Solution conditions can promote formation of either amyloid protofilaments or mature fibrils from the HypF N-terminal domain

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Abstract

The HypF N-terminal domain has been found to convert readily from its native globular conformation into protein aggregates with the characteristics of amyloid fibrils associated with a variety of human diseases. This conversion was achieved by incubation at acidic pH or in the presence of moderate concentrations of trifluoroethanol. Electron microscopy showed that the fibrils grown in the presence of trifluoroethanol were predominantly 3–5 nm and 7–9 nm in width, whereas fibrils of 7–9 nm and 12–20 nm in width prevailed in samples incubated at acidic pH. These results indicate that the assembly of protofilaments or narrow fibrils into mature amyloid fibrils is guided by interactions between hydrophobic residues that may remain exposed on the surface of individual protofilaments. Therefore, formation and isolation of individual protofilaments appears facilitated under conditions that favor the destabilization of hydrophobic interactions, such as in the presence of trifluoroethanol.

Keywords: Aggregation; amyloid fibrils; HypF N-terminal domain; protofilaments; trifluoroethanol

A number of human diseases, including Alzheimer’s disease and the spongiform encephalopathies, are associated with the formation of stable protein aggregates known as amyloid fibrils. In each of these pathological states, a specific protein or protein fragment changes from its native soluble form into insoluble aggregates which subsequently develop slowly into fibrils accumulating in a variety of organs and tissues (Kelly 1998; Dobson 1999; Bellotti et al. 1999; Rochet and Lansbury, 2000). This phenomenon frequently has serious consequences and usually leads to premature death. Despite the variation encountered in the amino acid sequences and native structures of proteins associated with these diseases, the amyloid fibrils from different pathologies display common structural features. They typically appear to be formed from 2–6 unbranched protofilaments 2–5 nm wide that associate laterally or twist together to form fibrils with a 4–13 nm width (Shirahama and Cohen 1967; Merz et al. 1983; Serpell et al. 1995, 2000). Formation of fibrils with such morphology has also been observed widely in vitro from the proteins associated with disease (Shirahama et al. 1973; Glenner et al. 1971; Goldsberry et al. 1997; Harper et al. 1997; Conway et al. 1998).

Recently, it has been found that proteins other than those associated with diseases are capable of forming amyloid fibrils under appropriate conditions in vitro (Guijarro et al. 1998; Litvinovich et al. 1998; Chiti et al. 1999; Konno et al. 1999; Krebs et al. 2000; Ramirez-Alvarado et al. 2000; Villegas et al. 2000; Yutani et al. 2000; Fandrich et al. 2001; Pertinhez et al. 2001). It has been proposed that amyloid fibril formation can occur when the native globular fold of a protein is destabilized under conditions in which noncovalent interactions still remain favourable (Chiti et al. 2001).
1999). The ability for rational design of conditions promoting amyloid formation has implications for understanding the origin of amyloid formation in vivo because the latter can no longer be assumed to arise from the unique physical properties of a limited number of protein sequences (Chiti et al. 1999; Dobson 1999; Rochet and Lansbury 2000). In addition, the possibility of producing amyloid fibrils under controlled conditions provides us with the opportunity to investigate the molecular basis of amyloid formation using a wide range of proteins.

In this work, we investigate the conditions favorable for amyloid fibril formation using the N-terminal domain of HypF, a newly cloned globular protein factor participating in the maturation of the prokaryotic [NiFe] hydrogenase that is involved in hydrogen metabolism (Friedrich and Schwartz 1993). In addition to showing that this domain is capable of forming in vitro amyloid fibrils of the type associated with diseases, we will describe how solution conditions can be designed to promote selective formation of either the mature fibrils typical of pathological states or their constituents, the protofilaments that normally associate laterally or twist together to form the mature fibrils.

