Structural Basis for the Inhibition of IAPP Fibril Formation by the Hsp60 Co-Chaperonin Prefoldin

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Abstract:

Chaperones, as modulators of protein conformational states, are key cellular actors to prevent the accumulation of fibrillar aggregates. Here, we integrated kinetic investigations with structural studies to elucidate how the ubiquitous co-chaperonin prefoldin (PFD) inhibits diabetes associated islet amyloid polypeptide (IAPP) fibril formation. We demonstrated that both human and archaeal PFD interfere similarly with the IAPP fibril elongation and secondary nucleation pathways. Using archaeal prefoldin model, we combined NMR spectroscopy with EM to establish that the inhibition of fibril formation is mediated by the binding of prefoldin’s coiled-coil helices to the flexible IAPP N-terminal segment accessible on the fibril surface and fibril ends. AFM demonstrates that binding of prefoldin to IAPP leads to the formation of lower amounts of aggregates, composed of shorter fibrils, clustered together. Linking structural models with observed fibrillation inhibition processes opens new perspectives for understanding the interference between natural chaperones and formation of disease-associated amyloids.

key words: Amylin; IAPP; chaperone; prefoldin; fibril formation; amyloid; protein structure.
Introduction

To date there are about 50 proteins or peptides identified which are implicated in amyloid diseases\(^1\). The hallmark of these pathologies is the formation of thermodynamically highly stable fibrils with a characteristic $\beta$-cross structure\(^1,2\) from native monomeric proteins. Most studied amyloidogenic proteins are involved in neurodegenerative diseases such as Alzheimer’s or Parkinson’s disease\(^3\). However, the cell-toxic formation of fibrillar aggregates can also be of systemic nature or localized in peripheral organs and can be associated with very different diseases such as arthritis or diabetes\(^4\). The 37 amino-acids long intrinsically disordered peptide hormone islet amyloid polypeptide (IAPP)\(^5\), also known as amylin, is found as amyloid aggregates surrounding $\beta$-cells in the pancreas in 90% of type II diabetes cases\(^6,7\). A variety of ways have been identified how amyloids can confer toxic activities, as for example the increase of the permeability of cell membranes by oligomers of amyloidogenic proteins\(^8\), and formation of porous structures\(^9\). Likewise, in the case of IAPP, not mature fibrils but small oligomeric species formed as intermediates during the fibrillation process have been observed to be the most toxic for $\beta$-cells\(^10\). Nevertheless, the mature amyloid fibrils, whose structures have recently been elucidated for IAPP\(^11–15\) are not innocuous as they can sequester proteins of the cell machinery\(^16,17\), exert mechanical stress on the cell\(^1\), and play a significant catalytic role during the fibril formation process\(^18\).

The fibrillation process starts with an oligomerization event, where a nucleus is formed, which rapidly elongates by templated incorporation of monomers. As soon as nucleation has occurred, the fibril growth proceeds exponentially since not only elongation occurs, but also
secondary processes, such as fibril fragmentation or secondary nucleation on fibril surfaces, which increase the rate of the fibrillation process. In vivo, under normal conditions, the protein quality control machinery composed of chaperone systems, ubiquitin-proteasome systems, and autophagy-lysosomal systems is able to prevent the formation of aggregates. However, in cases of misfolding diseases, these systems are either weakened due to a decrease of capacity during aging or overwhelmed due to increased concentrations or more-aggregation prone mutants of amyloidogenic proteins in hereditary conditions. A detailed mechanistic understanding of this natural quality control machinery allows for targeting its processes for medical intervention, either by targeted upregulation of its components or via functional replacement with small molecules. Chaperone molecules as modulators of protein conformational states are key factors acting on proteins during their transition from native state to the amyloid fold. Interacting with a multitude of species, chaperones are able to alter the fibrillation process by inhibition of aggregation, disaggregation, or detoxification. For instance, fibrillation inhibition can be mediated via interaction with misfolded monomers or small oligomers, which prevents the formation of seeding competent nuclei, but also interaction with fibril ends or decoration of the fibril surface is possible, which inhibits elongation and secondary nucleation processes respectively. Treatment of the diseases associated with amyloidogenic proteins whose native state is intrinsically disordered remains an important challenge for modern medicine, so a better understanding of the modulation of such amyloidogenic proteins by chaperones is of paramount interest.

Here we report the mechanistic study of IAPP fibrillation inhibition by the HSP60 type II co-chaperonin prefoldin (PFD). Homologues of this co-chaperonin are found in the cytosol.
of archaeal and eukaryotic cells. The heterohexameric PFD is a holdase with a characteristic “jellyfish” architecture, consisting of a β-barrel body and coiled-coil α-helix tentacles\(^\text{32}\).

Prefoldin binds substrates via a clamp-like mechanism and delivers it to HSP60 for refolding\(^\text{33}\).

PFD has previously been shown to interact with different disease-relevant amyloidogenic substrates and to inhibit their aggregation\(^\text{34–37}\), however no structural or mechanistic insights in this process have been described heretofore. We have integrated kinetic investigations with structural studies using atomic force microscopy (AFM), electron microscopy (EM) and nuclear magnetic resonance (NMR) spectroscopy to elucidate the different inhibition pathways and to provide a structural understanding of the prefoldin-amyloid interaction. The study of the highly dynamic complex between the 90 kDa PFD and intrinsically disordered 4 kDa IAPP enabled us to obtain insights into the chaperoning process at sub-molecular level. We show that inhibition of elongation and secondary nucleation is achieved by the interaction of PFD with IAPP fibril ends and surface, which subsequently leads to lower fibrillation rates and to the formation of clustered aggregates instead of individual fibrils.
**Results**

**PFD impedes IAPP aggregation.** IAPP aggregation kinetics in presence of PFD show a strong inhibitory effect of PFD on IAPP fibril formation *in vitro* (Fig. 1a and Supplementary Fig. S1a). For both *Pyrococcus horikoshii* and human PFD (denoted as PhPFD and hPFD, respectively), the overall tendencies are highly similar. The aggregation process is affected at low sub-stoichiometric PFD concentrations, suggesting an interaction between PFD and IAPP oligomers or fibril states, rather than a simple sequestration of IAPP monomers by PFD. In *de novo* aggregation assays monitored by thioflavin T (ThT) fluorescence, starting from monomeric IAPP without addition of any pre-formed aggregates, progressive addition of PFD slightly increases the duration of the lag phase (during which no aggregation is detectable by ThT fluorescence) and strongly reduces the fluorescence intensity (FI) in the final plateau phase. In addition, increased PFD concentrations result in a significant reduction of fibril growth rates associated with increased durations of the growth phase (the time span between lag phase and plateau phase) by more than an order of magnitude (Fig. 1a and Supplementary Fig. S1a).

