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FiberForge: enabling high-throughput simulations of the mechanical properties of helical fibrils

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The mechanical properties of amyloid-based materials are governed by fibril geometry, sequence, and polymorphism, yet systematic exploration of this vast design space has been limited by the lack of high-throughput modeling tools. Here we present FiberForge, an open-source workflow that automates construction of amyloid protofibrils, streamlines high-throughput simulations of amyloid deformation, and analyzes fibril trajectories to estimate mechanical properties and fracture mechanisms. Using 374 full-length amyloid crystal structures from the Protein Data Bank, FiberForge rebuilds fibrils with a mean per-chain RMSD of 1.7 Å (median 2.2 Å), demonstrating accurate structural recovery across wide sequence (18–420 aa) and symmetry ranges. Extensive SMD benchmarking on Aβ(1–42) (2MXU) yields a mean rupture force of 1.534 ± 0.164 nN from 232 replicas; bootstrapping analysis shows that three replicas suffice for converged elastic-modulus and strength estimates. High-throughput screening of the amyloid fiber dataset produces elastic moduli of 0.2–20 GPa and ultimate tensile strengths of 0.1–1 GPa. Comparison with four AFM-characterized systems shows agreement within an order of magnitude, underscoring the method's predictive capability. FiberForge's screening results also enable larger-scale sequence–structure–property analysis, revealing that mechanical behavior is correlated with molecular assembly geometry, especially hydrogen-bond density. While earlier work suggested the relevance of these features for particular systems, our results demonstrate their importance across diverse fibril architectures. FiberForge thus provides an end-to-end platform for molecular modeling and design of amyloid materials, enabling physics-based identification of sequences and polymorphs with targeted mechanical behavior.

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1 Introduction

The mechanical properties of amyloid fibers are remarkable.¹ They exhibit mechanical properties comparable to other protein materials such as microtubules,² actin filaments,³ and even spider silk.⁴ They possess substantial tensile strength, rivaling that of steel, due to hydrogen bonding between β-sheet structures along their length. Amyloid fibers are stiff with a tensile modulus ranging from 2 to 20 GPa.⁵ In addition to their impressive mechanical characteristics, amyloid fibers are typically more biocompatible due to their biosynthetic nature; this makes them particularly attractive for biomedical applications. Due to these extraordinary material properties, amyloid fibers have been the focus of numerous biomedical engineering

efforts, including hydrogels⁶ for tissue engineering, carbon capture⁷ materials, drug delivery,⁸ and so on. When these fibers form larger networks, such as gels or plaques, the properties of the network emerge from the collective behavior of the individual fibers. For this reason, understanding the behavior of individual fibrils is of fundamental importance to understanding the properties of the collective material. Mechanical properties of amyloid fibrils, such as stiffness and elasticity, can be characterized experimentally using atomic force microscopy (AFM). Computationally, steered molecular dynamics (SMD) offers atomic-level insight into deformation mechanisms by simulating the effects of applied forces. Together, these methods reveal the molecular interactions that govern amyloid stability and structural organization.

Despite the successful application of amyloids, the design of amyloid materials still faces several key challenges: the selection of the optimal fiber from the vast space of possible amyloid-forming proteins, prediction of the critical fiber length for the change of the fracture mechanism, differing properties exhibited by structural polymorphs, and lack of a curated dataset of amyloid structures relevant for mechanics investigation. On the issue of sequence selection, prior studies have shown that many proteins, even those not conventionally

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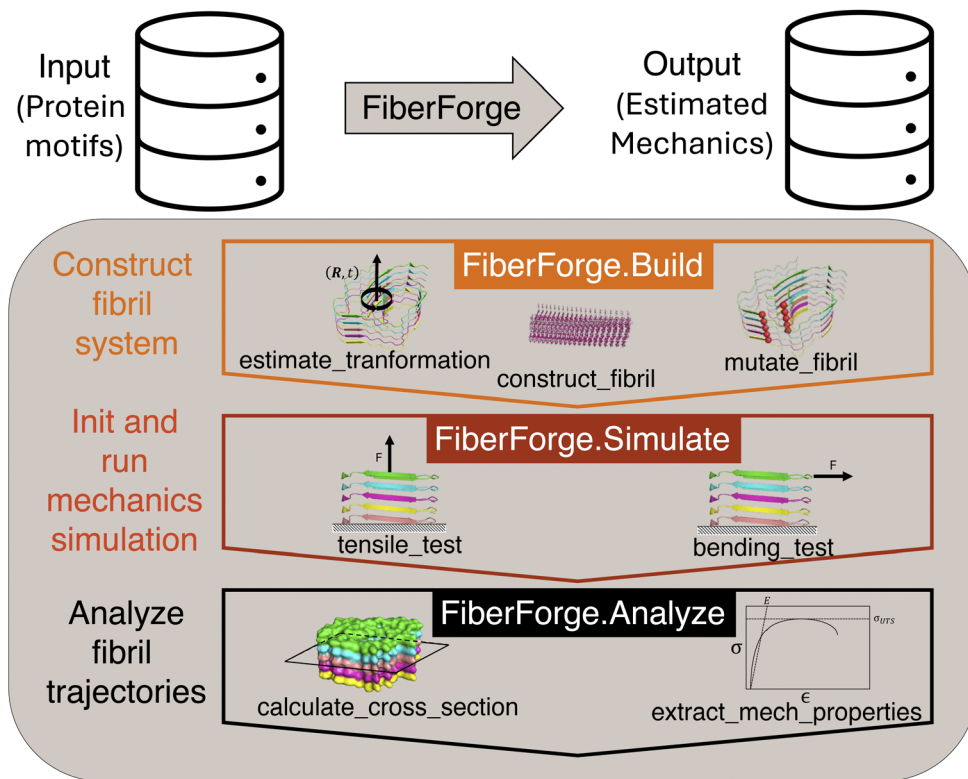


Fig. 1 Roadmap of FiberForge, an automated software suite for amyloid structural construction and mechanical characterization. FiberForge comprises three modules: FiberForge.Build, which supports geometrical characterization, fibril assembly construction, mutagenesis, and solvation; FiberForge.Simulate, which conducts tensile and bending tests using tailored SMD protocols; and FiberForge.Analyze, which processes simulation outputs to estimate mechanical properties like elastic and bending moduli. These modules enable the construction of an end-to-end workflow for simulating fracture mechanics in amyloid structures.

associated with amyloids, are capable of converting to amyloid fibers at high concentration under destabilizing conditions.⁹ The richness of the sequence space of possible amyloids necessitates a high-throughput (HTP), automated fiber construction, and simulation framework. In terms of structural polymorphisms, there can be a huge difference in properties between aggregate polymorphic structures for the same protein sequence.¹⁰ The sensitivity of mechanical properties to the underlying polymorphic structure highlights the importance of systematically sampling and characterizing different structural polymorphs. On the issue of fracture mechanism, insights from SMD modeling have revealed evidence of length-scale-dependent fracture mechanisms (stick-slip) that arise in both amyloids¹¹ and silk crystals.¹² As suggested by these studies, length-scale dependent mechanisms may be attributed to the fact that the fracture mechanism of a β -sheet-rich fibril is governed by the cooperative rupture of hydrogen bonds that stabilize the fibril structure. To investigate further cooperative rupture mechanisms arising in these materials, accurate construction of variable length amyloid structures is needed. Finally, we need a database that captures amyloid structural variations. Several amyloid databases have been curated, including AmyPro, AmyloGraph, and StAmP-DB.^{13–16} Although these databases are extremely helpful for the community interested in the pathological aspect of amyloids, they do not

capture sufficient structural differences that are relevant for the design of amyloids for their mechanical applications.

