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Fast Flow Microfluidics and Single-Molecule Fluorescence for the Rapid Characterization of α -Synuclein Oligomers

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- Supporting Information

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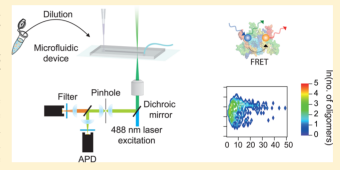
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ABSTRACT: α -Synuclein oligomers can be toxic to cells and may be responsible for cell death in Parkinson's disease. Their typically low abundance and highly heterogeneous nature, however, make such species challenging to study using traditional biochemical techniques. By combining fast-flow microfluidics with single-molecule fluorescence, we are able to rapidly follow the process by which oligomers of αS are formed and to characterize the species themselves. We have used the technique to show that populations of oligomers with different FRET efficiencies have varying stabilities when diluted into low ionic strength solutions. Interestingly, we have found that oligomers formed early in the aggregation



pathway have electrostatic repulsions that are shielded in the high ionic strength buffer and therefore dissociate when diluted into lower ionic strength solutions. This property can be used to isolate different structural groups of αS oligomers and can help to rationalize some aspects of αS amyloid fibril formation.

arkinson's disease (PD) is the second most common neurodegenerative disease after Alzheimer's disease, 26 affecting more than 1% of the world's population of those 27 aged over 65 years. Clinically, the disorder is characterized by a 28 loss of control of movement and the onset of resting tremors, 29 muscle rigidity, and bradykinesia. Additionally, there are 30 nonmotor characteristics, such as cognitive impairment, 31 depression, olfactory deficits, psychosis, and sleep disturbance 32 during the progression of the disease. Neuropathologically, PD 33 is characterized by the progressive loss of dopaminergic 34 neurons within the substantia nigra pars compacta of the 35 midbrain. One of the major pathological hallmarks of PD is the 36 presence of intraneuronal proteinaceous cytoplasmic inclusions, 37 referred to as Lewy Bodies (LBs). The finding that amyloid 38 fibrils of the protein α -synuclein (α S) are a major component 39 of LBs suggests that it is involved in the etiology of the disease.⁴ 40 LBs are also found in other neurodegenerative disorders such as 41 dementia with LBs, the LB variant of AD,⁵ and Down's 42 syndrome. Duplication and triplication of the chromosomal 43 region surrounding the gene coding for αS and a number of 44 point mutations⁸⁻¹³ lead to dominantly inherited PD, 45 providing further evidence for the link between the protein 46 and the diseases.

The finding that purified recombinant αS forms fibrils resembling those found in LBs¹⁴ has led to speculation that 49 aggregation plays a key role in the disease. When αS aggregates 50 in vitro, small soluble oligomers, which have been shown to be

cytotoxic, are generated, $^{15-17}$ such species can, in some cases, 51 progress directly into fibrillar structures. Despite the evidence 52 of their importance to the disease, the usually low abundance 53 and highly heterogeneous nature of the oligomeric species 54 formed during the aggregation reaction make them difficult to 55 study using traditional biochemical techniques. We have 56 previously used single-molecule fluorescence techniques to 57 characterize the aggregation and behavior of amyloid oligomers 58 of the SH3 domain of PI3 kinase, 18 the amyloid-beta 59 peptide, 19,20 and α S. 16 For the latter, we used single-molecule 60 Förster Resonance Energy Transfer (FRET) to show that α S 61 initially forms amorphous oligomers having a low FRET 62 efficiency, which consequently undergo a structural rearrange- 63 ment to form proteinase-K resistant, cytotoxic oligomers, 64 having a higher FRET efficiency. 16

The oligomers of αS generated during fibril formation 66 typically only make up a small fraction of the total protein 67 concentration, and so single-molecule techniques are needed to 68 detect their presence and characterize their structures. The 69 picomolar concentrations required for single-molecule detec- 70 tion, however, mean that a significant time is needed to collect 71 enough data to make useful conclusions. Through incorporat- 72 f1

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73 ing fast-flow microfluidics (Figure 1) to rapidly pass the low 74 concentration solutions through the confocal volume (at a

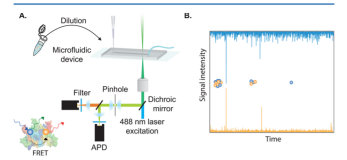


Figure 1. A. Instrumental setup. The microfluidic device is mounted onto a confocal microscope in which 488 nm radiation is focused down to a diffraction-limited spot within the 100 μ m wide channel. Fluorescence from Alexa Fluor 488 (AF488) and Alexa Fluor 594 (AF594) (excited indirectly via FRET) can then be collected and separated onto two separate Avalanche photodiode detectors (APDs). B. Example intensity traces. Bursts of intensity arise when fluorescently labeled α S transits the confocal volume. Oligomeric species have a burst in both the donor (blue trace) and acceptor (orange trace) channel, allowing them to be separated from the majority of monomeric protein (donor bursts only).

