Nucleated Conformational Conversion and the Replication of Conformational Information by a Prion Determinant

Tricia R. Serio,1* Anil G. Cashikar,2* Anthony S. Kowal,2 George J. Sawicki,2 Jahan J. Moslehi,2 Louise Serpell,4 Morton F. Arnsdorff,3 Susan L. Lindquist1,2†

Prion proteins can serve as genetic elements by adopting distinct physical and functional states that are self-perpetuating and heritable. The critical region of one prion protein, Sup35, is initially unstructured in solution and then forms self-seeded amyloid fibers. We examined in vitro the mechanism by which this state is attained and replicated. Structurally fluid oligomeric complexes appear to be crucial intermediates in de novo amyloid nucleus formation. Rapid assembly ensues when these complexes conformationally convert upon association with nuclei. This model for replicating protein-based genetic information, nucleated conformational conversion, may be applicable to other protein assembly processes.

In *Saccharomyces cerevisiae*, the prion element [PSI+] causes heritable changes in translational fidelity (1) through self-perpetuating changes in the state of Sup35 (2–4), a component of the translation termination factor. Together the glutamine-rich NH2-terminal (N) and highly charged middle (M) regions of Sup35 adopt distinct physical states (either soluble or insoluble higher order complexes), and these produce the distinct translational activities of [psi-] and [PSI+] strains (3, 4). Sup35 complexes replicate their conformational state, and thereby perpetuate [PSI+], by converting newly made Sup35 to the same state (5). In vitro, highly ordered NM complexes (fibers) promote the conversion of soluble NM to the same state, closely modeling the in vivo process (5). Lysates of [PSI+], but not [psi-], cells accelerate NM polymerization (3, 6), and mutations that enhance or inhibit [PSI+] propagation enhance or inhibit NM conversion (5, 7, 8) in a complementary way. But the conformational conversion mechanism that replicates the protein-based genetic information of [PSI+] is unknown.

The three most prominent models proposed for self-propagating changes in protein state fundamentally differ in the role that preexisting polymers play in converting soluble, S-state protein to assembled, A-state protein and in the nature of the rate-limiting step (Fig. 1). For templated assembly (TA) (9, 10), A-state complexes act as templates for S→A conformational conversion (the rate-limiting step) during assembly. Conformational conversion is also rate-limiting for monomer-directed conversion (MDC) (11), but S monomers convert to A monomers in solution, and assembly is a consequence, not a cause, of conversion. For nucleated polymerization (NP) (12–14), the S and A states are in equilibrium in solution, but the A state is rare and unstable. The rate-limiting step is not conformational conversion but chance association of a sufficient number of A monomers to form a stable, polymerization-competent nucleus. A-state monomers, generated by the S→A equilibrium, are stabilized by joining the nucleus, as in seeded crystallization.

**Nucleation by solid-phase A-state protein.** To test these models for NM conformational conversion and assembly, we first characterized the A state. During fibrillization, NM became resistant to cleavage with proteases: Chymotrypsin resistance appeared concomitant with assembly (Fig. 2A); a modest V8 resistance appeared later (Fig. 2B). Because chymotrypsin cleavage sites are restricted to N, and V8 sites to M, these observations support the hypothesis that N directs fiber formation with M on the outside (5). By x-ray diffraction, NM fibers had a prototypical amyloid, cross-β structure (Fig. 2C) (15). Fibers resisted solubilization in 8 M urea and 2% (w/v) SDS at room temperature (Fig. 2D), but were solubilized in 2% (w/v) SDS when boiled for 10 min. Fibers were also rigid and easily fractured into short pieces by sonication (Fig. 2, E and F).

Sonication greatly enhanced the nucleating activity of NM fiber solutions (Fig. 3 and Table 1) (16, 17). It did not do so by increasing the concentration of a soluble accelerator, because nucleating activity was removed by centrifugation at 100,000g for 10 min (17). Nor did it change the biochemical nature of the protein (18). Furthermore, fracturing the fibers did not simply keep nucleating material in suspension; even unassembled fibers remained in suspension during these experiments (17). When fresh NM was added, fragments converted to long, unclumped fibers (17). Thus, assembled protein is the species that accelerates conversion—eliminating MDC for NM—and fiber ends, rather than lateral surfaces, appear to be crucial for nucleation (seeding).