One methodology to navigate the multi-factor design landscape of amyloids is high-throughput computational physics-guided protein design.¹⁷ High-throughput computational design of amyloid materials requires a framework for the autonomous building of fibril structures, the initialization of simulation systems, and the analysis of simulation results. Unfortunately, prior software focused on amyloid properties covers a broad range of applications not specifically focused on this challenge; these software include experimental image analysis of fibril microscopy,¹⁸ *de novo* aggregate structure prediction,^{19,20} β -serpentine fibril structure prediction,²¹ and numerous examples of amylogenic region prediction.^{22–25}

In this work, we seek to address the issues mentioned above by creating a software, FiberForge, that automates the construction of amyloid protofibrils, streamlines the deployment of high-throughput simulations of amyloid deformation, and automatically analyzes the trajectory of amyloid fibril deformation to estimate mechanical properties and fracture mechanisms. FiberForge enables a deeper understanding of the structure–function relationships that dictate the mechanical properties of amyloid-based materials, providing critical insights for fundamental materials science. Moreover, it facilitates the *in silico* design and screening of amyloid mutants with



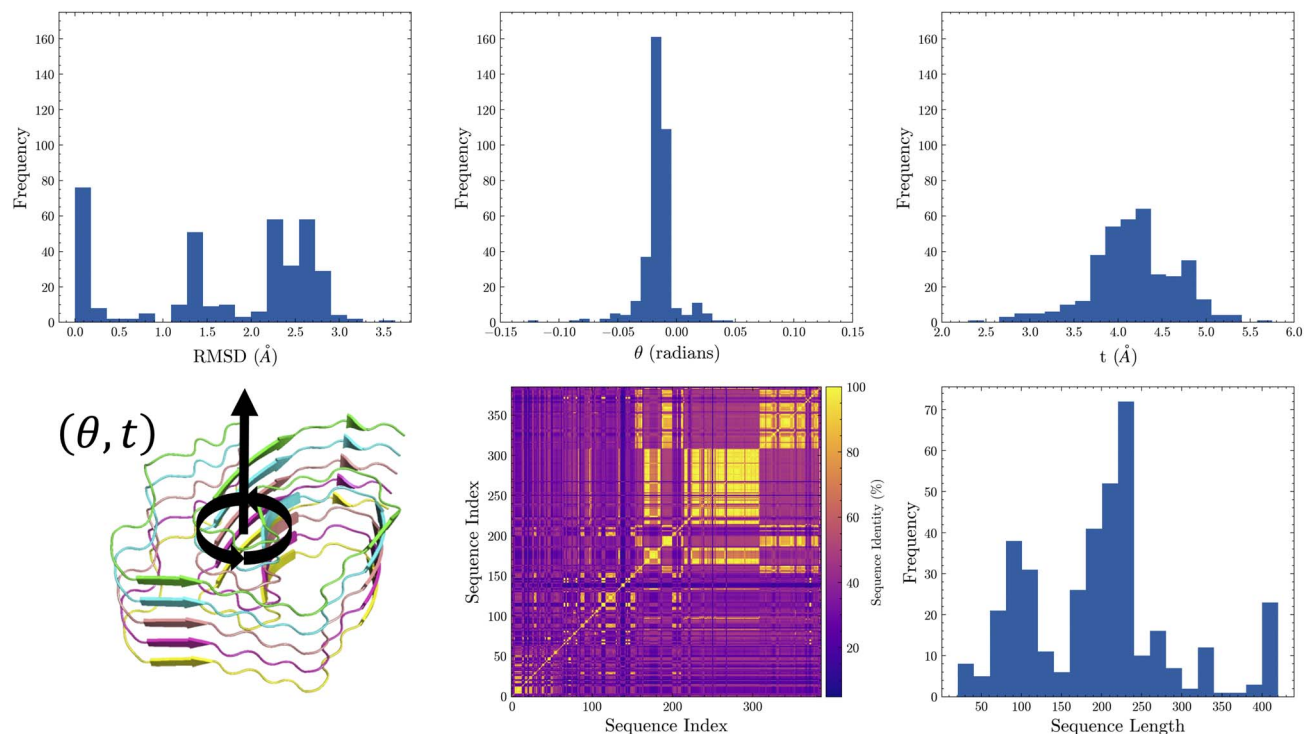


Fig. 2 Statistical distribution and structural features of the protein dataset. (Top left) Distribution of root mean square deviation (RMSD) values, showing a multimodal distribution indicative of structural variability in the dataset. (Top middle) Distribution of helical twist angles θ , centered near zero, suggesting minimal angular displacement between repeating units in most structures. (Top right) Distribution of helical rise values t . (Bottom left) Schematic illustration of the helical parameters (θ, t) , depicting the angular and translational symmetry along the fibril axis. (Bottom middle) Pairwise sequence identity matrix showing the percent identity between all pairs of protein sequences in the dataset. Diagonal and block patterns indicate groups of similar sequences. (Bottom right) Distribution of sequence lengths, indicating the dataset spans a diverse range of protein sizes.

tailored mechanical behaviors, accelerating the development of bioinspired materials for applications in nanotechnology, biomedical devices, and responsive materials.

1.1 Design and implementation

To address our goal of creating software to enable HTP computational design of amyloids, we need a data-structure that is as condensed as possible, but would still accurately describe the 3D geometry of amyloid protein assemblies. We created this data-structure by utilizing the underlying symmetry of amyloid structure: helical symmetry. This representation can describe amyloid fibrils using a 3-tuple: the growth axis vector, translation scalar, and rotation scalar. With this data structure, we were able to construct amyloid fibrils and molecular systems with high fidelity (See bottom left of Fig. 2 for depiction of data structure).

After designing our foundational data-structure for the description of amyloids, we built accompanying modules and functions to utilize this structure to perform mechanical property modeling tasks. Specifically, we designed our software suite to have 3 modules: Build, Simulate, and Analyze (See Fig. 1 for depiction of FiberForge). The Build module consists of functions that enable the geometrical characterization of structures containing helical symmetry, the construction of assemblies based on their helical geometrical parameters and sequence mutation information, and finally the construction of a fully

solvated amyloid fiber for simulation (see sample code in SI, Fig. S3). In our software, we are able to construct amyloids of different lengths by first learning the symmetry functions from multi-chain fibrils and then applying these symmetries to obtain the desired length fibril. The Simulate module consists of a tensile testing function and a bending testing function. These mechanics testing functions create an SMD protocol to conduct the application of an external force which depends on the testing conditions specified (see sample code in SI, Fig. S4). The Analyze module reads the output trajectories produced from the Simulate module and calculates mechanical properties of interest, such as elastic/bending modulus (see sample code in SI, Fig. S5). Together, these modules enable the construction of end-to-end workflows for fracture mechanics simulations of diverse amyloid structures.

2 Results and discussion

2.1 Amyloid dataset

As a prerequisite for building and testing FiberForge, we collected and curated a dataset of experimental amyloid fiber structures. We searched the PDB for entries associated with the keyword “fiber” (performed on February 4th 2024) and then selected all entries that were full sequence fibers, according to our visual inspection of the 3D structure. The compiled dataset consists of 374 structures, with sequence lengths ranging from



