Stepwise Organization of the β-Structure Identifies Key Regions Essential for the Propagation and Cytotoxicity of Insulin Amyloid Fibrils*

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Background: Oligomers and protofibrils have been observed in the early stages of fibrillation. Results: Fibrillation of insulin at a high salt concentration identified a new species of prefibrillar intermediate. Conclusion: Structural comparison of the intermediate and mature fibrils suggested regions responsible for self-propagation and cytotoxicity. Significance: The trapping of intermediate is an effective way of revealing molecular details of the organization of fibril structure.

Amyloid fibrils are supramolecular assemblies, the deposition of which is associated with many serious diseases including Alzheimer, prion, and Huntington diseases. Several smaller aggregates such as oligomers and protofibrils have been proposed to play a role in early stages of the fibrillation process; however, little is known about how these species contribute to the formation of mature amyloid fibrils with a rigid cross-β structure. Here, we identified a new pathway for the formation of insulin amyloid fibrils at a high concentration of salt in which mature fibrils were formed in a stepwise manner via a prefibrillar intermediate: minute prefibrillar species initially accumulated, followed by the subsequent formation of thicker amyloid fibrils. Fourier transform infrared spectra suggested the sequential formation of two types of β-sheets with different strength hydrogen bonds, one of which was developed concomitantly with the mutual assembly of the prefibrillar intermediate to form mature fibrils. Interestingly, fibril propagation and cellular toxicity appeared only after the later step of structural organization, and a comparison of β-sheet regions between the prefibrillar intermediate and mature fibrils using proteolysis led to the proposal of specific regions essential for manifestation of these properties.

Amyloid fibrils are β-sheet-rich protein assemblies that are often associated with more than 30 serious amyloidoses (1–3). Moreover, various proteins and peptides not related to amyloidosis have been shown to form amyloids or amyloid-like aggregates, which suggests the potential of amyloid fibrils as a common and unique form of polypeptides as well as native structures (4). Amyloid fibrils are generally formed through nucleation and growth phases. On the basis of this fundamental scheme, nucleation is one of the most important events controlling the pathogenesis and propagation of diseases: many previous reports have shown that nucleation progresses at an extremely low rate, whereas the growth phase proceeds promptly via the incorporation of monomers or oligomers into the ends of nuclei according to a mechanism of template-dependent propagation of the amyloidogenic conformation once nuclei are formed (2, 5). Furthermore, recent experimental studies have proposed the cytotoxic nature of oligomers; therefore, the role of prefibrillar intermediates in diseases has also been a focus (6–11). Given these findings, elucidating the mechanisms for the formation of amyloidogenic nuclei in detail is essential for clarifying the pathogenesis and propagation of amyloidoses at the molecular level. Nevertheless, much remains unknown regarding how and when precursor protein molecules assemble to convert themselves into the amyloidogenic conformations responsible for manifestation of cytotoxicity and propagation behaviors. Although the native protein folding process has been extensively studied, the process of fibrillation needs to be clarified in more detail to understand the exact mechanisms of cross-β assembly.

To address this issue, it is very beneficial to trap and characterize prefibrillar intermediates, which will provide valuable insights into the detailed mechanisms by which the cross-β structure is organized during nucleation as well as the role of oligomers in diseases. The presence of smaller aggregates such as oligomers and protofibrils formed during the nucleation process has recently been revealed experimentally for several proteins. Various techniques such as solid-state NMR (7), small angle x-ray scattering (12, 13), site-directed labeling of fluorescence probes (8), hydrogen/deuterium (H/D) exchange experiments (14), and immunological assays (6) have revealed the structural details of the prefibrillar oligomers involved in fibrillation. However, quantifying and characterizing their physico-
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chemical properties are often hampered due to their unstable and heterogeneous properties; therefore, it is very difficult to clarify how these species contribute to the formation of mature fibrils with a rigid β-sheet core.

Here, we investigated a fibrillation pathway of insulin in which oligomer- or protofibril-like prefibrillar intermediate species could be detected in the early stages of fibrillation. Insulin is a hormone protein associated with glucose metabolism and consists of two polypeptide chains (i.e. A-chain of 21 amino acid residues and B-chain of 30 amino acid residues) connected by two intermolecular disulfide bonds. Insulin is one of the most important subjects in the field of diabetes therapy, and it has also become one of the best models for investigating fibrillation mechanisms in the past few decades by readily forming amyloid fibrils in vitro (15). By incubating insulin at a high temperature and in the presence of a high concentration of salt, we identified a new pathway for the formation of insulin amyloid fibrils in which a large amount of immature intermediate fibrils accumulated prior to the formation of mature fibrils. The sequential formation of β-sheet structures with different strength hydrogen bonds was clarified in this pathway, and in conjunction with this stepwise β-sheet organization, self-propagating ability, and cellular cytotoxicity manifested only at a later stage of the process. By comparing β-sheet regions in prefibrillar intermediates with those in mature fibrils using proteolytic analysis combined with mass spectrometry, we identified the amino acid regions responsible for propagating ability and cytotoxicity, based on which the detailed molecular mechanisms of fibril formation have been discussed.