75 velocity of 2 cm/s), we have now reduced the data acquisition 76 time from 3 h, to just over 5 min per time-point, thereby 77 allowing us to increase drastically the time-resolution of our 78 technique and also to run several aggregations on one 79 instrument simultaneously. In this paper, we illustrate the 80 power of this technique by studying in more detail the 81 aggregation of α S.

The conformation of monomeric αS is affected by factors such as pH²¹ and ionic strength, 22 and we speculated that certain types of oligomers may be electrostatically destabilized to intermonomer electrostatic repulsion because of the high net charge of the C-terminal region. To test this hypothesis, we used the increased data-acquisition rate of this technique to study the effect of diluting the time-points from the aggregation into high and low ionic strength buffers. We found that the amorphous type oligomers, which are less cytotoxic, and more susceptible to proteinase-K digestion, dissociate at low ionic strengths, whereas the higher FRET species are stable to such a change in conditions.

■ EXPERIMENTAL SECTION

Protein Expression and Purification. BL21(DE3) Gold cells (Stratagene) were transformed with the Cys mutant A90C. Starters were diluted into Overnight Express Instant TB Medium (Novagen) supplemented with 1% glycerol and left to grow for 16–18 h at 30 °C. Following this, cells were harvested, and the protein was purified as previously published. 16

Device Fabrication. Microfluidic channels were fabricated using standard soft-lithography techniques into polydimethylsi-lo3 loxane (PDMS; Dow Corning) with SU-8 photoresist on silicon masters, as described previously. The channels were plasma-bonded to glass coverslides (V.W.R., thickness = 1) to create sealed devices. The channel height was 25 μ m. Each device was inspected on a white-light microscope (Nikon Ti-108 U), and only those without dust or aberrations were used.

Protein Labeling. The Cys variant of wild-type α S was labeled as reported previously. ¹⁶ Life Technologies Alexa Fluor H88 C₅ maleimide (AF488) and Alexa Fluor 594 C₅ maleimide

(AF594) were used in these reactions. Labeled protein was 112 isolated from unreacted dye using the method reported in 113 Cremades et al., 2012. The reaction yield was checked by 114 mass spectrometry for all reactions, and all labeling reactions 115 with a yield lower than 90% were discarded.

Single-Molecule Fluorescence Measurements of Ag- 117 gregating α S. A 70 μ M solution made up of equimolar 118 concentrations of AF488 and AF594 labeled α S was prepared 119 in 25 mM Tris-HCl, pH 7.4, 100 mM NaCl. The buffer was 120 freshly prepared before each experiment and passed through a 121 0.02 μ m syringe filter (Anotop, Whatman) to remove insoluble 122 contaminants. The aggregation mixture was supplemented with 123 0.01% NaN3 to prevent bacterial growth and was then 124 incubated at 37 °C with orbital shaking at 200 r.p.m. (25 125 mm orbital diameter). Regular time-points were taken 126 throughout the course of the aggregation and were immediately 127 diluted by a factor of 1:250,000 (in two stages) before being 128 loaded into a 200 µL gel-loading tip (Life Technologies, 129 Carlsbad, CA, USA) attached to the inlet port of a microfluidic 130 channel (25 μ m in height, 100 μ m in width, 1 cm in length) 131 mounted onto the single-molecule confocal microscope. The 132 confocal volume was focused 10 μm into the center of the 133 channel, and the solution was passed through the channel at an 134 average velocity of 2 cm/s by applying a negative pressure, 135 which was generated using a syringe pump (Harvard apparatus, 136 Holliston, MA, USA) attached to the outlet port via Fine Bore 137 Polyethylene Tubing (0.38 mm inner-diameter, 1.09 mm outer- 138 diameter; Smiths Medical International, Hythe, Kent, UK). 139 Before adding the gel-loading tip and withdrawing the sample, 140 the syringe, tubing, and microfluidic channel were first filled 141 with buffer, and any air-bubbles were purged from the system. 142 After the appearance of single-molecule bursts corresponding 143 to labeled α s passing through the confocal volume, the sample 144 was measured for 400 s. There was no noticeable decrease in 145 the event-rate over the duration of the measurement, 146 suggesting that any surface absorbance does not affect the 147 measurement.