A β (42), PDB: 2MXU

DAEFRHDSGYEVHHQKLVFFAEDVGSNKGAIIGLMVGGVVIA

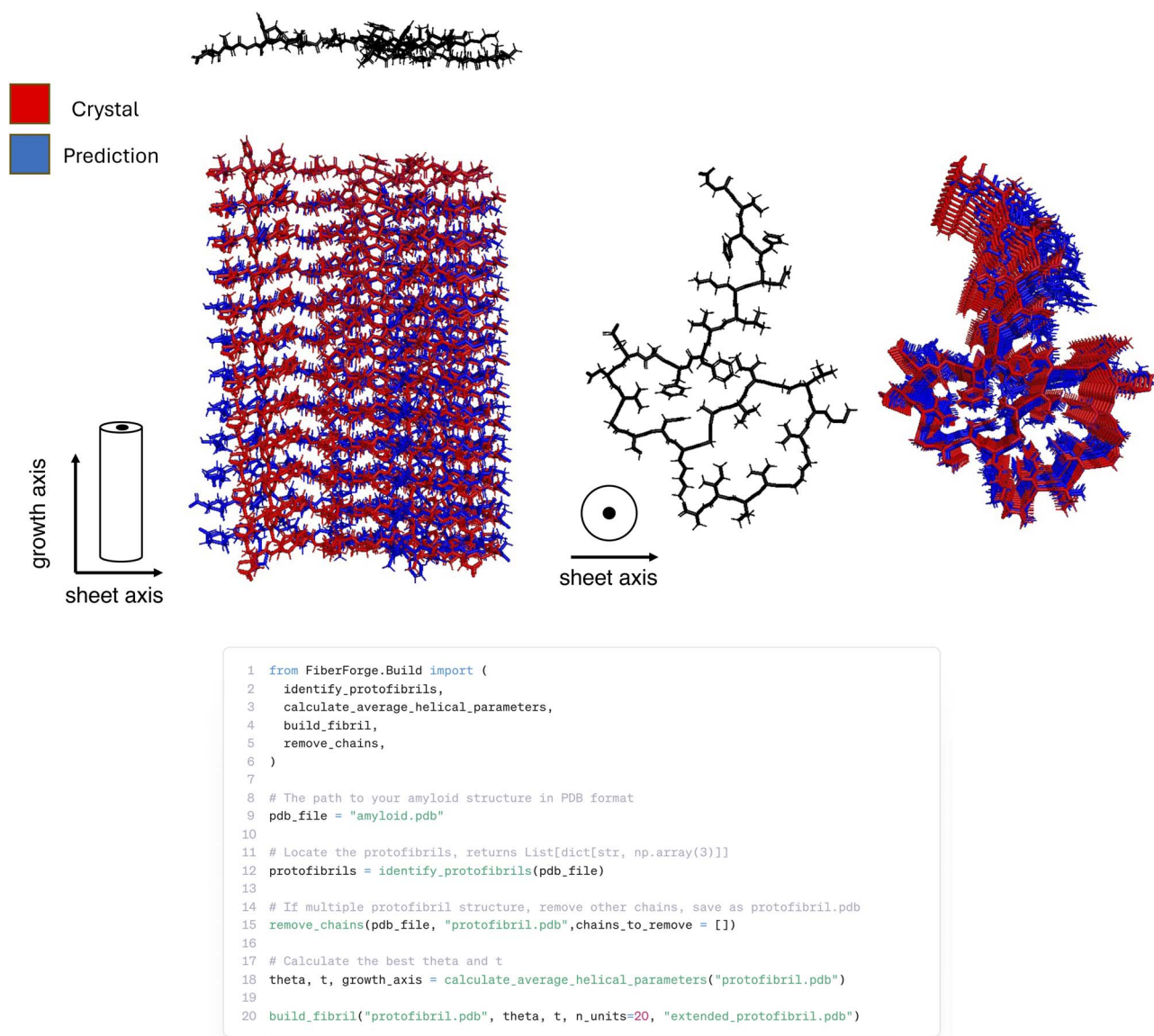


Fig. 3 Demonstration of construction results and accompanying code for creating amyloids of variables length. (Top) Benchmarking of amyloid crystal reconstruction for the A β (42) protein sequence. Black structures represent seed protein used for construction, blue structure represents predicted amyloid construction, and red represents experimental crystal structure. (Bottom) Code for estimating the helical symmetry parameters of the amyloid and the subsequent code for construction of the protein fibril.

a minimum of 18 amino acids (aa) to a maximum of 420 aa. The mean sequence length is 197 aa, with a median of 207 aa and a standard deviation of 88 aa. The resolution of the structures has a mean of 3.1 Å, a median of 3.1 Å, and a standard deviation of 0.5 Å.

2.2 Benchmarking of reconstruction from helical parameters

Accurate fibril construction affects the reliability of estimated mechanics from simulation. To validate our construction method, we applied our method to each structure in our curated

dataset, rebuilding each structure using only a seed chain and the helical parameters. After fibril construction, we calculated the geometric deviation of our reconstructed fibril structures from their respective experimental structures (See Fig. 3, FiberForge API for this process, and summary reconstruction histogram for our curated dataset). Reconstruction statistics are as follows: mean RMSD per chain: 1.7 Å, median RMSD per chain 2.2 Å, standard deviation of RMSD per chain 1.4 Å. As such, the structural models produced by FiberForge.Build shows high consistency with experimental structures for amyloids of various lengths and across a wide range of



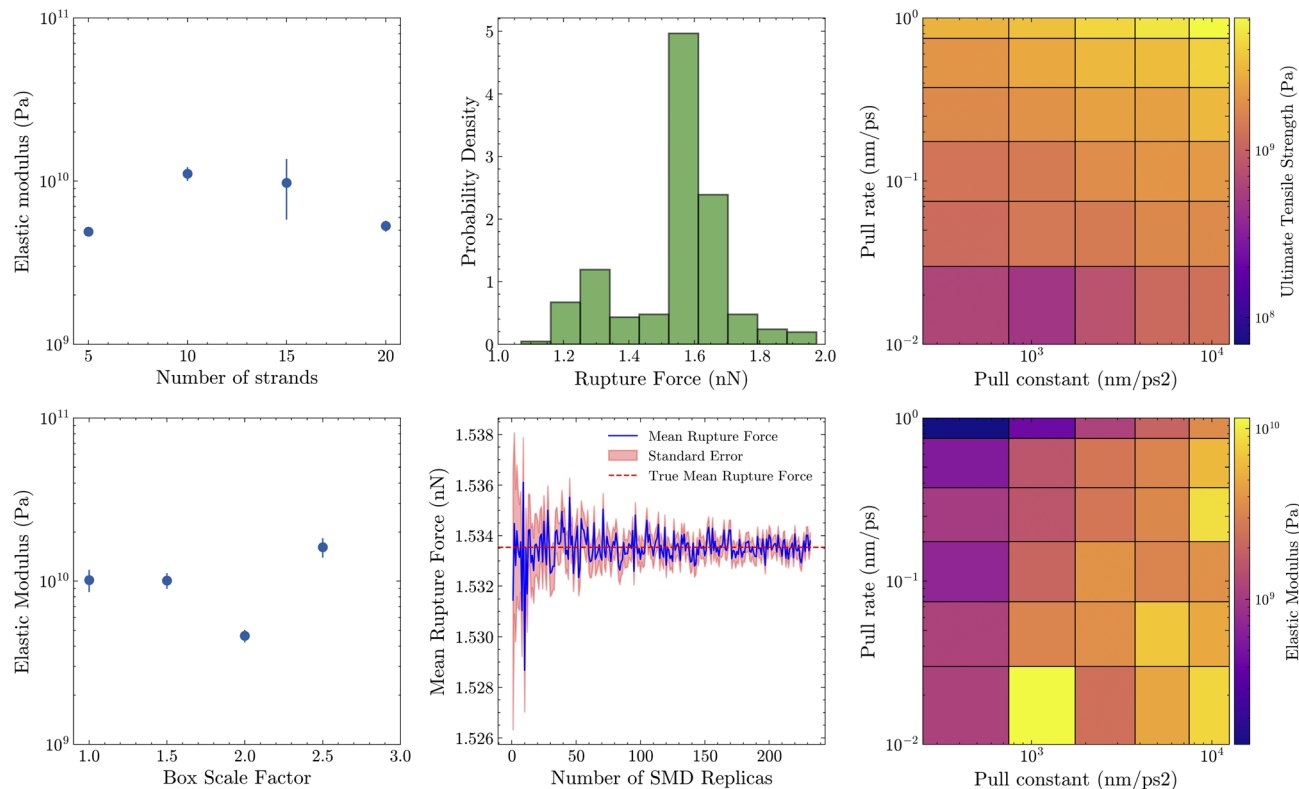


Fig. 4 Results of benchmarking simulation parameters on elastic modulus, rupture force, and ultimate tensile strength. SMD simulations were performed over 5 ns with 2 fs timesteps. (Top left) Elastic modulus (Pa) as a function of the number of β -strands in the amyloid fibril, showing how fibril size influences stiffness. (Top middle) Probability density distribution of rupture forces (nN) obtained from 232 steered molecular dynamics (SMD) replicas of the 2MXU system, with a peak near 1.600 nN. (Top right) Heatmap of ultimate tensile strength (Pa) as a function of pull constant (nm ps^{-2}) and pull rate (nm ps^{-1}), demonstrating how simulation parameters affect tensile strength estimates for the 2MXU system. (Bottom left) Elastic modulus (Pa) as a function of the box scale factor (i.e., number of water molecules), revealing the influence of simulation box size on measured stiffness. (Bottom middle) Convergence of the mean rupture force (nN) with the number of SMD replicas for the 2MXU system using bootstrapping. The blue line shows the running average, the red shaded area represents the standard error, and the dashed red line indicates the true mean rupture force. (Bottom right) Heatmap of elastic modulus (Pa) as a function of pull constant and pull rate, highlighting how mechanical stiffness estimates depend on SMD pulling parameters.

geometric constructs. A representative example of this group is a hIAPP polymorph (7YL7) which has a RMSD of 0.07 Å (see Fig. S1 for a depiction of this fibril). This is believed to be attributed to intra-chain rotation occurring between stacking of successive chains in an amyloid, particularly for residues occurring along the outer region, which has not been accounted for in our current building model of the amyloid symmetry. For some fibril structures our method deviated slightly from experimental structures. The major outlier is 6NZN, which we do not show in Fig. 2. This amyloid has an anti-parallel stacking occurring between chains in a fibril. The current version of our helical symmetry representation does not account for this behavior, but could be extended to handle this type of structure in the next version. Excluding the outlier, the largest RMSD per chain is 3.6 Å with the structure 6RTB (See Fig. S2 for depiction of reconstruction).

