How synonymous mutations alter enzyme structure and function over long timescales

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The specific activity of enzymes can be altered over long timescales in cells by synonymous mutations that alter a messenger RNA molecule’s sequence but not the encoded protein’s primary structure. How this happens at the molecular level is unknown. Here, we use multiscale modelling of three Escherichia coli enzymes (type III chloramphenicol acetyltransferase, D-alanine–D-alanine ligase B and dihydrofolate reductase) to understand experimentally measured changes in specific activity due to synonymous mutations. The modelling involves coarse-grained simulations of protein synthesis and post-translational behaviour, all-atom simulations to test robustness and quantum mechanics/molecular mechanics calculations to characterize enzymatic function. We show that changes in codon translation rates induced by synonymous mutations cause shifts in co-translational and post-translational folding pathways that kinetically partition molecules into subpopulations that very slowly interconvert to the native, functional state. Structurally, these states resemble the native state, with localized misfolding near the active sites of the enzymes. These long-lived states exhibit reduced catalytic activity, as shown by their increased activation energies for the reactions they catalyse.
energetic properties of this alternative state of proteins is a fundamental, unanswered question in biochemistry and molecular biology. Soluble, nonfunctional states do not just occur for enzymes; other protein functions can also be altered. The hormone‐transporting protein transthyretin, for example, can have 20% of its soluble fraction in a nonfunctional state. There are hints in the literature of some of the structural properties of this state. NMR‐derived structures of β/γ‐crystallin produced from two synonymous mRNA variants showed that it populates two soluble conformations differing in the formation of a disulfide bond. Disulfide bonds represent an energetically strong constraint on folding topologies and are uncommon: only 5 and 20%, respectively, of E. coli and human cytoplasmic proteins contain disulfide bonds. Furthermore, many enzymes that exhibit specific activity changes due to synonymous mutations do not contain disulfide bonds. Thus, the nature of these structurally altered subpopulations has not yet been resolved, although experimental data rule out misfolding involving quaternary structures. For example, gel separation and chromatography experiments rule out the formation of off‐pathway dimers and higher‐order oligomers for many enzymes that normally function in a monomeric state.

Here, we use a novel multiscale approach across the dimensions of time, space and energy to simulate the synthesis, post‐translational behaviour and function of enzymes under different translation rate schedules that arise from synonymous codons. We establish that our modelling approach can predict the experimentally measured changes in specific activity for CAT‐III, α‐alanine‐β‐alanine ligase B (DDLB) and dihydrolipoyl reductase (DHFR). Dissecting the structures and kinetics of co‐ and post‐translational folding that occur in our simulations, we show that synonymous mutations shift the folding pathways and populations of near‐native non‐covalent lasso entanglements, each of which has its own intrinsic activity (as measured by the reaction rate constant \( k_{\text{cat}} \)) and leads to long‐term changes in the enzymatic specific activity.

Results

Recapitulating experimental trends of CAT‐III

First, we applied our modelling protocol to the E. coli enzyme CAT‐III to test whether the method correctly predicts the influence of synonymous mutations on specific activity. CAT‐III has been shown to have a 20% decrease in specific activity when faster‐translating codons are introduced through synonymous mutations. We created both fast‐ and slow‐translating synonymous mRNA sequences (denoted, respectively, CAT‐III\(_{\text{fast}}\) and CAT‐III\(_{\text{slow}}\)) by replacing each wild‐type codon with its fastest or slowest synonymous variant (Fig. 1b); mRNA sequences are reported in Supplementary Methods Section 22). The resulting slow mRNA variant takes twice as long to translate than the fast variant (Fig. 1f). We simulated the synthesis and post‐translational behaviour of CAT‐III resulting from the fast and slow mRNA variants and calculated their respective specific activities (equation (4)) at the end of the post‐translational simulations. In our model, CAT‐III\(_{\text{fast}}\) exhibited 83.6% (95% confidence interval (CI) = 72.0–96.5%) of the activity of the fast variant (Fig. 1f). After synthesis was completed, DHFR\(_{\text{fast}}\) exhibited a specific activity that was 100% (95% CI = 100–100%) that of DHFR\(_{\text{slow}}\) (\( P = 1 \); that is, not statistically significant). Therefore, our model describes enzymes whose specific activity is either sensitive (DDLB) or insensitive (DHFR) to changes in translation speed over long timescales.