EXPERIMENTAL PROCEDURES

Heat-induced Fibril Formation—The spontaneous formation of insulin amyloid fibrils without seeds was performed by heating. A total of 1.0 mg/ml of bovine insulin (Sigma) was dissolved in 25 mM HCl (pH 1.6) containing 0, 0.1, or 1.0 M NaCl. 500-μl aliquots of the sample solutions were transferred to 1.5-ml polypropylene test tubes and heated from room temperature to 90 °C at a rate of ~2 °C/min using a heat block (Dry Thermo Unit DTU-1B; TAITEC, Nagoya, Japan). After reaching 90 °C, the samples were removed from the heat block and air-cooled back to room temperature with gentle mixing; the temperature dropped to below 30 °C within 5 min. After 20 min, a 7.5-μl aliquot of the sample was mixed with 1.5 ml of 5 μM thioflavin T (ThT)3 in 50 mM glycine-NaOH buffer (pH 8.5) at 25 °C, and the fluorescence intensity at 485 nm of this solution was measured with an excitation wavelength of 445 nm to assay the formation of amyloid fibrils (16). Samples were then heated again, a process that was repeated three times. The fibrillation time course was also examined by measuring ThT fluorescence at different time points immediately after increasing the temperature to various levels from 55 to 90 °C to initiate the fibrillation reaction. For this experiment, 500-μl aliquots of the sample solutions sealed in the 1.5-ml polycarbonate test tubes were immersed in the heat block, which had been preheated to the reaction temperature; under this condition, the temperature jump of sample solutions was completed within 5 min. Insulin concentrations were determined using an absorption coefficient of 1.0 for 1.0 mg/ml at 276 nm (17).

Atomic Force Microscopy (AFM)—A total of 10 μl of fibril samples was spotted onto a freshly cleaved mica plate. The residual solution was removed after 1 min by placing a piece of filter paper at the edge of the mica plate followed by drying. AFM images were obtained using a Nano Scope IIIa (Digital Instruments, Tonawanda, NY). The scanning tip used was a phosphorus (n)-doped Si (Veeco Instruments, Plainview, NY; spring constant = 20–80 N/m, resonance frequency = 245–289 kHz), and the scan rate was 0.5 Hz.

Fourier Transform Infrared (FTIR) Absorption Measurement—FTIR spectra were measured with a FT/IR-6100 model spectrometer equipped with a DLATGS detector (Jasco, Tokyo, Japan) and a water-circulating system connected to a thermoregulated water bath. Samples for the structural analysis of heat-induced fibrils were precipitated by centrifugation to concentrate the fibrils to ~10 mg/ml, and washed with a small amount of D2O solution containing 25 mM DCI and 0.1–1.0 M NaCl three times prior to the measurement to diminish the amount of contaminated water interfering with the observation of the amide I band region. Samples were sealed with a cell with CaF2 windows and a 50-μm polytetrafluoroethylene spacer, and the FTIR spectrum was monitored at room temperature by collecting 256 interferograms with a resolution of 2 cm−1. To time course monitor fibril formation, 10 mg/ml of native insulin solution dissolved in 25 mM DCI containing 1.0 M NaCl was used as a sample. Prior to the sample preparation, native insulin was preincubated in 25 mM DCI and at 37 °C for 24 h to complete the H/D exchange of amide protons. Samples were then sealed with the 50-μm optical path-length CaF2 cell embedded inside the water-circulating system, and the temperature of the sample liquid was immediately raised to the measurement temperature (i.e. 75 or 90 °C) within several minutes to initiate the fibrillation reaction. The first FTIR spectrum was monitored 5 min after the temperature increase and subsequent spectra were collected at an interval of 4 min at a constant temperature.

Seed-dependent Fibril Extension—The seed-dependent extension reaction was examined by adding sonicated fibrils as seeds to native insulin in a 25 mM HCl solution containing 100 mM NaCl at concentrations of 30 μg/ml and 0.3 mg/ml, respectively. To prepare seeds, fibrils were sonicated using a Microson sonicator (Misonix, Farmingdale, NY) at an intensity level of 2 and 20 1-s pulses. The extension reaction was carried out at 37 °C without agitation and the reaction was monitored at different time points by ThT fluorescence analysis.

Cytotoxicity Assay—Pheochromocytoma (PC12) cells were grown in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% heat-inactivated horse serum and 5% fetal bovine serum. PC12 cells were plated on poly-L-lysine-coated dishes for differentiation, and cultured in DMEM supplemented with 10 ng/ml of nerve growth factor. Aliquots of the prefibrillar intermediate or mature fibrils were then added to cell cultures at a final concentration of 8.5 μM, and cells were incubated in a humidified 5% CO2 at 37 °C for 48 h. Prior to the assay, fibril samples were subjected to 20 pulses of sonication using a Microsonic sonicator at an intensity level of 2 to improve

3 The abbreviations used are: ThT, thioflavin T; AFM, atomic force microscopy; FTIR, Fourier transform infrared; H/D, hydrogen/deuterium.
dispersion by breaking up fibril clumps. Cell death was evaluated by measuring the extent of release of lactate dehydrogenase intracellularly to the medium using a commercial CytoTox 96 Non-radioactive Cytotoxicity Assay (Promega, Madison, WI). All measurements were performed three times and the percentage of surviving cells was calculated by assuming the value obtained by the addition of solvent only (i.e. HCl and NaCl at final concentrations of 1.25 and 50 mM, respectively) to be 100%. Statistical analysis was performed by one-way factorial analysis of variance combined with Scheffe’s test for all paired comparisons, where p < 0.05 was considered statistically significant.

