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# Reversible Morphological Control of Cholecystokinin Tetrapeptide Amyloid Assemblies as a Function of pH

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ABSTRACT: Most amyloid assemblies are seen as irreversible and exhibit polymorphism because their assembly is kinetically controlled and different structures are trapped during the aggregation process. However, in the specific case of peptide hormones, formation of amyloid assemblies for storage purposes has been reported. This suggests a strict control of assembly and the ability to disassemble upon hormone secretion. In the present work, we have sought to test these assertions with a short peptide, the cholecystokinin (or gastrin) tetrapeptide (CCK-4), that has been found in both gastrointestinal tract and central nervous system, and whose sequence is shared by a large number of hormones. We have thus studied *in vitro* this peptide's self-assembling properties in dense phases at different pH levels, thus mimicking in vivo storage conditions. The solubility and morphology of the supramolecular assemblies have been shown to vary with the pH. At low pH, the tetrapeptide exhibits a low solubility and forms microcrystals. At higher pH levels, peptide solubility increases and above a high enough concentration, peptide monomers self-assemble into typical amyloid fibrils of 10-20 nm diameter. The physical network formed by these fibrils results in a birefringent hydrogel phase. Despite the different morphological features exhibited at different pH, structural analysis shows strong similarities. Both supramolecular assemblies – microcrystals and fibrils – are structured by  $\beta$ -sheets. We also show that all these morphologies are reversible and can be either dissolved or changed into one another by switching the pH. In addition, we demonstrate that a modification in the charge sequence of the peptide by amino acid mutation modifies its self-assembly properties. In conclusion, just as the CCK-4 sequence is the minimal sequence required for a complete biological activity at CCK<sub>B</sub> receptors in the brain, it is also sufficient to form amyloid fibers whose properties can be related to hormone storage and release purposes in vivo.

### INTRODUCTION

Until recently, the study of amyloid fibril assembly and structure has mainly been motivated by the will to understand the formation of pathological aggregations as observed in neurodegenerative diseases such as Alzheimer's, Parkinson's or Huntington's. The fibril formation proceeds from the misfolding of small proteins (or large peptides) which are usually constituted from a few tens up to a few hundreds of amino-acids.<sup>1</sup> The most studied models are amyloid- $\beta$  protein<sup>2</sup>, amylin or  $\alpha$ -synuclein.<sup>3</sup> Nevertheless, numerous examples were found in which proteins or peptides are assembled into amyloid-like fibrils that fulfill a beneficial role such as cytoprotection<sup>4,5,6</sup>, cell adhesion<sup>7</sup> or regulation of melanin biosynthesis.<sup>8</sup> As a consequence, these systems were named "functional amyloids".9 Moreover, these types of amyloid fibrils seem to be ubiquitous in the biological realm since they can be found in plants<sup>10</sup>, fungi<sup>11</sup> and bacteria<sup>12,13</sup> as well as in mammals.<sup>7,8,14,15</sup> All these examples highlight the potential of amyloids in the development of novel functional biomaterials. In contrast with pathological amyloids, functional amyloids are formed by amino acid sequences that have been selected during evolution. Hence we can expect that they follow robust assembly processes, have well-defined structures and that they are harmless. Additionally, they don't require destabilizing conditions (organic solvents or alcohol, extreme pH conditions etc...) to selfassemble, which makes them biocompatible.

Many of these functional amyloids result from the assembly of small proteins (most of them count well over 100 amino acids) that are difficult to purify and present complex co-aggregation phenomena, which render in vitro studies tedious. Thus, there is a strong incentive to study shorter peptides stemming from functional amyloid systems. By studying the family of pituitary peptide hormones, which were slightly shorter (9-41 amino acids, with an outlier at 199), Maji et al. opened new avenues.<sup>16</sup> Likewise, studies on peptide hormone analogues conducted by Paternostre & Artzner et al. have demonstrated the biophysical insight that can be retrieved from the study of this class of peptides.<sup>1</sup> Indeed, using a short peptide sequence makes it easier to identify the molecular determinants driving the assembly and controlling the supramolecular structures. The specific role of side chains and charges can be finely addressed and potential molecular dynamic simulation tractable. Moreover, the storage functionality of hormones amyloids suggests that their assembly is reversible upon well-defined triggers. Finally, peptide hormones self-assembly properties can be har-nessed for therapeutic applications.<sup>21–25</sup>

In the frame of the present study we have assessed the selfassembly properties of the cholecystokinin tetrapeptide (CCK-4) also known as gastrin tetrapeptide - as a function of pH. This sequence corresponds to the last four amino acids of the gastrin and the cholecystokinin C-terminus peptides that are respectively synthesized in the intestine and in the brain. In both cases it corresponds to the minimal sequence required for complete biological activity at the G-protein coupled receptor CCKB in both the central nervous system and gastrointestinal tract.<sup>26</sup> As such it is commonly used in experimental pharmacology to investigate the biological effects mediated by this receptor.<sup>27</sup> CCK-4 has been found in nerve terminals in pancreatic islets and reported to act as a potent releaser of insulin and other islet hormones.<sup>28</sup> There is also evidence suggesting that CCK-4 - along with CCK-33 and CCK-8 - acts as a neurotransmitter or neuromodulator in the central nervous system.<sup>29-31</sup> Indeed, CCK-related peptides are thought to participate in the central control of stress response and stress-induced memory dysfunctions.<sup>32</sup> It is also worth mentioning

that gastrin and cholecystokinin peptides have been found in dense secretory granules<sup>33,34</sup>. We can hypothesize that they are stored under an aggregated form – very likely amyloid structures, as other peptide hormones are.<sup>16</sup>