Accurate prediction of trends in specific activity

To experimentally test whether DDLB\(_{\text{slow}}\) has a reduced enzymatic activity compared with DDLB\(_{\text{fast}}\), we recombinantly expressed the fast and slow DDLB variants in E. coli using the same mRNA sequences as in the simulations. We then purified the enzyme and assayed the reaction kinetics (see Supplementary Methods Section 19). We observed that the fast variant had a higher level of protein expression 5 h after induction than the slow variant, consistent with the fast mRNA variant translating faster (Supplementary Fig. 11). The reaction rate constant \( k_{\text{cat}} \) was measured across five biological replicates of the fast and slow variants (Supplementary Table 1). We found that the specific activity of DDLB\(_{\text{slow}}\) was, on average, 88% (\( k_{\text{cat}} \) of DDLB\(_{\text{fast}}\): 95% CI = 81.3–94.8%) that of DDLB\(_{\text{fast}}\) (\( P = 0.0186 \); one‐tailed \( t \)‐test for \( k_{\text{cat}} \) of DDLB\(_{\text{slow}}\)/\( k_{\text{cat}} \) of DDLB\(_{\text{fast}}\) < 1). In contrast, we performed the same experiments to measure the \( k_{\text{cat}} \) of fast and slow DHFR variants (see Supplementary Methods Section 20). Consistent with our model prediction, the specific activity of DHFR\(_{\text{slow}}\) was indistinguishable from that of DHFR\(_{\text{fast}}\) (\( k_{\text{cat}} \) of DHFR\(_{\text{slow}}\)/\( k_{\text{cat}} \) of DHFR\(_{\text{fast}}\) = 91%; 95% CI = 68–115%; \( P = 0.5323 \); two‐tailed \( t \)‐test for \( k_{\text{cat}} \) of DHFR\(_{\text{slow}}\)/\( k_{\text{cat}} \) of DHFR\(_{\text{fast}}\) = 1). Thus, our modelling approach also successfully predicts the trends in changes in activity for DDLB and DHFR, indicating that the model is realistic.

Near‐native, lasso‐like entangled structures

Next, we identified in our simulations the structures, catalytic properties and folding pathways that cause these activity changes. First, we examined the structural distributions of CAT‐III, DDLB and DHFR in the post‐translational simulations using a clustering algorithm that employs both structural information and temporal interconversion rates between metastable states. Specifically, after numerous tests of different metrics, we performed structural clustering on the basis of the fraction of native contacts formed in the enzyme’s substrate‐binding regions (denoted as \( Q_{\text{act}} \); equation (3)) and the fraction of native contacts that exhibited non‐native topological entanglements (denoted as \( G \); equation (2); see Methods). The resulting metastable states are shown in Fig. 2a,b for CAT‐III, Fig. 3a,b for DDLB and Fig. 4a,b for DHFR.

Across the metastable states, we observed diverse misfolded structures for CAT‐III and DDLB (Supplementary Figs. 5 and 6). Most of the misfolded structures exhibited topological entanglements, and many of these entangled structures were near native, having >60% of native contacts formed (\( Q_{\text{act}} \geq 0.6 \) and \( G \geq 0.02 \); interactive visualization is provided at https://obrien‐lab.github.io/visualize_entanglements/). Misfolded, metastable states P9, P10, P11, P12 and P13 of CAT‐III exhibited \( Q_{\text{act}} \) and \( G \) values, respectively, of 0.66 and 0.18, 0.72 and 0.04, 0.80 and 0.02, 0.84 and 0.09 and 0.86 and 0.15, while the native state (P14) had values of 0.89 and 0 (Fig. 2g and Supplementary Fig. 5). The DDLB misfolded states P4, P6, P7, P8 and P9 exhibited values of 0.64 and 0.06, 0.83 and 0.19, 0.85 and 0.09, 0.91 and 0.05 and 0.92 and 0.15, respectively, while the native state (P14) had values of 0.89 and 0 (Fig. 2g and Supplementary Fig. 5). The DHFR misfolded states P6 and P9 exhibited values of 0.69 and 0.06 and 0.79 and 0.15, respectively, while the native state (P14) had values of 0.68 and 0 (Fig. 3g and Supplementary Fig. 5). In contrast, DHFR exhibited fewer misfolded states and only one entangled, misfolded state, with \( Q_{\text{act}} \) and \( G \) values of 0.73 and 0.05 compared with the values 0.93 and 0 for its native state (Fig. 4g and Supplementary Fig. 7). All of the topological entanglements that we observed had a non‐covalent lasso topology, where native contacts within a certain folded region established a closed loop along the protein backbone and another segment in the same chain threaded
Some entangled structures are long-lived kinetic traps