**Proteolytic Digestion of Prefibrillar Intermediate and Mature Amyloid Fibrils**—Intermediate and mature fibrils were prepared by incubating 5 mg/ml of insulin solution dissolved in 25 mM HCl containing 1.0 M NaCl at 75 °C for 1 and 3 h, respectively. The enzymatic digestion of these fibrils was performed by mixing 200 μl of 0.63 mg/ml of pepsin from porcine gastric mucosa with 500 μl of the sample solution and incubating at 37 °C for 24 h. After the proteolytic treatment, samples were centrifuged and the pellet was washed 3 times with 25 mM HCl containing 1.0 M NaCl. The pellet was then solubilized by suspending it in dimethyl sulfoxide and incubating at 25 °C for over 10 h. The reduction of disulfide bonds was performed by adding 50 times its volume of 20 mM Tris-HCl buffer (pH 7.0) containing 10 mM dithiothreitol (DTT), followed by incubation at 50 °C for 1 h. The free cysteine residues generated were carboxamidomethylated by adding 5 times its volume of 100 mM iodoacetamide dissolved in 100 mM Tris-HCl (pH 7.0), which was incubated by agitating at 25 °C for 40 min. Measuring the matrix-assisted laser desorption ionization-time-of-flight (MALDI-TOF) mass spectra of the obtained peptide fragments was performed with a Voyager DE-STR MALDI-TOF mass spectrometer (Applied Biosystems) using α-cyano-4-hydroxycinnamic acid as a matrix.

**RESULTS**

**Formation of Prefibrillar Intermediate Fibrils in the Presence of a High Concentration of NaCl**—We first investigated the effects of the concentration of NaCl on the formation of insulin amyloid fibrils, which was monitored with the ThT binding assay (Fig. 1, A–C). Earlier studies revealed the spontaneous formation of insulin amyloid fibrils when subjected to high temperatures under acidic conditions (18–20). In the present study, bovine insulin dissolved in 25 mM HCl was commonly used as a fundamental solution and three different concentrations of NaCl, i.e. 0, 0.1, and 1.0 M, were analyzed. Regarding the heating method, we employed a linear heat gradient from 25 to 90 °C at a rate of 2 °C/min. The results of NaCl concentrations of 0 M (A), 0.1 M (B), and 1.0 M (C) are shown. All measurements were performed three times and the error bars depict the S.D. ± mean. D–G, AFM images of insulin fibrils formed by heat-induced fibrillation. The amyloid fibrils formed at 0.1 and 1.0 M NaCl were sampled after the 1st and 2nd cycles of heat treatment in the presence of 0.1 M (D (1st), E (2nd), and 1.0 M (F (1st); G (2nd)) NaCl. The scale bars represent 500 nm.

1st cycle of heating and almost the same ThT fluorescence intensity was maintained after the 2nd and 3rd cycles of the heating treatment. AFM images verified the formation of typical amyloid fibrils with needle-like morphology, and fibril height or length appeared to be unchanged after the 2nd heating cycle, which was in accordance with the unaltered ThT fluorescence intensity (Fig. 1, D and E).

Interestingly, when insulin was heated in the presence 1.0 M NaCl, the intensity of ThT fluorescence was markedly higher than that observed at 0.1 M NaCl, and the intensity was subsequently decreased by half after the 2nd heating cycle (Fig. 1C). This two-step change in fluorescence intensity implied the transient accumulation of different conformations with distinct ThT binding properties from that of mature fibrils, considering that the intensity of ThT fluorescence is determined by the affinity of ThT-binding sites, number of ThT-binding sites, and ThT fluorescence strength for each bound state as well as the amount of fibrils. To test this possibility, the sample was subjected to AFM analysis. In line with our expectations, AFM
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images exhibited the dominance of very thin and short aggregates (Fig. 1F), which suggested the transient accumulation of a prefibrillar intermediate. Needle-like morphology appeared after the 2nd heat treatment (Fig. 1G). The average height values obtained by sampling 10 different fibrils were 2.5 ± 0.4 nm for the prefibrillar intermediate and 10.4 ± 3.1 nm for the mature fibrils. Therefore, we postulated that the second step accompanying the reduction in ThT fluorescence intensity was attributed to the maturation process toward longer and thicker amyloid fibrils.

When FTIR spectroscopy was performed to compare micromorphological structures between the intermediate-like species and mature fibrils formed after the 1st and 2nd cycles of heating at 1.0 M NaCl, respectively, the obtained spectra revealed a marked difference in shape in the amide I region of fibrils formed after the 1st (red) and 2nd (blue) cycles of heat treatment at 25 °C. The black dashed lines represent the spectra of native insulin without any heat treatment as a reference. The spectrum of amyloid fibrils formed by repeated self-seeding at 0.1 M NaCl was also shown in a green line for reference. Spectra were normalized so that the integrated intensity of the amide I band ranging from 1580 to 1750 cm⁻¹ was set to be equal. C and D, second-derivative infrared spectra at 1.0 M NaCl (D). The colors are the same as those in panels A and B.