In order to further investigate the inhibitory effects of both human and archaeal PFD on different processes contributing to IAPP aggregation, secondary nucleation and elongation assays were performed. Absence of agitation and addition of non-sonicated pre-formed fibrillar seeds enable observation of the surface-catalysed secondary nucleation process as the dominant contributor to the growth of fibrillar mass. Maximal growth rate drastically decreases for [PFD]:[IAPP] molar ratios above 1:113, and drops to almost zero for molecular ratios larger
than 1:17. For observation of PFD’s effect on elongation of IAPP fibrils, sonicated, short, pre-formed IAPP fibrils were added to IAPP monomers, offering a high number of fibril ends. The very first phase of linear growth reflects the pure elongation process, as it is the dominating mechanism of aggregation under this condition. Addition of PFD results in a distinct decrease of the initial growth rate (Fig. 1a). While the described effects on IAPP fibrillation are present for both archaeal and human PFD, the inhibitory effects of hPFD are more pronounced. The kinetic study of different modes of IAPP aggregation in presence of PFD shows that PhPFD and hPFD affect various steps of the fibrillation process. Sub-stoichiometric inhibition of both elongation and secondary nucleation implies interaction of PFD with IAPP fibrils, at the fibril ends and also along the fibril surface. Moreover, the final plateau values in all the kinetic experiments become significantly lowered if PFD is present.

**PFD induces formation of compact aggregates.** Atomic force microscopy (AFM) of the IAPP species present after the aggregation kinetics experiment showed changes in the appearance and the amount of IAPP aggregates in the presence of PFD (Fig. 1b and Supplementary Fig. S2). IAPP aggregation in absence of PFD results in a large amount of long, individually distinguishable fibrils (Fig. 1b left). In comparison, IAPP aggregates in presence of PFD are rare and rather clustered. The increase of PFD concentration lowers the overall amount of IAPP aggregates. Although the clustering gives rise to bigger assemblies of IAPP, some single fibrils are clearly visible at their perimeters (marked with arrows in Fig. 1b). Hence, this suggests that the observed clusters are not amorphous, but they represent rather bundles of shorter IAPP fibrils.
**IAPP binds PFD cavity in a dynamic manner.** In order to determine the binding site of IAPP on the PFD surface, NMR binding studies were performed with PhPFD, whose NMR assignment of backbone and methyl groups is available\(^{38}\) (Supplementary Fig. S3). Structurally, hPFD and PhPFD are similar\(^{39,40}\), but whilst hPFD is composed of six different subunits, the PhPFD complex contains 2 \(\alpha\)- and 4 \(\beta\)-subunits, which simplifies the investigation of this 86.4 kDa protein heterohexamer by solution NMR. The similar results obtained by AFM and ThT assays for the investigation of the effects of hPFD and PhPFD on IAPP fibril formation support the choice of this hyperthermophilic model system, which was used for all subsequent structural studies.

For the NMR investigation of the PhPFD-IAPP interaction, we have prepared highly deuterated complexes of PhPFD, selectively labelled on backbone (\(^{15}\)N\(^{-1}\)H probes) or end of side chains (\(^{13}\)CH\(_3\) probes) on either \(\alpha\)- or \(\beta\)-subunit. To determine the binding surface of IAPP on PhPFD, we prepared a spin-labelled IAPP construct, which allowed us to map the binding interface via paramagnetic relaxation enhancement (PRE)\(^{41-43}\). The disulfide bridge between cysteines two and seven on IAPP forbids spin labelling using standard thiol-reactive nitroxy derivatives, therefore, we linked a DOTA-cycle to the N-terminus using a two-\(\beta\)-alanine linker. The DOTA-functionalized IAPP was loaded with paramagnetic Gd\(^{3+}\) (Gd-IAPP) or diamagnetic Lu\(^{3+}\) cations (Lu-IAPP) (Supplementary Fig. S4). Labelled PhPFD was incubated at 30°C with either paramagnetic Gd-IAPP or diamagnetic Lu-IAPP reference (Fig. 2a-e). The residues involved in binding with IAPP were identified by comparing the intensities of PFD’s NMR signals in presence of Gd-IAPP or Lu-IAPP, respectively (Supplementary Fig. S5).
Mapping of the paramagnetically-broadened residues on the surface of PhPFD (PDB: 2ZDI, Fig. 2f) allows to locate the binding interface between IAPP and PhPFD (Fig. 2g-j). Whilst no PRE-broadening is observed for residues which are located on the top of the β-barrel body, broadening inside the cavity and along the sides of coiled-coil helices is detected. More specifically, strong interaction is observed in the middle part of the coiled-coil helices corresponding to residues 8-31 and 118-135 of the α-subunit. On the β-subunits the first 32 N-terminal residues are the most strongly broadened by the Gd-IAPP. These two regions face each other in the complex. All coiled-coil regions are especially enriched with glutamine. Interestingly, the regions strongly affected by IAPP paramagnetic relaxation present a globally negatively charged surface (-2) which can complement the net positive charge (+2) of the IAPP sequence. Also analysis of PFD’s electrostatic surface indicates that the regions strongly affected by IAPP paramagnetic relaxation are predominantly negatively charged or neutral (Supplementary Fig S6), suggesting that the electrostatic interaction may complement hydrophobic interactions, which presumably remain the main driving force for PFD-IAPP interactions. The large binding interface determined from the PRE-data suggests avidity-based binding to many lower affinity sites, which gives rise to the observed affinity. The NMR study suggests that monomeric IAPP is being incorporated in the chaperone cavity whilst binding and releasing the available interaction surfaces in a dynamic manner, as already observed for complexes involving unstructured proteins bound to other molecular chaperones\textsuperscript{41,44–46}.

