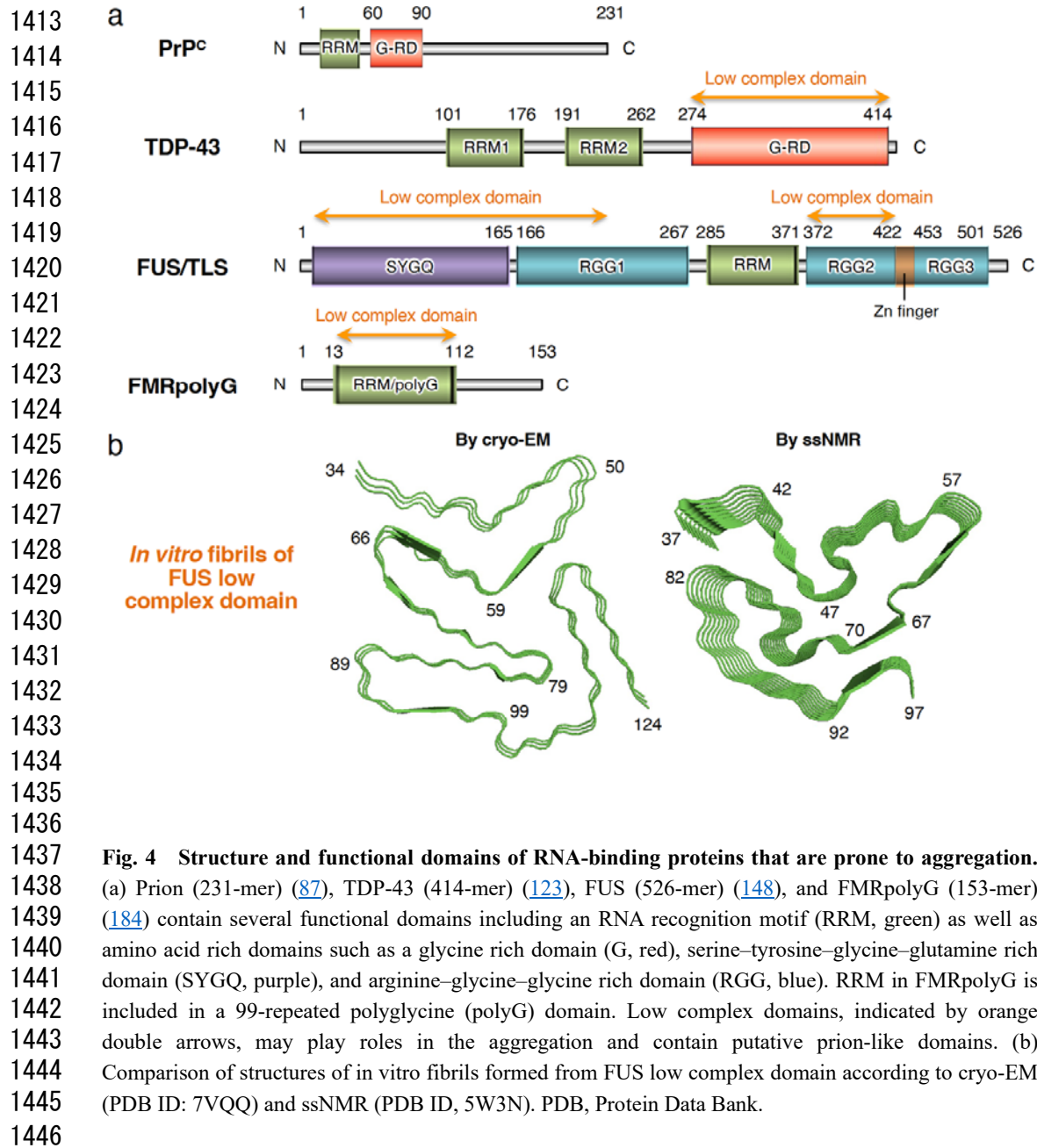


Fig. 3 Watson-Crick-type and Hoogsteen-type base pairing for natural nucleotides and G-quadruplex formation involved in protein aggregation. (a) Watson-Crick interactions have three hydrogen bonds between guanine and cytosine and two hydrogen bonds between adenine and thymine. Hoogsteen interactions have two hydrogen bonds between guanine and cytosine and two hydrogen bonds between adenine and thymine, which can be induced by rotating the adenine or guanine by 180° around the glycosidic bond. Although the latter is a minor base pairing of natural nucleotides compared with the former, it allows the formation of triplex and quadruplex structures of DNA or RNA that may contribute to its binding ability with functional proteins and protein aggregation. (b) G-quadruplex structure formed in DNA or RNA by guanine rich sequences induced from Hoogsteen-type interaction. Four guanine bases can form a square planar structure (G-tetrad), and the G-quadruplex is composed of two or more G-tetrads through a stacking process. An intramolecular parallel G-quadruplex that forms three separate parallel G-tetrads stacked 5' to 3' with three loops is shown. (c) Pseudoknot structure formed in DNA or RNA by Watson-Crick interaction and Hoogsteen interaction. R denotes the ribose-phosphate backbone.



1447 **Table 1. Characteristics of cross-seeding of amyloidogenic proteins.**

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Amyloid	Cross-seeding amyloid	Testing method	Effects	Year	Ref.
Aβ – tau					