The amino acid sequence of the CCK-4 fragment is Trp-Met-Asp-Phe-NH<sub>2</sub>. From now on it will be referred to as WMDF. Its detailed formula can be seen in Figure 1A. From a structural point of view, it is an amphiphilic peptide with two aromatic residues at the termini, respectively a tryptophan at the N-terminus and a phenylalanine at the C-terminus. The C-terminus is additionally amidified. In between, the two central residues are methionine, which can potentially be functionalized<sup>35,36</sup>, and aspartic acid, whose side-chain is negatively charged above pH 3.7. Another charge appears below pH 9.3 by protonation of the N-terminal. The resulting net charge of the peptide thus varies between -1 and +1 passing through zero at pH 6.1 (See Figure 1B). Between 3.71 and 8.76 the peptide adopts a zwitterionic form. This suggests that the backbone conformation and solubility strongly depends on the pH. To assess the role of the charge borne by the aspartic acid, we have also characterized the assembling properties of an analogue whose sequence Trp-Met-Asn-Phe-NH<sub>2</sub> (referred to as WMNF in the following text) in which the aspartic acid residue is replaced by an asparagine residue whose side chain is polar and uncharged. The only remaining charge is thus the N-terminus at acidic pH. Above pH 10, WMNF bears no charge at all.



Figure 1: A) WMDF in its zwitterionic form (e.g. at zero net charge pH~6.2). B) Calculated net charges for WMDF and WMNF sequences as a function of pH.

We will first show that the ionic state of the peptide plays a significant role in the morphology of the supramolecular assemblies and study their structure and conformation by small angle X-ray scattering (SAXS) and vibrational spectroscopies; the second section will establish the reversible nature of these structures. In a third section, we will further highlight the specific role of the charge borne by aspartic acid in studying the mutant peptide WMNF. Finally, we will discuss the biological relevance of the WMDF assembly diagram as a function of pH and concentration in term of hormone storage and release.

## **EXPERIMENTAL SECTION**

**Peptides.** The WMDF- $NH_2$  and WMNF- $NH_2$  peptides were purchased from Bachem (Switzerland). The WMDF-acetate and WMNF-TFA lyophilized salts were weighed and dissolved in the appropriate aqueous buffer at room temperature. The solutions were homogenized by vortexing.

**Transmission electron microscopy.** Transmission electron microscopy was performed on a Philips CM12 electron microscope operated at 80 kV. A drop of the solution at 3-5% w/w was deposited on a copper grid covered with a carbon film (Agar Scientific). After blotting off of the excess liquid, the material was stained with a 2% uranyle acetate solution or a 1.5% ammonium molybdate solution.

X-ray scattering and x-ray diffraction. SAXS-WAXS experiments were performed at the European Synchrotron Facility (ESRF), on the ID02 beamline. The wavelength of the incident beam was 0.995 Å with exposure times of the order of 0.1-0.5 s. The scattered X-ray were collected on two Rayonix 2D-detectors, covering respectively q-ranges of 0.0075 - 0.64 Å<sup>-1</sup> and 0.53 - 0.644.33 Å<sup>-1</sup>. The WAXS patterns reported in Figure 3D were obtained on an in-house set-up. The X-rays were produced by a Molybdene anode. The wavelength of the incident beam was of 0.722 Å. The scattered pattern was collected on a MAR detector. The sample-to-detector distance was 72 cm. The samples were inserted in 1.5 mm diameter quartz capillaries sealed with paraffin wax or spread on a kapton cell (10% WMDF in PBS and 5% WMNF in NaOH). The spectra were manipulated and analyzed using the SaxsUtilities (www.sztucki.de/SAXSutilities/) and SasView (http://www.sasview.org/) softwares. Attempts to index the diffraction patterns were made using the DICVOL06 program.<sup>37</sup> A series of cells was proposed by the software from analysis of the 19 peaks common in the two diffractograms. We used the following criteria to select the most relevant cells: (i) Cells must be compatible with the dimensions of the peptide ~9 Å x ~20 Å, V = 1153 Å<sup>3</sup>; (ii) All the peaks have to be indexed; (iii) Cells must not generate too many peaks that are not experimentally detected.

Attenuated Total Reflectance Fourier Transform Infrared (ATR-FTIR). ATR-FTIR spectra were recorded at a 2-cm<sup>-1</sup> resolution with a Bruker IFS 66 spectrophotometer equipped with a 45° N ZnSe ATR attachment. The spectra obtained resulted from the average of 30 scans and were corrected for the linear dependence on the wavelength of the absorption measured by ATR. The water signal was removed by subtraction of pure water spectrum multiplied by a coefficient chosen by superimposing the spectra in the 2500-4000 cm<sup>-1</sup> region. Afterwards, the spectra were normalized by setting the highest peak of the amide I band to 1. The spectra were then deconvoluted with PeakFit (Systat Software Inc) as a sum of Gaussian-Lorentzian components using the secondary derivative method. A linear baseline was used.

Fourier-Transform Raman Spectroscopy (FT-Raman). FT-Raman spectra were recorded at a 4-cm<sup>-1</sup> resolution with a Bruker IFS 66 spectrophotometer. The scattered light from a HeNe laser ( $\lambda$ =632 nm) was emitted (452 mW incident power) at the sample and was analyzed at a right angle. A nitrogen cooled CCD detector was used to collect Stokes Raman data. In order to check the possible degradation of the sample under the beam, we acquired series of 500 spectra. The plotted spectra are thus the average of 100×500 spectra, which correspond to an accumulated acquisition time of 20 hours. The samples were conditioned in quartz capillary tubes sealed with paraffin wax.