To disentangle a misfolded structure, it is often necessary to unfold some portion of the properly folded segments to attain the native state. This can be energetically costly. Therefore, we hypothesized that these near-native, entangled structures are long-lived kinetic traps. To test this hypothesis, we calculated the post-translational probability of being in each metastable state as a function of time. These results are reported in Figs. 2c, 3c and 4c. For CAT-III and DDLB, entangled states (P13 for CAT-III and P4 and P8 for DDLB) persisted with appreciable populations (>10%) until 60 s after nascent protein release from the ribosome. For DHFR, the single entangled state (P2) populated only 0.2% of the trajectories and disentangled by 10 s. As a further test, we examined whether any post-translational trajectory of CAT-III or DDLB that reached an entangled state ever converted to an unentangled state. For CAT-III and DDLB, we found that 92.7 and 100% of trajectories sampled an entangled state and, of these, 78.6 and 10.4% did not convert to a state that was not entangled by the end of the simulations. Thus, we conclude that many of these entangled structures represent long-lived kinetic traps that convert to the native state slowly.

Entangled structures have altered catalytic properties

The non-covalent lasso entanglement intermediates we observed are a form of misfolding, as they represent ordered structures that are not native. Therefore, we hypothesized that some of these entangled structures have catalytic properties different from those of the native ensemble. To test this hypothesis, we calculated the transition state barrier height of the enzyme reaction of each metastable state that formed a native or near-native active site (\(Q_{act} \geq 0.6\)) using a back-mapping procedure to an all-atom representation and subsequent quantum mechanics/molecular mechanics (QM/MM) umbrella sampling simulations of the catalytic reaction (see the potential of mean force plots in Supplementary Figs. 8–10). Thus, for each metastable state, we obtained the median activation free energy barrier \(\Delta G^\ddagger\) going from reactants to products. For all three enzymes, the specific activity values are normalized to the higher specific activity value found in fast and slow mRNA variants. The error bars in f and g represent 95% CIs about the mean, as estimated using bootstrapping over all simulation trajectories (sample sizes, \(n = 100\) trajectories for CAT-III and DDLB and \(n = 50\) trajectories for DHFR). P values characterize the statistical significance of the difference in specific activities between the proteins produced from the fast and slow variants. They were calculated using a one-tailed permutation test. The specific activities of CAT-III and DDLB are sensitive to translation speed changes, whereas that of DHFR is not.
Deep entanglements are usually long lived

Not all entangled structures are long-lived kinetic traps; otherwise, the entangled structure of DHFR (state P2 in Fig. 4g and Supplementary Fig. 7) would persist. Sliding a small number of residues out of the closed loop (Fig. 5; see also Supplementary Methods Section 11) tends to be easier than sliding a large number of residues. In addition, unfolding a small number of residues during disentanglement also tends to be easier than unfolding a large number of residues. Therefore, we hypothesized that the minimum number of residues involved in the threading segment, whose repulsion can cause disentanglement, and the minimum number of residues needed to unfold during the disentanglement process should correlate with the ability of entangled metastable states to interconvert to unentangled states. To test this hypothesis, we analysed the representative structures from each metastable state. The entanglement of DHFR involved only, on average, five residues in the threading segment, and sliding these segments through the loop should not cause any portion of the protein to unfold. For CAT-III and DDLB, entangled states that never disentangled in the
**Fig. 3** Slow translation partitions more DDLB into post-translational kinetically trapped entangled states. 

**a, b.**  Log probability surfaces ($-\ln[P]$), where $P$ is the probability of sampling particular $Q_{act}$ and $G$ values for the post-translational folding of fast-translating (a) and slow-translating (b) mRNA variants as a function of the order parameters $Q_{act}$ and $G$ (left) and the regions corresponding to different metastable states (right). The native state is located in the bottom right-hand corner of these plots. 

**c.** Time courses of gross state probabilities (soluble + insoluble; same colours as the metastable states in a and b) with error bars shown as transparent stripes for fast (left) and slow (right) variants. 