**Simultaneous acquisition of structure and assembly.** To investigate the relation between structural conversion and assembly, we simultaneously analyzed (i) assembly, by sedimentation, light scattering, transmission electron microscopy (TEM), and atomic force microscopy (AFM); (ii) structural change, by circular dichroism (CD), limited proteolysis, and dye binding; and (iii) stability, by SDS solubility. In unseeded reactions, the half-point of assembly, determined by centrifugation and light scattering, was reached at ~50 hours (Fig. 3). By ~50 hours, NM had gained 50% of its maximum ability to bind Congo red (CR) (Fig. 3); the random-coil signal in the CD spectrum (approaching 200 nm) was substantially reduced (17), fibers became abundant by microscopy (17), and nearly 50% of the protein had become resistant to cleavage with chymotrypsin (19) and insoluble in 2% (w/v) SDS (Fig. 3) (20). 8-Anilino-1-naphthalene sulfonate (ANS) fluorescence (21) also changed during fibrillation. It did not reveal, as it commonly does, a structural intermediate, but rather the accessibility of a hydrophobic environment in mature NM fibers (22). ANS fluorescence reached a maximum at the same time as other traits of the A state (Fig. 3).

A similar congruence between the acquisition of structure and assembly was observed in seeded reactions, both with unfractioned and with sonicated fibers (Fig. 3). The coincident acquisition of all A-state structural traits in reactions occurring over vastly different time frames indicates that the A structure is either attained concomitantly with polymerization, as predicted by TA, or is too rare to detect in solution during any of these time courses. **Kinetic analysis of the lag and assembly phases.** We next examined the concentration dependence of the lag phase, during which nuclei form, and the assembly phase, in which un polymerized protein joins these nuclei (23, 24). The NP and TA models make different predictions for the concentration dependence of both phases. Nucleation by NP depends on the number of A-state monomers that must come together.
to form a stable nucleus. The lag phase, therefore, varies exponentially with that number (4F). Nucleation by TA presumably depends on the rate of S→A conversion and thus should vary directly with concentration. The lag phase for polyhistidine-tagged NM varies little with protein concentration (5, 7). Because this lag affects polymerization (see below), we investigated the effects of concentration on the lag phase using unmodified NM. Over a 500-fold range (0.1 to 50 μM), a lag phase was detected (25), and even at the highest concentration this phase was ~20 hours, demonstrating a rate-limiting step in nucleus formation (Fig. 4A). However, the lag phase varied over this concentration range by less than 10-fold (Fig. 4A), a conundrum for either the NP or the TA model.

Assembly rates, according to NP, are determined by the rate at which soluble A-state protein is generated by the S→A equilibrium and collides with preformed nuclei (4F). The concentration of A and the rate of assembly should increase directly with total soluble protein. For TA, assembly is limited by the rate at which S-state protein, interacting with an A-state nucleus, is conformationally converted to regenerate the nucleating surface. Only in the extreme case where nucleating surfaces are in excess will assembly rates vary with the concentration of soluble protein (26).

Post-lag phase assembly times in unseeded reactions were constant over a broad range of concentrations but increased sharply below 5 μM (Fig. 4B). In these reactions, at the end of the lag phase, >90% of the protein was soluble and structurally unconverted (Fig. 3). Similarly, in seeded reactions containing a fixed quantity of soluble NM (5 μM) and different quantities of sonicated fibers, the rate of conformational change was proportional to the quantity of fibers added only until the seed concentration reached 1% (w/w) (Fig. 4C). At this point, soluble protein was in excess over fiber ends by ~50,000-fold (17). These experiments reveal a second conundrum for either NP or TA: Assembly rates showed little concentration dependence over a broad range but reached a limit when monomer concentrations remained in vast excess over polymerization surfaces.

**Evidence for an oligomeric intermediate in nucleation and assembly.** Evidence for an intermediate in fiber formation, which might explain these conundrums, was obtained when solutions of NM were preincubated for various times before the addition of fibers. When fibers (0.15% w/w) were added immediately to freshly solubilized NM, assembly proceeded as a first-order reaction (Fig. 4D); when they were added to NM solutions after an 8-hour preincubation, the reaction proceeded in two stages (Fig. 4D), with an initial phase 40-fold faster than the second phase. The contribution of the fast phase to the assembly process increased with the length of the preincubation, until the preincubating solutions themselves had initiated fiber formation (27). Thus, an intermediate accumulated during the lag phase that promoted rapid NM assembly and was quickly consumed once assembly began.