**Steady-state Fluorescence.** The influence of the pH on the fluorescence emission of the WMDF peptide was assessed by performing two titrations of two 1 mg/ml samples. The first sample was prepared in 0.5 M acetic acid and titrated with 1 M NaOH. The second sample was prepared in 0.2 M NaOH and titrated with 1 M HCl. The solutions were excited by a wavelength at 286 nm. Although some measurements were made with samples with absorbance values higher than 0.3 (as typically advised to avoid reabsorption effects by neighboring molecules), controls showed that the position of the peak maxima  $\lambda_{max}$  were not impacted by concentrations. Intensities are given in arbitrary units. This was in particular unavoidable in order to probe the effect of assembly on the fluorescence. ThT fluorescence assays have also been performed. These results are displayed and commented in details in the ESI section.

## **RESULTS AND DISCUSSION**

#### WMDF polymorphism as a function of pH.

WMDF is poorly soluble in acidic conditions (e.g. in acetic acid, between pH 2.4 and 5) and at intrinsic pH (i.e., when dissolved in pure water, pH~6) at concentration as low as 0.1 g/L. One obtains large aggregates that sediment to the bottom of the Eppendorf tube while the supernatant remains liquid. The concentration of peptide in the supernatant is estimated to be ~1.5 mM via UV absorption spectroscopy. When the pH is increased above pH 7, the solubility also increases. Macroscopically, the samples at concentrations below 5% w/w are clear liquids. Above 5-7% w/w, they form gels that are birefringent when observed between crosspolarizers (Figure 2A). At pH 7-8 (0.1 M PBS), however, the gels are whitish while at pH 13 (0.2 M NaOH) they are completely transparent. ThT fluorescence assays performed on liquid suspension (pH 6) and gel (pH 9) show an increase of fluorescence emission at 482 nm in both cases (See ESI, Figure S5), which is widely regarded as a diagnostic of amyloid formation.



Figure 2: A) Polarized optical microscope picture of a 10% w/w gel of WMDF solubilized in a phosphate buffer 0.1 M (pH 7.4) inserted in a quartz capillary. B) Transmission electron micrograph of a solution of WMDF solubilized in 0.5 M acetic acid (pH 2.5). C-D) Transmission electron micrographs of a 5% w/w solution of WMDF solubilized in a 0.1 M phosphate buffer (pH 7.4).

Transmission electron microscopy (TEM) gives access to the morphology of the objects formed in solution. In the suspensions of WMDF dissolved in acetic acid or water, straight and elongated crystals are observed (Figure 2B). Their length is of the order of the micron or more and their aspect ratio is about 1:10. In contrast, solutions prepared at higher pH appear to contain several micron-long unbranched fibrils, with an estimated diameter of  $19\pm7$  nm, very similar to amyloid fibrils as they are classically described<sup>38</sup> (Figure 2C-D). At intermediate pH (e.g. 6-7), a coexistence between the microcrystal and fibril morphologies can be observed. The proportion of the two species seems to depend on the pH since at pH 6 the crystals are predominant, while at higher pH fibrils take over. It is also noticeable that the crystals formed at higher pH seem less sharp than those formed at pH 2.5-5.

The structural features of the WMDF basic solutions are further probed by small angle X-ray scattering (SAXS). Figure 3A-B exhibit the 2D-scattering patterns of a 10% w/w WMDF solution solubilized in NH<sub>4</sub>Ac 0.2 M (pH 11) collected the SAXS and WAXS detectors of the ID02 beamline, ESRF respectively. These oriented patterns clearly show the typical cross-ß pattern characteristic of amyloid fibers. Three main reflections are indicated by arrows labelled *mer* (for meridional), *eq* (for equatorial) and *a*, at respectively 1.34 Å<sup>-1</sup> (*d*=4.7 Å), 0.36 Å<sup>-1</sup> (*d*=17.5 Å) and 0.11 Å<sup>-1</sup> (d=57.11 Å). The meridional 4.7 Å reflection corresponds to the spacing between hydrogen bonded β-strands that run perpendicular to the fibril axis. The equatorial 17.5 Å reflection corresponds to a β-sheet network resulting from an alternated stacking of antiparallel peptides (spacing between the sheets).<sup>17</sup> This inter-sheet distance is about twice that which is traditionally observed in typical amyloid fibers (8-11 Å). The reflection a is actually the first oscillation generated by the scattering from a cylindrical shape as shown in Figure 3C, which displays the SAXS pattern of 10% WMDF in 0.2 M NaOH and the corresponding fit. The fit was obtained by assuming a cylinder shape with a 47 Å radius.

Figure 3D displays the azimuthal profiles of WMDF solutions in 0.2 M NaOH (pH 13) at concentrations ranging from 5 to 20% (traces 1-5). These scattering patterns were collected on the single detector of an in-house set-up. At concentrations below 10%, the traces are barely distinguishable from the solvent (trace 0), except for a very weak bump around 0.2 Å<sup>-1</sup>. From 10% and higher, the scattering strongly increases at the smallest angles, indicating the formation of large objects. As described above, the typical amyloid cross- $\beta$  fiber reflections are clearly visible. The correlation length  $\xi$  estimated from the full width at half-maximum (FWHM) of the meridional and axial reflections with the relation  $\xi=2\pi$ /FWHM is about 100 Å, which is of the order of the diameter of the fibrils observed in TEM (Figure 2D) and with the diameter of the cylinders used to fit the SAXS pattern (Figure 3C).



Figure 3: Small Angle X-Ray Scattering (SAXS) of WMDF gels solubilized in basic conditions. A-B) Oriented scattering pattern of 10% WMDF in NH<sub>4</sub>Ac (pH 11) on SAXS detector (A) and WAXS detector (B). The capillary and hence the fibrils were oriented horizontally. C) SAXS pattern of a 10% WMDF gel solubilized in 0.2 M NaOH (blue line) and the corresponding fit assuming flexible cylinders (purple line). D) SAXS-WAXS patterns of WMDF solutions in NaOH 0.2 M at different concentrations – 0: blank NaOH 0.2 M; 1: 5%; 2: 7%; 3: 10%; 4: 15%; 5: 20%. The arrows indicate the reflections corresponding to the meridional (*mer*) and equatorial (*eq*) orientations in respect with the fibrils' axis.