**d.** Aggregation, degradation and Hsp70 binding propensities of each metastable state, calculated using Supplementary Equation (19). 

**e, f.** Percentage of soluble protein in each metastable state, calculated using Supplementary Equation (17). 

**g.** Crystal structure of DDLB showing the native state and the entangled state P4. The entanglement in state P4 is represented in the same way as in g. The native conformation corresponding to the entanglement is highlighted in the native state structure. The proteinase K site residue 192 is shown as a green ball. It is more exposed to solvent in the entangled state P4. In g, the entangled states are labelled in red, the native state (that is, state P10) is labelled in green and the others are labelled in black. All error bars represent 95% CIs of the statistics calculated by bootstrapping over all simulation trajectories/representative structures (sample sizes, $n = 1,000$ trajectories in c, $n = 5$ representative structures in d and e and $n = 1,000$ frames in f). Each of the bar charts in d and e is overlaid with the data points of the five representative structures. The structures of P4 and P8 can be explored interactively at https://obrien-lab.github.io/visualize_entanglements/.

**h.** Recommended state structure. The proteinase K site residue 192 is shown as a green ball. It is more exposed to solvent in the entangled state P4. In h, the entangled states are labelled in red, the native state (that is, state P10) is labelled in green and the others are labelled in black. All error bars represent 95% CIs of the statistics calculated by bootstrapping over all simulation trajectories/representative structures (sample sizes, $n = 1,000$ trajectories in c, $n = 5$ representative structures in d and e and $n = 1,000$ frames in f). Each of the bar charts in d and e is overlaid with the data points of the five representative structures. The structures of P4 and P8 can be explored interactively at https://obrien-lab.github.io/visualize_entanglements/.
were extrapolated to 298 K. We are $k = 2.24 \times 10^{-3} \text{s}^{-1}$, extrapolated to 0 M urea at 0°C. We then carried out temperature jump (from DHFR state P2) to an all-atom representation (see Supplementary Methods Section 13). We also simulated and calculated the unfolding time of native DHFR ensembles, we also simulated and calculated the unfolding time of native DHFR at these temperatures and compared it with its experimentally measured unfolding rate ($k_{\text{uf,exp}} = 2.24 \times 10^{-3} \text{s}^{-1}$, extrapolated to 0 M urea at 298 K)$^{12a}$. From these simulations, the disentangling times ($k_{\text{de}}^{\text{sim}}$) were extrapolated to 298 K. We found that the all-atom force field accelerated the unfolding of DHFR by 144-fold at 298 K. By correcting $k_{\text{de}}^{\text{sim}}$ for this force field bias using this acceleration factor (that is, $k_{\text{de}}^{\text{rescale}} = 144/k_{\text{de}}^{\text{sim}}$), we estimated that for the deep entanglements in CAT-III and DDLB, the values of $k_{\text{de}}^{\text{rescale}}$ are $6.2 \times 10^{3} \text{s}^{-1}$ (95% CI = 29 s–$1 \times 10^{3} \text{s}$), and 1.3 $\times 10^{4} \text{s}^{-1}$ (95% CI = 28 s–$4 \times 10^{3} \text{s}$), respectively. The shallow entanglement of DHFR becomes disentangled with a timescale of 71 s (95% CI = 4 $\times 10^{3}$–2 $\times 10^{4} \text{s}$), which is two