To investigate whether the FTIR peaks at 1621 and 1634 cm⁻¹ are assigned to cross-β structures, x-ray diffraction was analyzed supplementarily for the prefibrillar intermediate and mature fibrils. The diffraction patterns clearly supported the presence of cross-β structure both for the intermediate and mature fibrils by exhibiting two intense reflections at ~4.8 and ~11 Å, although the poor quantitativity of powder diffraction data hampered to verify an increased amount of the cross-β structure for the mature fibrils compared with that for the prefibrillar intermediate species (data not shown). Consequently, at least the FTIR intermolecular β-sheet band at 1621 cm⁻¹ was assigned to the cross-β structure, failing to the exact assignment of the peak at 1634 cm⁻¹.

Time Course for the Accumulation of an Intermediate Structure—To further explore the kinetic features for the formation of intermediate fibrils, the time course of fibril formation at 1.0 M NaCl was monitored by ThT fluorescence at varying temperatures ranging from 55 to 90 °C. A marked increase in ThT fluorescence was promptly observed after initiating the reaction at 75 °C and above, which then gradually decreased, representing the transient accumulation of intermediate fibrils (Fig. 3A). The two-phase change in ThT fluorescence specifically appeared to be observed at high temperatures only: the population of intermediate fibrils with high ThT fluorescence intensity in the range of 60 to 70 °C was not as large as that observed at higher temperatures, and the pattern appeared to migrate toward a typical sigmoidal pattern with a lag phase followed by extension as the incubation temperature became lower (Fig. 3B). No increase in ThT fluorescence intensity was observed within the experimental period at 55 °C (Fig. 3B). When spontaneous fibril formation was monitored at 0.1 M NaCl, the time course showed a typical sigmoidal pattern even at 90 °C, without the accumulation of prefibrillar intermediate species with high ThT fluorescence intensity (Fig. 3C).

We further investigated the β-sheet structure organization using real-time monitoring of FTIR spectroscopy forming insulin fibrils inside an optical cell (Fig. 4). Amide I absorption bands shifted slightly toward shorter wavenumbers to be represented as amide I’ because all of the amide protons of the insulin sample were substituted by deuterium prior to initiation of the fibrillation reaction for this measurement (see “Experimental Procedures”). When monitored at 75 °C, development of the β-sheet peak at 1619 cm⁻¹ was observed immediately after elevating the temperature, which confirmed the abrupt formation of prefibrillar intermediate fibrils with a partial amount of β-sheet structures (Fig. 4, A, C, and E). However, the subsequent development of another β-sheet peak at 1628 cm⁻¹ was significantly delayed, failing to be observed within the present experimental time period. The retardation of this later mat-
uration process may have been due to adsorption of the prefibrillar intermediate on the surface of CaF₂ windows with which the sample liquid contacts in a large area due to the very short path length (50 μm). When the reaction speed was accelerated by raising the temperature to 90 °C, the appearance of the peak of the β-sheet structure at 1628 cm⁻¹ was successfully observed, although the β-sheet structure at 1619 cm⁻¹ already formed within a dead time before the first scan of the FTIR spectrum. Overall, the combination of experiments with two different temperatures revealed the sequential formation of these two types of β-sheets with different strength hydrogen bonds: as was seen in the time-depen-
FIGURE 5. Effect of protein concentrations on accumulation of the prefibrillar intermediate of insulin amyloid fibrils. Time course for the spontaneous formation of insulin amyloid fibrils in the presence of 1.0 M NaCl was monitored at 75 °C using ThT fluorescence at different concentrations of protein, i.e. 1 (black), 5 (red), 10 (green), and 20 mg/ml (blue). ThT fluorescence intensity was divided by protein concentration for plots of measurements at 5, 10, and 20 mg/ml of insulin to bring their level in line with that at 1 mg/ml.

with a slow growth rate would be diminished as a result of competitive propagation (24). The synchronous propagation of these two FTIR peaks thus also supports the idea that the prefibrillar intermediate fibrils in the presence of 1.0 M NaCl was monitored at 75 °C using ThT fluorescence at different concentrations of protein, i.e. 1 (black), 5 (red), 10 (green), and 20 mg/ml (blue). ThT fluorescence intensity was divided by protein concentration for plots of measurements at 5, 10, and 20 mg/ml of insulin to bring their level in line with that at 1 mg/ml.

FIGURE 6. Trapping of the prefibrillar intermediate of insulin by cooling to room temperature. Intermediate fibrils were formed by incubating insulin at a concentration of 1 mg/ml dissolved in 25 mM HCl containing 1.0 M NaCl at 75 °C, and after accumulation of the prefibrillar intermediate, which was indicated by a markedly higher ThT fluorescence intensity, the reaction temperature was decreased to 25 °C and then the population of prefibrillar intermediate was monitored using ThT fluorescence intensity (gray circles). The time point of the temperature decrease is marked by an arrow. The result without a temperature decrease (i.e. the result of constantly incubating at 75 °C throughout the experiment; Δ) is also represented for comparison.