Single-Molecule Instrumentation. A Gaussian laser 149 beam at a wavelength of 488 nm was first attenuated using 150 neutral density filters and passed through a spatial filter before 151 being directed through the back-port of an inverted microscope 152 (Nikon Ti-U). A dichroic mirror (Di01-R405/488/594 153 Semrock) reflected the laser light through an oil-immersion 154 objective (Nikon CFI Plan Apochromat VC 60X Oil N2 NA 155 1.4, W.D 0.13 mm), which focuses it to a diffraction-limited 156 confocal spot within the sample being studied. The emitted 157 fluorescence was collected by the same objective and passed 158 through the dichroic, before being focused by a tube lens within 159 the microscope body through a 50 μ m pinhole (Thorlabs). A 160 second dichroic (585DRLP Horiba) then separates the 161 fluorescence from the two different fluorophores; the longer 162 wavelength passes through the dichroic and was focused by a 163 lens (Plano apo convex, focal length = 50 mm, Thorlabs) 164 through a band-pass filter (FF01-629/53) onto the Avalanche 165 Photodiode (APD) detector. The shorter wavelength was 166 reflected by the dichroic and is focused through a second set of 167 filters (535AF55 Horiba, 540LP Omega) onto the second APD. 168 Outputs from the two APDs are connected to a custom- 169 programmed field-programmable gate array, FPGA (Colexica), 170 which counts the signals and combines them into time-bins, 171 which are selected according to the expected residence time of 172 molecules traveling through the confocal volume.

Figure 2. A. Device design. The microfluidic chip consists of a single-channel, 100 μ m in width, and 25 μ m in height, red scale bar is 500 μ m in length. A gel-loading tip containing the analyst is added to the sample inlet, and the outlet is attached to a syringe within a syringe pump to pull sample through the device. Zoom: Brightfield image of the microfluidic device channel (scale bar is 100 μ m in length). B. and C. Cumulative frequency histogram of monomeric and oligomeric events, respectively, over a 400 s measurement of 280 pM aggregated αS. The histograms are fitted to a straight line yielding R^2 = 0.999 and R^2 = 0.998 for monomeric and oligomeric events, respectively. This shows that the event detection rate remains constant over the measurement time.

Stability in Different Ionic Strength Solutions. To determine the stability of different samples of αS oligomers with respect to varying ionic strength, the solution of αS was diluted by a factor of 1:250,000 into a 5 mM tris buffer (pH 7.4) containing a range of NaCl concentrations (from 0 to 100 mM). The diluted sample was then immediately analyzed on the single-molecule confocal instrument.

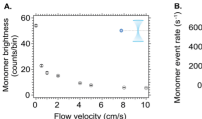
RESULTS AND DISCUSSION

Microfluidic Device Design. In order to determine 182 183 whether an increase in the data acquisition rate for αS 184 aggregation samples could be achieved using fast-flow, a simple 185 microfluidic device was designed and made from PDMS. The 186 device consists of a single channel, 100 μ m in width, 25 μ m in 187 height, and 1 cm in length. The width and height were selected to prevent larger fibrils blocking the device, and this is demonstrated by the constant monomer and oligomer 190 detection rate over the duration of the measurement, even at 191 later time-points when fibrils are present (Figure 2B and C). 192 The aggregation samples were diluted before being loaded into 193 a low-binding gel-loading tip, which was inserted into the entry port of the device (Figure 2A.). Tubing connected to a syringe within a syringe pump was attached to the exit port, allowing for accurate control of the flow-velocity by application of a negative pressure. By withdrawing sample in this way, rather 197 than infusing sample directly from a syringe, the dead volume was minimal, and the exposure of the samples to hydrophobic surfaces, such as the inside of plastic tubing, was minimized.

