# Short hydrogen bonds enhance non-aromatic protein-related fluorescence

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#### 28Abstract

29Fluorescence in biological systems is usually associated with the presence of aromatic 30groups. Here, we show that specific hydrogen bonding networks can significantly affect 31fluorescence employing a combined experimental and computational approach. In particular, 32we reveal that the single amino acid L-glutamine, by undergoing a chemical transformation 33leading to the formation of a short hydrogen bond, displays optical properties that are 34significantly enhanced compared to L-glutamine itself. *Ab initio* molecular dynamics 35simulations highlight that these short hydrogen bonds prevent the appearance of a conical 36intersection between the excited and the ground states and thereby significantly decrease non-37radiative transition probabilities. Our findings open the doors for the design of new 38photoactive materials with biophotonic applications.

# 43Significance statement

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44Intrinsic fluorescence of non-aromatic amino acids is a puzzling phenomenon with an 45enormous potential for technological and biomedical applications. The physical origins of 46this effect, however, remain elusive. Herein, we demonstrate how specific hydrogen bond 47networks can modulate fluorescence. We highlight the key role played by short hydrogen 48bonds in the networks on the ensuing fluorescence and we provide a detailed molecular 49mechanism to explain this unusual non-aromatic optical properties. Our findings should 50benefit the design of novel optically active biomaterials for applications in biosensing and 51imaging.

## 53Introduction

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55Short peptides void of any aromatic residues have been shown to display an intrinsic 56fluorescence in the visible range (1, 2). This has primarily been observed in fibrillar protein 57structures linked to neurodegenerative diseases, such as Alzheimer's, Parkinson's and 58Huntington's diseases (3-6). Furthermore, optical properties of double amino acid based 59nanowires have also been reported, consisting either of two non-aromatic or two aromatic 60amino acids (2, 7, 8). We have previously suggested that the fluorescence of non-aromatic 61short crystal structures forming part of the amyloid-beta protein is enhanced by proton 62delocalisation (5). We have hypothesised that one of the prerequisites for this fluorescence 63observed in either amyloid structures or short peptide nanowires is related to hydrogen 64bonding or aromatic interlocks which, for the latter, decreases the bandgaps down to the 65semiconductive regions (9).

67Despite our previous suggestion that proton delocalisation is strongly coupled to this intrinsic 68fluorescence, its direct role on putative fluorescing states has not been elucidated. We have 69thus searched for a model system, such as a single amino acid-based structure, that displays 70similar optical properties to amyloid fibrils and is permissive to more sophisticated 71computational approaches. We have been inspired by the small peptide nanostructures that 72have been pioneered by the Gazit laboratory (9) and by the fact that there are several 73neurodegenerative diseases that have been connected with an increased level of glutamines 74produced as part of a protein, as for example Huntingtin in Huntington's disease which 75renders the protein more aggregation prone (10). It has been known that the amide group in

76L-glutamine (L-glu) is highly labile and thus can rapidly hydrolyse. We show here that the 77single amino acid L-glu upon heating in water can form a nanostructural material with optical 78properties similar to the ones observed in other amyloid fibrils such as in fibrils of amyloid-79beta, alpha-synuclein or tau (4, 11, 12).

81Using X-ray diffraction (XRD), we show that L-glu dissolved in water and upon heating 82becomes cyclysed forming a previously unreported structure which resembles L-83pyroglutamine and which has been reported to be a component of amyloid-beta in the brain 84(13), but involves a low-barrier hydrogen bonded anionic dimer with an ammonium 85counterion. We have termed the new structure, i.e. L-pyroglutamine complexed with an 86ammonium ion, L-pyro-amm. L-pyro-amm has a microcrystalline plate morphology as shown 87by scanning electron microscopy (SEM). The newly formed solid has a unique hydrogen 88bond network formed by very strong hydrogen bonds (SHB) with a length of approximately 892.45 Å which is confirmed by using terahertz time-domain spectroscopy (THz-TDS).

91By employing a combination of electronic structure calculations and *ab initio* molecular 92dynamics in the ground and excited state we provide an interpretation of the reported 93experiments. We illustrate the important contribution of different vibrational distortions on 94the optical properties. Furthermore, our simulations identify the origins of the non-aromatic 95fluorescence in L-pyro-amm, demonstrating the key role played by the SHBs, which prevent 96the appearance of a conical intersection, significantly reducing the chances of non-radiative 97relaxation towards the ground state and hence increasing the optical properties.

## 99Methods and Materials

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101Experimental

102Sample preparation of L-glutamine

103L-glutamine (L-glu) (#G3126, #G8540, Sigma-Aldrich, Gillingham, UK) and L-104pyroglutamine (L-pyro) (#83160, Sigma-Aldrich) were dissolved in  $18.2\Omega$  MilliQ H<sub>2</sub>O at a 105concentration of 0.3 M or 1 M. Aliquots were placed in a 65°C oven, since heating up 106proteins to 65°C increases the formation of amyloid structures as reported previously (*14*). 107Each aliquot was rotated to dissolve the powder once a day. Samples were either analysed in 108liquid form or dried on a glass or quartz cover slip (#043210.KG, Alfa Aesar, Lancashire,

109UK) either at room temperature (RT) or on a heat block set to 50°C. L-pyro-amm formed 110translucent crystals when dried.

112Emission and excitation wavelength scans

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113Emission and excitation spectra were taken on a Hitachi F-4500 FL spectrophotometer 114(Hitachi High-Technologies Corporation, Tokyo, Japan) at room temperature (RT) in a quartz 115cuvette. For measurements, the excitation slit resolution was 5 nm or 10 nm and the emission 116slit resolution was 20 nm. The PMT voltage was set at 950 V and the scan speed set at 240 117nm/min. The excitation scan was measured between 250 - 400 nm and the emission filter set 118to the emission maxima of the sample stated in the figure legend, with a slit resolution of 20 119nm. The emission scan was measured between 380 - 560 nm and the excitation filter set to 120the excitation maxima of the sample stated in the figure legend, using a slit resolution of 5 121nm. Four measurements were taken for each sample which were repeated at least three times 122and the background (air or  $H_2O$ ) was subtracted from the average.

124Absorption measurements

125Absorption measurements were taken on a UV-Vis-NIR Spectrophotometer, UV-3600 Plus 126(Shimadzu, Kyoto, Japan) and Cary 6000i (Agilent, Santa Clara, USA). 1 M or 0.3 M L-glu, 127L-pyro or L-pyro-amm solutions were measured in 10 mm QX cuvettes (Hellma Analytics, 128Müllheim, Germany) or dried on quartz coverslips. Measurements were taken between 129wavelengths 200 – 800 nm using 1 nm steps at a slow scan speed and a 1 nm resolution. The 130light source change wavelength was set at 393 nm and the grating change wavelength set at 131750 nm. Both direct and integrating sphere measurements were taken and showed little 132difference in results, direct measurements are shown in the manuscript. Samples were 133measured at least three times and the experiments repeated at least three times, measurements 134were then averaged and H<sub>2</sub>O or coverslip only control was subtracted.