In Figure 4, the powder diffraction patterns of WMDF in acetic acid (trace 1) and PBS (trace 2) are displayed in comparison with the scattering pattern of 20% WMDF in NaOH (trace 3). The diffraction patterns confirm the crystalline organization of the objects observed in Figure 2B. In addition to reflections attributed to amyloid structures, about 20 diffraction peaks are clearly visible in both patterns. Most peaks are common to both spectra, however there are still some differences that lead to slightly different cell parameters. Peak indexation trials led to the conclusion that both samples are polymorphic.In both cases, all cells are monoclinic and their parameters are rather close and consistent with the dimensions of the peptide monomer (See Table S1 in ESI). Their volumes ( $V \sim 2300 \text{ Å}^3$ ) suggest that they contain two peptides. These cells could not be linked to specific morphologies observed in TEM. Nevertheless, the correlation length  $\xi$  estimated from the FWHM of the first four peaks is of the order of 1800 Å, which is about the lateral size of the elongated crystals observed in TEM (cf. Figure 2B).



Figure 4: Diffraction patterns of crystallized WMDF. 1 (red line): Dried sample of WMDF prepared in acetic acid. 2 (green line): Crystals contained in a 10% WMDF gel prepared in PBS (pH 7.38) 3 (blue line): Concentrated solution (20% w/w) of WMDF fibrils prepared in NaOH 0.2 M (pH 13). The arrows labelled "eq" and "mer" indicate the position of the amyloid reflections. The overlap between the SAXS and WAXS detectors of the ID02 beamline, ESRF is seen in the 0.55-0.65 Å<sup>-1</sup> range.

Attenuated Total Reflection Fourier Transform Infrared (ATR-FTIR) vibrations give information on the conformation of the peptide backbone in their assembled form. Figure 5A presents the area of the spectra corresponding to the amide vibrations which are the major characteristic bands of a peptide infrared spectrum. The amide I band, between 1600 and 1700 cm<sup>-1</sup> is mainly associated with the C=O stretching vibration and is directly related to the backbone conformation and the hydrogen bonding pattern. The amide II bands, between 1510 and 1580 cm<sup>-1</sup>, result from the N-H bending vibrations and from the C-N stretching vibrations.<sup>39</sup>

After water spectrum subtraction and normalization, the amide I region is the part of the ATR-FTIR spectra that is the most characteristic of the peptide assembly (Figure 5A). At any pH

tested, the spectra exhibit very thin and intense bands in the 1631-1645 range corresponding to a strong  $\beta\text{-sheet}$  structure.  $^{40,41}$  A satellite peak of lower intensity in the 1676-1684 cm<sup>-1</sup> is indica-tive of an antiparallel organization.<sup>42</sup> The fit of this region by Gaussian-Lorentzian peaks makes the change of conformation explicit as the pH increases (Figures S2 and S3). At acidic pH levels the vibration at 1643 cm<sup>-1</sup> is the strongest and the vibration at 1627 cm<sup>-1</sup> is barely detectable. As the pH increases, an inverse of the situation is observed as is illustrated in Figure S3. This shift of the main vibration is indicative of stronger hydrogen bonds and could be liken to the shift observed between transthyretin in native (1630-1643) and in fibril form (1611-1630).<sup>43,44</sup> In both cases, these differences can be ascribed to the longer strand formation in fibrils versus native/crystal form.<sup>44</sup> Yet, the increase in the intensity of bands assigned to unordered structures in 1650-1660 cm<sup>-1</sup> suggests that amyloid fibrils are less structured than the microcrystals.

The amide II spectral region (Figure 5A) is rather independent of pH, with a large band around 1550 cm<sup>-1</sup> which mostly originates in N-H vibrations. In each case, this part of the spectrum can be broadly deconvoluted by Gaussian functions (not shown) centered at 1497±2, 1528±4, 1546±2 and 1558±4 cm<sup>-1</sup> whose respective intensities can vary. The 1497±2 cm<sup>-1</sup> band is very the likely ring vibrations of amino acids containing aromatic rings.<sup>45</sup> Vibrations at 1528 and 1546 cm<sup>-1</sup>, which become prominent at basic pH levels, are attributed to the dominance of the bending of hydrogen bonded NH groups in antiparallel β-sheet structures.<sup>46</sup>



Figure 5: Vibrational spectroscopic characterization of WMDF solutions at different pH (amide I and amide II regions). A: ATR-FTIR spectra. B) FT-Raman spectra. 1: WMDF solubilized in acetic acid 500 mM (pH 2.4); 2: WMDF solubilized in pure water (pH 6.1); 3: WMDF solubilized in 5% Mannitol (pH 6);4 and 4': 10% w/w WMDF solubilized in a 100 mM phosphate buffer (pH 7); 5: 10% WMDF solubilized in a 200 mM NaOH solution (pH 13.3). Raman peak attribution is shown in Supplementary Materials.