**Results**

Figure 1A shows the far-UV circular dichroism (CD) spectra of the purified HypF N-terminal domain acquired in the presence and absence of 5 M GdnHCl (guanidine hydrochloride). Unlike the CD spectrum acquired in the presence of GdnHCl, the spectrum of the protein acquired under native conditions is consistent with a well-structured protein with an α/β topology. Figures 1B, C show the change of mean residue ellipticity at 222 nm on the increase of urea concentration and temperature, respectively. The urea and heat-denaturation curves reveal single sharp transitions, indicating that this protein fragment has a globular and well-defined three-dimensional structure under native conditions. The analysis of the urea denaturation curve yielded values of 29±3 kJ/mol, 7±1 kJ/mol per mole, and 4.2±0.2 for the $\Delta G(H_2O)$, $m$, and $C_m$ values, respectively. Unlike the urea-denatured protein, the heat-denatured domain was not capable of recovering native-like ellipticity when cooled to room temperature, indicating that the thermal denaturation process is irreversible. Therefore, determination of the thermodynamic parameters characterising the unfolding thermal reaction was not attempted.

To induce formation of amyloid fibrils, the HypF N-terminal domain was incubated under different solution conditions including (i) 20 mM citrate buffer (pH 3.0), (ii) 50 mM acetate buffer and 15–40% (v/v) trifluoroethanol (TFE) at pH 5.5, and (iii) 50 mM acetate buffer (pH 5.5) at a temperature of 50°C, which was chosen as the highest temperature at which the protein is completely native. Protein samples incubated under non-denaturing conditions were also prepared as controls. The samples, containing the protein at concentrations ranging from 0.1 to 0.8 mg/ml, were left at room temperature (except those incubated at 50°C) for 1–4 weeks. Electron micrographs acquired from samples incubated at pH 3 reveal the presence of many fibrils (Fig. 2A–C). Analysis of $\sim 100$ fibrils from three different specimens identified three different types of fibrils, each displaying a characteristic width: 3–5 nm, 7–9 nm, and 12–20 nm. Figure 2B shows an area of a specimen in which the three different types are all present (numbered 1, 2, and 3, respectively). An 8 nm fibril shown in the figure (num-
Fig. 2. Electron micrographs of amyloid fibrils formed from the N-terminal domain of HypF. Fibrils were grown for one month at room temperature by incubating the domain, at a concentration of 0.3 mg/mL, in 20 mM citrate buffer (pH 3.0) (A–C) and in 50 mM acetate buffer (pH 5.5) in the presence of 30% (v/v) TFE (D–F). The numbers in (B) and (F) indicate fibrils with diameters of 3–5 nm (1), 7–9 nm (2) and 12–20 nm (3). The pictures obtained from samples incubated at low pH (A–C) show that the 7–9 and 12–20 nm fibrils are abundant, whereas the 3–5 nm fibrils are rare. By contrast, the 3–5 nm and 7–9 nm fibrils are highly populated in samples containing TFE (D–F), whereas those with larger widths are virtually absent.

The aggregates of the HypF N-terminal domain formed in the presence of TFE were analysed further with two colorimetric assays specific for amyloid fibrils. Samples incubated for one month in the presence of moderate concentrations of TFE cause the enhancement of fluorescence of Thioflavine T (ThT). This enhancement is more than 30 times higher than that observed on addition of a protein sample incubated in the absence of TFE (Fig. 3). The absorption peak of Congo red also undergoes a red-shift when aliquots of the protein solutions incubated under these conditions are added to the dye (Fig. 4). An increase of the absorbance values throughout the whole range of wavelength investigated here is also observed when the protein aggregates are added to the dye. This increase clearly originates from the light scattering by the protein aggregates; indeed, the spectrum of the aggregated protein in the absence of Congo red yields absorbance values higher than zero (Fig. 4, dotted line). Subtraction from the experimentally measured spectrum of the aggregates in the presence of Congo red of spectra of the aggregates alone and Congo red alone yields a difference spectrum with a maximum at 530–540 nm (Fig. 4, inset). Such a result is expected only if ordered amyloid-like aggregates are present in the solution (Klunk et al. 1989). Figure 5 shows the time-course of the change of ThT fluorescence as the protein aggregates form at pH 3 or in the presence of TFE. Whereas at acidic pH the aggregates develop after a lag phase of several days, in the presence of TFE the maximum ThT fluorescence is reached after a few minutes (Fig. 5, inset). The observed kinetics...
indicate that the prevalence of fibrils with larger diameters in the samples incubated at low pH cannot be attributed to a more rapid rate of the fibril formation process.