136SEM (scanning electron microscopy)

137SEM was performed using a FEI Magellan 400 HR-SEM at an acceleration voltage of 2 kV. 138L-pyro-amm samples were lyophilised by freezing in liquid nitrogen and freeze drying in a 139LyoQuest 85 (Telstar, Terrassa, Spain) and imaged on a glass coverslip.

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141X-ray diffraction

142L-pyro-amm was dried on a glass coverslip in a 50°C oven and then at RT until crystals 143formed. Single crystal X-ray diffraction (SCXRD) measurements were performed at 180 °K 144with a Bruker D8-QUEST PHOTON-100 diffractometer, which utilised a Cu Kα radiation (λ 145= 1.54 Å), and an APEX-II CCD. Absorption corrections were made using SDABS, and data 146integration and reduction were performed with SAINT+. All non-hydrogen atoms were 147refined isotropically and anisotropically, followed by inclusion of the hydrogen atoms 148(determined using the excess electron density) and refinement isotropically. The structure is 149deposited in The Cambridge Crystallographic Data Centre, CCDC No. 1981551.

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151Terahertz Time-Domain Spectroscopy

152All THz-TDS spectra were acquired using a commercial Terapulse 4000 spectrometer 153(TeraView Ltd, Cambridge, UK). Samples were prepared for THz-TDS measurements by 154diluting the solid air dried L-pyro-amm with polyethylene (~ 10% w/w concentration) by 155gentle mixing using an agate mortar, followed by pressing into 2 mm thick, 13 mm diameter 156pellets using a hydraulic press. All THz-TDS spectra shown are a result of division of sample 157and blank datasets, with the blank dataset represented the THz-TDS response of a pellet of 158pure polyethylene.

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160Theoretical

161Density functional theory (DFT)-THz Calculations

162Calculations were performed using both the CRYSTAL17 (15) and Quantum Espresso (16) 163software packages. Geometry optimisations and vibrational analyses performed with the 164CRYSTAL17 code utilised the atom-centred split-valence triple-zeta 6-311g(2d,2p) basis set 165for all atom types. Based on a previous study related to ionic molecular crystals (17), the 166range-corrected WB97-X (18) functional was used. The vibrational analysis was performed 167within harmonic approximation, and infrared intensities were determined using the Berry 168Phase method (19). Energy convergence criteria were set to  $\Delta E < 10^{-8}$  and  $10^{-11}$  hartree for the 169geometry and frequency calculations, respectively.

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171Periodic time dependent (TD)-Density Functional Theory (DFT) Excited State Calculations
172Simulations were performed using the fully periodic Quantum Espresso software package.
173The Becke-Lee-Yang-Parr (B3LYP) hybrid density functional was used with an energy
174cutoff of 40 Ry. The calculations of the excited state were performed within the framework of
175TDDFT using the Liouville-Lanczos formalism implemented in the freely available

176Quantum-Espresso package (20). In this approach, the optical spectra are obtained directly 177over the wide spectral range without taking into account the numerically complex 178calculations of the single exited states. We used plane wave basis set and the electron-ion 179interactions were taken into account via norm conserving Martins-Troullier pseudopotentials 180(21). To determine the ground state wave function, we used the gamma point of the Brillouin 181zone. All the periodic calculations employed the computationally demanding B3LYP (22) 182hybrid functional, the kinetic energy cutoff of 40 Ry was used for the wave functions. The 183intrinsic band width for the spectra was set to 0.003 Ry (~0.0408 eV).

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185Periodic Structure Geometry Optimisation

186The structures obtained from the experiments were first geometrically optimized at 0°K using 187the Broyden-Fletcher-Goldfarb-Shanno (BFGS) minimisation algorithm implemented in 188CP2K (23, 24) package. A convergence criterion for the wave function optimisation was used 189as 5x10<sup>-7</sup> a.u.. Applying the method of the Gaussian and plane wave, the wave function was 190expended in the Gaussian double-zeta valence polarised (DZVP) basis set. The cutoff for the 191electronic density was set to 300 Ry. We used the gradient correction to the local density 192approximation and the core electrons were treated via Goedecker-Teter-Hutter 193pseudopotentials (25). In all the calculations, we used the Becke-Lee-Yang-Parr (BLYP) (26) 194functional with the D3(0) Grimme (27) dispersion corrections for the van der Waals 195interactions.

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197Ab Initio Molecular Dynamics Simulations

198Ab initio Molecular Dynamics simulations (AIMD) were performed using Quickstep 199algorithm implemented in CP2K. In these calculations, the propagation of the nuclei was 200taken into account within the framework of the Born-Oppenheimer approximation. The 201simulations were performed in the NVT ensemble and the temperature was controlled during 202the simulations by using the velocity-rescaling thermostat (28). We used the time step of 0.5 203femtosecond to update the nuclear coordinates and velocities while the total length of the 204simulations for each system is 50 picoseconds.

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206Excited State Cluster Calculations

207A set of excited state calculations were performed on glutamine clusters in order to 208understand the role of the environment on the optical properties. Specifically, the optical

209properties of L-pyro-amm were investigated using various isolated cluster models with the 210Gaussian09 software package. The clusters were extracted directly from the crystal structure 211and used in various combinations (dimers, trimers, tetramers) to perform TD-DFT 212calculations. A split-valence triple-zeta 6-311g(2d,2p) basis set was used for all atom types 213together with the hybrid B3LYP functional. Some benchmark simulations, comparing the 214optical properties obtained from the periodic calculations using B3LYP to range corrected 215hybrid functionals like CAM-B3LYP, were also performed with these clusters.

217We also performed a series of excited state optimisations on various model systems built 218from L-pyro-amm in order to examine the nature of the geometrical distortions that occur on 219the lowest electronic excited state. These calculations were also performed with the 220Gaussian09 software package. All clusters were surrounded with a continuum dielectric 221constant of 80, representing pure H<sub>2</sub>O. The 6-311G(2d,2p) basis set was used for all atoms 222together with the range corrected hybrid functional CAM-B3LYP (29). The clusters were 223first optimised in the ground state after which they were optimised on the first electronic 224excited state.

226Non-adiabatic Decay Probabilities Using Excited State Molecular Dynamics

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227Excited state AIMD was employed, as implemented in the LIO quantum-chemical package 228(https://github.com/MALBECC/lio) (30–33), to analyse the influence of three key factors 229determining the optical properties of L-pyro-amm: (i) the formation of a SHB, (ii) the 230presence of ammonium, and (iii) the combination of ring deplanarisation and carbonyl 231stretching. Our model system for this study was the L-pyro-amm dimer with a single SHB. In 232order to assess the influence of the ammonium ion on the optical properties of L-pyro-amm, 233AIMD was performed both on the L-pyro-amm with and without the ammonium ion. 234Analogously, in order to shed light on the role of the SHB in the L-pyro-amm fluorescence, 235in addition to performing simulations of L-pyro at the natural SHB distance of 2.5 Å, we 236performed three replicas constraining the HB distance to 3.0 Å, 3.5 Å and 4.5 Å, respectively.