Increase in Data Acquisition Rate with Fast Flow 202 Microfluidics. Many steps in protein aggregation reactions are likely to take place on time-scales that are shorter than the 3 h time points used in our previous study on the oligomers formed during the aggregation of α S. ¹⁶ We sought to increase the time resolution by using fast flow microfluidics. It has previously been shown that it is possible to increase the rate of data acquisition in single-molecule confocal experiments by flowing the molecules through the probe volume at high speeds (see Figure 1).²³ There is, however, an upper limit to the velocity at which molecules can flow through the confocal volume while still enabling the collection of meaningful data. At higher speeds, there is an increase in the rate at which fluorescent 214 species transit the probe volume; in parallel, there is, however, a 215 decrease in the number of excitation-emission cycles of the 216 fluorophores, reducing event brightness. Thus, there is an 217 optimum velocity at which the event rate is high, but the 218 brightness of the fluorescent bursts is not sufficiently low to 219 make them undetectable. With two-color excitation, the rate at which it is possible to flow and still get meaningful data is 220 higher than for the FRET experiments in which only one color 221 is directly excited. For the work presented here, only the donor 222 dyes are directly excited, and the presence of signal in the 223 acceptor channel due to FRET is used to identify the oligomers 224 in the presence of an excess of monomers. The oligomers, 225 however, typically have more than one donor fluorophore 226 present, and so it is possible to flow at higher velocities than 227 those used for detecting species with only one directly excited 228 donor dye.

To determine the optimum flow velocity for our experiment, 230 an aliquot of αS solution was removed after incubation under 231 aggregation conditions for 24 h and diluted to 300 pM, before 232 being flowed through a simple one-channel device mounted on 233 the single-molecule confocal instrument at a range of velocities. 234 With increasing flow velocity, it was necessary to increase the 235 laser intensity, as under these conditions, the molecules spend 236 less time in the confocal volume. The optimum intensities were 237 determined previously²³ by selecting those laser powers that 238 gave the highest brightness at each velocity, up to the maximum 239 power achievable on our instrumental setup. The brightness of 240 the bursts is also dependent on the flow velocity, and so it was 241 necessary to vary the thresholds (see the Supporting 242 Information for details of thresholding and data analysis) in 243 the donor and acceptor channels for each flow velocity. To 244 achieve this in an unbiased way, we have previously²³ shown ²⁴⁵ that the thresholds at each flow velocity can be selected to give 246 the maximum fraction of coincident fluorescent bursts in both 247 channels using a dual labeled DNA duplex as an example. 24 The 248 flow velocities, time bin-widths, and automatically selected 249 thresholds are presented in SI Table 1. Figure 3 shows the 250 f3 effect of varying flow speed on the measured monomer 251 brightness (A) and the donor channel burst-rate (B). Even with 252 the increase in laser power that can be used at higher flow 253 velocities, the monomer brightness decreases due to the 254 molecules spending less time in the probe volume and 255 therefore completing fewer excitation-emission cycles. How- 256 ever, with increasing flow velocity, the donor burst-rate initially 257 increases, reaching a maximum of 530 ± 115 burst s⁻¹ at a rate 258 of 5 cm/s. At higher velocities, there is a decrease in the 259 number of monomers detected, as the dye molecules are unable 260 to emit enough photons to be detected above the applied 261 thresholds.

As the oligomers are likely to contain more than one donor- 263 labeled molecule, they are brighter than the monomer alone 264 and have a greater probability of emitting sufficient photons to 265 be detected above the threshold level at higher flow velocities. 266 64



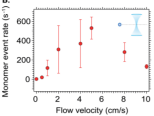


Figure 3. Effect of flow velocity on A. the measured mean monomer brightness and B. the monomer burst rate (mean \pm SD, n=3) when 280 pM of α S was passed through the microfluidic channel at varying flow velocities. With increasing flow velocity, the intensity of the bursts decreases due to fewer excitation-relaxation cycles in the shortened residence time in the probe volume. By contrast, the donor burst-rate initially increases, due to a faster encounter rate with the probe volume, but then decreases as a larger proportion of the molecules become insufficiently bright to detect.

267 Figure 4A shows the number of oligomers detected as a 268 function of the mean flow velocity. As with the rate of 269 monomer detection, there is an initial increase in the number of 270 events detected as the flow velocity increases, before a slight 271 decrease at higher velocities.