Synthetic A β 42	Tau (mice)	A β -injected P301L-tau Tg mice	NFT \uparrow	2001	(37)
A β (mice)	Tau (mice)	JNPL3 x Tg2576 mice	NFT \uparrow , A β plaques \uparrow	2001	(38)
Brain A β	Brain tau	AD patients	A β -tau complex in NFT	2006	(36)
Aβ – αSyn					
A β (mice)	α Syn (mice)	hSYN x J9 Tg mice	Motor deficit \uparrow , α Syn inclusion \uparrow	2001	(44)
Brain A β	Brain α Syn	AD, PD, DLB patients	Colocalization	2006	(45)
Brain A β	Brain α Syn	DLB patients, Tg mice	Interaction in membrane	2008	(47)
Fibrillar, oligomeric A β 40, A β 42	Fibrillar, oligomeric α Syn	Th-T, TEM (<i>in vitro</i>)	Fibril \uparrow	2012	(48)
Tau – αSyn					
Tau (mice)	Synthetic α Syn	α Syn-injected P301S-tau Tg mice	Tau inclusion \uparrow	2013	(49)
Brain tau	Synthetic α Syn	Injection of α Syn to 5xFAD Tg mice	P-tau and P- α Syn ^a inclusion \uparrow , A β plaques \uparrow	2020	(50)
Aβ – IAPP					
Synthetic A β 42	IAPP (mice)	A β -injected IAPP Tg mice	Colocalization	2015	(56)
Oligomeric A β 42	Oligomeric IAPP	Th-T, TEM, SDS-PAGE, ^b MTS (<i>in vitro</i>)	Fibril \uparrow , oligomer \uparrow , cytotoxicity \uparrow	2020	(55)
αSyn – IAPP					
Brain α Syn	Brain IAPP	PD, DLB patients	α Syn deposits in pancreatic β cells	2018	(57)
Synthetic α Syn	IAPP (mice)	α Syn-injected IAPP Tg mice	Amyloid deposits in pancreatic β cells	2020	(58)
Synthetic α Syn fragment	Synthetic IAPP	Th-T, AFM, MTT (<i>in vitro</i>)	Fibril \uparrow , cytotoxicity \downarrow	2022	(59)

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^aphosphorylated-tau and phosphorylated- α Syn