238The initial structures for this analysis were extracted from the ground state AIMD described 239above. Subsequently, 3 ps ground state AIMD at 300K was performed to equilibrate the 240system, followed by 1ps of excited state AIMD. The non-radiative decay probability (NRP) 241was computed every time step using the TDDFT-based Trajectory Surface Hopping 242algorithm without permitting decays to the ground state (*34–36*). The non-adiabatic coupling

243elements between *S0* and *S1* (Equation 1) can be estimated analytically employing the 244method introduced by Tapavicza, *et al.*, (*37*).

$$\sigma_{S0,S1} = \langle \Psi_{S0}(r,R(t)) | \nabla_R | \Psi_{S1}(r,R(t)) \rangle \tag{1}$$

246In this context, the non-radiative decay probability (NRP) from *S1* to *S0* (Equation 2) is 247expressed as:

$$NRP(t) = -2 \int_{t}^{t+\Delta t} dt' \frac{Re[c_{S1}c_{S0}^{*}\sigma_{S0,S1}(t')]}{c_{S0}c_{S0}^{*}}, \qquad (2)$$

249where C coefficients satisfy (Equation 3)

$$i\hbar \frac{\partial c_i(t)}{\partial t} = C_i(t)\omega_i - i\hbar \sum_{j=0}^{N_{states}} C_j \sigma_{ij}.$$
 (3)

251NRP(t) is the probability for the system to relax from SI to SO without fluorescent emission at 252time t. Hence, the accumulated non-radiative decay probability (Equation 4) accounts for the 253total relaxation probability from the excitation instant (t=0) up to time t.

$$ANRP(t) = \int_0^t dt' NRP(t') \tag{4}$$

256As mentioned above, this methodology was applied to five different initial conditions of the 257dimer at SHB, 3.0, 3.5, and 4.5 Å of hydrogen distance and for the dimer-amm at SHB 258distance. In order to avoid introducing large structural deformations, the distance constrains 259in the HB were introduced in combination with position restrains on the nitrogen and the  $C_{\beta}$  260of the pyroglutamic ring.

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262The excited state AIMDs were performed employing the density functional theory (DFT) 263using the hybrid functional PBE0 level (38), with a 6-31G basis set and a time step of 0.5 fs. 264The computation of excitation energies, excited-state gradients, and non-adiabatic coupling 265vectors, the Linear Response TDDFT method was used within the Tamm-Dancoff 266approximation (37, 39).

# 268Results and Discussion

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270It has long been known that poly-glutamine can form amyloid-like fibrillar structures *in vitro*. 271The more glutamine residues in the poly-glutamine polymer, the faster the aggregation 272propensity of the polypeptide chain. This led us to investigate whether L-glu on its own, 273under conditions which normally promote fibril formation, such as an increase in temperature

274(14), was able to form structures with similar optical properties, as recently observed for 275amyloid fibrils (5, 40, 41).

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277We first investigated the structure of L-pyro-amm, which formed after incubation of L-glu for 2788 days at 65°C, using SEM and observed crystal structures shown in Fig. 1a. However, in 279order to investigate whether L-glu had indeed changed its crystal structure arrangement we 280performed XRD analysis of the resulting material. In Fig. 1b we show the crystal structure of 281the heated L-glu structure, which we termed L-pyro-amm, and the published crystal 282structures of L-glu and L-pyroglutamine (L-pyro) in Suppl. Fig. 1. Note, the L-pyro structure 283was analysed as it displayed structural similarities to the newly formed L-pyro-amm. Figures 284were obtained from geometry optimisations of the nuclear positions of the atoms using 285experimental densities. L-pyro-amm consists of 8 pyroglutamine groups and 4 ammonium 286ions (144 atoms) complexed within the crystal (see Fig. 1b). In contrast, as shown in Suppl. 287Fig. 1a, L-glu consists of 4 glutamine molecules (80 atoms) in the unit cell which form 288hydrogen bonds involving the termini and side chain. Furthermore, as shown in Suppl. Fig. 2891b, L-pyro consists of 12 pyroglutamine molecules (192 atoms) in the unit cell forming 290hydrogen bonds involving the NH and COOH groups.

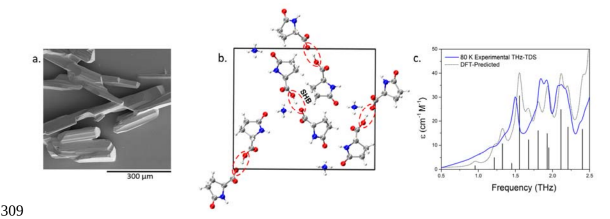
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292L-pyro-amm has a rather unique hydrogen bond network structure since four of the 293pyroglutamine molecules are deprotonated and hence have a nominal negative charge, while 294the other four molecules are neutral. One of the important implications of this difference is 295that L-pyro-amm contains a very strong hydrogen bond. The red circled regions in Fig. 1b 296correspond to a short hydrogen bonds (SHBs) with a length of 2.45 Å, while those in L-glu 297and L-pyro range between ~2.55-2.85 Å, Suppl. Fig. 1.

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299The structural change was further confirmed using THz-TDS measurements, as this technique 300is strongly dependent on the bulk packing arrangement as well as on the internal covalent 301structure of the molecules (42). The THz-TDS spectrum of the resulting solid, as well as the 302solid-state DFT predicted spectrum based on the single crystal XRD (SCXRD)-determined 303structure, is shown in Fig. 1c (full spectral assignment available in Suppl. Fig. 2). The 304agreement between the experimental and theoretical spectra further supports that full 305conversion of the sample occurs and thus enables additional investigations into the structural 306and electronic properties of the material. The agreement is also indicative of the ability of the

307theoretical model to not only reproduce the experimental structure, but also the weak forces 308found in solid structures.



310Figure 1. L-glu forms L-pyro-amm upon heating.

311 (a) SEM image of crystals of L-pyro-amm dried. (b) XRD analysis of heated L-glu sample 312show the newly formed structure, L-pyro-amm. Geometry optimisations show that 8 313pyroglutamine groups and four ammonium ions (144 atoms) are complexed in the crystal and 314a SHB of 2.45 Å (within red dashed lines) is present near the ammonium ion (white-315hydrogen, red-oxygen, blue-nitrogen, grey-carbon). (c) Experimental (blue line) and 316theoretical (grey line) THz-TDS of the L-pyro-amm sample are in agreement and confirm the 317presence of the new L-pyro-amm structure.

319We first investigated whether there were any differences in the optical properties associated 320with the three crystal structures. Comparing the absorption of L-glu, L-pyro and L-pyro-amm 321in water, we show that only L-pyro-amm has a significantly red-shifted absorption which lies 322in the 275-320 nm range, whereas both L-glu and L-pyro primarily absorb in the deep UV 323(<250 nm) (see Fig. 2a).