**IAPP binds PFD with two binding segments.** To determine IAPP residues involved in binding of PhPFD, we recorded 2D NMR $^{15}$N-$^1$H-correlation spectra of U-$^{15}$N-labelled IAPP
at 30°C after addition of increasing ratios of PFD. Quantifications of the IAPP backbone amide signal broadening and chemical shift perturbations (CSPs) in the NMR titration experiment were used to map the binding sequence (Fig. 3a-d). Strong CSPs were observed for IAPP residues 10-19 and 24-28 (Fig. 3e). Peak broadening, resulting from apparent decreased tumbling rates upon binding of the 86.4 kDa prefoldin complex to small flexible IAPP, was located towards the N-terminus (Fig. 3d). C-terminal residues 34-37 are almost not impacted by interaction with PFD and are therefore most likely not involved in direct binding. Interestingly, binding is not restricted to the region 20-29, which is strongly involved in fibril formation and found buried in recently elucidated cryo-EM structures11–15. The spectral distribution in the proton dimension remains narrow, which suggests that IAPP does not fold upon binding, but remains essentially disordered. This confirms the PRE-results obtained upon monitoring the interaction of paramagnetically tagged IAPP on PFD which show an extended binding surface (Fig. 2). More quantitative analysis of binding strength was performed with the TITAN lineshape fitting program47, assuming a 1:1 stoichiometry. Line-shapes and change of signal frequencies were fitted for Ala13, Val17, His18, Ser19, Leu27, Phe23, Ala25, and Ser28, as these residues are important for the binding and analysis is not hampered by peak overlap. The apparent $k_{off}$ rate was estimated by line shape fitting to be in the order of 6000 s$^{-1}$, corresponding to a residence time of IAPP on PFD in the order of 100-200 µs. The apparent dissociation constant $K_D$ (61 ± 3.2 µM - error determined by bootstrap analysis) shows that IAPP has an affinity for PFD in the low to medium range. Estimation of the dissociation constant $K_D$ by biolayer interferometry (BLI), for which monomeric biotinylated IAPP was
immobilized on a streptavidin sensor, gave a result in the same order of magnitude (16 ± 2.4 μM) (Fig. 3f).

**Inhibition is mediated by PFD binding to IAPP fibril ends and surface.** To understand the observed effect of substoichiometric inhibition of elongation and secondary nucleation observed in the fibrillation assays, we have investigated PFD binding to IAPP fibrils by electron microscopy (EM). The major polymorph of IAPP fibrils used for this study has a pitch of ~48 nm (Fig. 4a-c), consistent with previous cryo-electron microscopy (cryo-EM) structure determination of IAPP fibrils (polymorph 1 in Röder et al. 2020, Fig 4d). Preformed IAPP fibrils were incubated with PFD for 30 min, transferred to a carbon grid, stained with SST and then visualized by electron microscopy. In the electron micrographs, strong decoration of the fibril with PFD is observed (Fig. 4 e-h). Visual analysis shows that on the fibril surface approximately every 80 nm one PFD molecule is found, that is about 10% of the surface (Fig. 4e). Upon investigation of the fibril ends, a density was repeatedly observed towards the end of the fibrils (Fig. 4h), which suggests a binding of PFD also to the fibril termini - about 40% of ends are accompanied by a density.

To characterize this interaction further at a sub-molecular level, we have docked PFD (PDB: 2ZDI) on the previously determined cryo-EM structure of IAPP fibrils, based on experimental PRE-broadening on PFD and chemical shift perturbation detected by solution NMR between PFD and IAPP (Fig. 5d,e), using the software HADDOCK\(^{48,49}\). The docking procedure was either directed to the fibril surface or to the fibril ends, thereby creating two representative complex-models (Fig. 5d,e). Analysis of these models indicates that IAPP-PFD
binding is strongly mediated by the N-terminal part of IAPP, which remains flexible and solution-exposed also in the fibrillar state, and that protruding N-termini of several neighbouring IAPP molecules can interact with the same prefoldin (up to six). This model rationalizes the pattern observed in the negative stain micrographs, as well as the inhibitory effect of PFD on IAPP fibrillation in the kinetic assays.
Discussion

Previously, the co-chaperonin prefoldin was reported to inhibit the fibrillation of highly toxic amyloidogenic proteins \textit{in vivo} and \textit{in vitro}\textsuperscript{34–37}, but no mechanistic or structural information was obtained so far. Here we show that prefoldin inhibits IAPP fibrillation at substoichiometric concentrations and interacts with multiple IAPP species, such as monomers and mature fibrils, and propose models for these interactions. We gained structural and mechanistic insights into the interaction of PFD with two different states of amyloidogenic IAPP (monomeric and fibrillar) by combining solution NMR, EM, AFM, and kinetic measurements.

NMR interaction studies report on binding between monomeric IAPP and PFD by chemical shift perturbations and peak broadening on IAPP. Analysis of these effects allowed us to identify two segments on IAPP, one towards the N-terminus (10-19) and one in the middle segment (24-28) to be important for the interaction (Fig. 3). Extensive deuteration and methyl labelling enabled us to study the interaction from the perspective of the 86.4 kDa PhPFD. PRE-mapping with a paramagnetic-labelled IAPP construct on PFD allowed to determine the sites involved in IAPP binding (Fig. 2). As it has been previously observed for chaperone-client complexes, such as the one between Hsp90 and tau protein, the complex is characterized by a broad interaction surface\textsuperscript{46}. IAPP induced PRE-broadening effects are observable inside the whole PFD cavity. Especially strong broadening is observed towards the middle of the coiled-coil helices of the α-subunit, suggesting that this is an important region for binding. The apparent $K_D$ was determined to be of intermediate strength (between 16 and 61 µM, according to BLI and NMR investigation). The extensive binding interface and multiple binding regions
on IAPP suggests avidity-based binding to multiple lower affinity contacts which can be sampled by IAPP. The spectra are characteristic of fast exchange between bound and free form with an apparent lifetime of the bound state on the order of 100 to 200 µs. As typical for chaperone-client complexes\textsuperscript{45}, the interaction between monomeric IAPP and prefoldin cannot be described as one static complex, but as a structural ensemble where IAPP dynamically binds and unbinds the available interaction sites inside the prefoldin cavity. Possible representative models of this complex were obtained by docking of IAPP on PFD driven by NMR derived structural restraints (one of the computed models is shown in Fig. 5a,b). \textit{De novo} ThT-fluorescence assays show that this interaction between IAPP monomers and prefoldin does only lead to a weak effect on the lag-time. Probably, neither restructuring effects on IAPP, nor sequestration of enough IAPP monomers in PFD cavity occurs, to strongly impact the primary nucleation step, and therefore the PFD-monomer interaction would play a minor role in the inhibition mechanism of IAPP fibril formation. However, the NMR studies between monomeric IAPP and PFD bring important information about residues of IAPP and PFD susceptible to be involved in the interaction with different species during the IAPP fibrillation pathway.