Finally, FT-Raman spectroscopy also offers fingerprints of the different conformations encountered in each condition (Figure 5B and Section 4of the ESI). The strong amide I peak at ~1675 cm<sup>-1</sup> once more confirms the formation of antiparallel  $\beta$ -sheet. However, because the peaks arising from tryptophan, phenylalanine and amide bands may overlap, exact attribution can be difficult. We

propose attributions in the ESI (Table S2). Nevertheless, some qualitative conclusions may be drawn regarding the tryptophan environment.<sup>47,48</sup> First, the W3 vibration is located at 1549 cm<sup>-1</sup> in both acidic and basic conditions. The W3 vibration can be related to the torsion angle  $\chi^{2,1}$  and defines the orientation of the indole ring with respect to the C- $\alpha$  atom of the amino acid backbone<sup>49</sup> and in both acidic and basic conditions  $\chi^{2,1} \approx 90^{\circ}$ . Likewise, the W17 vibration at 876 cm<sup>-1</sup> at both pH (ESI, Figure S4C), indicates that the indole N-H is involved in a medium-strength hydrogenbond as is formed with solvent water. The intensity ratio of the W7 Fermi doublet ( $I_{1360}/I_{1340}$ ) is also reported to depend on the hydrophilic/hydrophobic tryptophan environment. With ratios of 1.5 and 1.26 at acidic and basic conditions respectively, it can be inferred that tryptophan environment is only moderately hydrophobic.

The tryptophan environment can also be assessed by steadystate fluorescence spectroscopy. Figure S5 (ESI) shows that the tryptophan environment is rather polar, even when the peptides are aggregated into fibrils or microcrystals. In the latter cases the average  $\lambda_{max}$  is about 350 nm, which is similar to the values given for an exposed tryptophan in a protein. Values reported for buried tryptophan residues are closer to 330 nm.<sup>50</sup> However, this value is significantly shifted with respect to the ones measured for the monomers in solutions (366 nm) which are similar to those reported for the amino acid alone in solution.<sup>51</sup> As for the intensity of the emission peak, it decreases as the proportion of assembled peptides increases.

Overall, there is a set of evidence pointing towards an antiparallel arrangement of the peptides in the different morphologies found. Despite an expected hydrophobic-driven self-assembly, it appears that the tryptophan residue is not fully buried in the supramolecular structure but may be involved in hydrogen bonding rather than in  $\pi$ -stacking. Calculations of contact potentials between two adjacent peptides are also in favor of antiparallel structures where the backbone would be staggered by three amino acids (See ESI, Figure S1), an arrangement that allows hydrophobic interactions between phenylalanine residues, but not for tryptophan residues as suggested by Raman and fluorescence spectra. X-ray diffraction analyses also suggest that in spite of a polymorphism at the molecular level, molecules are always packed in a similar kind of unit cell; the conformational differences detected as a function of pH would then only arise from different lateral chain organizations. Additional characterization with solid-state nuclear magnetic resonance<sup>52,53</sup> could probably confirm the proximities between residues.

Here, we have demonstrated that even a small fragment of Cholecystokinin are able to self-assemble into amyloid structures even though aggregation propensities predictions are null for this specific amino acid sequence for most algorithms<sup>54–57</sup> (see Figure S6A-D in ESI). Nevertheless, both Aggrescan<sup>58</sup> and Zyggregator<sup>59</sup> predict aggregation (Figure S6E-F). Prior experimental work on gastrin related peptides by Oakley et al. had reported fibril formation at 0.5% w/w in saline solution.<sup>60</sup> Notably, they also mentioned that "fibrils not only disaggregate completely after mild alkali treatment but the peptide also fails to form fibrils at a high pH". We have clearly demonstrated that this is not the exact case. The critical aggregation concentration simply increases with pH. It is probably the sheer shortness of the sequence that makes the morphology extremely sensitive to pH, especially at pH levels close to physiological conditions. However, as Hu et al. have pointed out in the case of Pmel17:RPT, functional amyloids do not necessarily require a specific structure or morphology.<sup>61</sup> In the present case, if we assume a storage end, there is indeed no need for a very regular structure; only packing density and reversibility are required.

## Assessment of the reversible nature of the amyloid assemblies of WMDF.

One of the primary hypotheses about functional amyloids fibrils that are assumed to serve as storage means is that this type of supramolecular assembly must be reversible. In this respect, WMDF hydrogels were shown to be easily solubilized upon dilution. However, given the detection threshold of our analytical techniques it is difficult to prove that the fibrils themselves are disassembled and not simply dispersed in a solution of larger volume. To keep the peptide concentration constant during the putative disassembly process, basic solutions of WMDF were submitted to acid vapors (either glacial acetic acid or hydrogen chloride vapors). Conversely, acidic or neutral solutions were submitted to (basic) ammonia vapors.

In most cases, it is macroscopically clear that the sample is undergoing drastic changes: initially whitish and light scattering acid samples containing microcrystals and larger aggregates rapidly clear up (Figure 6 A-B), while liquid (at concentrations below 10% w/w, Figure 6 C-D) or gel basic samples (at concentrations above 10% w/w, Figure 6 E-F) precipitate. ATR-FTIR and SAXS characterizations confirm the optical observations. Figure 6A shows an initially well-structured spectrum of an acidic sample (red line) transformed into a weak disordered spectrum (blue line) after said sample has been submitted to basic ammonia vapors. Corresponding SAXS spectra show diffraction peaks (red line) disappearing (blue line) as well as intensity decreasing at smaller angles indicating the disappearance of large objects in the solution (Figure 6B). However, the scattering pattern is indicative of cylindrical shapes (fibrils) in solution. Conversely, Figure 6C-D show non structured spectra of a low concentration basic sample (blue lines) transformed into structured spectra (red lines) as pH is decreased with acetic acid vapors. The characteristic vibration bands or diffraction peaks correspond to those described in previous section for crystal structures obtained directly in water or acetic acid. SAXS spectra also clearly shows an increase in scattering intensity corresponding to the formation of large supramolecular objects (Figure 6D). The non-structured spectra correspond to solutions containing non-assembled solubilized WMDF monomers. Finally, Figure 6E illustrates the transition from a structured FTIR spectrum (blue line) corresponding to a fibrillar assembly to a structured spectrum (red line) corresponding to a crystalline structure. Here again, the spectral modifications are in line with the differences observed in Figure 5 for both amide I and amide II bands. The changes in SAXS patterns (Figure 6F), in this case, are not drastic although we observe a clear increase of diffraction peak intensity indicating growth of the crystalline phase as pH level decreases.