325We next compared the experimental absorption spectra of L-glu, L-pyro and L-pyro-amm 326with the ones obtained from time dependent density functional theory (TDDFT). We 327highlight here, that the small size of the systems permitted us to determine the spectra using a 328hybrid functional, thereby not only advancing the quality of our theoretical predictions from 329previous studies (5, 40, 41) but also coupling the optical properties directly to different 330vibrational modes.

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332Fig. 2b illustrates the absorption spectra obtained for the TDDFT calculations on the 3 333periodic systems in the ground state (i.e. at 0 °K). Panel b.i) shows the relative oscillator 334strength as a function of the frequency while panel b.ii) illustrates the second derivative of the 335oscillator strength permitting the positions of the maxima in the spectra to be more easily 336identified. The spectra reveal some striking differences between the different systems. 337Interestingly, we observe that L-pyro is essentially dark throughout the frequency range up to 338~ 6eV. On the other hand, L-pyro-amm shows the presence of more structure in the spectrum. 339Specifically, it is the only system for which the spectrum features a low energy excitation at 340226 nm (5.5 eV) and subsequently other peaks slightly above 220 nm (5.625eV) and 216 nm 341(5.75eV). While L-glu exhibits a peak at 222 nm (5.58eV), it is dark up to 206 nm (~6eV).

343We have previously shown that thermal fluctuations and in particular nuclear vibrations, such 344as proton transfer, have a large impact on the absorption spectra of peptide structures as 345compared to the 0 °K behavior (5, 43–45). In Fig. 2c we show that, compared to the 0 °K 346spectra in Fig. 2b, thermal fluctuations cause a large red shift to around 3.4 eV (365 nm) for 347L-pyro-amm, close to what is observed experimentally. These spectra were computed 348averaging over 25 frames sampled from the molecular dynamics simulations. Interestingly, 349no such effect is observed for L-glu which remains weakly absorbing up to more than 5 eV 350(247 nm) as seen at 0 °K.

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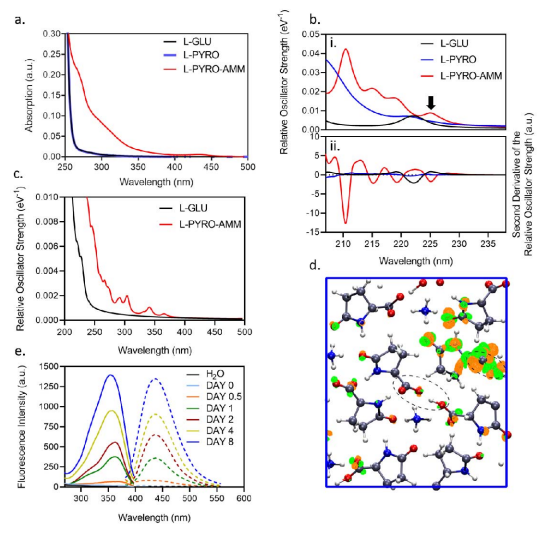
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352In order to understand better the physical origin of the low energy excitation at 226 nm (~5.5 353eV) in L-pyro-amm, we computed the electron response density at this frequency. This is 354illustrated in Fig. 2d, where we observe that most of the electron response is localised in 355regions around the pyroglutamine rings as well as regions near the SHB (see dashed circle in 356Fig. 2d). The optical response thus entails a charge reorganisation involving several parts of 357the molecular crystal. However, since L-pyro contains the same pyroglutamine rings as L-358pyro-amm but absorbs more in the UV-spectrum, we conclude that the structural changes in 359the crystal in the presence of the SHB, which are the main features that distinguish L-pyro 360from L-pyro-amm, are responsible for the large Stokes shift observed in L-pyro 361experimentally and computationally.

363We next investigated whether the above structures also display fluorescence excitation and 364emission properties as has been observed for amyloid-like structures reported previously (5,

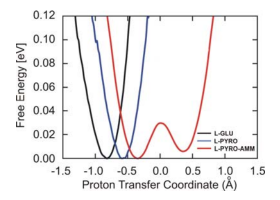
36540, 41). Fig. 2e shows the excitation scan from 250-400 nm (solid lines) with the emission set 366at 430 nm of L-glu in water at day 0 to 8 after incubation at 65°C. We observe an excitation 367peak at around 360 nm which is similar to what we have measured previously for amyloid 368proteins (5). The corresponding emission scan (dashed lines) with the excitation set at 360 369nm and emission from 380-560 nm showed an emission peak around 430 nm, again lying in 370the same visible range as for amyloid fibrils. When the L-pyro-amm solution was dried the 371excitation and emission peaks were slightly blue shifted (Suppl. Fig. 3a) which may be due to 372a change in the molecular environment in the dried state. Importantly, we do not see any 373fluorescence in L-glu (without heating, i.e. at day 0 Fig. 2e.). To determine the importance of 374the ammonium ion experimentally, L-pyro was incubated in water and heated at 65°C for 8 375days, and only a very weak fluorescence has been detected (Suppl. Fig. 3b).



377Figure 2. Optical properties of L-pyro-amm are distinct from L-glu and L-pyro.

378(a) Absorption spectra of 0.3 M L-glu (black), L-pyro (blue) and L-pyro-amm (red) (L-glu 379incubated for 8 days at 65°C) in water taken between 200 – 500 nm shows primarily features 380of L-pyro-amm. (b) Absorption spectra of L-glu, L-pyro and L-pyro-amm obtained from 381 periodic density functional theory calculations with the B3LYP functional. L-pyro-amm 382 features the lowest lying excited states which are characterised by the largest oscillator 383strengths. (c) Absorption spectra for L-glu and L-pyro-amm obtained from periodic 384simulations at room temperature. The spectra were computed by averaging over 25 frames 385randomly sampled from the ab initio molecular dynamics simulations. (d) The excited state 386electron density computed for L-pyro-amm from the optimised structure computed at the first 387peak (arrow in panel b). The lowest excited state density shows a response from various parts 388of the crystal structure including the pyroglutamic acid ring and the SHB region (see dashed 389circle). The orange and green surfaces correspond to regions involving a decrease and 390increase in electron density respectively, shown at an iso-value of 1x10<sup>-5</sup>. (e) 1 M L-glu was 391incubated at 65°C and the excitation and emission spectra were measured over time. 392Excitation spectra were measured between 250-400 nm with emission set at 420 nm and 393emission spectra were measured between 380-560 nm with the excitation set at 360 nm.

395As alluded to earlier, one of the factors that distinguishes L-pyro-amm from the other systems 396is the presence of the SHB (highlighted by red circles in Fig. 1b) and the presence of the 397ammonium ion. In order to characterise the behaviour of the SHB, we conducted *ab initio* 398molecular dynamics simulations of the three systems at 300 °K and examined the proton 399transfer coordinates defined as the difference in distance between the proton (H) and the two 400oxygen atoms (O1 and O2) that sandwich it and is commonly referred to as the proton 401transfer coordinate (d<sub>O1-H</sub>-d<sub>O2-H</sub>) as shown in Fig. 3 for different types of hydrogen bonds in 402the crystals. It is clear that the SHB in L-pyro-amm is characterised by a double-well 403potential. The barrier associated with this proton transfer is on the order of thermal energy, 404indicating that zero-point energy (ZPE) would make the proton transfer barrierless (*46*). An 405examination of similar proton transfer coordinates for hydrogen bonds in L-glu and L-pyro 406show that they are characterised by only single-well potentials.