The ThT-fluorescence assays (Fig. 1) show inhibition of IAPP fibrillation at substoichiometric PFD ratios, which suggests interaction between PFD and high molecular weight IAPP species (e.g., oligomers or fibrils). Further investigation of fibrillation pathways by seeded ThT assays showed that both elongation and secondary nucleation pathways were inhibited by the presence of PFD (both human and archaeal). The subsequently performed EM analysis allowed to image PFD bound to ends and surface of IAPP fibrils (Fig. 4), which
reveals that the observed inhibition of the elongation and secondary nucleation pathways is achieved by these interactions (Fig. 5c). Only a low sub-stoichiometric PFD:IAPP ratio is required to obtain a major effect on the fibrillation kinetics, when inhibition is mediated by binding to fibrillar species\(^{50}\). The structures of fibrillar IAPP were elucidated recently by cryo-EM\(^{11–15}\) One major polymorph observed contains two protofilaments with three \(\beta\)-strands each and has a pitch of 48 nm\(^{13–15}\). The fibril core is composed of residues 13-37 with residues 1-12 protruding from the fibril core which are therefore not resolved in the cryo-EM map due to their dynamic behaviour. This polymorph (PDB: 6Y1A) was used to model the PFD-IAPP-fibril interaction as observed by EM, utilizing interacting IAPP and PFD residues identified by NMR to dock PFD and the fibril. This allowed us to get a sub-molecular insight into the interaction observed at low resolution by negative stain EM. From the fibril structure, it is clear that only the N-terminal binding epitope determined on IAPP is available for PFD binding when a fibril is formed. The PFD-IAPP fibril interaction through the N-terminal epitope was confirmed by the model obtained by NMR-guided docking\(^{48,49}\) (Fig. 5d,e). Yet, ca. up to six N-termini of IAPP in its fibrillar state can fit into the PFD cavity when PFD is binding to the fibril surface or end, possibly leading to a higher affinity as compared to monomer binding. On the fibril end, the second binding site in the middle IAPP segment is additionally available to stabilize the interaction. According to EM, the PFD density at the fibril end was higher than on the lateral fibril surface. This suggests that the fibril end has a particularly high affinity for PFD, which can be explained by the fact that both IAPP segments important for interaction are exposed at the fibril end. The preference of PFD to bind to fibril ends correlated with the strong inhibition of fibril elongation observed in IAPP aggregation assays (Fig. 1).
A striking effect of PFD on the IAPP fibrillation curves monitored by ThT FI was a decrease of the plateau height with addition of PFD (Fig 1). This could be due to a change in the fibrillar structure of IAPP leading to different binding of ThT molecules and subsequently decreased ThT FI, shielding of the surface due to PFD decoration, or due to the formation of less IAPP fibrils. A control experiment revealed that pure shielding of the ThT-IAPP fibril surface interaction by PFD is not sufficient to explain the amplitude of the observed effect (see material and methods section). However, AFM samples taken directly after termination of fibrillation assays show indeed decreased amounts of fibrils, but simultaneously, a morphological change in the observed aggregates is noticed. The fibrils which have formed in presence of PFD are shorter and cluster together, forming larger aggregates. The height profile and observed pitch, which can be measured on fibrils protruding from these aggregates in the AFM images, however are similar to the major polymorph formed in the absence of PFD. It seems that both decreased formation of aggregates and the formation of bigger clusters of aggregates, possibly also decorated by PFD molecules, lead to the observed effect in the ThT assays. The presence of PFD changes the amount and the length of fibrils and leads to their clustering. It is beyond the scope of this work to study changes in the toxicity of the aggregates formed in the presence of PFD, but it is worth noting that in the literature reduction of toxicity was observed upon decrease of the total exposed hydrophobicity by increase of aggregate size and surface shielding\textsuperscript{51,52}.

In conclusion, in this study we demonstrated that both human and archaeal PFD are able to inhibit fibril formation of the amyloidogenic protein IAPP. Solution NMR enabled us to
identify two binding regions on IAPP (from residues 10 to 19 and 24 to 28), and to map the corresponding binding sites within the PhPFD cavity. We established, using IAPP fibrillation assays and electron microscopy, that the sub-stoichiometric inhibition of IAPP fibrillation by PFD is mainly due to binding to fibril surface and ends, thereby inhibiting both secondary nucleation and elongation. Binding to the fibril is mediated by the N-terminal regions of IAPP of which up to six can be enclosed in the PFD cavity, interacting with PFDs coiled-coil helices. Binding to fibril ends is possibly supported by the second binding region, located in the middle segment of IAPP. The presence of PFD leads to the formation of lower amounts of aggregates, composed of shorter fibrils and clustered into formations of bigger size, which could be a potential detoxifying mechanism.
Material and Methods

Protein preparation

IAPP. Human IAPP (H-KCNTATCATQ RLALNFVHSS NNFGALSST NVGSNTY-NH₂; molecular mass 3903.4 Da; purity 93.2%), biotinylated IAPP (biotinyl-[β-Ala]-[β-Ala]-KCNTATCATQ RLALNFVHSS NNFGALSST NVGSNTY-NH₂; molecular mass 4271.9 Da; purity 99.2%), and DOTA (2,2′,2″,2‴-(1,4,7,10-Tetraazacyclododecane-1,4,7,10-tetrayl)tetraacetic acid) labelled IAPP (DOTA-[β-Ala]-[β-Ala]-KCNTATCATQ RLALNFVHSS NNFGALSST NVGSNTY-NH₂; molecular mass 4432.4 Da; purity 95.9%), all with an amidated C terminus and a disulfide bond between Cys2 and Cys7 were custom synthesized (Pepscan, Lelystad). Identity and purity were confirmed by RP-HPLC and mass spectrometry.

For kinetic experiments, to ensure monomeric starting material, the IAPP peptide powder was dissolved at 2 mg ml⁻¹ in 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP), incubated at room temperature for 1 h and lyophilized. Then 1 mg peptide powder was dissolved in 0.5 ml aqueous 6 M guanidine hydrochloride solution, and size-exclusion chromatography was performed on a Superdex 75 Increase 10/300 column (GE Healthcare) equilibrated with 10 mM 2-(N-morpholino)ethanesulfonic acid (MES)/NaOH buffer, pH of 6.0 using an NGC liquid chromatography system (Bio-Rad). The monomer peak fraction was collected, aliquoted, flash frozen in liquid nitrogen, stored at −80 °C, and thawed on ice straight before further use.
DOTA-functionalized IAPP was loaded with either paramagnetic Gd$^{3+}$ or diamagnetic Lu$^{3+}$ cations, respectively. Lyophilised DOTA-[β-Ala]-[β-Ala]-IAPP-NH$_2$ (400 µg, 90.2 nmol) was dissolved in 500 µl 20 mM sodium acetate/acetic acid buffer pH 4.53 and 18 µl 500 mM aqueous GaCl$_3$ solution was added (9 µmol, i.e., a 100-fold molar excess). The solution was incubated at 25 °C and 300 rpm shaking. Time dependent Gd$^{3+}$ loading was monitored by analysing the reaction mixture from withdrawn 10 µl aliquots at different time points by RP-HPLC on a 9.4 mm x 250 mm Zorbax 300SB-C8 column connected to a 1260 HPLC system (both Agilent) operated at a column temperature of 80 °C, a flow rate of 4 ml min$^{-1}$ and UV absorbance monitoring at 214 nm. A linear gradient from aqueous 8% (v/v) acetonitrile (ACN), 0.1% (v/v) trifluoroacetic acid (TFA) to 60% (v/v) ACN, 0.1% (v/v) TFA within 40 min was applied. Gd$^{3+}$ loading was completed after ~2 h and the remaining reaction mixture was subsequently purified by two to three consecutive preparative RP-HPLC runs (each 30 to 50 nmol Gd$^{3+}$-DOTA-IAPP) under elution conditions described above.