2.3 Benchmarking of steered molecular dynamics

Essential to validating the reliability of mechanical property prediction is the expected variability of values predicted from SMD. Previous SMD studies of amyloid fracture involved very

few replicates, as little as a single SMD simulation. This is likely because the primary focus was to qualitatively investigate fracture mechanism rather than predicting specific mechanical properties.^{11,26} Here, we assessed how much rupture-force modeling results fluctuate across SMD simulations to judge their accuracy and reproducibility (Fig. 4). Specifically, we carried out a case study on the A β (42) amyloid (PDB: 2MXU), the same structure used in our construction benchmark, to quantify this inherent variability. We took the same equilibrated system and the same pulling process under different random seeds (a total of 232 replicates). The results show that the mean rupture force is 1.530 nN, the standard deviation of the rupture force is 0.164 nN, and the standard error of the rupture force is 0.0108 nN. Across the entire replica dataset, the mean value fluctuates between 1.526 nN and 1.538 nN—a total spread of only 0.012 nN (12.0 pN), or one percent of the mean. This means that, even though the curve looks “noisy”, the absolute variation is small. As such, a handful of trajectories already gives a reasonable estimate: after just 1–3 replicas the mean typically sits within a few $\times 10^{-3}$ nN of the large-sampling-averaged mean (red dashed line). Thus, for qualitative comparisons or screening studies, running only a couple of replicas is usually



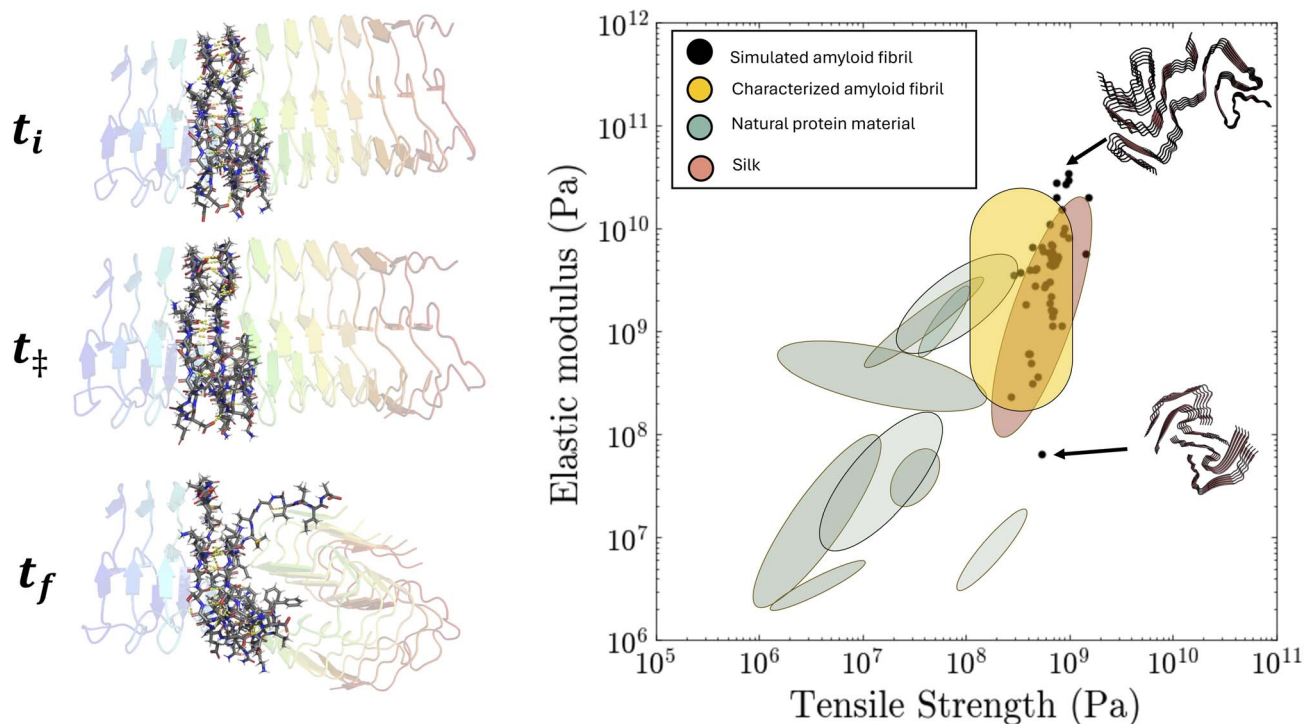


Fig. 5 Summary of virtual screening study of the mechanical properties of our curated dataset of amyloids. (Left) A visual of a tensile test of a $A\beta(42)$ protofibril (2MXU). Initial state of the fibril, transition state, and final state, from top to bottom respectively, can be visualized. (Right) Ashby plot of our screening study, inlaid black points are our results, where each point represents a unique protofibril structure.

sufficient. For applications that demand quantitative convergence to the second decimal place (± 0.010 nN, *i.e.*, ± 5 pN), the shaded standard-error band shows that roughly 30–40 independent replicas are needed. Beyond this point the residual uncertainty falls below 0.005 nN, ensuring the mean is stable to 0.010 nN precision.

Next, we quantified how simulation settings affect the mechanical constants extracted from SMD pulling (Fig. 4). Raising the spring constant from $5 \times 10^2 \rightarrow 1 \times 10^4$ nm ps $^{-2}$ and the pulling rate from 0.01 \rightarrow 1 nm ps $^{-1}$ drives a clear, monotonic stiffening: the ultimate tensile strength climbs from around 5.3×10^9 Pa to 1.1×10^{10} Pa and the elastic modulus from around 3.0×10^9 Pa to 9.5×10^9 Pa, matching the behavior suggested by prior theory of SMD.²⁷ In contrast, box size and number of strands can also perturb the calculated elastic modulus, but no significant change was observed on the scale of the estimated elastic modulus.

Lastly, we compared our estimated mechanical properties to those obtained from experimental methods (see Fig. 5 for results across the entire curated dataset). Our simulated values fall within the expected range for amyloid fibrils (roughly 10^9 to 10^{11} Pascals for elastic modulus and 10^8 to 10^9 Pascals for tensile strength).⁵ Furthermore, we compared our estimated properties with a single experimental study (see Table 1). Although our results showed reasonable agreement with experimental values, for example, for the insulin amyloid with PDB ID 8SBD, we obtained 4.7 GPa for the simulated elastic modulus compared to the reported experimental value of 3.2 GPa, our simulated estimates were consistently higher. This

discrepancy is expected: theoretical predictions and atomistic simulations typically produce substantially larger elastic moduli than experiments because the deformation rates used in SMD are several orders of magnitude higher than those employed in AFM and related experimental techniques.²⁸ Additionally, experimental measurements exhibit substantial variability across studies due to differences in fibril preparation, mechanical testing protocols, deformation rates, and solution conditions. For these reasons, and to mitigate the variability inherent in experimental datasets, comparison with experiment was limited to a single group. To test the generality of this approach, future work will focus on expanding comparisons across multiple experimental systems, despite the challenge posed by the relatively small number (on the order of 10) of available single-fibril mechanical measurements and their substantial inter-study heterogeneity.

Table 1 Comparison of individual experimentally determined elastic modulus³⁴ and SMD determined (simulated) elastic modulus. In experiments, the protein solution was diluted to 0.1 wt% in Milli-Q water at pH 2. In simulations, amyloid fibrils were placed in explicit water, and counterions were added to maintain overall charge neutrality

Common name	PDB entry	$E_{\text{exp}} \pm \text{SE}$	$E_{\text{sim}} \pm \text{SE}$
Insulin	8SBD	3.2 ± 0.6	4.7 ± 0.2
τ -protein	7YPG	3.4 ± 0.7	11.3 ± 3
β -lactoglobulin	6GK3	3.7 ± 0.8	8.6 ± 1
$A\beta(1-42)$	2MXU	3.2 ± 0.8	4.8 ± 0.9



In addition to the benchmarks, we should note the intrinsic stochasticity underlying the rupture force measurement from single-molecule AFM experiments. Early physical theories were developed to describe rupture force distributions in single hydrogen bonded systems²⁹ (Bell–Evans model), these theories were later extended to protein unfolding, *i.e.* multi-hydrogen bonded system, then eventually the connection between AFM and SMD techniques.^{27,28} These theories were applied to describe amyloid fibril rupture.¹¹ Essentially, amyloids are held together by a network of hydrogen bonds occurring between proteins which result in induced force being distributed across a network of bonds. The rupture of this network is a non-equilibrium, stochastic process; as such, the outcome of SMD mechanical property estimates inherently depends on many random factors, making statistical analysis essential.