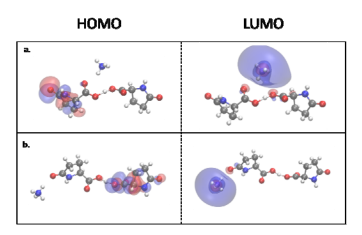


409Figure 3: Free energy profiles along the proton transfer coordinate show only L-pyro-410amm displays a double well potential. Free energy profiles along the proton transfer 411coordinate are displayed for L-glu (black), L-pyro (blue) and L-pyro-amm (red) at room 412temperature. The L-glu and L-pyro display single well potentials, while the L-pyro-amm 413system is the only one the exhibits a double well potential implying that there is proton 414transfer from one side to the other.

416The nature of the optical properties is sensitive to the environment in which the glutamine 417molecules reside. It has previously been reported that charged amino acids already display an 418absorption in the range of 250-350 nm that is significantly red shifted (47, 48). The origins of 419the low energy absorption were attributed to charge transfer excitations. The simulations of 420these systems were performed in the gas phase, rather than considering the protein 421environment such as shown for L-pyro-amm in Fig 2d. In comparison to the results presented 422in Fig. 2d, data presented in Fig. 4, show that the origins of the electronic transitions equally 423 arise from a charge transfer (CT) between the highest occupied molecular orbital (HOMO) on 424the anionic dimer, and the lowest unoccupied molecular orbital (LUMO) centred on the 425ammonium cation when performed in the gas phase. Interestingly, the correct transition 426energy is only predicted when the ammonium cation is spatially near the centre of the dimer, 427which corresponds to the delocalisation of the negative charge and the SHB. Two generalised 428geometries, with the ammonium cation near the SHB (as seen in Fig. 4a) and away from the 429SHB (Fig. 4b), with the corresponding HOMO and LUMO orbitals are shown. The results 430predict a transition of 304 nm for dimer (a) and 669 nm for dimer (b), with dimer (a) most 431 closely resembling the chemical environment present within the crystalline material.

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433The results show that charge transfer is capable to lead to absorption in the near UV when 434investigated in the gas phase, i.e. neglecting the direct protein environment. However, 435including the protein environment in the molecular crystal (Fig. 2d) results instead in the 436excitation being a charge reorganisation involving several different molecular groups of the 437crystal. Indeed, by shuffling the protons along the SHBs in the ground-state, we observe an 438electronic response involving the entire structural units of L-pyro-amm including both the 439hydrogen bonded regions and the pyroglutamic acid rings when the protein environment is 440accounted for (Suppl. Fig. 4).



442Figure 4. Comparison of the HOMO and LUMO orbitals on two L-pyro-amm models.

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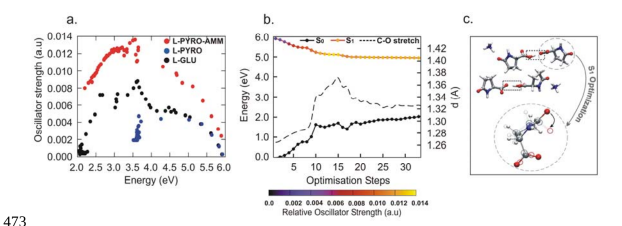
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443L-pyro-amm structures are presented where the ammonium cation is located directly near the 444hydrogen bond (a), and where the ammonium cation is located away from the hydrogen bond 445(b). While both models predicted charge-transfer HOMO-LUMO states, only in the case of 446(a) is the transition predicted to be in the vicinity of the experimentally observed peak, 304 447nm compared to 669 nm for (b).

449Up to this point, we have shown that proton transfer along SHBs is an important part of the 450structural fluctuations in the ground state structure of L-pyro-amm. In order to characterise 451the nuclear relaxation upon photoexcitation, we conducted both excited state optimisations as 452well as excited state molecular dynamics simulations. We first show the results from the 453excited state optimisations obtained from clusters carved out from the different glutamine 454crystals and surrounded by a continuum dielectric constant of 80. Fig. 5a shows a scatter plot 455of the difference between the first excited state and ground-state energies as well as the 456corresponding oscillator strengths for the three systems, L-glu, L-pyro and L-pyro-amm. The 457scatter plots were obtained over the course of the excited state optimisation. We observe that

458the L-pyro-amm system is characterised by the largest oscillator strengths peaking at 459approximately 3.5eV (354 nm), which is consistent with our experimental findings. Although 460L-glu and L-pyro also have a peak at around 3.5 eV it is much weaker than the one of L-pyro-461amm. We thus decided to focus on a series of excited state optimisations for various clusters 462of L-pyro-amm.

464Fig. 5b shows the evolution of the first excited state energy for one of the L-pyro-amm 465clusters (Fig. 5c) across the optimisation steps. We find that the excited state energy drops by 4660.5 eV combined with an enhancement of the oscillator strength while the ground state 467energy, shown in solid-black, rises by about 1.5eV. In this case, the closing of the energy gap 468to approximately 3eV is associated with an increase of the carbonyl oxygen bond as observed 469in previous studies (48–50), as well as a de-planarisation of the pyroglutamine ring (see Fig. 4705c). Similar features are also observed in other clusters including one with the ammonium ion 471closer to the short hydrogen bond (Suppl. Fig. 5).



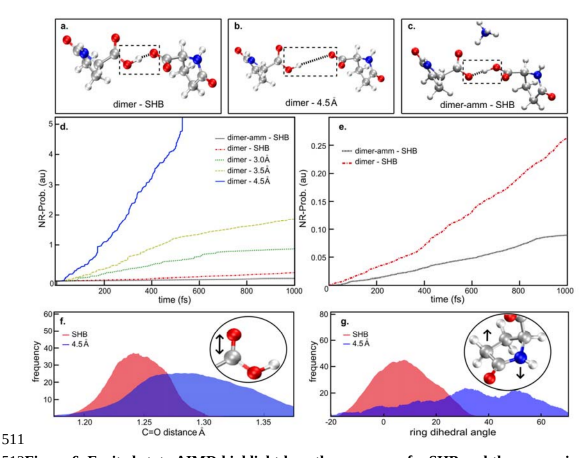
474Figure 5. L-pyro-amm optical properties as seen through excited state optimisations.

475(a) Scatter plot of the oscillator strengths versus the emission energy (defined as the 476difference between the first excited state and the ground-state) during the excited 477optimisations of L-glu (black), L-pyro (blue) and L-pyro-amm (red). (b) Ground and excited 478state energies are plotted as a function of the excited state optimisation of the system shown 479in c. The curves on the excited state are colour-coded with the oscillator strengths. S<sub>0</sub> and S<sub>1</sub> 480refer to the ground and excited state energies, respectively. (c) A snapshot of the optimised 481excited state cluster also showing the lengthening of the carbonyl bond and the 482deplanarisation of the ring.