Time dependent RP-HPLC analytics are shown in Supplementary Fig. 4a. Loading of DOTA-IAPP with Lu$^{3+}$ was performed in an analogous way (Supplementary Fig. 4b).

U-[${}^{15}$N]- or U-[${}^{15}$N, ${}^{13}$C] isotopically labelled human IAPP was recombinantly expressed in *Escherichia coli* in M9 medium supplemented with either 2 g l$^{-1}$ [${}^{15}$N]-NH$_4$Cl and 3.2g l$^{-1}$ nonlabeled glucose or with 2 g l$^{-1}$ [${}^{15}$N]-NH$_4$Cl and U-[13C] glucose and purified according to established protocols$^{53}$. Purified recombinant IAPP from preparative RP-HPLC as final purification step was batch lyophilized, dissolved in HFIP, aliquoted and lyophilized again. Identity and purity (93%) were confirmed by analytical RP-HPLC under isocratic conditions with 28.5% (v/v) ACN, 0.1% (v/v) TFA as mobile phase on a 4.6 mm x 250 mm Zorbax
300SB-C8 column (Agilent) at 80 °C column temperature and 1 ml min⁻¹ flow rate (Supplementary Fig. S4c). For quality control a 2D $^{15}$N-TROSY spectrum of U-$[^{15}$N]-IAPP was recorded at 10 °C on a NMR spectrometer with a $^1$H frequency of 850 MHz (Supplementary Fig. S4d).

**Human Prefoldin (hPFD – 97.0 kDa).** *E. coli* Rosetta2 (DE3) pLysS cells (Novagen) were transformed with pET-21a plasmids encoding for PFD1, PFD3, PFD5 and PFD6 or pET-41 (Genecust) encoding for PFD2 and PFD4 subunits of hPFD. The N-terminal 37 residues of PFD3 and four residues of PFD4 were deleted, PFD2 and PFD4 contained N-terminal hexahistidine tags followed by a thrombin cleavage site (Supplementary Fig. S7a). Bacteria were either transformed with the plasmids for PFD1, PFD3, PFD5 and PFD6 individually or co-transformed with the plasmids for PFD2 and PFD3. The cells were grown at 30 °C in LB-medium with the required antibiotics (Chloramphenicol, Ampicillin, or Kanamycin). At OD$_{600\text{nm}}$ of 0.6 protein production was induced by IPTG (1 mM) and the expression was performed at 20 °C for 8 h (PFD1, PFD5), 30 °C for 4 h (PFD2, PFD3, PFD4) or 30 °C for 2 h (PFD6). All purification steps were carried out at 4 °C. Cells expressing the different subunits were mixed (100 ml for PFD1, PFD5 or PFD2/PFD3, 150 ml for PFD6 and 200 ml for PFD4). Cells were sonicated in 50 ml of 50 mM sodium phosphate buffer (pH 7), containing 300 mM NaCl, 5 mM β-mercaptoethanol and complemented with 0.025 mg/ml RNase (Euromedex), 0.025 mg/ml DNAse (Sigma Aldrich) and 1 anti-protease tablet (cOmplete™). Affinity purification, his-tag removal, cation exchange and size exclusion chromatography were performed as previously described. SEC-MALS and SDS-PAGE followed by Coomassie staining was used to monitor the purification process (Supplementary Fig. S6b,c).
Prefoldin from *Pyrococcus horikoshii* (PhPFD – 86.4kDa). *E. coli* BL21 (DE3) cells transformed with pET23c plasmids encoding either for the α- or β-subunit of PFD from *Pyrococcus horikoshii* (point mutation on α-subunit S98G) were used for protein expression (Supplementary Fig. S8a). To obtain unlabelled subunits, cells were grown at 37 °C in LB-medium, at OD<sub>600nm</sub> of 0.8 protein production was induced by IPTG (1 mM) and the expression was performed at 37 °C for 3 h. For production of U-[<sup>2</sup>H, <sup>13</sup>C, <sup>15</sup>N]-PhPFD samples, cells were progressively adapted to M9/2H2O media in three stages over 24 h. In the final culture the bacteria were grown at 37 °C in M9 media prepared with 99.85% 2H2O (Eurisotop), 2 g/l of U-[<sup>2</sup>H,<sup>13</sup>C] D-glucose and 1 g/l <sup>15</sup>NH<sub>4</sub>Cl (Cambridge Isotope Laboratories). For production of U-[<sup>2</sup>H, <sup>12</sup>C, <sup>15</sup>N], A-[<sup>13</sup>CH<sub>3</sub>]<sup>β</sup>, I-[<sup>13</sup>CH<sub>3</sub>]<sup>β</sup>, L-[<sup>13</sup>CH<sub>3</sub>]<sup>β</sup>, V-[<sup>13</sup>CH<sub>3</sub>]<sup>γ</sup>, T-[<sup>13</sup>CH<sub>3</sub>]<sup>γ</sup> labelled subunits, 2 g/l of U-[<sup>2</sup>H] D-glucose (Sigma Aldrich) was used as carbon source. When the OD<sub>600nm</sub> reached 0.7, a solution containing 240 mg/l of 2-[<sup>13</sup>C]methyl-4-[<sup>2</sup>H<sub>3</sub>]acetolactate (NMR-Bio) was added in M9/<sup>2</sup>H<sub>2</sub>O media for the stereoselective labelling of pro-S Leu<sup>δ2</sup> and Val<sup>γ2</sup> methyl groups<sup>55</sup>. 40 min later 3-[<sup>13</sup>C]-2-[<sup>2</sup>H]-L-Alanine, (S)-2-hydroxy-2-(2'-[<sup>13</sup>C],1’-[<sup>2</sup>H<sub>3</sub>])ethyl-3-oxo-4-[<sup>2</sup>H<sub>3</sub>]-butanoic acid (NMR-Bio) and 2,3-(<sup>2</sup>H) 4-(<sup>13</sup>C)-L-Threonine (NMR-Bio) were added to a final concentration of 250 mg/l, 100 mg/l and 50 mg/l respectively<sup>56,57</sup> for the simultaneous labelling of Ile<sup>δ1</sup>, Ala<sup>δ</sup>, Thr<sup>γ2</sup> methyl groups. Protein production was induced by IPTG (1 mM) and protein expression was performed at 37 °C for 3 h. With such protocol, the level of deuteration of protein is higher than 97%, while selectively labelled methyl probes are estimated to be protonated at more than 95%.
The purification protocol of archaeal prefoldin subunits was adapted from Okochi and coll. as previously published. SDS-PAGE followed by Coomassie staining, SEC-MALS and cryo-EM was used to control purity and homogeneity of produced samples (Supplementary Fig. S7b-d). Mass spectrometry was performed on the α-subunits (data not shown). For interaction studies, the buffers of PhPFD-α2β4 samples were exchanged for 25 mM MES/NaOH buffer (pH 6.5), 25 mM of MgCl₂ using Amicon Centrifugal Filter Units (Merck).