**Fig. 4** | No kinetically trapped states arise in synonymous variants of DHFR. a, b, Log probability surfaces (−ln[P]), where P is the probability of sampling particular Q$_{\text{act}}$ and G values) for the post-translational folding of fast (a) and slow (b) mRNA variants as a function of the order parameters Q$_{\text{act}}$ and G (left) and the regions corresponding to different metastable states (right). The native state is located in the bottom right-hand corner of these plots. c, Time courses of gross state probabilities (soluble + insoluble; same colours as the metastable states in a and b) with error bars shown as transparent stripes for fast (left) and slow (right) variants. d, Aggregation, degradation and Hsp70 binding propensities of each metastable state, calculated using Supplementary Equation (19). e, Percentage of soluble protein in each metastable state, calculated using Supplementary Equation (17). f, Median transition state barrier heights ($\Delta G$) for the near-native metastable states calculated from the QM/MM simulations. g, From left to right: representative structures of the shallow-entangled state P2 (the closed loop and threading segment of the entangled region are coloured in red and blue, respectively), the native state P3 and the crystal structure (4KJK), with the substrate shown in magenta. The Q$_{\text{act}}$ and G values of the most probable cluster (microstate) for each state are reported below the structure in the format (Q$_{\text{act}}$, G value, $\Delta G$ value). In a–g, the native state (that is, state P3) is labelled in green and the others are labelled in black. All error bars represent 95% CIs of the statistics calculated by bootstrapping over all simulation trajectories/representative structures (sample sizes, $n = 500$ trajectories in c, $n = 5$ representative structures in d and e and $n = 1,000$ frames in f). Each of the bar charts in d and e is overlaid with the data points of the five representative structures. The structure of P2 can be explored interactively at https://obrien-lab.github.io/visualize_entanglements/.
orders of magnitude faster than the deep entanglements. These results suggest that the deep-entangled metastable states we have identified are likely to be long-lived kinetic traps, persisting for hours.

Native-like entangled states are likely to remain soluble
The proteostasis machinery in cells has the potential to catalyse the folding (repair) or degradation (clearance) of these entangled structures. The entangled structures could also potentially aggregate and be removed from the pool of soluble enzymes. We accounted for the effects of these processes by estimating the aggregation, degradation and Hsp70 binding propensities of the different metastable states (see Supplementary Methods Section 14.2 and Supplementary Equation (19)).

We observed, as expected, that the less structured metastable states have a higher propensity to aggregate, be degraded or interact with chaperones. For example, relative to the native state, states P1–P8 of CAT-III (see structures in Supplementary Fig. 5) are much more likely to experience one of these processes, as indicated by the larger magnitude of the bar plots in Fig. 2d. However, states P9–P13 exhibit similar propensities to the native state (P14) to aggregate, be degraded or become chaperone substrates. Similar results were observed for DDLB (Fig. 3d) and DHFR (Fig. 4d). Thus, our model indicates that some of these entangled structures do not interact with the proteostasis machinery any more than the native state does. Therefore, the altered specific activity we observe is likely to persist inside cells over long timescales. This prediction was confirmed by our enzyme kinetics assays (see the section ‘Accurate prediction of trends in specific activity’), since the fast and slow variants of DDLB were expressed in E. coli cells possessing the full complement of the proteostasis machinery (see Supplementary Methods Section 19).

We estimated the percentage of proteins in each metastable state that are likely to remain soluble by accounting for the aggregation, degradation and Hsp70 binding propensities within each metastable state (see Supplementary Methods Section 14.2 and Supplementary Equation (17)). We found that for both CAT-III and DDLB, many of the near-native entangled structures remained soluble. At least 97% of the proteins in entangled states P10, P11, P12 and P13 for CAT-III were estimated to remain soluble (Fig. 2e), while at least 89% of the proteins in entangled states P4, P6, P7, P8 and P9 of DDLB were expected to remain soluble (Fig. 3e). Thus, many of the long-lived, near-native entangled states are likely to remain soluble and free from aggregation, degradation and catalysed folding by chaperones.

The reason for this is that these near-native structures sequester the residues and sequence motifs that promote these processes to an extent similar to that in the native state. For example, state P13 of CAT-III exposes a similar amount of hydrophobic surface area as the native state ensemble (36.2 versus 34.7 nm²), so too with the exposed surface areas of residues that promote aggregation (21.6 versus 18.9 nm²) and interactions with the chaperone Hsp70 (40.0 versus 38.6 nm²).