The fast phase of assembly was not observed when fibers were added to NM solutions preincubated for 8 hours in the same buffer but with 1 M NaCl (27). The CD spectra of NM solutions was similar in 1 and 0.15 M NaCl, suggesting that secondary structure was not substantially altered (27). However, solutions scattered more light in 0.15 M than in 1 M NaCl immediately after NM was solubilized (Fig. 4E).

This early signal was due not to the formation of fibers (Fig. 3) but to oligomeric complexes (17). Notably, salt and other conditions (4°C, His10 tag) that increased or decreased initial light scattering (Fig. 4E) increased or decreased the rate of fibrillization in a complementary way (Fig. 4F), providing independent evidence that nucleus formation and assembly are mediated by an oligomeric intermediate.

If fibrillization were mediated by a complex composed of a defined number of subunits that interact through specific surfaces, the rate should be exponentially dependent on soluble protein concentration. Curiously, the equilibrium between monomers and oligomers was not affected by concentration in the range of our experiments. Initial light scattering from NM in physiological salt increased in simple proportion to concentration over a 100-fold range (0.3 to 35 μM) (27), and oligomers were always much less abundant than monomers (28).

**Structural and morphological characterization of NM oligomers.** During assembly, spherical, variably sized oligomeric complexes were detected by several types of microscopy (Fig. 5). By scanning transmission electron microscopy (STEM), their mass was 1.6 × 106 daltons (SD = 0.8, n = 359), corresponding to complexes of ~20 to 80 NM monomers (Fig. 5, B and C). Notably, fibers, which appeared smooth by TEM (Fig. 2E), were shown by AFM to be composed of segments with an average height of ~4 nm (Fig. 5, A, D, and E). Most complexes were in the range of 2 to 4 nm

---

**Table 1.** Time course reactions with 5 μM NM alone (unseeded) or in the presence of 2% (w/w) intact (seeded) or sonicated fibers either unrotated or rotated as for Fig. 2, quantified by CR binding as for Fig. 3. L½, lag time; A½, 50% assembly time.

<table>
<thead>
<tr>
<th></th>
<th>Unseeded</th>
<th>Unsonicated seed</th>
<th>Sonicated seed</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>L½ (hours)</td>
<td>A½ (hours)</td>
<td>A½ (hours)</td>
</tr>
<tr>
<td>Unrotated</td>
<td>33</td>
<td>29</td>
<td>3</td>
</tr>
<tr>
<td>Rotated</td>
<td>0.33</td>
<td>0.2</td>
<td>0.2</td>
</tr>
</tbody>
</table>

**Fig. 1.** Models for conversion and assembly. Smooth circles, A-state protein; jagged circles, S-state protein; and open jagged circles denote possible conformational differences between soluble monomers and oligomers. See text for explanations of the models.
Observations indicate that oligomers are less rigidly structured than fibers.

Rotation accelerates both the lag and assembly phases. An on-pathway role for oligomers is supported by the effects of modest agitation. Both the lag phase and the assembly phase were accelerated ~100-fold when reactions were rotated at 60 rpm (Table 1). Rotation at a slower speed produced a more modest increase in rate, but again both phases were affected similarly (8). Rotation did not fracture fibers to create new ends (30), as previously postulated (7). We suggest that rotation accelerates polymerization by dissociating overly large complexes and/or increasing the collision of eligible complexes with each other and with fiber ends. Indeed, rotation eliminated the strong concentration dependence of assembly below 5 μM (Fig. 4B).

Rate-limiting conversion. Because rotation and seeding accelerate NM assembly by different mechanisms (altering fluid dynamics versus providing preformed ends for polymerization), combining the two might be expected to produce an even greater increase in rate. Rotation did increase assembly rates in seeded reactions, but only by a factor of 15, not 100 (Table 1). Substituting sonicated fibers for unfractured fibers produced no further increase (Table 1). Together with the data described above, these observations strongly suggest that assembly rates under these conditions approached a limit imposed by the time required for conformational conversion.