The samples incubated at high temperatures show the presence of visible protein aggregates that increase considerably the fluorescence of ThT. However, such a ThT fluorescence enhancement was observed only at protein concentrations >0.3 mg ml⁻¹ and was about one third of that observed at low pH or in the presence of TFE. Moreover, these aggregates do not have a fibrillar appearance under electron microscopy, appearing as large protein aggregates devoid of any geometric form. Nor do they cause the characteristic red shift of the Congo red dye. Therefore, the aggregated material observed under these conditions was not investigated in any further detail. The ability of the amorphous aggregates to bind to ThT indicates that ThT binding is not completely specific for amyloid fibrils as found for Congo red (Khurana et al. 2001) We do not exclude, however, the possibility that fibril formation can occur at temperatures >50°C or at this temperature value under different conditions of pH or ionic strength. The samples used as negative controls and containing the N-terminal domain of HypF in acetate buffer (pH 5.5) did not show the presence of any aggregates under electron microscopy, nor did they modify the fluorescence spectrum of ThT or the absorption spectrum of Congo red.

Discussion

As with other protein systems investigated so far (Guijarro et al. 1998; Litvinovich et al. 1998; Chiti et al. 1999; Konno et al. 1999; Goda et al. 2000; Krebs et al. 2000; Ramirez-Alvarado et al. 2000; Villegas et al. 2000; Yutani et al. 2000; Fandrich et al. 2001; Pertinhez et al. 2001), amyloid formation by the HypF N-terminal domain in vitro was achieved by destabilizing the native state of the protein under conditions in which non-covalent interactions still remain favorable. This indicates that solution conditions promoting the formation in vitro of amyloid fibrils can be designed rationally and support the hypothesis that the abil-
Amyloid formation by the HypF N-terminal domain

Amyloid formation by the HypF N-terminal domain is a property of many, if not all, polypeptide chains (Chiti et al. 1999). Furthermore, using the HypF N-terminal domain we have shown that it is possible to select solution conditions to promote formation of either amyloid fibrils or their constituent protofilaments. Moderate concentrations of TFE destabilize the interactions promoting the assembly of protofilaments formed from this protein domain. It was reported previously that long-term incubation of acylphosphatase in the presence of moderate concentrations of TFE also generates relatively long protofilaments that remain dissociated for periods longer than those observed for other fibril formation processes (Chiti et al. 1999). Furthermore, addition of TFE to fibrils obtained from the peptide corresponding to the 10–19 region of the transthyretin sequence disrupts the formation of TFE to fibrils obtained from the peptide corresponding to the 34–42 region of the sequence of the Aβ fragment associated with Alzheimer’s disease generates protofilaments, rather than complete fibrils, when the hydrophobic residues that are thought to be exposed on the surface of the protofilament are replaced with hydrophilic ones (Lazo and Downing 1999). Finally, protofilaments are also more stable when the L55P mutant of transthyretin undergoes aggregation (Lashuel et al. 1999).

All these findings, in addition to the results reported here with the HypF N-terminal domain, indicate that protofilament association into larger fibrils is likely to be guided, at least in part, by interactions between the side chains of hydrophobic residues that remain solvent-exposed on the surface of individual protofilaments and that mild destabilization of such interactions makes it possible to form and isolate the individual protofilaments. The ability of TFE to stabilize mainchain hydrogen bonds within individual protofilaments may also contribute to the relative stability of these structures in solutions containing TFE, as β-sheet structure is probably an important force driving the assembly of individual protein or peptide molecules into protofilaments. Protofilaments can grow and be stable independently of any further supramolecular organization. Solutions containing moderate concentrations of TFE are very favorable for the stabilization of individual protofilaments as TFE is known to weaken hydrophobic interactions without disrupting the backbone hydrogen bonds within individual protofilaments. Although we cannot make the generalization that TFE leads universally to protofilament formation (as opposed to other partially denaturing conditions such as low pH, high temperatures, and destabilizing mutations), we propose that dissociation of fibrils into their constituent protofilaments requires conditions destabilizing hydrophobic interactions that might exist at the interface between interacting protofilaments.