We also analyzed the cytotoxicities of intermediate and mature fibrils using cell viability assays with PC12, a rat adrenal pheochromocytoma cell line. An increasing number of studies have suggested that early stage aggregates in the fibril formation process are more toxic than mature fibrils (6–10). Although the severe toxic nature of prefibrillar intermediates was initially expected based on the findings of these reports, cytotoxicity was observed only for the mature fibrils and the intermediate species was nearly non-toxic to a similar level as that of non-aggregated intact insulin (Fig. 7B). Slight excess over 100% observed for the intact insulin and prefibrillar intermediate can be attributed to insulin-enhanced division of neuronal cells. Although the result for the cellular toxicity of the intermediate fibrils was contrary to what was expected, it is clear that structural differences between the intermediate and mature amyloid fibrils led to their distinctive toxicity and self-propagation properties. Based on the two-step change in the FTIR spectrum in the amide I′ region, the β-sheet structural feedback reaction was confirmed for the second maturation step was confirmed to be the key region responsible for the self-propagation and cytotoxicity of insulin amyloid fibrils.

Identification of the β-Sheet Regions Responsible for Propagation and Cytotoxicity—We performed a limited proteolysis experiment to specify the β-sheet region(s) organized during the maturation process. Proteolysis is one of the most promising methods that can be used to elucidate the β-sheet core regions of amyloid fibrils, and several studies have demonstrated that the flexible regions of amyloid fibrils were suscep-
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The prefibrillar intermediate and mature fibrils formed by heat treatment in the presence of 1.0 m NaCl were used as seeds. A, the time course for the seed-dependent extension reactions performed in a 25 mM HCl solution containing 0.1 m NaCl. In these reactions, 30 μg/ml of prefibrillar intermediate (○) and mature fibrils (▲) were seeded. The time courses in the absence of seeds (open inverted triangles) and presence of conventional insulin fibrils obtained by repeated self-seeding at 0.1 m NaCl (■) were also plotted for negative and positive controls, respectively. B, cytotoxic effect in PC12 cells upon the addition of native insulin without heat treatment for prefibrillar intermediate, mature fibrils, and conventional insulin fibrils obtained by the repeated self-seeding of fibrils originally formed at 0.1 m NaCl. The concentration of fibrils added was 8.5 μM and the incubation time was 48 h. Fractions of surviving cells, which were calculated from the activity of lactate dehydrogenase released from dead cells, are represented in this figure. All measurements were performed three times and the error bars depict the S.D. ± mean.

4. *p < 0.0001.

DISCUSSION

A New Species of Prefibrillar Intermediate Detected by Salt-induced Perturbation of the Fibrillation Pathway—The transient formation of several oligomer- or protofibril-like immature species have so far been reported for the fibrillation of insulin (12, 23, 31–35). Their population and properties appeared to be highly dependent upon the reaction conditions such as the concentrations of proteins, solvent composition, and incubating temperature or pressure, which suggested that the phase boundary between monomers and prefibrillar intermediates was sensitively perturbed by these external factors. In agreement with these results, a theoretical study recently proposed that even a slight perturbation in solution conditions can drive the fibrillation pathway toward the system in which oligomers are present as a stable intermediate phase (36). Here, we identified a new pathway for the formation of insulin amyloid fibrils by combining a high concentration of NaCl and high temperature, in which characteristic prefibrillar intermediate fibrils accumulated prior to the formation of mature fibrils. One of the most characteristic properties of this intermediate species is its markedly higher ThT fluorescence intensity, which might have resulted from a larger surface area of the prefibrillar intermediate per weight concentration compared with that of mature fibrils and then served as a clear sign of the presence of intermediate species with an unconventional time-dependent pattern (Figs. 1C and 3A); however, the exact reasons why ThT fluorescence intensity for the prefibrillar intermediate with an

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The main effect of salts has generally been considered to be a combination of screening of the electric charge of protein molecules through direct interactions between ions and proteins according to the electroselectivity series and solvent interactions perturbing protein hydration and/or bulk water conditions through indirect water-solute interactions according to Hofmeister series (34, 37). Through these effects, salts have been expected to modulate a balance between the electrostatic and hydrophobic interactions critical for fibrillation (37, 38), and tilting its balance toward stronger hydrophobicity by the presence of high concentrations of salts often results in the formation of worm-like or protofibril-like aggregates (39, 40). In light of the conceptual proposal that the presence of lag time is an essential characteristic by which crystal-like amyloid fibrils are distinguished from glass-like amorphous aggregates (38), the elimination of lag time in the present stepwise pathway (Fig. 3A) suggests that the prefibrillar intermediate fibrils assume an intermediary property between crystalline and amorphous structures. It is plausible that salt-induced increments in hydrophobic interactions among insulin molecules promoted the rapid accumulation of metastable prefibrillar intermediate fibrils with less organized immature β-structures.