Consistency with limited proteolysis mass spectrometry experiments
To experimentally test whether these entangled states exist, we carried out limited proteolysis mass spectrometry (LiP-MS) in which E. coli lysates were globally unfolded through incubation in 6 M guanidinium chloride, then refolded by dilution, and the conformations of the resulting refolded proteins were assessed by their susceptibility to proteolysis with proteinase K. With liquid chromatography tandem mass spectrometry, tens of thousands of fragments were identified and quantified, of which a number were from DDLB and DHFR, enabling an assessment of whether or not their refolded conformations were similar to their native forms. In these experiments, proteins were allowed to refold for 1 min, 5 min or 2 h following dilution from denaturant, then pulse proteolysis was conducted, providing a snapshot of their structural ensemble at distinct timescales. Peptide fragments that contained a cleavage site arising from proteinase K were interpreted as demarcating sites within a protein that were solvent exposed or unstructured; hence, if such a fragment was present in greater abundance in the refolded samples (relative to untreated, native samples), it implied that a population of the protein failed to form the native-like structure at that site. Additional details can be found in Supplementary Methods Section 21. For DDLB, we found one statistically significant peptide fragment (abundance ratio greater than twofold; P < 0.01 by Welch’s t-test), with a proteinase K cleavage site at residue 192, whose abundance was at least 2.6 times greater in the refolded samples at all three time points (Fig. 3h; the full dataset is provided in Supplementary Table 13), indicating that residue 192 is more exposed to solvent than in the native state. Cross-referencing this site against the long-lived metastable states in our simulations (Fig. 3g), we found that state P4 contains an entanglement in which residue 192 is part of the threading segment and more exposed to solvent (Fig. 3h). This misfolding results in a 12-fold increase in the solvent-accessible surface area for residue 192 compared with the native state, in which it is part of a β sheet. In addition, states P4 and P8 were sampled by DDLB when refolding was commenced from a thermally denatured state in our model (Supplementary Fig. 15), indicating that many of the same misfolded states populated during synthesis are also populated through refolding in the absence of the ribosome. Thus, comparison of the LiP-MS data to co- and post-translational folding is reasonable. Therefore, our simulations provide a molecular interpretation of LiP-MS refoldability experiments and are consistent with the existence of protein entangled states.
In contrast, we predicted that DHFR does not exhibit long-lived misfolded states. Indeed, in the LIP-MS data there was a peptide that exhibited a significant abundance difference in the refolded samples after 5 min of refolding time; however, it was not present at the 1 min and 2 h time points (Supplementary Table 14), indicating that the refolded protein is indistinguishable from its native conformation on long timescales. Thus, any structural distortions in DHFR during refolding were short-lived. The single significant peptide indicates that a region spanned by residues 77–98 is more exposed to solvent in the misfolded state than in the native state. Indeed, in the short-lived entangled state, P2, we found that this region had a 3% larger solvent-accessible surface area than in the native state (a 1.05-fold increase with a 99% CI of 1.02–1.07; two-tailed t-test, \( P = 6 \times 10^{-5} \)). Therefore, this negative control provides further evidence of the accuracy of our model’s predictions.

**Synonymous mutations alter the entangled state populations**

Next, we examined how translation speed changes affected the post-translational populations of entangled states. We observed that for CAT-III, 60 s after translation termination, 76.3% (95% CI = 73.6–78.9%) of the structures were entangled when synthesis was fast, while 71.6% (95% CI = 68.8–74.3%) were entangled when synthesis was slow. Likewise, the entangled population of DDLB accounted for 35.7% (95% CI = 32.7–38.7%) and 30.2% (95% CI = 27.3–33.0%) of the total under slow and fast translation, respectively. In contrast, the population of entangled structures for DHFR remained zero regardless of the translation speed 10 s post-termination. Thus, synonymous mutations can alter the population distributions of entangled states over long timescales for deep entanglements.

**Synonymous mutations cause a divergence of folding pathways**

We hypothesized that the changes in translation speed alter the co-translational folding pathways of the protein. First, we assessed how different the nascent chain structural distributions were under fast and slow translation by applying the Jensen–Shannon divergence metric (Supplementary Equation (21)) to the population of microstates identified as part of our Markov state modelling. A Jensen–Shannon value of 0 means that there is no difference between the distributions, while a value of In2 means the distributions are completely different. We found that for all three proteins, the structural divergence induced by synonymous mutations was small at short nascent chain lengths and tended to increase as the nascent chain length increased (Fig. 6b,c,h). The maximum divergence occurred at or near the longest nascent chain length before the nascent chain was released from the ribosome. For DHFR, the structural ensembles arising from fast and slow synthesis started to diverge at approximately 110 residues in length, while for CAT-III and DDLB, the two distributions started to diverge at 190 and 210 residues, respectively. Thus, the fast and slow translation rates altered the co-translational distributions of conformations for all three enzymes.