The association quotient, Q (defined in the Supporting 2.72 273 Information), indicates the fraction of molecules that 274 simultaneously exhibit bursts of fluorescence above the applied thresholds in both detection channels. Figure 4B shows how Q 276 varies as a function of flow velocity for the oligomeric sample 277 solution. Increasing the mean flow velocity from 0 to 0.5 cm s 278 leads to a slight increase in Q₁ as observed previously for dual-279 labeled DNA duplexes. 23 At higher velocities, there is a 280 decrease in the Q value, due to the reduction in the brightness of the fluorophores, which are unable to complete sufficient 282 excitation-emission cycles within the probe volume. In 283 performing experiments under fast-flow, it is desirable that 284 the flow velocity is high enough that many events are detected 285 but not so high that the number of photons detected limits the 286 quality of the data. To take both of these factors into account in 287 selecting the optimum flow velocity, the product of the event rate and Q against the varying flow speed was determined and 289 is plotted in Figure 4C. There is an increase in the product up 290 to 2 cm/s, followed by a plateau at higher velocities indicating 291 that a mean flow velocity of 2 cm/s is ideal for performing 292 measurements of the oligomeric species under investigation

here, since the event rate is increased (events are detected 293 \sim 150 times more frequently than under static conditions), 294 while the Q value is still reasonably high.

A further advantage of using microfluidic delivery to the 296 confocal volume is that the flow velocity of the individual 297 species is independent of their diffusivity/size, as the molecules 298 are under laminar flow. In stationary measurements, relying on 299 diffusion only, larger species are able to occupy the probe 300 volume for a prolonged period of time, which can lead to 301 multiple counting of events. In addition, the path taken through 302 the confocal volume is more constant under flow (i.e., the 303 molecules flow in one-dimension, rather than traveling through 304 via Brownian motion), meaning that the transit time remains 305 constant, and any intensity changes due to varying residence 306 times are removed. Also, since the flow is constant regardless of 307 the species size, then species that are larger than the confocal 308 volume diameter, such as fibrils, will occupy a greater number 309 of time-bins than smaller species such as oligomers, as it takes a 310 longer period of time for them to transit fully the confocal 311 volume. This effect allows for fibrillar species to be separated 312 from the smaller oligomeric events, which do not occupy 313 multiple bins (see the Supporting Information).

Following the Kinetics of an in Vitro Aggregation of 315 α **5.** To determine the ability of the enhanced methodology to 316 follow the in vitro formation of α S oligomers, solutions of α S 317 were incubated under conditions favoring aggregation, and 318 aliquots were taken and analyzed at regular time intervals using 319 single-molecule fluorescence. As monomers become incorporated into oligomers and fibrils, their numbers decrease, and 321 this effect can be observed in Figure 5, which shows the 322 fs number of donor fluorescent bursts detected above the applied 323 threshold within the 400 s measurement time for the 50 h time-course of the aggregation reaction.

For each event detected, the FRET efficiency and 326 approximate size of the oligomer giving rise to it can be 327 determined (see the Supporting Information for details) and 328 then binned into histograms (Figure 6A). As time progresses, 329 66 the range of the sizes increases; after 6 h of incubation, there 330 are a few events attributable to an oligomer containing more 331 than 10 monomer units, whereas after 48 h of incubation, a 332 significant number of events have larger sizes of up to 50 333 monomer units. This is expected as the oligomers grow in size 334 through monomer addition. It is also evident that there is a 335 transition from species with a low FRET efficiency to those 336

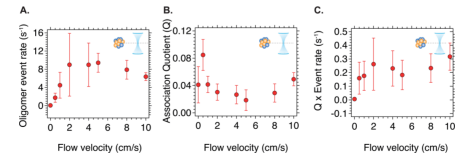


Figure 4. A. The effect of increasing flow velocity on the oligomer event rate when a 280 pM sample of aggregated α S (starting monomer concentration) was passed through the microfluidic channel at varying flow velocities. With increasing flow rate, the event rate increases, making a plateau level after 2 cm/s, before decreasing slightly (mean \pm SD, n = 3). B. The association quotient as a function of flow velocity (mean \pm SD, n = 3). There is a slight increase in the association quotient in changing the flow velocity from 0 to 0.5 cm/s; however, it decreases at higher velocities due to the fluorophores being less bright and therefore less easily detectable. At higher velocities still (>5 cm/s), the association constant slightly increases; however, this is due to the significant decrease in the monomer event rate (Figure 1B). C. The product of the association quotient and event rate increases from 0 to 2 cm/s before plateauing (mean \pm SD, n = 3).