**Thioflavin T (ThT)-fluorescence assays**

IAPP aggregation kinetics were monitored by ThT fluorescence intensity (FI). All the experimental solutions contained 5 µM monomeric IAPP, 10 µM ThT, and 6 mM NaN₃ in 50 mM MES/NaOH (pH 6.5), 25 mM MgCl₂. Each assay contained a row of IAPP:PFD ratios: a sample without PFD and 10 samples with different PFD concentrations corresponding to logarithmically equidistant molar ratios PFD:IAPP between 1:5000 and 1:1. *De novo* assays had no additional components. For seeded assays, pre-formed mature IAPP fibrils were added (8 to 9% of overall IAPP concentration in monomer equivalent): either non-sonicated in the secondary nucleation assay, or sonicated (Sonopuls MS 72 microtip sonotrode, Bandelin; 10% amplitude, 4 cycles of 1 s pulse and 5 s break) in the elongation assay. For all assays, the samples were prepared as triplicates in protein LoBind tubes (Eppendorf) on ice and then transferred into a 96-well half-area polystyrene non-binding surface (NBS) microplate (3881, Corning). ThT FI values were recorded in a BMG FLUOstar Omega microplate reader (BMG LABTECH, Ortenberg) using excitation wavelengths of 448-10 nm and emission of 482-10 nm, each data point corresponds to an orbitally averaged value over 3 mm with 20 flashes per
well. In the *de novo* assays, 20 s orbital shaking at 300 rpm before each cycle was applied as additional agitation.

Kinetic analysis was performed and parameters such as growth rate and growth acceleration were extracted by numerical approximation of first and second derivatives with respect to time of the ThT FI curve (Fig. 1a, Supplementary Fig. S1a). For the analysis of the elongation assays only the first linear parts of the curves were considered, since under the chosen conditions other aggregation mechanisms besides elongation start contributing to IAPP fibrillation from relatively early time-points.

Since in all kinetic assays the resulting final ThT FI plateau values were significantly lowered upon increasing PFD concentration, a control experiment was performed: after the completion of an aggregation assay, PFD concentrations were equalized in all the wells and then ThT FIs were re-measured. The difference in the plateau heights was preserved as before this manipulation. Hence, the change in the final plateau values cannot be simply attributed to PFD preventing ThT alignment on the fibrillar surface and its following fluorescence enhancement, but it rather reflects the amount of available fibrillar IAPP in solution (data not shown). For this reason, the kinetic analysis shown in Fig. 1a was performed on non-normalized data. However, the comparison of the changes in kinetic parameters extracted from normalized and non-normalized data showed highly similar tendencies (Supplementary Fig. S1a), indicating the reliability of the conclusions on PFD inhibitory effects on IAPP aggregation.

Experimental data shown in Fig.1 for *de novo* and secondary nucleation assays were lag-time-corrected before averaging over the triplicates. Lag time, defined as the time before the first order time derivative reaches its 5th percentile, varied in triplicates on average about 6%.
Exemplary comparison of modified and non-modified data is shown in Supplementary Fig. S1b.

**AFM imaging**

For the AFM imaging, after recording *de novo* aggregation assays, the samples were taken out of the plate (96-well Half Area Black/Clear Flat Bottom Polystyrene NBS Microplate, Corning). The solution was extensively mixed in order to include the aggregates potentially sticking to the walls of the well. 2 µl of each sample were put onto a freshly cleaved muscovite mica surface and dried during the incubation over 10 minutes under the clean bench. Subsequently, the samples were washed 3 times with 100 µl Milli-Q H₂O and dried with a steam of N₂ gas. Imaging was performed in intermittent contact mode (AC mode) in a JPK NanoWizard 3 atomic force microscope (JPK, Berlin) using a silicon cantilever with silicon tip (OMCL-AC160TS-R3, Olympus) with a typical tip radius of 9 ± 2 nm, a force constant of 26 N/m and resonance frequency around 300 kHz. The images were processed using JPK DP Data Processing Software (version spm-5.0.84). For the presented height profiles a polynomial fit was subtracted from each scan line first independently and then using limited data range. For each sample a net area of 2500-18400 µm² was scanned for the overview, while part of it was additionally imaged with a higher resolution.

**Biolayer interferometry (BLI)**

The BLI measurements were conducted using BLItz-System (FortéBio, Sartorius), at room temperature. Super Streptavidin (SSA) biosensors (Sartorius) were hydrated in 50 mM
MES/NaOH (pH 6.5), 25 mM MgCl$_2$ buffer, loaded with biotinyl-[β-Ala]-[β-Ala]-IAPP-NH$_2$, quenched with biotin to eliminate the nonspecific binding of PhPFD to biosensors, and equilibrated in buffer. The sensors were plunged in solutions containing different concentrations of PhPFD from 0.5 µM to 150 µM and the association curves were recorded over 180 s, followed by the dissociation in the according buffer for 240 s. The resulting binding kinetics were corrected using the blank reference, then a steady state analysis was performed using $R_{eq} = [A]R_{max}/([A] + K_D)$ fitting equation, where $R_{eq}$ and $R_{max}$ stay for equilibrium and maximum responses, and $[A]$ is analyte (PhPFD) concentration.