2.4 Virtual screening results

We applied our FiberForge workflow to model the mechanical properties of 72 PDB-resolved amyloid fibrils spanning 18 peptide sequences to probe how sequence and geometry shape mechanical performance and to demonstrate the pipeline's throughput. For each structure, we prepared the workflow code to parse the helical rise and twist, build a 40 nm-long, 6–12 nm-wide atomistic model of about 50 000 atoms, equilibrate it, and ran a 10 ns constant-velocity pull at 0.1 nm ps^{-1} , with post-processing handled automatically; the full set finished in 36 h on a single GPU node. The resulting Ashby plot (Fig. 5) contains 190 simulated data points: elastic moduli range from 7.0×10^7 to $1.1 \times 10^{10} \text{ Pa}$ (median $3.2 \times 10^9 \text{ Pa}$) and ultimate tensile strengths from 9.0×10^7 to $1.0 \times 10^{10} \text{ Pa}$ (median $7.8 \times 10^8 \text{ Pa}$). These values align with experimentally reported amyloid properties (the Young's modulus: 10^8 – 10^{10} Pa ; tensile strength: 10^8 – 10^9 Pa) and sit well above those of typical structural proteins such as collagen or keratin. Wider multiprotofilament fibrils occupy the upper-right of the plot, while slimmer two-strand assemblies fall lower, reflecting the expected dependence on cross-section.^{5,30,31} The match to experiment and the ability to complete dozens of simulations overnight support both the reliability and scalability of our software for large-scale mapping of sequence–structure–property relationships in amyloids.

2.5 Sequence–structure–property relations

To identify the key structural and sequence-derived determinants of amyloid mechanical performance, we examined correlations between the geometric descriptors extracted from PDB structures, sequence hydrophobicity metrics, and the mechanical properties computed *via* SMD (see SI, Fig. S6). Ultimate tensile strength (UTS) exhibited a moderate positive correlation (0.62) with elastic modulus, consistent with the expected coupling between stiffness and failure load in stiff materials. Elastic modulus showed moderate correlation with Hydrogen-bond density (HBD) and sequence hydrophobicity, (0.44 and 0.26, respectively), suggesting a dependence of geometric and sequence properties on elasticity. UTS showed a strong correlation with HBD (0.68) and a moderate negative

correlation with the translation distance between chains (-0.23), suggesting the HBD greatly affects the maximum force that the amyloid can withstand before fracturing. Reducing the translation distance between chains also increase the UTS, which may be a causal effect or simply a result of stronger interactions between chains.

Notably, HBD showed a strong positive correlation with both elastic modulus and UTS, reinforcing earlier theories that interchain HBD is the primary factor affecting deformation resistance. Though earlier studies demonstrated this behavior in specific amyloid systems,^{5,11,12} our results imply that it is a general design principle of planar amyloid systems. Helical twist and rise generally showed weak correlations with mechanical properties, suggesting that local geometric offsets alone do not strongly dictate mechanical response. These two properties showed moderately strong negative correlation with each other, suggesting that a greater degree of twist in an amyloid reduces the spacing between between chains. Sequence mean hydrophobicity displayed modest (0.26) but statistically significant correlations with elastic modulus, indicating that hydrophobic packing may stabilize inter-chain interfaces and thereby alter load-bearing capacity.

Having established key correlations among the structural and sequence descriptors, we next asked whether these features could predict mechanical behavior. To evaluate their predictive power, we performed an automated screening of machine-learning algorithms using AutoML,³² testing dozens of regression models across multiple random train–test splits. The random forest regressor consistently achieves good performance and demonstrates strong robustness to dataset resampling. Based on these results, we selected the random forest as the base model for all subsequent analyses, and assessed how different feature groups contribute to predictive accuracy for elastic modulus and UTS (see SI for the parity plots of cross-validation and hold-out test, Fig. S7).

For elastic modulus, feature sets composed solely of helical parameters (magnitude of translation and rotation), sequence–distance metrics (HBD, cross-sectional area, and H-bond count), hydrophobicity metrics (fraction of hydrophobic residues, mean residue hydrophobicity, standard deviation of residue hydrophobicity), or evolutionary-sequence metrics yield near zero or even negative R^2 , indicating that no single class of descriptors is sufficient to predict elasticity variations. Notably, inclusion of evolutionary sequence embeddings³³ alone decreases predictive accuracy relative to the geometric features, suggesting that sequence evolutionary information does not meaningfully contribute to the elastic behavior of planar amyloids. The strongest predictive performance is achieved when geometric descriptors were combined with hydrophobicity features ($R^2 = 0.377$ on the holdout set), highlighting the interplay between amyloid structure and amino acid properties on elasticity. For UTS, hydrophobicity or helical descriptors alone produced weak predictive performance, and ESM embeddings again performs poorly in isolation. The highest R^2 values are achieved using the geometric, hydrophobicity, and ESM feature set ($R^2 = 0.516$ on the holdout set). These results emphasize that amyloid UTS is influenced by the geometry of



interchain organization, particularly features such as HBD, and evolutionary sequence information. However, the gap in R^2 values between cross-validation and the hold-out test suggests that the model may be overfitting due to the limited size of the training dataset. We anticipate that this issue will diminish as additional molecular simulation data become available.

Overall, both properties are driven predominantly by geometric organization—most notably HBD, which emerges as the single most informative descriptor for elastic modulus and a significant predictor for UTS. Elastic modulus depends largely on HBD and hydrophobicity with only minor gains from any additional features and little to no improvement from evolutionary embeddings. By contrast, UTS also relied heavily on HBD but benefited substantially from the inclusion of evolutionary sequence information, indicating that tensile strength is more sensitive than elasticity to sequence-encoded nuances once the primary geometric scaffold (HBD) is accounted for. This establishes a path toward rapid, simulation-free prediction and optimization of amyloid-based materials, helping to define general design principles and highlighting the promise of integrating interpretable structural features with machine learning for next-generation biomolecular material design.

3 Conclusion

We have developed FiberForge, an end-to-end, high-throughput software suite that automates the construction, simulation, and analysis of amyloid protofibrils for mechanical characterization. By encoding fibril geometry with a helical-symmetry tuple, FiberForge.Build rebuilds experimental structures with good accuracy (median RMSD 2.2 Å across 374 PDB fibrils) and readily generates user-defined lengths and sequence variants. FiberForge.Simulate deploys tailored SMD tensile and bending protocols, while FiberForge.Analyze extracts mechanical observables—elastic modulus, rupture force, and ultimate tensile strength—directly from the resulting trajectories.

Benchmarking shows that (i) reconstruction fidelity remains high across diverse polymorphs, (ii) rupture-force calculations converge within ± 0.010 nN after 30–40 replicas, and (iii) the expected rate- and spring-constant-dependent stiffening follows classical non-equilibrium theory. Using FiberForge, we screened 72 experimentally resolved fibrils (18 sequences), completing 190 tensile tests in 36 h on a single GPU node. The simulated elastic moduli and tensile strengths overlap closely with single-fibril AFM measurements.

Our structure–property analyses reveal that amyloid mechanics arise from the coupled effects of molecular assembly geometry and residue-level interactions. Across fibrils, HBD emerges as the strongest determinant of elasticity and a strongly correlated with UTS. Evolutionary sequence information does not appear to aid in the prediction of elasticity but does appear to aid in the prediction of UTS. This establishes a route toward rapid, simulation-free property prediction and materials optimization.

While FiberForge provides a robust computational platform for mapping sequence–structure–property relationships in amyloids, several limitations remain. The current framework is primarily

optimized for parallel, helical-symmetry protofibrils and does not yet capture anti-parallel, mixed-symmetry, or irregular architectures. Moreover, it is best suited for protofibril-scale fracture characterization, whereas the multi-stranded fibril bundles observed in mature fibers remain beyond its present modeling scope. From a simulation standpoint, the accessible timescales of classical molecular dynamics inherently limit the treatment of rate-dependent mechanical behavior.

Future developments will address these limitations by (i) extending FiberForge.Build to support anti-parallel, mixed-symmetry, and bundled fibril assemblies, (ii) coupling coarse-grained or multi-scale fracture models to capture longer timescales and larger systems, and (iii) integrating automated mutation, ranking, and machine-learning-based property prediction modules to enable high-throughput screening and accelerated design.