Figure 6: Structural characterization of WMDF samples before and after pH change. A-B) 5% WMDF solubilized in water (red line) then submitted to ammonia vapors (blue line). C) 5% WMDF solubilized in 0.2 M NaOH (blue line) then submitted to glacial acetic acid vapors (red line). D) 9% WMDF solubilized in 0.2 M NaOH (blue line) then submitted to glacial acetic acid vapors (red line). E) 13% WMDF solubilized in 0.2 M NaOH (blue line) then submitted to glacial acetic acid vapors. F) 15% WMDF solubilized in 0.2 M NaOH (blue line) then submitted to glacial acetic acid vapors. The spectra have been shifted for clarification. The stars (\*) in the FTIR spectra indicate the peak arising from acetic acid in the solution. In the middle panels pictures of the corresponding samples stored in Eppendorf tubes are shown.

In Figure 7 we show examples of microscopic investigations with TEM. Initially, the sample is typical of a poorly solubilized WMDF solution with aggregates and crystals (Figure 7A1), which is consistent with the light scattering seen by the naked eye. When basified with ammonia vapor (pH > 11), fibrils appear (Figure 7A2), in agreement with observations made with samples directly prepared in NaOH, albeit some remaining aggregates can be found. Conversely, Figure 7B exhibits a gelated sample initially solubilized in NaOH. When submitted to hydrochloride vapors, the sample becomes more liquid and finally turns blue, which is possibly due to oxidation of part of the sample by the strong acid. In contrast, samples acidified with acetic acid vapours remain uncolored. Transmission electron micrographs show that the fibrils initially structuring the gel (Figure 7B1) have completely disappeared and been replaced by large aggregates of crystals (Figure 7B2), which is coherent with other samples characterized with FTIR and SAXS that did not turn blue.



Figure 7: Transmission Electron Micrographs of A1) A trouble solution of WMDF solubilized in PBS; A2) the same sample after 10 minutes in ammonia vapors; B1) a fibrillar gel of WMDF solubilized NaOH 0.2 M; B2) the same sample after 10 minutes in HCl vapors. Scalebars = 200 nm (Panels A) and 1  $\mu$ m (Panels B).

Overall, these observations prove that both morphologies described in this study are reversible with pH and can be turned into one another. Additionally, the formation of microcrystals from fibrils refutes the idea that the crystals observed in the samples directly prepared at low pH are simply the unsolubilized remains of the powder or the result of nonspecific aggregation that would compete with fibril formation. Given the higher solubility of the peptide at high pH, increasing the pH can also lead to complete dissolution of supramolecular assemblies.

This reversible feature is not so common. It is indeed usually reported that amyloid fibrils are insoluble, very resistant and only disappear under very harsh denaturing conditions. Moreover, after that treatment, no newly formed fibrils would form again. Previously, Kurouski *et al.*<sup>62</sup> have observed that apo- $\alpha$ -lactalbumin fibrils spontaneously refold from one polymorph to another as a result of salt removal and drop of temperature. However, the WMDF system is more comparable with other self-assembling peptides whose sequence also comes from hormones and that also simply disappear upon dilution. Such behavior was observed with Lanreotide (a synthetic analogue of somatostatine) nanotubes that disassemble upon dilution or melt as temperature is increased while retaining their ability to reform nanotubes<sup>63</sup> or  $\beta$ –endorphin amyloid fibrils.<sup>64</sup> Triptorelin (a synthetic analogue of GnRH) also exhibits different morphologies with pH.<sup>19</sup>

# Charge modification by amino-acid exchange and further discussion on the role of charges.

As other studies have illustrated, charges and counterions play a crucial role in supramolecular assemblies (see Gobeaux *et al.* and references herein).<sup>65</sup> It is also anticipated that this role is all the more crucial here that the sequence is short. As expected, modifying one charge, i.e. replacing the aspartic acid (D) in the third position – whose side chain is negatively charged at basic pH - by an uncharged asparagine residue (N), has a direct effect on solubility and assembling properties of the peptide. Indeed, at basic pH levels, where the net charge of the WMNF is zero (Figure 1B), the peptide is now poorly soluble. On the other hand,

at acidic pH levels, WMNF is much more soluble that WMDF despite the fact that they exhibit the same net charge. As can been seen on TEM images (Figure 8), the modification also has a direct effect on the morphology of the objects formed at all pHs. In pure water, WMNF spontaneously forms long, unbranched fibrils of about 12 nm wide (Figure 8A). With a better resolution (Figure 7B), one can perceive a substructure suggesting that the fibrils are composed of two 6 nm wide fibrils. For solutions prepared in 150 mM acetic acid, the supramolecular assemblies take the form of long twisted ribbons (Figure 8C-D) whose width may vary between 25 and 80 nm. A plot of the twist period as a function of the width of the ribbons, taken from 83 individual measurements, suggests a correlation (see inset in Figure 8D). Macroscopically, the entanglement of these ribbons form birefringent gels. At acidic pH levels the fibrils and ribbons all exceed ten microns in length. Whereas at neutral or basic pH levels, the objects formed appear more thickset (Figure 8E-F for 100 mM PBS). They rarely exceed one micron and have an aspect ratio ranging from 5 to 15. Interestingly, these objects are also twisted (black arrows, Figure 8F), albeit not more than one twist is observed on each object.