**NMR spectroscopy**

**IAPP-PFD titration.** The NMR titration samples were prepared by adding unlabelled PhPFD into solutions of U-[¹⁵N]-labelled IAPP. Lyophilized, purified monomeric IAPP was suspended in an ice-cold buffer (25 mM MES/NaOH (pH 6.5), 25 mM MgCl$_2$) at a concentration of 64 µM and aliquoted. The aliquots were flash-frozen in liquid nitrogen and stored at -80 °C. For every titration point the aliquots were thawed on ice; different amounts of PFD, buffer and 4 µl of $^2$H$_2$O were added, so that the final concentration of IAPP was set to 29 µM (total volume 44 µl). The ratios: 1:0 (apo), 1:0.4, 1:0.75, 1:1, 1:1.25, 1:2, 1:3, 1:4 and 1:8 were tested. Each sample was transferred into a 1.7 mm NMR tube and 2D [¹⁵N]-SOFAST-HMQC spectra$^{59}$ were acquired at 30 °C on a Bruker Avance III HD spectrometer equipped with a 1.7 mm cryogenic probe and operating at a $^1$H frequency of 850 MHz. Data were processed using nmrPipe/nmrDraw$^{60}$ and the $K_D$ and $k_{off}$ were extracted using TITAN global lineshape fitting$^{47}$.
using NMR signals of residues A13, V17, H18, S19, F23, A25 and L27, assuming a 1:1 stoichiometry, therefore using the two-state binding model.

**PhPFD assignment.** Backbone atoms and methyl groups resonances of both α- and β-PhPFD subunits were previously assigned\textsuperscript{38}, using a combination of (i) 3D BEST-TROSY HNCO, HNCA, HN(CA)CB, HN(CO)CA, HN(COCA)CB and HN(CA)CO experiments\textsuperscript{61}; (ii) structure-based analysis of inter-methyl NOEs\textsuperscript{62}; (iii) mutagenesis\textsuperscript{63} and (iv) 3D HCC-relay experiments\textsuperscript{64,65} connecting backbone to methyl moieties. List of assigned chemical shifts is available online (BMRB accession code 50845) and annotated 2D \textsuperscript{13}CH\textsubscript{3}- and \textsuperscript{15}N-TROSY spectra (70°C in 50 mM Tris (pH 8.5), 100mM NaCl) are presented on Supplementary Fig. S3.

**PRE-experiments.** 200 µM \textsuperscript{15}N- or \textsuperscript{13}CH\textsubscript{3}-labelled U-[\textsuperscript{2}H]-PhPFD sample was combined at a ratio of 1:1 (or 1:2 (for methyl labelled samples) with DOTA-modified IAPP loaded with Lu\textsuperscript{3+} (diamagnetic) or Gd\textsuperscript{3+} (paramagnetic) and transferred in 4 mm Shigemi tubes (200 µl)). For each sample 2D \textsuperscript{15}N-BEST-TROSY\textsuperscript{66} or 2D SOFAST-METHYL-TROSY spectra\textsuperscript{63} were recorded at 30 °C for ~0.5 day/sample on a Bruker Avance III HD spectrometer equipped with a cryogenic probe and operating at a \textsuperscript{1}H frequency of 950 MHz. As a control experiment to exclude false positive interaction of PFD with the DOTA-cycle, we verified that addition of Gd\textsuperscript{3+}-loaded DOTA to methyl-labelled PFD does not lead to specific residue broadening (Supplementary Fig. S9).

**Electron microscopy (EM)**

IAPP fibrils were formed by incubation of 64 µM monomeric IAPP at room temperature for a minimum of one week in 10 mM MES/NaOH (pH 6.0). Concentrated purified PFD in 25 mM
MES/NaOH (pH 6.5) 25 mM MgCl₂ was added in a ratio of 1:100 (PFD:IAPP-monomer concentration) and incubated for 30 min. Samples were adsorbed to the clean side of a carbon film on mica, stained with sodium silicotungstate (SST) Na₄O₄SiW₁₂ at 1% (w/v) in distilled water (pH 7-7.5) and transferred to a 400-mesh copper grid. The images were taken under low dose conditions (<30 e⁻/Å²) with defocus values between -1.2 and -2.5 µm on a FEI Tecnai 12 LaB6 electron microscope at 120 kV accelerating voltage, 30,000x nominal magnification using CCD Camera Gatan Orius 1000 and analysed with Gwyddion software.

For cryo-EM imaging of PhPFD, 3.5µl of PFD solution in 50 mMTris, 100 mM NaCl pH=8.5 buffer at 100µM was blotted on a Quantifoil R2/2300 mesh gold grid coated with carbon and ionized by glow discharge. A FEI Vitrobot (automated vitrification machine) MARK IV was used at 6.5 sec blotting time, 100% humidity at 20 °C. About 1000 movies were recorded with a Thermo Scientific GLACIOS 200kV FEG with a Falcon II electron counting direct detection camera with EPU (automatic data collection) with a pixel size of 1.206 Å. About 700 000 particles were extracted using guided particle picking, using the PhPFD crystal structure (PDB: 2ZDI). 2D classification was performed using relion standard procedures.

Modelling of IAPP-PhPFD complexes

Interaction models were created by HADDOCK rigid model docking, using the HADDOCK 2.4 web server. For the model of the complex between monomeric IAPP and PhPFD, the basic docking protocol was used with the interaction residues on IAPP defined as the residues which showed above average chemical shift displacement upon PFD addition for most of the
ratios (residues 6, 10-13, 17-19, 24, 26-28). The PhPFD interaction site was defined as residues which showed more than 50% of signal intensity loss upon interaction with paramagnetic-labelled IAPP (α-subunit: 8, 9, 12, 20, 22, 24, 26, 29, 31, 37, 41, 65, 118, 125, 129, 132, 135, 146; β-subunit: 4, 12, 15, 17, 22, 24, 26, 34, 46, 58, 81; β’-subunit: 4, 12, 15, 17, 22, 24, 26, 34, 58, 61, 81, 107; Supplementary Fig. S5). The solution structure of IAPP in SDS micelles (PDB: 2L86) determined by NMR was used as the starting IAPP monomer conformation. Residues on the N-terminus of β-subunits missing in the PhPFD crystal structure (PDB: 2ZDI) were obtained from molecular modelling (MODELLER). For the complex involving PhPFD and IAPP fibril, the cryo-EM structure of the polymorph 1 of IAPP fibrils (PDB: 6Y1A) was used. The 12 missing N-terminal residues were added using molecular modelling (MODELLER), whereby a disulfide bond between residue two and seven was enforced. The same residues in PhPFD-IAPP monomer docking were used as restraints in PhPFD-IAPP fibril docking. For the fibril end interaction model, an IAPP fibril construct of 16 chains was used, and all the according residues on all the chains were given as docking restraints. To direct the docking towards the middle of the fibril, a longer IAPP fibril construct of 26 chains was used, and the interaction restraints were only set on the 18 chains in the middle of the construct.
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References


Figures Legends:

Figure 1. Inhibition of IAPP aggregation by Prefoldin.