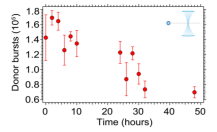


Figure 5. Change in the number of donor bursts measured over time during the aggregation of α S in 150 mM tris buffer with 100 mM NaCl (mean \pm SD, n = 3). The donor burst rate decreases as monomers are incorporated into oligomers and fibrils, because their concentration in solution is lower.

337 with a high FRET efficiency as the aggregation progresses. After 338 6 h of incubation, 95.9% \pm 6.2% (mean \pm SD, n = 3) of the 3—

150mers are within the lower FRET efficiency population, whereas 339 at 48 h, only $37.8\% \pm 4.2\%$ (mean \pm SD, n=3) of them are 340 within the lower FRET population.

By generating FRET efficiency histograms from events 342 corresponding to dimers, and species greater in size than 343 trimers, it is possible to observe a change in the FRET 344 efficiency more easily (Figure 6B). For the smaller apparently 345 dimeric oligomers, only one FRET population is observed, 346 although this could be a mixture of the two populations 347 observed for larger species (it also has a FRET efficiency in 348 between those of the two populations of larger species), but 349 since the events must have a greater number of photons than 350 the applied thresholds in both the donor and acceptor channel, 351 events with either very high or low FRET efficiencies and few 352 emitted photons may not be detected, and this may prevent the 353 two peaks from being resolved for the smaller sized oligomers. 354 For the larger oligomers, two peaks can be clearly resolved and 355

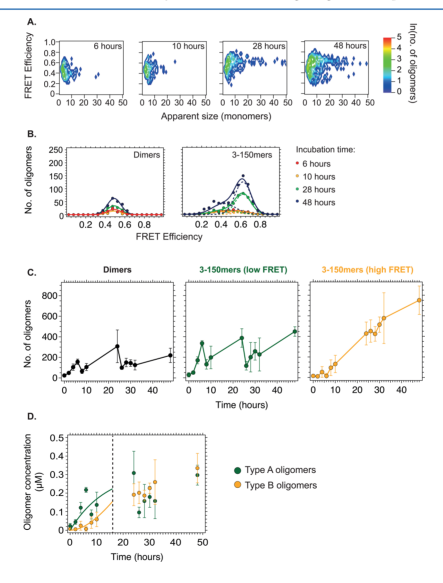


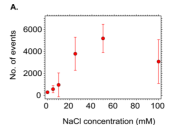
Figure 6. A. 2D contour plots of size and FRET efficiency of various times from solutions of aggregating α S. The size and FRET efficiency both increase over time. B. FRET efficiency histograms for various time-points taken during the aggregation of α S. The oligomers are separated into two different size groupings for which the populations were identified. C. Kinetic traces for the three different populations of α S aggregates (mean ± SD, n = 3). There is an increase in the number of oligomers over time; however, the major increase for the type A oligomers occurs in the first 24 h, whereas for the type B species, the increase is in the second 24 h. D. Results of the global fitting of the kinetics of formation of the type A and type B oligomers (solid green and yellow lines, respectively). The fitting is performed on the first six data-points (up to 10 h), before fibrils can be identified by the presence of a pellet after centrifugation at 15,000 r.p.m. for 15 min.

356 fitted to Gaussian distributions. After integrating the fitted 357 FRET histograms, the changes in the number of oligomers 358 within each population can be plotted (Figure 6C). All three 359 populations show an increase in the number of oligomers over 360 time; for the low FRET form of the larger oligomers 361 (corresponding to the "type A" oligomers identified pre-362 viously 16), the increase appears to occur rapidly after 2 h of 363 incubation and continues at a high rate during the first 10 h. 364 For the higher FRET larger species (corresponding to the "type 365 B" oligomers), the increase does not occur until after 8 h of 366 incubation and then continues at a steady rate for the rest of the 367 time-period. This delay in their presence can be attributed to the fact that they are formed from the conversion of type A oligomers. The apparent dimers show a gradual increase 370 throughout the time-course, as this population is formed from a mixture of the low and high FRET species that cannot be 372 separated due to low photon counts.

In our previous study, 16 the single-molecule data were fit to a nucleation growth model with a conformational conversion step, and we have used the same kinetic analysis here to analyze the analogous data. The model assumes that primary nucleation results in the creation of oligomers of type A from monomeric protein molecules and that type A oligomers can grow through 379 monomer addition or can convert into type B oligomers, resulting in several parallel pathways for the formation of type B oligomers of a given size (details of the early time analytical and exact numerical solutions of the model are presented in the Supporting Information). Only the first six time-points here are fit with this model (Figure 6D), since at later times, additional events, such as the formation of large fibrillar species, the 386 contribution of the reverse processes involving conversion of 387 type-B oligomers into type-A oligomers, and the dissociation of 388 type-A oligomers, become significant and result in under-fitting 389 of the data.