Together with our structure–property analysis—showing that mechanical behavior emerges from the interplay of supramolecular geometry and residue-level chemistry, and that features such as hydrogen-bond density, while important in specific amyloid systems, are not universally predictive across all fibrils, FiberForge provides a unified computational foundation for disentangling the multiscale determinants of amyloid mechanics.

FiberForge provides a computational platform for mapping sequence–structure–property relationships in amyloids and accelerates the *in silico* discovery of bio-inspired, high-performance protein materials. Combined with emerging single-fibril experimental techniques, FiberForge promises to bridge atomic-scale insight and macroscopic material design, paving the way for next-generation biomedical and sustainable engineering applications built upon amyloid architectures.

4 Methods

4.1 Molecular modeling protocol

The general procedure for the simulations involves the following steps: First, specify the PDB ID of the starting structure to ensure the correct protein is selected. For example, 2MXU of A β 42 was used in our demo. Next, place the corresponding PDB file in the designated directory. The file is stripped of non-protein molecules. The (θ, t, \mathbf{a}) parameters are then estimated (see parameter estimation section for details). After the parameters are estimated, the system is constructed (see system construction section for details).

In SMD, an external force $F_{\text{ext}}(t)$ is applied to the system according to the equation:

$$F_{\text{ext}}(t) = -k(x(t) - x_0(t))$$

where k is the spring constant, $x(t)$ is the position of the particle at time t , and $x_0(t)$ is the reference or target position at time t .

4.2 Sequence–structure–property relations

4.2.1 Dataset curation. Structures corresponding to incomplete simulations, missing stress–strain data, or those included in a predefined blacklist (*e.g.*, entries with known formatting or geometric anomalies) were excluded from



analysis. For each retained fibril, the corresponding atomic coordinates (PDB format) were parsed using BioPython's PDBParser.³⁵ The amino-acid sequence of the first complete chain was extracted directly from standard residue records.

4.2.2 Descriptor calculations. Inter-chain hydrogen bonds (H-bonds) were quantified using a distance-based criterion: all nitrogen and oxygen atoms were enumerated for each chain, and an H-bond was counted when a donor–acceptor atom pair from distinct chains occurred within 3.5 Å. The final value was normalized by the number of chain pairs to yield an average per-interface H-bond count. Cross-sectional area was computed using the `calculate_cross_sectional_area` routine in FiberForge. In this procedure, the fibril structure is first loaded with MDTraj³⁶ and restricted to protein atoms. A van der Waals surface is generated using the Shrake–Rupley algorithm³⁷ with a 0.6 Å probe radius. The fibril axis is obtained from the helical symmetry estimation procedure, wherein the optimal rotation angle, translational rise, and symmetry axis are determined by minimizing the structural deviation between adjacent chains (See Section 4.4.2 for details on this calculation). Each solvent-accessible surface point is orthogonally projected onto a plane perpendicular to this symmetry axis by removing its axial component. The projected surface elements are summed to yield the geometric cross-sectional area. Helical twist and translational rise values were taken directly from the symmetry parameters estimated for each protofibril. HBD was defined as the total inter-chain H-bond count divided by the computed cross-sectional area.

To assess sequence contributions to mechanics, we calculated Kyte–Doolittle hydrophobicity³⁸ values for each residue in the extracted sequence. Mean, standard deviation, and the fraction of residues with positive hydrophobicity score were used as sequence-level hydrophobicity features. To examine whether protein sequence information, beyond simple hydrophobicity metrics, improves prediction of fibril mechanical performance, we extracted primary sequences for each simulated amyloid structure and embedded them using a state-of-the-art protein language model. First, amino-acid sequences were parsed from the first chain of each PDB file using BioPython. For each valid fibril, we also retrieved precomputed structural descriptors including inter-chain hydrogen-bond count, cross-sectional area, helical rotation and translation offsets, elastic modulus, and ultimate tensile strength (UTS). To capture higher-order, nonlocal sequence information, we computed deep sequence embeddings using the ESM-2 protein transformer model (`esm2_t33_650M_UR50D`).³³ Each sequence was batch-converted into token format, passed through the pretrained model, and the 33rd-layer per-residue representations were averaged to obtain a fixed-length embedding vector for each fibril sequence.

Finally, sequence embeddings were paired with inter-chain hydrogen-bond density, cross-sectional area, Kyte–Doolittle hydrophobicity statistics, as well as helical parameters, to construct an extended feature vector representing sequence, structure, and inter-chain interaction characteristics.

4.2.3 Descriptor–predictor correlations. We computed pairwise Spearman correlation coefficients among elastic

modulus, UTS, helical rotation, inter-chain translation, HBD, and mean hydrophobicity. Spearman *p*-values were obtained using functions adapted from SciPy,³⁹ with safeguards implemented to avoid undefined statistics for constant or short vectors. Correlation matrices and corresponding *p*-value heatmaps were visualized using Seaborn,⁴⁰ with upper-triangle masking applied to avoid redundancy (See in SI, Fig. S6).

4.2.4 Property prediction. Each mechanical target (elastic modulus or UTS) was assembled into a response matrix, and feature vectors were concatenated into a feature matrix. The resulting dataset was used to evaluate how different feature groups influence predictive performance.

Before performing cross-validation analyses, we first identified an appropriate regression model using an automated model-selection procedure. We applied an AutoML workflow³² that screened a broad set of machine-learning algorithms across randomized train–test splits. We performed this operation several times on different train–test partitions to obtain model which we believed was robust to the train–test partitioning. This initial step enabled systematic comparison of dozens of candidate models under repeated resampling, providing a robust estimate of algorithm stability rather than performance on any single data partition. Across all tested regressors, the random forest consistently achieved strong and reproducible performance, demonstrating resilience to dataset reshuffling and minimal sensitivity to outlier splits. Based on these findings, the random forest model was selected as the base estimator for all subsequent analyses.

We trained random forest models under two complementary evaluation schemes. First, we performed 10-fold cross-validation on the full dataset to quantify model robustness and assess the relative predictive value of specific feature subsets. Feature groups included: (i) geometric offsets only, (ii) hydrogen-bonding and cross-sectional metrics, (iii) hydrophobicity features, and (iv) full sequence embeddings. Both features and targets were standardized within each training fold, and performance was assessed *via* the coefficient of determination R^2 . Second, to evaluate generalization, we performed a strict 80/20 train–holdout split, retrained models exclusively on the training portion (including normalization), and computed hold-out R^2 on unseen fibrils. For both evaluation schemes, parity plots were generated for each target–feature-set combination to visualize predictive accuracy and systematic bias (See SI, Fig. S7).

4.3 Data structure

Successful methodologies^{41,42} to reduce the structural complexity of protein aggregates involve understanding and utilizing the symmetries that govern their structure. Prior work has focused on exhaustively characterizing the symmetries that can exist in amyloid structures.⁴³ The symmetries that exist within a protofibril can be characterized by a translational (distance between proteins) and rotational (parallel *vs.* anti-parallel and equifacial *vs.* antifacial) symmetries which characterize the helical structure. In a structure with helical symmetry, a rotation around an axis followed by a translation



along that axis leaves the structure invariant. If R_θ is a rotation by angle θ and T_d is a translation by distance d along the same axis, the symmetry operation can be represented as a combined transformation $R_\theta T_d$. More formally, the helical group H is defined:

$$H = \{R_\theta^k \cdot T_d^k | k \in \mathbb{Z}\} \quad (1)$$

The “classical” pathogenic amyloid structure is characterized by helical symmetry (See eqn (1) for formal description). The rotational symmetry about the growth axis can be represented by a single scalar ($\theta \in \mathbb{R}$), the translational symmetry by a translation scalar ($t \in \mathbb{R}$), and the axis about which these parameters are applied (the growth axis) ($\mathbf{a} \in \mathbb{R}^3$).

The tuple (θ, t, \mathbf{a}) characterizes the symmetry of the protein sheets that make up the protein fibril. It should be noted that functional amyloids, amyloids characterized by solenoid fibrils such as CsgA, can also be described using helical symmetry. In this case θ is simply 1 *i.e.*, only translational symmetry.