a) IAPP aggregation kinetics in absence and presence of various concentrations of PhPFD (left column) and hPFD (middle column) with [PFD]:[IAPP] molar ratios ranging from 1:5000 to 1:1, followed by ThT fluorescence intensity (FI). Top row: de novo aggregation assay, where the aggregation starts from monomeric IAPP only, in presence of agitation. Middle row: secondary nucleation assay, where the aggregation starts from the mixture of monomeric IAPP and non-sonicated preformed IAPP fibrils (seeds), with no agitation. Bottom row: elongation assay, showing non-agitated aggregation of monomeric IAPP in presence of shorter sonicated seeds, where the initial linear parts of the curves (inset) are reflecting pure elongation of fibril seeds. The right column shows extracted kinetic parameters of changes in IAPP aggregation assays: growth phase duration, maximal growth rate, and initial growth rate for the according aggregation assays. Further kinetic analysis is presented in Supplementary Fig. S1a. b) AFM images of the samples at the end of the aggregation kinetic assays shown in section a (top row). Fibrils of IAPP (5 μM) aggregated in absence of PFD, rare clustered aggregates of IAPP formed in presence of 0.76 μM and 5 μM of PhPFD or hPFD (1:6.6 and 1:1 [PFD]:[IAPP] ratios accordingly). Each square image represents an area of 2x2 μm². In order to visualize objects decidedly differing in height, two colour code scales were used: the gradient from dark brown to white represents either 0 nm to 60 nm height, or 0 nm to 1 nm (highlighted in grey). The full AFM overview is presented in Supplementary Figure S2, including inter alia the control sample containing PFD alone and illustrating that the carpet-like coverage of the
surface with small roundish particles at increased PFD concentration is the appearance of PFD itself.

**Figure 2: Binding sites of IAPP on PhPFD structures**

a) PRE-labelled IAPP was obtained by fusion of a DOTA-macrocycle to the N-terminus of IAPP and subsequent loading of DOTA with Lu$^{3+}$ (diamagnetic control) or Gd$^{3+}$ (paramagnetic). Line broadening resulting from the proximity of Gd-IAPP (red spectra), compared to the control Lu-IAPP (blue spectra), is visible in 2D $^{15}$N-TROSY spectrum of β-subunit (b), 2D $^{13}$CH$_3$-TROSY spectrum of β-subunit (c), 2D $^{15}$N-TROSY spectrum of the α-subunit (d), and $^{13}$CH$_3$-TROSY spectrum of the α-subunit of PFD (e). A partial peak doubling is observed for the β subunits, due to their position in the PFD heterohexameric complex, some of which could be assigned to β and β$^+$ (panels b and c), while the two α-subunits are spectroscopically equivalent (panels d and e). f) Model of PhPFD (PDB: 2zd) with α-subunits in blue, β-subunits in green. g) No line-broadening was observed for residues on the top of the PFD complex. Conversely a strong effect was observed inside the cavity (j) and along the elongated coiled-coil helices of (h) β/β$^+$-subunits (two views related by 180° rotation) and (i) α-subunits dimer (side view).

**Figure 3: Interaction of monomeric IAPP with Prefoldin**

a) Extracts of 2D $^{15}$N-SOFAST-HMQC spectra of IAPP titration by addition of PhPFD. [PhPFD]:[IAPP] molar ratios ranging of 0, 0.3, 0.75, and 1.25 are shown. b) Fit of K_D signals of A13, V17, H18, S19, F23, A25 and L27. c) Summary of detected chemical shift
perturbations (CSPs) $\Delta \delta$ (calculated as $\Delta \delta = \sqrt{\Delta N^2 + \Delta H^2}$) induced by addition of a two-fold excess of PFD on uniformly $^{15}$N-labelled sample of IAPP. d) Peak broadening of IAPP signals upon addition of PhPFD, calculated as the intensity ratio of NH signals in IAPP/PhPFD complex (I) and in absence of PhPFD ($I_0$). Asterisk denotes values obtained at 10°C due to peak overlap at 30°C. e) IAPP sequence with colour coding of detected CSPs at ratio 1:2 (residue with above average $\Delta \delta$, that is $\geq 0.12$ ppm in red) and line broadening at ratio 1:2 (residue with $I/I_0 \leq 0.26$ in yellow). f) Steady state analysis of biolayer interferometry (BLI) experiment on PhPFD binding to monomeric IAPP: measured values (dark blue) and fit (light blue)

**Figure 4: EM images of PhPFD bound to IAPP fibrils**

a) Electron microscopy images of IAPP fibrils, stained with sodium silicotungstate (SST) at 30,000x magnification. b-c) Close up of main fibril species. d) Molecular model of polymorph 1 of the IAPP fibril (PDB: 6Y1A$^{13}$). e) Preformed fibrils (67 µM) were incubated with PhPFD in a ratio of 100:1 (IAPP monomer equivalent concentrations) for 30 min and imaged under the same conditions as IAPP fibrils (a). Decoration of the fibril surface with PhPFD is observed, remaining PhPFD particles are observed in the background. f-h) Close-ups of the fibrils show the decoration of the fibril surface (pink arrows) and end (white arrows). i) Molecular model corresponding to the observed fibril decoration. Horizontal white and black bars correspond to a length of 100 nm and 10 nm, respectively. Simulated TEM images of (d) and (i) are shown in Supplementary Fig. S10.
Figure 5: Structural Model of inhibition of IAPP fibril formation by PFD

Schematic model describing the inhibition mechanism of PFD on IAPP fibril formation. PFD interacts with monomeric IAPP but this transient interaction does not lead to a significant decrease of the lag-phase. (a) and (b) present docking models of the complex between monomeric IAPP (PDB: 2L86) and PhPFD (PDB: 2ZDI) based on NMR derived interaction information (Fig. 2, 3). c) Inhibition of secondary nucleation and elongation results from coverage of fibril surface and ends by PFD (Fig. 4). The presence of PFD leads to a decreased steady phase fibril mass (Fig. 1a), which results from the formation of less aggregates with an altered morphology (Fig. 1b). The inset zoom represents a model of IAPP fibril structure with unfolded residues 1 to 12 in yellow and the structured fibril core is represented in purple (from residues 13 to 37). (d) and (e) present docking models of PhPFD (PDB: 2ZDI) on IAPP fibril (PDB: 6Y1A) surface and extremities, respectively, integrating structural information obtained by NMR (Fig. 2, 3) and EM (Fig. 4); the black arrows indicate the fibril axis with the tips pointing towards the fibril ends.
Figure 1

**a**

- **PhPFD**
  - *De novo* (+ agitation, - seeds)
  - Growth phase duration

- **hPFD**
  - Secondary nucleation (- agitation, + non-sonicated seeds)
  - Elongation (- agitation, + sonicated seeds)

**b**

- 5 μM IAPP
- +0.76 μM PhPFD
- +0.76 μM hPFD
- +5 μM PhPFD
- +5 μM hPFD
Figure 2
Figure 3
Figure 5