This method of fitting generates two well-defined micro-391 scopic parameters; the primary nucleation rate constant for 392 oligomers of type A was found to be 1.05×10^{-7} s⁻¹, and the 393 conversion rate constant from type A to type B oligomers as 394 2.02×10^{-5} s⁻¹. These values are similar to those determined 395 previously, ¹⁶ suggesting that the information obtained using 396 single-molecule fluorescence and fast-flow microfluidics is fully 397 consistent with the data taken without flow.

Effect of Varying Ionic Strength on the Stability of 399 **Oligomers.** In order to probe the nature of the interactions within the oligomeric species, we studied the effect of ionic strength on their stability. We examined two different samples, 402 one formed after 14 h of incubation in which there are mainly 403 type A oligomers and the other after 38 h in which there are 404 mainly the type B oligomers. These samples were diluted into a solution of 5 mM tris buffer at pH 7.4 containing varying concentrations of NaCl to change the ionic strength. The FRET efficiency histograms are shown in SI Figures 3 and 4, and Figure 7 shows the number of oligomers detected from 409 integrating the fitted FRET efficiency peaks. For the sample 410 collected after 14 h of aggregation, there is a noticeable increase 411 in the number of detected events as the ionic strength of the 412 dilution buffer is increased, suggesting that there are 413 destabilizing charge repulsions in the earlier formed oligomers. 414 Moreover, the destabilization (likely involving disaggregation) 415 of the type A oligomers upon dilution into low ionic strength 416 buffer conditions is rapid and occurs during the time it takes for 417 the sample to be added to the instrument, since the same 418 number of oligomeric events and FRET histograms are found



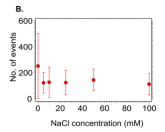


Figure 7. Effect of ionic strength on the number of events detected for aggregating samples of α S after 14 h (A.) and 38 h (B.) of aggregation (mean \pm SD, n=3). The aggregation was done in 150 mM tris buffer with 100 mM NaCl, before being diluted into varying concentrations of NaCl. After 14 h, there is a significant increase in the number of events detected with increasing sodium chloride concentration. After 38 h, the effect of the varying ionic strength on the number of oligomers detected is not pronounced.

when the sample is analyzed at later times. For the sample 419 taken after 38 h of aggregation, however, there is no significant 420 effect of the ionic strength on the detected oligomers. The 421 more compact, higher FRET efficiency oligomers are less 422 susceptible to changes in ionic strength than the earlier formed 423 oligomers, which appear to be destabilized by charge repulsion. 424 In addition to shielding the charge interactions, the higher ionic 425 strength can also increase the stabilizing hydrophobic 426 interactions, which could also account for the oligomers 427 dissociating at lower salt concentrations.

Lowering the pH of the solution, 21 increasing the salt 429 concentration, ²⁵ or removing the C-terminal region of the ⁴³⁰ protein has been demonstrated to drastically increase the rate 431 of α S aggregation. Hover et al.²⁶ and Dedmon et al.²⁷ have 432 suggested that the C-terminus interacts with the N-terminus in 433 the monomeric form of the protein, shielding the NAC region 434 and preventing its self-assembly. At high ionic strengths, this 435 interaction will be diminished, leading to an increased 436 association of the monomers. However, the findings here 437 suggest that it is not only the monomer conformation that is 438 affected by varying the ionic strength of the solution but also 439 the stability of the early formed oligomers is affected. At low 440 ionic strengths, fewer early oligomers will be formed, and 441 therefore a decrease in the aggregation rate of the protein is 442 expected as a consequence of a significant destabilization of 443 these oligomers under those conditions.

In a second series of experiments, αS was incubated in 25 445 mM Tris buffer with 100 mM NaCl under conditions favoring 446 protein aggregation, and regular time-points were diluted into 447 either 25 mM Tris with 100 mM NaCl or into distilled water 448 (pH 7) before being analyzed. Figure 8A shows the resulting 449 f8 2D contour plots of size and FRET efficiency, FRET efficiency 450 histograms are presented in Figure 8B, and the numbers of 451 oligomers for each time-point when diluted into either Tris 452 buffer or distilled water are shown in Figure 9. The lower FRET 453 f9 efficiency oligomers (type A) were found to almost entirely 454 dissociate on dilution into distilled water, whereas the high 455 FRET efficiency oligomers (type B) are stable to dilution into 456 distilled water and do not dissociate.