A model for NM amyloid formation. Our investigations of the S→A transition and its relation to assembly suggest a mechanism for the replication of NM’s conformational information, which is the basis of its capacity to serve as a genetic element. This model, nucleated conformational conversion (NCC) (Fig. 1), incorporates aspects of NP and TA with additional features. Conformational conversion is nucleated by a complex of A-state proteins and is driven by assembly. Nuclei form by conformational rearrangements of less structured protein within the context of a structurally molten, oligomeric intermediate. When nuclei are present assembly proceeds rapidly, with less structured NM acquiring the A state through a templating or induced-fit mechanism at fiber ends. Other species might be competent for polymerization, but we suggest that assembly is most efficient with oligomers (31).

We propose that NM oligomers are initially micelle-like: They form through still fluid associations, involve variable stoichiometries, and are in equilibrium with monomers as well as smaller and larger complexes. Several lines of evidence directly support an on-pathway role for these unusual intermediates. Moreover, they provide a simple explanation for two aspects of the kinetic analysis that are otherwise baffling: (i) because overly large ineligible complexes increase proportionately with eligible complexes at higher concentrations, both the lag and the assembly phases show little variation in rate over a broad range of protein concentrations; and (ii) because oligomers are the limiting species for rapid assembly, as the number of nuclei increases, assembly rates are limited even when total soluble protein is in vast excess to fiber ends.
Fig. 4. Analysis of the lag and assembly phases. NM (5 μM) resuspended in CRBB in the absence (A, B, E, and F) or presence (C and D) of sonicated fibers. (A) Inverse lag time (1/L) versus log of monomer concentration ([M]). (Left) Ellipticity at 220 nm was monitored in reactions stirred with a magnetic bead in 1-cm quartz cuvettes containing 2 ml of NM solutions at 0.1, 0.15, 0.2, 0.5, 1, and 5 μM in a Jasco J-715 Spectropolarimeter. Slope = 0.4. (Right) CR binding for reactions at 1.1, 2.5, 5, 10, 20, and 50 μM NM, quantified as in Fig. 3. Slope = 0.7. (B) Assembly time (Ay) versus concentration for undisturbed reactions (hours) (O) or reactions rotated as in Fig. 2 (min) (C). (C) Ellipticity at 220 nm, monitored for undisturbed NM (5 μM) with 0.1, 0.15, 0.2, 0.5, 1, 2, 4, and 7% (w/w) sonicated fibers added just before the start of data collection. Rates were calculated by fitting to a single exponential equation with GraFit 4 (Erithacus Software Limited, Surrey, UK, 1998). (D) Ellipticity at 220 nm, monitored as in (C) with sonicated fibers (0.15% w/w) added immediately (left) or after 8 hours at 25°C (right). Rate = 0.017 ± 0.001 mdeg/min (left), 0.527 ± 0.060 mdeg/min, and 0.013 ± 0.007 mdeg/min (right). Data were collected immediately after addition of fibers. (Top) Kinetics were fit to single or double exponential equations as in (C). (Bottom) Residuals (Res) from the fitting. (E) Light scattering as in Fig. 3 immediately after dilution into buffer. Lane 1: NM, 25°C, CRBB; lane 2: NM, 4°C, CRBB; lane 3: NM, 25°C, 1 M NaCl, 5 mM potassium phosphate (pH 7.4); and lane 4: His10-tagged NM, 25°C, CRBB. (F) Unseeded reactions were performed and analyzed by CR binding as in Fig. 3. Conditions: ( ) NM, 4°C, CRBB; ( ) His10-tagged NM, 25°C, CRBB; ( ) NM, 25°C, CRBB; ( ) NM, 25°C, 5 mM potassium phosphate (pH 7.4), 1 M NaCl. Each experiment was performed three times with similar results.