Candidates of Specific Regions Contributing to the Propagation and Cytotoxicity of Amyloid Fibrils—As an advantageous property of the prefibrillar intermediate found in the present study, it was very stable at room temperature (Fig. 6), which allowed its application to a wide range of structural investigations. This stable population of intermediate fibrils has been attributed to a kinetic trap with an elevated energy barrier that separated the subsequent maturating process at room temperature, as was previously observed for the soluble intermediates of Aβ(1–40) that survived for a much longer time by lowering temperatures (41) and for annealing of the β-sheet core structure of mammalian prion fibrils that only proceeded at high temperatures (42). Through detailed comparison of β-sheet regions between the prefibrillar intermediate and mature fibrils as well as their seeding ability and cellular toxicity, particular peptide regions strongly associated with the propagation and cytotoxicity of insulin amyloid fibrils, have been specified successfully (Fig. 8C).

The growing ends of seed fibrils are generally considered to behave like a template in the template-dependent self-propagation mechanism of amyloid fibrils, onto which precursor proteins attach and successively change their conformation to replicate the same template structure, referred to as the dock-lock model (43, 44). Previous pre-steady state kinetic studies performed with Trp fluorescence (45) and H/D exchange comparisons of mature fibrils and prefibrillar intermediate species are shown on the upper and lower sides of each panel, respectively, and blue and magenta labels on the assigned peaks represent peptide fragments found only in the prefibrillar intermediate and in both the prefibrillar intermediate and mature fibrils, respectively. Black labels represent peaks derived from an uncleavaged fraction of insulin molecules. Also see Tables 1 and 2 for numerical data on the assigned peaks representing peptide fragments found only in the prefibrillar intermediate and in both mature fibrils, respectively. Also see Tables 1 and 2 for numerical data on the assigned peaks representing peptide fragments found only in the prefibrillar intermediate and mature fibrils, respectively. Regarding peptide fragments in which two possible identifications were assigned on the basis of the mass values, both of these possibilities were represented by dashed double-headed arrows with asterisks at the corresponding cleavage sites on the primary sequence. The regions sensitive to proteolytic digestion only in the intermediate fibrils are colored green.

FIGURE 8. Identification of amino acid regions responsible for manifesting the propagation ability and cytotoxicity. A and B, MALDI-TOF mass spectra of peptide fragments of prefibrillar intermediate and mature fibrils. In this study, peptides were identified under two conditions: with (A) and without (B) the reduction of disulfide bonds inside the insulin molecule. The spectra of mature fibrils and prefibrillar intermediate species are shown on the upper and lower sides of each panel, respectively, and blue and magenta labels on the assigned peaks represent peptide fragments found only in the prefibrillar intermediate and in both the prefibrillar intermediate and mature fibrils, respectively. Black labels represent peaks derived from an uncleavaged fraction of insulin molecules. Also see Tables 1 and 2 for numerical data on the primary sequence of the A-chain and B-chain. The blue and magenta arrows represent peptide fragments found only in intermediate fibrils and those found in both intermediate and mature fibrils, respectively. Cleavage sites are summarized on the primary sequence of insulin, in which blue and red dashed lines represent the sites cleaved in the intermediate and mature fibrils, respectively. Regarding peptide fragments in which two possible identifications were assigned on the basis of the mass values, both of these possibilities were represented by dashed double-headed arrows with asterisks at the corresponding cleavage sites on the primary sequence. The regions sensitive to proteolytic digestion only in the intermediate fibrils are colored green.
TABLE 1
Assignment of the digestive fragments of mature and prefibrillar intermediate fibrils
Fibrils were depolymerized into monomers by the addition of dimethyl sulfoxide after the proteolytic treatment, and the disulfide bonds of the sample were reduced and carboxamidomethylated before the measurement.