^bsodium dodecyl sulfate–polyacrylamide gel electrophoresis

1453 **Table 2. Characteristics of cross-inhibition of amyloidogenic proteins.**

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Amyloid	Cross-seeding amyloid	Testing method	Effects	Year	Ref.
TTR – Aβ					
TTR (mice)	Aβ (mice)	mAb(anti-TTR)-injected Tg2576 mice	Aβ plaques↑, tau phosphorylation↑	2004	(62)
TTR (mice)	Aβ (mice)	APP ^{swe} /PS1Δ9 x TTR ^{+/-}	Insoluble Aβ↑, Aβ plaques↑	2007	(63)
Recombinant TTR	Recombinant Aβ40	Th-T, TEM (<i>in vitro</i>)	Nucleation↓	2018	(64)
Recombinant TTR	Recombinant Aβ40	AFM, DLS (<i>in vitro</i>)	Nucleation↓, cytotoxicity↓	2020	(65)
TTR – IAPP					
Recombinant TTR	Synthetic IAPP	Th-T	Nucleation↓, elongation↓	2021	(66)
TTR (mice)	Aβ (mice)	HFD ^a -treated <i>App</i> ^{NL-F/NL-F} mice	TTR expression↓, Aβ plaques↑	2021	(67)
BRICHOS – Aβ					
Bri2, Bri3 (recombinant, mice)	Aβ (synthetic, mice, human)	TgAβPPare mice, AD brain, Th-T	Fibril↓, Bri 2, Bri3 bound Aβ (mice, human)	2018	(69)
Expressed BRICHOS (Drosophila)	Expressed Aβ (Drosophila)	Coexpressed Drosophila	Longevity↑, Aβ plaques↓, locomotor loss↓	2014	(70)
Recombinant Bri2, Bri3	Recombinant Aβ42	In silico, DLS, SDS-PAGE	Oligomer↓ (Bri2 > Bri3)	2020	(74)
Recombinant proSP-C	Recombinant Aβ42	Th-T, TEM, cryo-EM, electrophysiology	Oligomer↓, cytotoxicity (gamma oscillation)↓	2015	(72)
Recombinant Bri2	Recombinant Aβ42	TEM(3D), SDS-PAGE	Oligomer↓	2017	(73)
Expressed proSP-C and Bri2 (Drosophila)	Expressed Aβ (Drosophila)	Coexpressed Drosophila	Longevity↑, locomotor loss↓, impaired eye phenotype↓	2016	(71)
BRICHOS – IAPP					
Expressed Bri2 (Drosophila)	Expressed IAPP (Drosophila)	Coexpressed Drosophila, Bri2-siRNA-treated EndoC-βH1 cell	Longevity↑, cell death↑	2018	(75)

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^ahigh fat diet

1458 **Table 3. Binding characteristics of amyloidogenic proteins and nucleic acids.^a**

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Amyloid	Nucleic acid	K_D (nM)	Testing method for K_D	Binding region of amyloid	Year	Ref.
Prion						
murine PrP ^C (FL ^b)	plasmid DNA	250	light scattering	N.D. ^c	1999	(93)
murine PrP23-231	short dsDNA	25	light scattering	N- and C-terminal domains	2001	(94)
hamster PrP23-231	short dsDNA	90	fluorescence polarization	N- and C-terminal domains	2006	(98)
ovine PrP (FL)	D12 DNA (G4)	62	SPR	N- and C-terminal domains	2013	(96)
Human PrP ^C (FL)	tRNA	1,700	fluorescence polarization	N- and C-terminal domains	2018	(104)
Human PrP ^C (FL)	Cm47 RNA (pseudoknot)	1,500	fluorescence polarization	N- and C-terminal domains	2018	(104)
TDP-43						
human TDP-43 (FL)	(UG) ₆	27	EMSA	RRM1	2005	(128)
TDP101-261	RNA34nt-(UG) ₆	5.3	EMSA	RRM1	2013	(130)
human TDP43 (FL)	ssDNA-(TG) ₁₂	90	FCS ^d	N.D.	2018	(132)
TDP-43 (RRM1+RRM2)	ssDNA-(TG) ₁₂	51	ITC	RRM1 loop3, RRM2 pocket around V220	2021	(131)
human TDP43 (FL)	RNA14nt	32	ITC	N.D.	2022	(129)
FUS/TLS						
TLS (FL)	ggugRNA25nt	250	EMSA	RGG ^h repeats in RRM	2001	(158)
FUS (FL)	mRNA200nt	56	EMSA	RRM, RGG rich domain (Zn finger)	2015	(162)
TLS/FUS (FL)	r(UUAGGG) ₄ (G4)	6.2	EMSA	RGG rich domain	2018	(163)
RRM in FUS	hnRNP A2/B1 stem-loop RNA	9,200	ITC	RRM, RGG rich domain (Zn finger)	2019	(166)
TLS/FUS (FL)	PSD-95 ⁱ GQ2 (G4)	28	steady-state fluorescence spectroscopy	RGG rich domain	2020	(164)
FUS (FL)	PSD-95 RNA(G4)	3.2	SPR	RGG rich domain	2021	(165)
FMRpolyG						
FMRpolyG	RNA	N.T. ^e	N.T. ^e	(CGG) ₉₉	2021	(184)

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^aOnly amyloidogenic proteins whose K_D values were determined are shown when many examples were investigated.

^bfull length

^cnot determined

^dfluorescence correlation spectroscopy

^enot tested