4.4 Core algorithms

4.4.1 Protofibril isolation. As noted above, amyloid fibers often have a distinctive hierarchical structure. As such, experimentally determined crystal structures of amyloids commonly contain a fiber bundle composed of several protofibrils. The presence of fiber bundles necessitates the algorithmic isolation of protofibrils that compose the fiber bundle. Specifically we analyze the protein structure from a PDB file to group chains into potential protofibrils based on their spatial proximity.

$$\min_{\theta, t, \mathbf{a}} \sum_{i=1}^N \sum_{j=1}^{M_i} \|\mathbf{r}_{ij}^{(2)} - (\mathbf{R}(\theta, \mathbf{a})\mathbf{r}_{ij}^{(1)} + t\mathbf{a})\|^2$$

$$\mathbf{R}(\theta, \mathbf{a}) = \begin{bmatrix} \cos \theta + a_x^2(1 - \cos \theta) & a_x a_y(1 - \cos \theta) - a_z \sin \theta & a_x a_z(1 - \cos \theta) + a_y \sin \theta \\ a_y a_x(1 - \cos \theta) + a_z \sin \theta & \cos \theta + a_y^2(1 - \cos \theta) & a_y a_z(1 - \cos \theta) - a_x \sin \theta \\ a_z a_x(1 - \cos \theta) - a_y \sin \theta & a_z a_y(1 - \cos \theta) + a_x \sin \theta & \cos \theta + a_z^2(1 - \cos \theta) \end{bmatrix}$$

We created the function `identify_protofibrils` to identify groups of protein chains forming protofibrils based on spatial proximity within a given PDB file. The PDB structure was first parsed using PDBParser from Biopython,³⁵ and for each chain, the atomic coordinates were extracted. The center of mass for each chain was then computed by averaging the Cartesian coordinates of all its atoms.

To group chains into protofibrils, a clustering approach based on a predefined distance threshold was applied. Each chain's center of mass was compared against those in existing protofibril clusters. If the distance between a chain's center and any chain already assigned to a protofibril was below the threshold, the chain was incorporated into that protofibril. If no

existing protofibril met the distance criterion, a new protofibril cluster was initiated.

This procedure resulted in a list of protofibrils, where each protofibril was represented as a dictionary containing chain identifiers and their corresponding centers of mass. This method facilitated the automated identification of spatially grouped chains for further structural analysis.

For a formal description, let the protein structure contain C chains, each with a center of mass $\mathbf{r}_i \in \mathbb{R}^3$, $\mathcal{P} = \{P_1, P_2, \dots, P_k\}$ as the set of protofibrils where each P_j is a subset of chains, $d_{ij} = \|\mathbf{r}_i - \mathbf{r}_j\|$ as the Euclidean distance between chain centers \mathbf{r}_i and \mathbf{r}_j , and δ as the distance threshold for protofibril membership.

The goal is to find a partitioning of chains into protofibrils that minimizes the maximum intra-protofibril distance:

$$\begin{aligned} & \min_{\mathcal{P}} \max_{P_j \in \mathcal{P}} \max_{\mathbf{r}_i, \mathbf{r}_j \in P_j} d_{ij} \\ & s.t. \quad d_{ij} \leq \delta \\ & \quad \forall \mathbf{r}_i, \mathbf{r}_j \in P_j. \end{aligned}$$

4.4.2 Parameter estimation. After the protofibrils have been isolated within a given fibril bundle, the helical structural parameters of the protofibrils can be identified. Protofibrils are composed of an assembly of proteins. Each protein takes the shape of an approximately flat sheet in the case of pathogenic amyloids, or a cylindrical solenoid in the case of functional amyloids. Given the coordinates of a protofibril we can estimate the parameters characterizing the helical symmetry by minimizing the sum of squared deviation of each protein in the fibril:

where θ is the rotation angle (in radians), t is the translation along the symmetry axis, \mathbf{a} is the axis of symmetry (as a unit vector), $\mathbf{r}_{ij}^{(1)}$ is the coordinates of atom j in chain 1 of pair i , $\mathbf{r}_{ij}^{(2)}$ are the coordinates of atom j in chain 2 of pair i , N is the number of chain pairs, M_i is the number of atoms in the i -th chain pair, $\mathbf{a} = [a_x, a_y, a_z]$, $\|\mathbf{a}\| = 1$, and $\mathbf{R}(\theta, \mathbf{a})$ is the rotation matrix about axis \mathbf{a} by angle θ .

4.4.3 System construction. Once the structural parameters of a protofibril are determined or estimated, protofibrils of any length can be generated by iterative application of (θ, t, \mathbf{a}) to a single “seed” protein sheet (see Fig. 1 for details). Formally, the transformation applied at step n can be written as:

$$\mathbf{r}_n = \mathbf{R}(\theta, \mathbf{a})\mathbf{r}_{n-1} + t\mathbf{a} \quad (2)$$



The protofibril structure can be constructed using either the mBuild⁴⁴ or Pymol⁴⁵ software packages to enable direct conversion for molecular dynamics simulations.

A key challenge in SMD of amyloids is the large system size required, given the nanometer scale of fibrils. To automate system construction efficiently, we leveraged the amyloid's helical parameters to optimize its orientation in the simulation box. Specifically, we aligned the fibril along its identified growth axis—a 3D vector—placing it in a rectangular box with its longest dimension parallel to the growth direction. Once the fibril is correctly oriented we can place solvent molecules using PackMol.⁴⁶

4.4.4 Fracture simulations. Due to the orthotropic nature of amyloid fibrils, two directions of deformation fully determine their mechanics: the axis parallel to the growth direction of the fibril and the axis perpendicular to the growth direction of the fibril.

The FiberForge package offers two modes of deformation: tensile and shear. Tensile deformation refers to a pulling force applied parallel to the growth axis, while shear deformation involves a pulling force applied perpendicular to the growth axis (see Fig. 1 for a depiction of these mechanical tests).

To initialize these simulations without human input, we utilize pre-defined data structures. Specifically, we used the translational component *t* of the helical symmetry to orient the fibrils in the simulation box. For tensile tests, the fibrils are aligned parallel to the applied force, and for shear tests, they are aligned perpendicular to the applied force.

Springs are attached to the center of mass of the end chains of the amyloid for a tensile simulation. For a shear simulation restraints are placed on one end of the fibril and a spring is attached to the other end.

SMD simulations are performed using GROMACS.⁴⁷ Systems are solvated using Packmol. After the fibril is placed in the correct orientation in the simulation box, energy minimization is performed. Following minimization, the system undergoes equilibration under an NVT ensemble for 50 ps, followed by further equilibration under an NPT ensemble for another 50 ps. Finally, SMD pulling is performed for 500 ps.

Parameters affecting SMD simulations, such as the pull rate and the force constant, are determined manually. The optimal pull rate is estimated to be 0.01 nm/ps and the optimal force constant is 1000 kJ (mol⁻¹ nm⁻²). Temperature coupling is performed using the Nose–Hoover extended ensemble under standard conditions, and pressure coupling is performed using the Parrinello–Rahman extended ensemble.

4.4.5 Trajectory analysis. The final method we developed for this project is the automated analysis of trajectories to estimate mechanical properties. In its current form, FiberForge can estimate 4 mechanical properties: elastic modulus, shear modulus, ultimate tensile strength, and ultimate shear strength.

The elastic modulus is a measure of a material's resistance to elastic deformation and is generally defined as

$$E = \frac{\sigma}{\varepsilon}$$

where σ is the applied stress and ε is the resulting strain within the linear elastic region.

Similarly, the UTS is defined as

$$\text{UTS} = \max(\sigma)$$

that is, the maximum stress a material can withstand before failure during a tensile test.

These properties are estimated by analyzing the geometry of the protofibril and the SMD trajectory. The F_{ext} and the corresponding deformation distance produced from the SMD simulations are extracted from the output files. To reduce the noise we apply a moving average filter to the stress–strain curve. To further process the simulation data, automated routines were implemented to smooth and fit the stress–strain relationships using cubic spline interpolation. Next, we estimate the linear elastic region of the stress–strain curve by calculating the second derivative of the stress with respect to strain and then extracting the inflections points. The elastic modulus E is then obtained as the slope of the fitted curve within the linear elastic region, which is identified between the initial point and the first inflection point of the second derivative of the spline. This inflection point also defines the yield point, corresponding to the transition between elastic and plastic deformation. To estimate the tensile and shear strength, the maximum F_{ext} obtained during the SMD simulation is used.

Author contributions

KNP and ZJY designed the study. KNP implemented the software, performed SMD simulations, analyzed and gathered computational data. KNP wrote the manuscript, and KNP and ZJY revised the manuscript.

Conflicts of interest

The authors declare no competing financial interest.