Figure 8: Transmission electron micrographs of WMNF solubilized in water (A-B), 150 mM acetic acid (C-D) and 100 mM PBS (E-F). Inset in panel D shows the plot of the period between two twists (as indicated by arrows in panel D) as the function of the ribbon width in acidic conditions. The linear regression yields a slope of 7.7 with a coefficient of determination  $R^2$  of 0.6824.

The ATR-FTIR spectra (Figure 9A) indicate a strong and uniform structure and organization into  $\beta$ -sheets: all spectra exhibit a strong peak around 1640 cm<sup>-1</sup> with a FWHM lower than 20 cm<sup>-1</sup> (the fitted Lorentzian-Gaussian peaks yield values between 11 and 14 cm<sup>-1</sup>). At all pHs tested up to 7.4, the main peak is located at 1641.8 cm<sup>-1</sup>. In NaOH (pH ~13), it shifts to 1639.2 cm<sup>-1</sup>. Other peaks traditionally attributed to  $\beta$ -sheets are also observed at 1627 and 1685 cm<sup>-1</sup>, which again suggests an antiparallel organization of the  $\beta$ -sheets. Finally, another peak is detected at 1662 cm<sup>-1</sup> that can be attributed to disorder. The peaks in the amide II region (1500-1600 cm<sup>-1</sup>) do not change with pH and differences with those observed in the WMDF spectra are minor.

The SAXS characterization of the WMNF solutions also points towards a  $\beta$ -sheet organization. At both pHs tested, the equatorial and meridional reflections of the cross β-sheet are detected. However, the structures formed at a low pH are a lot more organized than those formed at higher pHs. Here, the scattering curve corresponding to the sample prepared in PBS only shows a few reflections and an increasing scattering at low q indicating the presence of large, polydisperse scattering objects. The meridional reflection at 1.25 Å<sup>-1</sup>, indicates an average interstrand spacing of about 5 Å, a slightly higher value than the 4.7 Å commonly reported. The 11.7 Å spacing between the sheets corresponding to the equatorial reflection is a more typical value. By contrast, the scattering curves of WMNF in acetic acid suggest a more regular structuration. The meridional reflection at  $1.31 \text{ Å}^{-1}$  (4.79 Å) indicates a tighter packing of the peptide strands. The strong correlation peak 0.0296 Å<sup>-1</sup> (~212 Å) is characteristic of interactions between the scattering objects while the oscillations indicate their size monodispersity. The red curves are taken along the capillary and show that the sample is not homogeneous, probably because of sedimentation, and that the correlation peak disappears readily as the fibril concentration decreases. The curves without the correlation peak can be reasonably fitted by a cylindrical form factor. The fitted radius is about 5.2 nm, which is of the same order of magnitude than the radius of the objects observed by TEM in Figure 8A-B.



Figure 9: Characterization of the WMNF gels. A) Amide I & II regions of the ATR-FTIR spectra of 10% w/w WMNF solutions solubilized in 0.2 M acetic acid (1), water (2), 0.1 M PBS (3) and 0.2 M NaOH (4). B) SAXS-WAXS patterns of 10% WMNF gels in acetic acid (red lines) and in PBS (green line). A fit of the SAXS part assuming a cylindric shape is plotted in purple. The noise in the 0.2-0.8 Å<sup>-1</sup> range appeared with the subtraction of the kapton cell scattering. The insert is a blow-up of the interstrand amyloid peak plotted on a linear scale.

#### Phase Diagram.

From the multiple conditions that we have tested, it emerges that there is a fine balance between the solubility of the peptide and its self-assembly properties. For example, at pH 11, the critical assembly concentration is lower than at pH 13 (~3% vs ~10%) where the WMDF monomer is highly soluble because of its net charge. Likewise, the assembly/gelification process is faster at pH 11. In the case of the WMNF peptide, it is interesting to note that the excellent solubility of the asparagine amino acid (which is 5 times higher than that of aspartic acid in water) is transferred to

the whole peptide sequence. However, as emphasized by Do *et al.*<sup>66</sup>, the terminal net charge is very important in regulating solubility and assembly. In the present case, it appears to be essential. Indeed, when the N-terminus is deprotonated at basic pH levels, the WMNF solubility is poor, while in the case of WMDF the deprotonation of the aspartic acid balances the high hydrophobicity of the aromatic-rich sequence.

Additionally, the peptides themselves contribute to the solutions buffering and there is a complex feedback between the pH of the solutions, the state of charge of the peptides and their state of assembly.<sup>65</sup> As a consequence, at high peptide concentrations the pH might be slightly different than at lower concentrations. As we have seen, the morphology of the WMDF and WMNF supramolecular assemblies are sensitive to the pH of the solution. This explains why in some cases the morphology observed in TEM does not exactly match that inferred from SAXS measurement for the same buffer. Indeed, the concentration of solutions needed for TEM observations are significantly lower than that necessary to obtain a good X-ray scattering signal. Likewise, the polymorphism and variable solubility observed when WMDF is solubilized in PBS (pH 7.4) may be attributed to its complex zwitterionic nature. Indeed, at high concentrations, it is possible that the peptides compensate their charges two-by-two. The resulting dimers would then further associate in many different ways in absence of uncompensated charges to direct the assembly with charge repulsion. This mechanism would then lead to the formation amorphous aggregates.6

Our observations are summarized in the phase diagram Figure 10 and in Table 1. There are basically four zones: (a) a microcrystal phase at low pH, (b) a non-assembled phase at high pH/low concentration, (c) a fibrillar phase at high pH/high concentration and (d) coexisting fibrils and crystals at intermediate pH/intermediate concentrations and high pH/very high concentrations. Table 1 compares the solubility, morphology and net charge of WMDF and WMNF at three different pH levels. For simplicity, we have only discussed the influence of the pH. However, we have used different buffers and we cannot exclude that the different type of ions can play a role in the assembly properties.