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484To further explore the preceding mechanisms, we next turn to deploying non-adiabatic decay 485 estimations using excited state ab initio molecular dynamics simulations, as implemented in 486the LIO quantum-chemical package (https://github.com/MALBECC/lio) (30–33). This 487approach enables investigating the S0←S1 de-excitation probability providing a clear 488interpretation of the ensuing optical properties of L-pyro-amm (see Methods for details). The 489 nature of the S0←S1 transition is mainly a  $\mathbb{Z}$ ''=0 ←  $\mathbb{Z}$ '=0 (where  $\mathbb{Z}$ ' and  $\mathbb{Z}$ '' are the 490vibrational quantum numbers in the electronic ground and excited state respectively) with a 491FC factor between ground vibrational levels of ~0.82 (see Suppl. Fig. 6), therefore the excited 492state simulation was initiated by a vertical excitation to the S1 state. We performed two sets 493 of excited state simulations: (i) varying the strength of the SHB to study its influence on the 494fluorescence of L-pyro dimer (Fig.6 a-b), and (ii) comparing L-pyro with L-pyro-amm 495dimers, to assess the role of ammonium ion on the transient excited state dynamics (Fig. 6c). 496Each trajectory was propagated for 1 ps from which the non-radiative decay probability 497(NRP) was determined by the fewest-switches trajectory surface hopping approach (34–36). 498Fig. 6d shows the time evolution of the accumulated NRP for various hydrogen bond lengths. 499The accumulated NRP (ANRP) represents the total probability for  $S_0 \leftarrow S_1$  non-radiative 500relaxation. The larger value of the NRP implies that the system has a higher likelihood of 501decaying non-radiatively to the ground-state. On the other hand, a lower NRP would involve 502a longer excited-state lifetime increasing the fluorescence probability. Fig. 6d demonstrates 503unequivocally that the presence of the short hydrogen bond significantly reduces the chances 504of non-radiative relaxation towards the ground state. Examining the energy gap between the 505excited and ground state for this system shows that it occurs approximately ~3 eV (413 nm) 506(see Suppl. Fig. 7a) consistent with fluorescence in the blue/green visible regime. The 507presence of the ammonium ion further enhances, albeit by a subtle amount, the probability of 508trapping the system in the excited state.

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512Figure 6. Excited state AIMD highlight how the presence of a SHB and the ammonium 513ion is enhancing fluorescence. Excited state AIMD performed for the (a) L-pyro dimer, (b) 514the L-pyro dimer constraining the SHB distance, and (c) the L-pyro-amm. (d) Accumulated 515non-radiative  $S_0 \leftarrow S_1$  decay probability (ANRP) for L-pyro dimer (dimer-SHB) and the 516constrained SHB L-pyro (dimer-) where the SHB distance was fixed at 3.0 Å, 3.5 Å and 4.5 517Å and for the L-pyro-amm dimer (dimer-amm-SHB). (e.) Accumulated non-radiative decay 518probability for L-pyro (red discontinuous line) and L-pyro-amm (black dashed line). (f) 519Carboxyl C=O distance histogram for the L-pyro-amm dimer at the SHB distance (red curve) 520or constraining the SHB bond to 4.5 Å (blue curve). (g) Ring dihedral angle histogram for the 521L-pyro-amm dimer at the SHB distance (red curve) or constraining the SHB bond to 4.5 Å 522(blue curve).

524The excited state optimisations shown in Fig. 5, illustrate the important role played by the 525vibrational distortions in the excited state that on the ensuing optical property. In agreement 526with this, Figs. 6f and g show the distribution of the C=O bond length and the ring de-527planarisation (computed as the sum of its internal dihedral angles) for the dimer systems with

528the SHB and weaker (longer) hydrogen bonds in the left and right panel, respectively. 529Interestingly, we see that the trajectories that undergo non-radiative decay are characterised 530by both, a significant increase in the carbonyl stretch (C=O bond length) by about a tenth of 531an Angstrom, and a large de-planarisation of the dihedral angle. The combined distortion of 532these vibrational coordinates leads to non-radiative decay as observed in the dimer system 533with longer hydrogen bonds. In the system with the short and strong hydrogen bonds, these 534modes are hampered which essentially prevents the system from easily accessing conical 535intersections. This increases the likelihood of observing fluorescence in the L-pyro-amm 536structure characterised by short-hydrogen-bonds. Therefore, both our experiments and 537simulations demonstrate the crucial role played by the SHB and in determining the 538fluorescence of L-pyro-amm.

# 540Conclusions

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542The experimental and theoretical findings presented here, elucidate a rather complex 543molecular mechanism associated with the non-aromatic intrinsic fluorescence in protein-like 544structures. In the case of L-glutamine, a chemical reaction creates a newly formed structure 545involving a cyclised pyroglutamic acid ring. This new structure features absorption in the UV 546and emission in the visible range very similar to the chemically distinct amyloid fibrils (4, 5, 54711, 12, 49–51). The structural chromophore responsible for the optical properties in this new 548protein-related structure arises from a hydrogen bond network associated with structures 549involving short hydrogen bonds. Indeed, we have shown previously that similar to L-pyro-550amm, the crystallised structure of the hydrophobic core of amyloid beta, 2Y3J, L-pyro-amm 551contains a SHB (52) leading to a double well potential in the ground state. The presence of 552strong hydrogen bonds along which proton transfer occurs and secondly, specific ionic 553interactions in close proximity, such as involving the ammonium ion of L-pyro-amm, affect 554the optical properties. Although the fluorescence observed in these systems is much weaker 555compared to conventional fluorophores, the physical and chemical properties of the hydrogen 556bond networks reported here maybe a generic feature across many other peptide structures.

558Our non-adiabatic molecular dynamics simulations demonstrate that the presence of short 559hydrogen bonds with specific environmental conditions hinder vibrational deformations that 560can access conical intersections, as was previously speculated (5). The role of carbonyl 561groups is also consistent with a recent experimental study showing their importance for non-

562aromatic fluorescence (53). SHBs have recently been observed in different biological systems 563which have long been associated with either intrinsic fluorescence, such as NADP/NAD (54), 564FAD/FMN (55), the light-sensing chromophore in photoactive yellow protein (56), or in the 565active site of many enzymes, such as hydrolases and oxidoreductases (57, 58), many of which 566consist of a highly complex H-bond structures similar to amyloid proteins. Furthermore, it 567has been recently reported that 1 in every 16 hydrogen bonds in over 1600 proteins are 568characterised by short hydrogen bonds (58). Thus, the mechanisms we espouse here may be 569more general. Our findings offer the possibility of designing novel biomaterial for 570applications in optical sensing or the design of novel biocompatible catalysts.