| Assignment                                                                 | Mature fibrils | Prefibrillar intermediate |
|---------------------------------------------------------------------------|----------------|---------------------------|
| Observed m/z                                               Theoretical m/z | Assignment                          | Observed m/z                                               Theoretical m/z | Assignment                          |
| 2339.049                                                            2338.981  | GIVEQCCASVCSLYQLENYC (A:1–21)     | 1027.491                                                            1027.503  | LVCGERGFF (B:17–25)         | 2544.280                                                            2544.249  | FNQLHGCGLHEALYLVCGERGF (B:1–23) and/or FNQLHGCGLHEALYLVCGERGF (B:2–25) |
| 2360.988                                                            2360.963  | GIVEQCCASVCSLYQLENYC (A:1–21) [M + Na]  | 1190.564                                                            1190.566  | YLVCGERGFF (B:16–25) and/or YLVCGERGFF (B:17–26) | 2619.344                                                            2619.317  | FNQLHGCGLHEALYLVCGERGF (B:1–24) and/or FNQLHGCGLHEALYLVCGERGF (B:2–24) |
| 2392.874                                                            2392.981  | Unmatched                                        | 1254.607                                                            1254.605  | FNQLHGCGLHEALYLVCGERGF (B:1–11) | 2745.272                                                            2745.216  | Unmatched                                        |
| 2500.268                                                            2500.272  | Unmatched                                        | 1303.655                                                            1303.650  | LYLVCGERGFF (B:15–25) | 2838.416                                                            2838.385  | FNQLHGCGLHEALYLVCGERGF (B:1–25) |
| 2691.387                                                            2691.317  | FVNQLHGCGLHEALYLVCGERGF (B:5–24) and/or FNQLHGCGLHEALYLVCGERGF (B:1–25) | 2392.938                                                            2392.981  | GIVEQCCASVCSLYQLENYC (A:1–21) | 2500.268                                                            2500.272  | FVNQLHGCGLHEALYLVCGERGF (B:5–24) |
| 2838.416                                                            2838.385  | Unmatched                                        | 2360.997                                                            2360.963  | GIVEQCCASVCSLYQLENYC (A:1–21) [M + Na]  | 2392.938                                                            2392.981  | GIVEQCCASVCSLYQLENYC (A:1–21) |
| 2500.268                                                            2500.272  | Unmatched                                        | 2544.280                                                            2544.249  | FNQLHGCGLHEALYLVCGERGF (B:1–23) and/or FNQLHGCGLHEALYLVCGERGF (B:2–25) | 2745.272                                                            2745.216  | Unmatched                                        |
| 2500.268                                                            2500.272  | Unmatched                                        | 2691.344                                                            2691.317  | FNQLHGCGLHEALYLVCGERGF (B:1–24) and/or FNQLHGCGLHEALYLVCGERGF (B:2–24) | 2838.416                                                            2838.385  | FNQLHGCGLHEALYLVCGERGF (B:1–25) |
| 2892.322                                                            2892.322  | Unmatched                                        | 2560.959                                                            2561.140  | GIVEQCCASVCSLYQLENYC (A:1–13) plus FNQLHGCGLHEALYLVCGERGF (B:1–25) | 2745.272                                                            2745.216  | Unmatched                                        |
| 2360.997                                                            2360.963  | GIVEQCCASVCSLYQLENYC (A:1–21) and/or FNQLHGCGLHEALYLVCGERGF (B:1–25) | 2745.272                                                            2745.216  | Unmatched                                        | 2392.938                                                            2392.981  | GIVEQCCASVCSLYQLENYC (A:1–21) |
| 2892.322                                                            2892.322  | Unmatched                                        | 2561.140                                                            2561.140  | GIVEQCCASVCSLYQLENYC (A:1–13) plus FNQLHGCGLHEALYLVCGERGF (B:1–25) | 2838.416                                                            2838.385  | FNQLHGCGLHEALYLVCGERGF (B:1–25) |
| 3251.622                                                            3251.613  | Unmatched                                        | 2500.268                                                            2500.272  | GIVEQCCASVCSLYQLENYC (A:1–21) plus FNQLHGCGLHEALYLVCGERGF (B:1–25) | 3251.622                                                            3251.613  | FNQLHGCGLHEALYLVCGERGF (B:1–25) |
| 3398.681                                                            3398.681  | FVNQLHGCGLHEALYLVCGERGF (B:1–30)         | 3398.681                                                            3398.681  | FVNQLHGCGLHEALYLVCGERGF (B:1–30)         | 3398.681                                                            3398.681  | FVNQLHGCGLHEALYLVCGERGF (B:1–30) |

* Observed and theoretical m/z values were represented as monoisotopic masses.

TABLE 2
Assignment of the digestive fragments of mature and prefibrillar intermediate fibrils monitored without the reduction of disulfide bonds
Fibrils were depolymerized into monomers by the addition of dimethyl sulfoxide after proteolytic treatment and then used for measurement.

| Assignment                                                                 | Mature fibrils | Prefibrillar intermediate |
|---------------------------------------------------------------------------|----------------|---------------------------|
| Observed m/z                                               Theoretical m/z | Assignment                          | Observed m/z                                               Theoretical m/z | Assignment                          |
| 5174.1                                                            5174.0  | GIVEQCCASVCSLYQLENYC (A:1–21) plus FNQLHGCGLHEALYLVCGERGF (B:1–25) | 2560.959                                                            2561.140  | GIVEQCCASVCSLYQ (A:1–13) plus FNQLHGCGLHEALYLVCGERGF (B:1–11) |
| 2561.140                                                            2561.140  | GIVEQCCASVCSLYQ (A:1–13) plus FNQLHGCGLHEALYLVCGERGF (B:1–11) | 2789.168                                                            2789.251  | GIVEQCCASVCSLYQ (A:1–13) plus FNQLHGCGLHEALYLVCGERGF (B:1–13) |
| 2789.251                                                            2789.251  | GIVEQCCASVCSLYQ (A:1–13) plus FNQLHGCGLHEALYLVCGERGF (B:1–13) | 2852.156                                                            2852.262  | GIVEQCCASVCSLYQ (A:1–15) plus FNQLHGCGLHEALYLVCGERGF (B:1–11) |

* For the mature fibrils, observed and theoretical m/z values are represented as averaged masses because mass resolution of the observed peak is insufficient for accurate determination of its monoisotopic mass. For the prefibrillar intermediate, observed and theoretical m/z values are represented as monoisotopic masses. The m/z value for intact insulin (i.e., A:1–21 plus B:1–30) was excluded from the list.

Set with NMR analysis (46), and $R_e$ relaxation dispersion and transferred cross-saturation NMR experiments (47) have suggested the presence of short amino acid regions involving the formation of a transient monomer-seed complex within precursor proteins. Considering these findings, our present results appear to have identified such specific sequences that play an essential role as a strong candidate for “template” coding a characteristic fibril structure of insulin.