Data availability

Archived code for FiberForge can be found on GitHub <https://github.com/ChemBioHTP/FiberForge/tree/v1.0.0> with DOI: <https://doi.org/10.5281/zenodo.18202973>. The most up to date version of FiberForge can be found at <https://github.com/ChemBioHTP/FiberForge>. All additional code and data used to produce figures can be found on Zenodo with DOI: <https://doi.org/10.5281/zenodo.17782766>.

Supplementary information (SI) is available. See DOI: <https://doi.org/10.1039/d5dd00307e>.

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Notes and references

- M. Schleegeer, T. Deckert-Gaudig, V. Deckert, K. P. Velikov, G. Koenderink, M. Bonn, *et al.*, *Polymer*, 2013, **54**, 2473–2488.
- M. Kikumoto, M. Kurachi, V. Tosa and H. Tashiro, *Biophys. J.*, 2006, **90**, 1687–1696.
- J. M. Ferrer, H. Lee, J. Chen, B. Pelz, F. Nakamura, R. D. Kamm and M. J. Lang, *Proceedings of the National Academy of Sciences*, 2008, **105**, 9221–9226.
- F. Vollrath and D. P. Knight, *Nature*, 2001, **410**, 541–548.
- T. P. Knowles and M. J. Buehler, *Nat. Nanotechnol.*, 2011, **6**, 469–479.
- G. Bhak, S. Lee, J. W. Park, S. Cho and S. R. Paik, *Biomaterials*, 2010, **31**, 5986–5995.
- D. Li, H. Furukawa, H. Deng, C. Liu, O. M. Yaghi and D. S. Eisenberg, *Proceedings of the National Academy of Sciences*, 2014, **111**, 191–196.
- S. K. Maji, D. Schubert, C. Rivier, S. Lee, J. E. Rivier and R. Riek, *PLoS Biol.*, 2008, **6**, e17.
- L. Goldschmidt, P. K. Teng, R. Riek and D. Eisenberg, *Proceedings of the National Academy of Sciences*, 2010, **107**, 3487–3492.
- G. Yoon, M. Lee, J. I. Kim, S. Na and K. Eom, *PLoS One*, 2014, **9**, e88502.
- B. Choi, G. Yoon, S. W. Lee and K. Eom, *Phys. Chem. Chem. Phys.*, 2015, **17**, 1379–1389.
- S. Keten, Z. Xu, B. Ihle and M. J. Buehler, *Nat. Mater.*, 2010, **9**, 359–367.
- M. Varadi, G. De Baets, W. F. Vranken, P. Tompa and R. Pancsa, *Nucleic Acids Res.*, 2018, **46**, D387–D392.
- N. Louros, K. Konstantoulea, M. De Vleeschouwer, M. Ramakers, J. Schymkowitz and F. Rousseau, *Nucleic Acids Res.*, 2020, **48**, D389–D393.
- M. Burdukiewicz, D. Rafacz, A. Barbach, K. Hubicka, L. Bkakala, A. Lassota, J. Stecko, N. Szymanska, J. W. Wojciechowski, D. Kozakiewicz, *et al.*, *Nucleic Acids Res.*, 2023, **51**, D352–D357.
- N. Louros, R. Van Der Kant, J. Schymkowitz and F. Rousseau, *Bioinformatics*, 2022, **38**, 2636–2638.
- Q. Shao, Y. Jjiang and Z. J. Yang, *J. Chem. Inf. Model.*, 2022, **62**, 647–655.
- I. Usov and R. Mezzenga, *Macromolecules*, 2015, **48**, 1269–1280.
- I. André, P. Bradley, C. Wang and D. Baker, *Proceedings of the National Academy of Sciences*, 2007, **104**, 17656–17661.
- C. W. Wood, M. Bruning, A. A. Ibarra, G. J. Bartlett, A. R. Thomson, R. B. Sessions, R. L. Brady and D. N. Woolfson, *Bioinformatics*, 2014, **30**, 3029–3035.
- S. A. Bondarev, O. V. Bondareva, G. A. Zhouavleva and A. V. Kajava, *Bioinformatics*, 2018, **34**, 599–608.
- S. Maurer-Stroh, M. Debulpaep, N. Kuemmerer, M. L. De La Paz, I. C. Martins, J. Reumers, K. L. Morris, A. Copland, L. Serpell, L. Serrano, *et al.*, *Nat. Methods*, 2010, **7**, 237–242.
- S. O. Garbuzynskiy, M. Y. Lobanov and O. V. Galzitskaya, *Bioinformatics*, 2010, **26**, 326–332.
- A. W. Bryan Jr, M. Menke, L. J. Cowen, S. L. Lindquist and B. Berger, *PLoS Comput. Biol.*, 2009, **5**, e1000333.
- C. Kim, J. Choi, S. J. Lee, W. J. Welsh and S. Yoon, *Nucleic Acids Res.*, 2009, **37**, W469–W473.
- R. Paparcone and M. J. Buehler, *Biomaterials*, 2011, **32**, 3367–3374.
- H. Lu and K. Schulten, *Proteins: Struct., Funct., Bioinf.*, 1999, **35**, 453–463.
- B. Isralewitz, M. Gao and K. Schulten, *Curr. Opin. Struct. Biol.*, 2001, **11**, 224–230.
- E. Evans and K. Ritchie, *Biophys. J.*, 1997, **72**, 1541–1555.
- U. G. Wegst and M. F. Ashby, *Philos. Mag.*, 2004, **84**, 2167–2186.
- M. F. Ashby, L. Gibson, U. Wegst and R. Olive, *Proceedings of the Royal Society of London. Series A: Mathematical and Physical Sciences*, 1995, **450**, 123–140.
- S. Pandala, *LazyPredict*, GitHub repository, <https://github.com/shankarpandala/lazypredict>.
- Z. Lin, H. Akin, R. Rao, B. Hie, Z. Zhu, W. Lu, N. Smetanin, A. dos Santos Costa, M. Fazel-Zarandi, T. Sercu, S. Candido *et al.*, bioRxiv, 2022, preprint.
- J. Adamcik, C. Lara, I. Usov, J. S. Jeong, F. S. Ruggeri, G. Dietler, H. A. Lashuel, I. W. Hamley and R. Mezzenga, *Nanoscale*, 2012, **4**, 4426–4429.
- P. J. Cock, T. Antao, J. T. Chang, B. A. Chapman, C. J. Cox, A. Dalke, I. Friedberg, T. Hamelryck, F. Kauff, B. Wilczynski, *et al.*, *Bioinformatics*, 2009, **25**, 1422.
- R. T. McGibbon, K. A. Beauchamp, M. P. Harrigan, C. Klein, J. M. Swails, C. X. Hernández, C. R. Schwantes, L.-P. Wang, T. J. Lane and V. S. Pande, *Biophys. J.*, 2015, **109**, 1528–1532.
- A. Shrake and J. A. Rupley, *J. Mol. Biol.*, 1973, **79**, 351–371.
- J. Kyte and R. F. Doolittle, *J. Mol. Biol.*, 1982, **157**, 105–132.
- P. Virtanen, R. Gommers, T. E. Oliphant, M. Haberland, T. Reddy, D. Cournapeau, E. Burovski, P. Peterson, W. Weckesser, J. Bright, *et al.*, *Nat. Methods*, 2020, **17**, 261–272.
- M. L. Waskom, *J. Open Source Softw.*, 2021, **6**, 3021.
- F. DiMaio, A. Leaver-Fay, P. Bradley, D. Baker and I. André, *PLoS One*, 2011, **6**, e20450.
- J. L. Watson, D. Juergens, N. R. Bennett, B. L. Trippe, J. Yim, H. E. Eisenach, W. Ahern, A. J. Borst, R. J. Ragotte, L. F. Milles, *et al.*, *Nature*, 2023, **620**, 1089–1100.
- D. S. Eisenberg and M. R. Sawaya, *Annu. Rev. Biochem.*, 2017, **86**, 69–95.
- C. Klein, J. Sallai, T. J. Jones, C. R. Iacovella, C. McCabe and P. T. Cummings, *Foundations of molecular modeling and simulation, Select papers from FOMMS 2015*, 2016, 79–92.
- W. L. DeLano, *et al.*, *CCP4 Newsl. Protein Crystallogr.*, 2002, **40**, 82–92.
- L. Martínez, R. Andrade, E. G. Birgin and J. M. Martínez, *J. Comput. Chem.*, 2009, **30**, 2157–2164.
- M. J. Abraham, T. Murtola, R. Schulz, S. Páll, J. C. Smith, B. Hess and E. Lindahl, *SoftwareX*, 2015, **1**, 19–25.