## **572Supporting Information**

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573Supporting data include; Supplementary Figure 1. Clusters of L-glu and L-pyro. 574Supplementary Figure 2. Full spectral assignment of THz data. Supplementary Figure 3. L-575pyro-amm has blue shifted fluorescence when dried and displays higher fluorescence 576intensity than L-pyro. Supplementary Figure 4. The optical properties of L-pyro-amm are 577sensitive to the environment and involves the electronic response of the entire structure. 578Supplementary Figure 5. L-pyro-amm undergoes vibrational distortions upon excitation as 579seen through excited state optimisations. Supplementary Figure 6. Potential energy surface 580along the SHB proton transfer coordinate in the ground (S0, panels A and D) and the excited 581(S1, panels B and C) states. Supplementary Figure 7. The energy gap between excited and 582ground state of L-pyro-amm is consistent with the experiments and eventually deactivates via 583a conical intersection back to the ground state. SCXRD data are available at the Cambridge 584Crystallographic Data Centre, CCDC No. 1981551. Raw data are available at the University 585of Cambridge Repository https://doi.org/10.17863/CAM.57945.

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# **592Author Contributions**

593G.S.K.S and A.H conceptualised the manuscript. \*A.D.S and M.N.Q contributed equally. 594A.D.S prepared samples for all experimental data. P.J.W. performed SEM experiments. 595A.D.B and M.T.R performed XRD measurements. M.T.R. performed THz experiments and

596DFT-THz calculations. A.D.S. performed excitation and emission measurements. A.D.S and 597S.T.J. performed absorption measurements. M.T.R and E.M.K performed TD-DFT cluster 598calculations. M.N.Q, E.P, L.G, R.G and A.H performed AIMD, Periodic TD-DFT Excited 599State Calculations and Periodic Structure Geometry Optimisation calculations. G.D.M, 600U.N.M, and M.C.GL performed excited state molecular dynamics. A.D.S, M.N.Q, M.T.R, 601S.T.J, L.G, G.D.M, U.N.M, M.C.G.L, J.A.Z, D.C, A.H and G.S.K.S contributed to 602manuscript writing. All authors have given approval to the final version of the manuscript.

## 604Notes

605The authors declare no competing financial interest.

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# 619**Abbreviations**

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620SEM, scanning electron microscopy; XRD, X-ray diffraction; THz-TDS, terahertz time 621domain spectroscopy; DFT, density functional theory; TD-DFT, time dependent DFT; SHB, 622short hydrogen bond; CT, charge transfer; L-glu, L-glutamine; L-pyro, L-pyroglutamine; L-623pyro-amm, L-pyroglutamine complexed with an ammonium ion.

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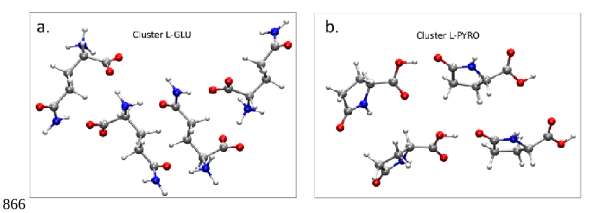
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836Supplementary Information for 838Short hydrogen bonds enhance non-aromatic protein-related fluorescence 840Amberley D. Stephens, Muhammad Nawaz Qaisrani, Michael T. Ruggiero, Gonzalo Diaz Miron, Uriel 841N. Morzan, Mariano C. González Lebrero, Saul T.E. Jones, Emiliano Poli, Andrew D. Bond, Philippa 842J. Woodhams, Elyse M. Kleist, Luca Grisanti, Ralph Gebauer, J. Axel Zeitler, Dan Credgington, Ali 843Hassanali, Gabriele S. Kaminski Schierle 845Ali Hassanali, Gabriele S. Kaminski Schierle 846Email: ahassana@ictp.it, gsk20@cam.ac.uk 



# 867Supplementary Figure 1. Clusters of L-glu and L-pyro

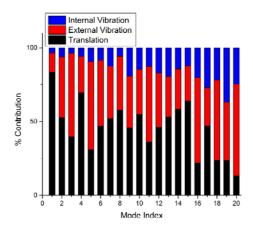
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868Structural representations of (a) L-glutamine (L-glu) and (b) L-pyroglutamine (L-pyro) 869clusters from the published crystal structure.



Mode Number	Frequency (THz)	Intensity (km raok	-1) Assignment
1	0.955	0.22	Antisymmetric transletion in b
2	1.215	0.71	Coupled translation in a and rotation about be
3	1.325	1.52	external rotation about b
4	1.450	0.36	Asymmetric translation in c
5	1.554	4.37	External rotation about a
5	1.676	1.76	Translation in a and external asymmetric relation about H-band coordinate
7	1.805	2.28	External esymmetric rotation of dimer pairs
8	1.929	2.11	External symmetric rotation of dimer pairs
9	1.950	1.28	Asymmetric rotation about h-bond coordinate, ommontum translational motion
10	2.113	3.53	Translation and retation (breathing) ground ammonium cation
11	2.208	2.5	Symmetric external rotation of entire formula units
12	2.404	2.87	In phase external notation perpendicular to h-band coordinate
13	2.502	0.45	Out of phase external rotatation perpendicular to h-bond coordinate
14	2.564	0.73	External out of phase rotatation of h-bonded chains about b
15	2.565	8.91	Out of phase external rotation of individual pyroglutarnic molecules with translation of ammonium cation
16	2.654	18.09	External vibration coupled with tension of the COOH group
17	2.579	0.29	External rotation and torision of pyrogiutamic ring
15	2.776	4.94	External rotation and torsion of ring and carboxyl group
19	2.988	2.68	External rotataion and translation of ammonium

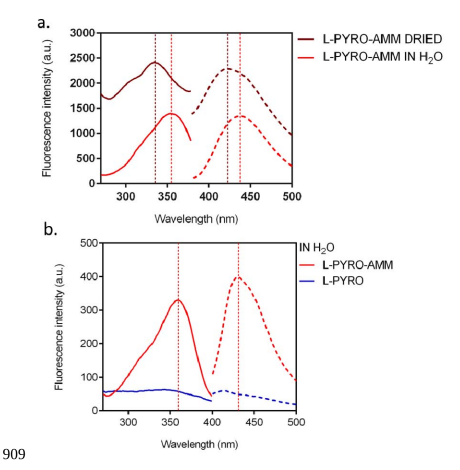
# 894Supplementary Figure 2. Full spectral assignment of THz data

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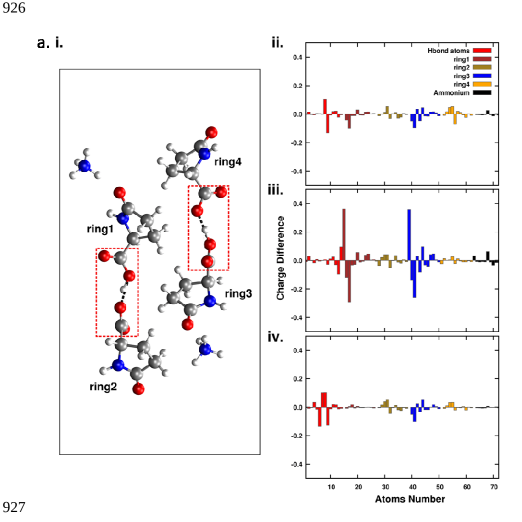
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895The top chart shows the contribution to each IR-active mode including external translations 896and hindered rotations, and internal vibrational motions (i.e. torsions), while the bottom table 897lists the detailed assignment for each mode.