Figure 10: Phase diagram of WMDF. As pH level increases, the peptide solubility increases. Additionally, morphology changes from microcrystals at low pH to classic amyloid fibrils at higher pHs. The possible scenarios for hormone release, either by dilution (path (i)) or pH change (path (ii)) are illustrated by yellow arrows. Path (ii) corresponds to the experiment presented in Figure 6A-B and Figure 6C-D and path (iii) to the experiment presented in Figure 6E-F.

Table	1: Summary comparing WMDF and WMNF	assembly
	properties at three pH levels.	

		WMDF	WMNF
		(+)WMDF	(+)WMNF
2.5	Net charge	+1	+1
Ηd	Solubility	Bad	Good
	Morphology	nanocrystals	Fibrils
		(+)WMD(-)F	(+)WMNF
7.4	Net charge	0	+1
μd	Solubility	Mediocre	Good
_	Morphology	Nanocrystals/fibrils	Twisted ribbons
		WMD(-)F	WMNF
13	Net charge	-1	0
μd	Solubility	Good	Mediocre
	Morphology	Fibrils	Twisted aggregates

#### **Biological implications.**

Although most of the conditions explored in our *in vitro* study can be considered to be far from physiological conditions, some lessons can be drawn from our observations.

First, as observed with other protein hormones<sup>16,68,69,70</sup>, the CCK-4 aggregates are very likely to be used as storage means in secretory vesicles in which they undergo post-translational processing.<sup>71</sup> There are indeed several advantages to storage into reversible amyloid structures. The most obvious advantage is to reach high concentrations in small volumes while keeping an iso-osmotic relationship with the surrounding cytoplasm. Furthermore, an assembled form might be able to oppose the lysis or at least to slow it down. Indeed, it has been measured that the CCK-4 half-life in human plasma is 13 minutes.<sup>72</sup> As noted by Dannies, aggregation is also an economical means of sorting because self-association removes the hormones from other possible pathways.<sup>68</sup>

Secondly, the influence of concentration and pH on the aggregation may provide explanations on how CCK-4 neuro-hormones are released in vivo. Indeed, it has been proposed that vesicles containing aggregated hormones fuse with plasma membrane, so their dense-cores are released and dissolve, causing a burst of hormone in the bloodstream on a timescale much faster than could result from increasing synthesis.<sup>68</sup> If we consider the approximate phase diagram drawn from our observations (Figure 10), two dissolution mechanisms are possible: either by simple dilution (i) or by a small shift in pH that would allow shifting from aggregated form to the unassembled form at a fixed concentration (ii). This would be consistent with the literature that reports slightly acidic pH levels in secretory granules (low c.a.c) and higher pH level in the plasma in which the hormones are released (higher c.a.c). Admittedly, we see no biological relevance to path (iii) except that it demonstrates the reversibility of both the assembled forms. Even though the microcrystal morphology may at first seem unlikely to be encountered in vivo' it has been reported that in certain conditions insulin is stored as microcrystals in densecore secretory granules.<sup>73,74</sup> As mentioned above, there is no monomorphism requirement for a storage purpose. What is more, microcrystal suspensions can be envisioned as an easily syringeable drug formulation that would release monomers when dissolved in physiological conditions.75

Thirdly, by working with solutions reaching up to 20 % w/w (> 200 mg/mL) we are closer to the crowding in cells<sup>76,77</sup> than most *in vitro* studies that use solution concentrations around 2 mg/mL. Concentrations in dense core granules might even reach higher values.<sup>78,79</sup> Further studies should try to mimic the *in vitro* condition in terms of salt composition (multivalent ions) and cosolutes like glycoaminoglycans that have been reported to increase aggregation and could also have an effect on supramolecular aggregates morphology. However, in this study we have favored "extreme" and somewhat unrealistic salt and pH conditions. Indeed, since we hypothesized that the different morphologies were guided by the state of charge, we tried to reduce polymorphism by working at pH levels far from the peptide pK<sub>a</sub>s.

Finally, although the present study does not deal with kinetics (we will report thorough kinetic measurements in a forthcoming article), we have observed that aggregation and gelification usually happens within a few hours if not minutes which is also coherent with what would be a functional behavior *in vivo*. For example prolactin, whose secretory pathway has been studied in details, is detected in aggregates only 30 minutes after synthesis.<sup>68</sup>

# CONCLUSION

In conclusion, the reported results show that even a short peptide sequence can hold well-defined and controllable assembly properties. With only four amino-acids in its sequence the Cholecystokinin/Gastrin tetrapeptide (aka WMDF or CCK-4) can attain different conformations that lead to polymorphic supramolecular assemblies. By using rather extreme pH conditions, we show that it is possible to choose one or the other morphology. Furthermore, the formation of these morphologies is reversible upon pH modification. We also show that the charge modification obtained by replacing the aspartic acid by asparagine is able to modify the morphology of the amyloid. Finally, although we have little in vivo data specific to CCK-4 to compare with our in vitro results, the behavior and phase diagram of CCK-4 aggregation are compatible with the requirements for a storage application. Further studies will examine the kinetics of formation and release of these thermodynamically-driven fibrils structures and hopefully bring new information on the generic mechanisms of amyloid assembly.

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# ASSOCIATED CONTENT

**Supporting Information Available.** Diffraction peaks indexation of WMDF in acetic acid and PBS with corresponding cell parameters. Contact potential estimations. WMDF FTIR fits. Full FT-Raman plot and peak attribution table. ThT fluorescence assays. Steady state fluorescence study. Amyloidogenicity predictions with online tools. WMNF FTIR amide I fit.

This information is available free of charge via the Internet at http://pubs.acs.org.

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