Fig. 5. Microscopy analysis of NM assembly. (A) AFM deflection image of a carbon-coated grid prepared for TEM as in Fig. 2. The sample is from an unseeded reaction of His10-tagged NM after 20 hours of incubation in CRBB at 25°C, imaged in contact mode with a Digital Instruments Nanoscope III by using standard 200-μm thin-featured Si3N4 cantilevers. (B and C) STEM images of His10-tagged NM 4 hours after resuspension in CRBB obtained as described (42). (D) AFM deflection image of uncleaved reactions of untagged NM after overnight incubation at 25°C. Fifty microliters of a 1:500 dilution of 5 μM untagged NM in CRBB was spotted onto 15-mm mica disks and adsorbed for 30 s. Disks were washed with 100 μl of CRBB and then 100 μl of water (three times). Samples were air-dried and imaged as in (A). (E) AFM surface plot of the height data in (D). (F) TEM image of His10-tagged NM 3 hours after resuspension in NM as in Fig. 2. Bars, 100 nm.

References and Notes
1. B. Cox, Hereditas 20, 505 (1965).
16. The quantity of fibers added to the assembly reactions as seed was undetectable by the methods we used.
17. T. R. Serio et al., data not shown.
18. Sonicated fibers remained resistant to SDS solubilization and chymotrypsin, sedimented, bound Congo red and ANS, retained β-sheet-rich CD spectrum, and scattered light (17).
19. 125I-immunoblotting allowed quantitation of chymotrypsin sensitivity; however, the resistance of NM to digestion with chymotrypsin was confirmed by Coomassie Brilliant Blue staining (17).
20. In three separate unseeded reactions, CR binding reached the half-maximal point at 53, 45, and 51 hours. In each case, the other traits appeared coincidentally with CR-binding capacity (17).
22. The existence of an ordered hydrophobic surface has also been detected by birefringence of CR bound to NM fibers (A. G. Cashikar and S. L. Lindquist, unpublished observation). It is unclear whether the CR and ANS binding sites are the same or different.
23. L3 is defined as the period of time in which CR binding is less than 10% maximal. A4 is defined as the interval of time required for CR binding to proceed from 10 to 90% maximal levels.
25. It could not be tested below this concentration because protein was lost to the sides of the tubes (17).
Hydrological Impact of Heinrich Events in the Subtropical Northeast Atlantic

Edouard Bard,¹,² Frauke Rostek,¹ Jean-Louis Turon,² and Sandra Gendreau²

Reconstructing the impact of Heinrich events outside the main belt of ice rafting is crucial to understanding the underlying causes of these abrupt climatic events. A high-resolution study of a marine sediment core from the Iberian margin demonstrates that this midlatitude area was strongly affected both by cooling and advection of low-salinity arctic water masses during the last three Heinrich events. These paleoclimatic time series reveal the internal complexity of each of the last three Heinrich events and illustrate the value of parallel studies of the organic and inorganic fractions of the sediments.

High-resolution paleoclimatic records show that the glacial climate was far more variable than previously thought. Studies of North Atlantic sediments show that the circulation of the surface and deep waters was repeatedly perturbed by massive surges and melting of icebergs. Although most researchers would agree that these so-called Heinrich (H) events originated from the Laurentide ice sheets (1–6), there is growing controversy about the precise behavior of other ice sheets during these events [see recent works on lithological indices (7, 8) and studies based on the Sr-Nd isotopic signatures of ice-rafted detritus (IRD) (9, 10)].

The influence of Heinrich events has sometimes been invoked to explain unusual high-frequency climatic events at low latitudes [e.g., (11)]. Abrupt decreases of sea surface temperature (SST) in phase with Heinrich events were detected in sediments of the Bermuda Rise (12) and the Mediterranean Sea (13). Further south, the tropical western Atlantic apparently warmed during H1 and the Younger Dryas (YD) (14). The strong, direct effect of melting icebergs is generally thought to be reflected alterations in the ratio of the prion protein to Hsp104 and other cellular factors, rather than in the intrinsic folding and conversion properties of Sup35.

Reconstructing the impact of Heinrich events outside the main belt of ice rafting is crucial to understanding the underlying causes of these abrupt climatic events. A high-resolution study of a marine sediment core from the Iberian margin demonstrates that this midlatitude area was strongly affected both by cooling and advection of low-salinity arctic water masses during the last three Heinrich events. These paleoclimatic time series reveal the internal complexity of each of the last three Heinrich events and illustrate the value of parallel studies of the organic and inorganic fractions of the sediments.