The ability for rational design of suitable solvent conditions for the formation of individual protofilaments has implications for the investigation of the structure and the mechanism of formation of amyloid fibrils. The possibility of preparing isolated protofilaments to order will allow structural investigations to be carried out on materials with a lower degree of complexity than mature fibrils. It will also facilitate the biophysical investigation of the steps associated with aggregation as protofilament formation is likely to be a less complex process than full fibril formation. More importantly, experimental procedures directed to the formation of protein aggregates and fibrils of various types will help in the identification of the aggregate species responsible for cell damage.

Materials and methods

Cloning, expression, and purification

The genomic DNA for the HypF N-terminal domain was isolated from Escherichia coli DH5α cells by Genomic Prep™ (Pharmacia). The DNA fragment corresponding to residues 993–1263 of the whole hypF gene was amplified by PCR using suitable primers containing the restriction sites for BamHI and EcoRI, respectively. Samples (1.0 μM) of each primer were added to 100 ng of template DNA, 10X PCR buffer, 200 μM dNTPs, and 2.5 units of Taq DNA polymerase (Finnzymes) in a volume of 50 μL. The fragments resulting from PCR amplification (94°C for 15 s, 52°C for 1 min, 72°C for 1 min, 25 cycles) were digested with BamHI and EcoRI, ligated into pGEX-2T downstream and in frame with glutathione S-transferase, and sequenced in their entirety. Expression of protein in the E. coli DH5α cells and its subsequent purification were carried out as described previously for muscle acylphosphatase (Modesti et al. 1995). Protein purity and quality were checked by SDS-PAGE, ES-MS, and amino acid analysis. The resulting sequence of the domain is AKNTSCGVLRRQKGVGFRPFVWQLAQLNLHDGVDGVYRLEPDTELTVQLYQHCPLLDRISVERFPWLSQPLTEFTIR. A Gly-Ser dipeptide is also present at the N terminus as a result of the cloning in pGEX-2T.

Circular dichroism

Far-UV CD spectra were acquired using a Jasco J-720 spectropolarimeter with a thermostat (Great Dunmow, Essex, UK). The CD spectra were recorded at a protein concentration of 0.2 mg mL⁻¹ in 50 mM acetate buffer (pH 5.5), 25°C. To acquire the CD spectrum of the domain under denaturing conditions, guanidinium chloride was added to a final concentration of 5.5 M.

Equilibrium urea- and heat-denaturation

25 samples containing 0.2 mg/mL protein in 50 mM acetate buffer (pH 5.5) were equilibrated at 25°C in the presence of different concentrations of urea ranging from 0 to 8 M. The mean residue ellipticity at 222 nm was plotted versus urea concentration and the resulting curve analyzed using the method described by Santoro and Bolen (1988). For acquiring a heat-denaturation curve, a 0.2 mg/mL solution of the N-terminal HypF domain, incubated in acetate buffer (pH 5.5), was placed in the CD cell holder and the temperature increased slowly at a rate of 0.5°C/min. A thermocouple was used to monitor the temperature inside the cuvette and
the CD signal at 222 nm was acquired at 1.0°C intervals from 20° to 76°C.

Electron microscopy

Electron micrographs were acquired using a JEM 1010 transmission electron microscope at 80 kV excitation voltage. In each case, a 2 μL sample of protein solution was placed on a formvar- and carbon-coated grid. The sample was then negatively stained with 30 μL of 2% uranyl acetate and observed at a magnification of 12–30,000×. Morphological investigation and size determination of the fibrils were performed by analysing ~30–40 fibrils per specimen and 3 specimens for each experimental condition (to achieve a total number of ~100 fibrils for each experimental condition).

Thioflavine T assay

The protein domain was incubated for either one week or one month at a concentration of 0.8 mg/mL in 50 mM acetate buffer, 30% (v/v) TFE (pH 5.5) or in 20 mM citric acid (pH 3). At regular time intervals, aliquots of the protein samples were withdrawn to perform the ThT assay as described above.

Congo red assay

The protein was incubated for either one week or one month at a concentration of 0.1 mg/mL in 50 mM acetate buffer, 30% (v/v) TFE (pH 5.5) or in 20 mM citric acid (pH 3). At regular time intervals, aliquots of the protein samples were withdrawn to perform the ThT assay as described above.

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