By diluting the samples into distilled water, it is therefore 458 possible to separate the high FRET efficiency oligomers from 459 the lower FRET efficiency ones. These are the oligomers that 460 were previously identified as being the cytotoxic species, ¹⁶ and 461 this method highlights their greater stability and allows them to 462 be distinguished from the second population not only from 463

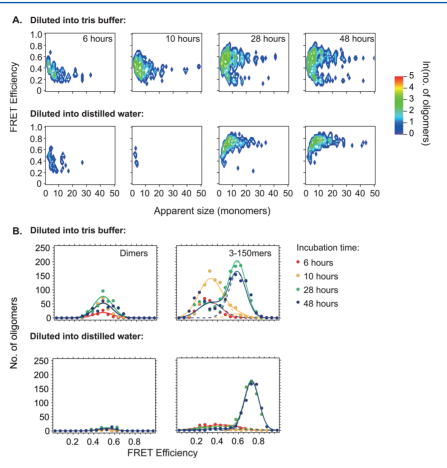


Figure 8. A. Representative 2D contour plots of size and FRET efficiency showing the effect of diluting the samples from an aggregating sample of α S into either 25 mM tris buffer with 100 mM NaCl or into distilled water. B. Representative FRET efficiency histograms of dimers and larger oligomers generated when the samples were either diluted into either 25 mM tris buffer with 100 mM NaCl or distilled water.

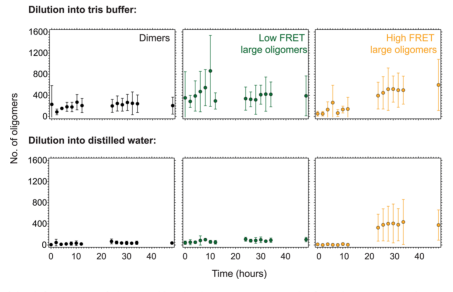


Figure 9. Kinetic traces of the different types of oligomer diluted either into 25 mM Tris buffer with 100 mM NaCl or into distilled water (mean \pm SD, n = 3). The low FRET efficiency oligomers can be seen to be highly destabilized in low ionic strength solutions.

464 their FRET efficiency but also from their sensitivity to ionic 465 strength.

466 CONCLUSION

467 Through the use of a microfluidic device and fast-flow, it is 468 possible to collect single-molecule data 150 times faster than using stationary measurements. At the shorter time accessible 469 by this approach, two different populations of oligomers are 470 observed during the aggregation, one exhibiting a lower FRET 471 efficiency and one resulting from the conversion of these into 472 more compact oligomers having a higher FRET efficiency. The 473 data generated using this method has been further analyzed by 474

475 fitting it to a simple kinetic model in which type A oligomers, 476 are first formed before being converted to type B oligomers, 477 both of which can grow in size via monomer addition. The 478 kinetic parameters determined from this fitting are analogous to 479 those previously determined. Overall, the developments have 480 reduced the time required to analyze a data set from 3 h to just 481 over 5 min, allowing for greater time resolution, and also 482 several experiments to be run simultaneously on one 483 instrument, since numerous samples can be measured back-484 to-back within a short period of time.

The reduced time allows for further biophysical measure-486 ments to be made, and we have shown that by diluting the oligomeric samples into distilled water, the earlier formed lower than the straightful properties of the sociate, possibly because of the charge repulsions destabilizing the oligomers being less shielded. This finding allows for the isolation of the higher FRET efficiency, cytotoxic oligomeric species from a solution of poligomers, simply by changing the ionic strength of the buffer. Our findings substantiate the finding that there are two forms of oligomeric species generated when αS aggregates; not only do they differ in FRET efficiency and susceptibility to proteinase-K digestion but also they behave differently when diluted into low ionic strength buffers.

8 ASSOCIATED CONTENT

499 Supporting Information

500 The Supporting Information is available free of charge on the 501 ACS Publications website at DOI: 10.1021/acs.anal-502 chem.5b01811.

Justification for choice of dyes, the measurement parameters, details regarding the data analysis and modeling, and individual oligomer FRET histograms at different ionic strengths (PDF)

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515 Notes

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516 The authors declare no competing financial interest.

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