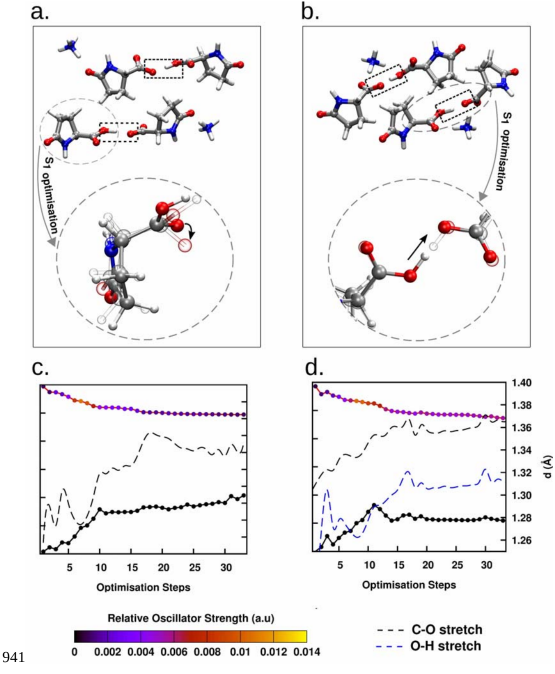
# 910Supplementary Figure 3. L-pyro-amm has blue shifted fluorescence when dried and 911displays higher fluorescence intensity than L-pyro

912(a)The excitation peak of L-pyro-amm when dried (solid dark red line) is blue shifted with a 913peak maximum  $\sim$  340 nm compared to L-pyro-amm in H<sub>2</sub>O (solid red line) which has a peak 914maximum  $\sim$  360 nm. The emission peak of L-pyro-amm when dried (dashed dark red line) is 915also blue shifted, with a peak maximum  $\sim$  420 nm, compared to L-pyro-amm in H<sub>2</sub>O (dashed 916red line) with a peak maximum  $\sim$  430 nm. (b) 1 M L-glu and 1 M L-pyro (blue) were 917incubated in H<sub>2</sub>O for 8 days at 65°C. After 9 days, the L-glu had converted into the L-pyro-918amm structure (red). L-pyro-amm has a clear excitation peak maximum at  $\sim$ 360 nm and 919emission peak maximum at  $\sim$ 430 nm, while L-pyro (blue), although not completely dark, has 920no clear excitation or emission peak.



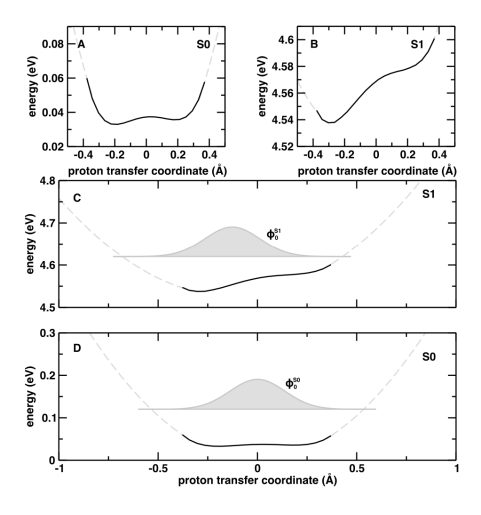
928Supplementary Figure 4. The optical properties of L-pyro-amm are sensitive to the 929environment and involves the electronic response of the entire structure

930(a)(i) L-pyro-amm cluster used to examine the sensitivity of the optical response on different 931parts of the cluster upon moving different protons. (ii) Charge differences between the ground 932and excited state are computed using restrained electrostatic potential atomic partial charges 933(RESP) for the cluster shown in i). Note, the electronic response involves all the atoms of the 934cluster. (iii) The two protons in the rectangle regions are displaced to be in the centre of the 935hydrogen bond and the charge differences are then computed. As illustrated, the proton 936displacement leads to a larger change in magnitude of the charges. (iv) The charge 937differences are computed for another nuclear configuration for which the protons are kept 938fixed but the O—O distance is increased from 2.45 to 3.2 Angstroms. The charge differences 939obtained here are quite similar to the original condition shown in (ii).



942Supplementary Figure 5. L-pyro-amm undergoes vibrational distortions upon 943excitation as seen through excited state optimisations

944(a-b) correspond to two additional clusters to the one reported in the main text that were built 945from the L-pyro-mm crystal structure. c) and (d) correspond to the evolution of the excited 946state energy and ground-state energy over the course of the optimisation. Also shown, are the 947changes in the C-O bond lengths (dashed black) which increases on the excited state surface. 948In (d) we show the O-H bond stretch where a proton transfer is observed on the excited state. 949

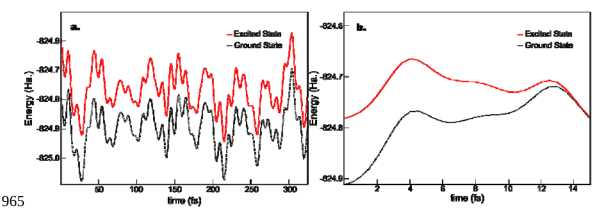


952Supplementary Figure 6. The S0 and S1 nuclear wavefunctions associated to the SHB 953proton transfer coordinate reveal a vibronic transition dominated by a  $0 \leftarrow 0$  character.

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954The calculated potential energy surfaces (black continuous lines) are prolongated with a 955quadratic fitting (gray dashed lines). Panels C and D illustrate the ground-state vibrational 956wavefunction in the S0 and the S1 electronic states ( and respectively). The resulting 957Franck-Condon factor between and is 0.82, showing that the nature of the S1  $\leftarrow$  S0 958transition is mainly a  $\square$ ''=0  $\leftarrow$   $\square$ '=0 (where  $\square$ ' and  $\square$ '' are the vibrational quantum numbers 959in the electronic ground and excited state respectively).



966Supplementary Figure 7. The energy gap between excited and ground state of L-pyro-967amm is consistent with the experiments and eventually deactivates via a conical 968intersection back to the ground state

969(a) Shows the energy of the excited and ground state obtained from the excited state 970molecular dynamics simulations where non-adiabatic decay probabilities are estimated. We 971see clearly that the energy difference between the excited and ground state is approximately 3 972eV (413 nm) consistent with experimental data. (b) Shows the same excited state and ground 973state energy for another simulation where the system de-activates very quickly within 10s of 974femtoseconds through a conical intersection where the ground and excited state energies 975intersect.