Interestingly, the specified peptide regions corresponded well with positions with high fibrillation propensity to form the cross-$\beta$ spine core forming rigid intermolecular steric zippers facing two self-complementary sheets, which has been reported by screening for amyloidogenic short peptide segments (48, 49). By identifying such peptide segments, a novel amylose analysis has recently been developed to predict the amyloidogenicity of proteins (50). The deep involvement of the steric zipper structure in the self-propagating ability has been supported by the inhibition of insulin fibrillation by an octapeptide coding amino acid sequence that forms a steric zipper (LVEALYLYL) (51) and the effectiveness of the structure-based design of the peptide inhibitors of amyloid fibril formation for favorable affinity with the steric-zipper structure (52). Regarding the structural basis of cytotoxicity, the steric zipper structure is also more likely to intrude into biomembranes, resulting in cellular dysfunction, although the detailed roles of the steric zipper structure have not yet been elucidated. Curiously, the peptide regions specified in this work also correspond approximately to cleavage sites by insulin degrading enzyme (53), suggesting that insulin degrading enzyme suppresses the deposition of insulin in vivo by preferentially cleaning up these key regions with strong amyloidogenic features.

Proposed Model for the Organization of $\beta$-Sheet Structures and Implication for the Molecular Basis of Fibril Architecture—On the basis of the present results, a schematic model representing the stepwise formation of the $\beta$-sheets of amyloid fibrils with different strength hydrogen bonds is illustrated in Fig. 9. Many prefibrillar intermediates and some amounts of $\beta$-sheet structures were observed initially in this pathway. The intermediate formed mutually assembled, accompanying the organization of additional $\beta$-sheet structures important for fibril functionality (Figs. 8C and 9, green regions). This scheme assumed the direct involvement of the prefibrillar species focused on in
Structural Analysis of Insulin Prefibrillar Intermediate

The present study as an on-pathway intermediate playing the role of a conformational unit for the formation of mature amyloid fibrils. When the thickness of the prefibrillar intermediate is postulated to be 2.5 nm on the basis of average height determined from the AFM image, the conjugation of two prefibrillar intermediate fibrils is roughly comparable with the dimensions of a single insulin protofilament (3 × 4 nm) that has been proposed in the previous study of cryo-electron microscopy (54). It is thus plausible that two prefibrillar intermediates assemble mutually to form a protofilament during the maturation process (Fig. 9). Additionally, the average height of mature fibrils (i.e., 10.4 nm) suggests that roughly 4 protofilaments further bunch up to form the hierarchical structure of one matured insulin fibril. The organization of two types of β-sheet structures at distinct regions appears to bear a strong resemblance to a recent experimental finding for the fibrillation mechanism of the Sup35NM prion, in which interactions outside the amyloid core drive oligomer formation and subsequently led to the organization of cross-β structures inside the amyloid core region (55).

Whether oligomeric species bear the primary responsibility as an on-pathway intermediate or merely an off-pathway product remains a controversial issue in the mechanisms of amyloid formation (9). Intermediates have been regarded as dead-end products for some proteins (23) or, alternatively, as by-products acting indirectly as a kind of scaffold that assists secondary nucleation (31). On the other hand, the direct conversion of oligomeric species to mature amyloid fibrils has also been proposed for several proteins such as the amyloid-β peptide (7), β-lactoglobulin (56), insulin (12), and several amyloidogenic peptides (57), in which oligomers were shown to evolve themselves into amyloidogenic nuclei via direct coil-β conformational conversion (7, 56, 57) or mutual fusion (58). A mechanism has been reported in which oligomers instead of monomers were shown to be involved as a unit of docking for fibril elongation (59). In the present study, the prefibrillar intermediate species have been considered an on-pathway intermediate given the sequential change in the FTIR spectrum and very low amount of residual insulin molecules coexisting with the prefibrillar intermediate (see “Results”); however, it is still difficult to completely rule out the possibility that prefibrillar intermediates are indirectly converted to mature fibrils via monomers produced by the dissociation of intermediates.

Through the present work, we propose that salts are expected to be a useful factor for fine-tuning the energy landscape of protein fibrillation and identifying the stepwise aggregating pathways of amyloid fibrils via a prefibrillar intermediate by the effects of salt will be very profitable for determining the key regions essential for the propagation and cytotoxicity of many amyloidogenic proteins. Although the fibrillation pathway of insulin proteins was investigated under acidic conditions in this study because neutral pH strongly inhibits the amyloid fibril formation through the formation of stable native hexamers (15), it will be important to explore prefibrillar intermediate species at physiological pH and temperature as a future challenge to specify the regions taking a pivotal role in dictating the pathogenesis and transmission of amyloidoses. The molecular design or amino acid substitution targeting those regions will become a promising therapeutic strategy at the molecular level. Furthermore, recognizing the key regions controlling fibril functions further implies the structural bases underlying amyloid polymorphism, and the discovery of different amino acid sequences and/or stereoscopic patterns of side chain docking will improve our understanding of the molecular mechanisms underlying the prion strain phenomenon (55). Intensive studies aimed at such key regions will open a new perspective for the clarification of fibril formation mechanisms, and furthermore, attain a breakthrough for the development of therapeutic strategies against a wide range of amyloidosis and neurodegenerative disorders.

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