The post-translational (metastable state) structural distributions (Supplementary Equation (22)) for DHFR, which were initially different between fast and slow synthesis, rapidly converged to the same distribution (that is, they had Jensen–Shannon values equal to zero; Fig. 6h). For CAT-III and DDLB, the two distributions did not reconverge 60 s after synthesis (0.02 for CAT-III and 0.27 for DDLB; see Fig. 6b,c). Thus, the changes in conformation caused by synonymous mutations are quickly forgotten by DHFR due to its rapid equilibraition, while CAT-III and DDLB retain a memory of those co-translational differences due to kinetic trapping.

To characterize changes in co- and post-translational folding pathways, we calculated the populations and pathway probabilities of transitions between metastable states occurring co- and post-translationally under the different translation schedules (see Supplementary Methods Section 17). As shown in Fig. 6, black arrows between metastable states indicate that a transition was observed between those states in both translation schedules in the top 80% of populated pathways. Blue arrows indicate that the transition was seen only in the slow translation schedule and orange arrows indicate that the transition was seen only in the fast translation schedule.

We identified nine co- and three post-translational metastable states for DHFR (representative structures shown in Fig. 6g). The co- and post-translational folding pathways were very similar, as most of the transitions were seen in both translation schedules (black arrows in Fig. 6g). There were some differences though. Transitions from state C7 to C9 and from C9 to C8 were observed only for the slow schedule. The pathway probabilities, however, demonstrated that even though the initial post-translational structural distribution differed, all states converged into the native state by the end of the post-translational simulations (Fig. 6i). For example, the pathway probabilities of C8→P3 and C9→P3 were 100% for both fast and slow translation. An exception was the transition probability of C6→P3 that occurs in the fast variant but not in the slow variant, because in the slow variant state C6 is never populated. However, because C6 quickly converts to the native state P3, the effect on the time-dependent native state population is negligible. Thus, the co-translational folding of DHFR was slightly affected by differences in translation speed; however, these differences quickly disappeared due to rapid folding from all post-translational metastable states.

In contrast, CAT-III exhibited a co- and post-translational folding network (Fig. 6a) and pathway probabilities that were sensitive to translation speed changes and whose resulting differences persisted post-translationally. Nine co-translational and 14 post-translational metastable states were identified for CAT-III (Fig. 6a). Seven of these post-translational metastable states (P5, P6, P9, P10, P11, P12 and P13) exhibited entangled structures, while no co-translational states exhibited entanglement. Thus, entanglement is a post-translational process for CAT-III (see Supplementary Videos 1 and 2 visualizing, respectively, the process of entanglement versus correct folding).

Very different co-translational folding pathways were observed starting at a nascent chain length of approximately 195 residues, where the divergence metric exhibited a large increase (Fig. 6b), and transitions into and out of co-translational metastable state C9 started to differ between the fast and slow translation schedules (as indicated by the blue and orange arrows in Fig. 6a). For example, only in the slow schedule could C9 transition to C6, while only in the fast schedule could C5 transition to P1. These differences in allowed transitions persisted post-translationally as well, with state P13 being an effective sink during the post-translational simulations (that is, allowing no direct or indirect transitions to the native state P14 once it was reached). This sink had deep-entangled structures, consistent with our earlier observation that the deep-entangled states tend to be kinetic traps (as indicated by the red border around metastable state P13 in Fig. 6a). Finer-grained consideration of the folding pathways showed that post-translationally, CAT-III fast partitioned 5.0% (95% CI = 1.8–8.9%) more protein into sink P13, while CAT-III slow partitioned 4.7% (95% CI = 0.8–8.6%) more protein into native state P14. The transitions towards the native state P14 were enhanced for CAT-III slow whereas the transitions towards the kinetic trap P13 were enhanced for CAT-III fast. This is also indicated by the pathway probabilities shown in Fig. 6c. Synonymous mutations also led to smaller changes in other states, including the entangled states P6, P10 and P12. Thus, the change in translation speed causes differences in the CAT-III co-translational folding pathways once at least 195 residues have been synthesized, and these differences lead to changes in the populations of post-translationally entangled states, thereby altering the transition state barrier for the reaction this enzyme carries out (Fig. 2f) and affecting the specific activity of CAT-III over long timescales (Fig. 1g).

Similar to CAT-III, DDLB also exhibited co- and post-translational folding pathways that were sensitive to changes in translation speed across its 11 co- and ten post-translational metastable states. However,
Fig. 6 | Co- and post-translational folding pathways of CAT-III, DDLB and DHFR. a–d. The most probable folding pathways of CAT-III (a), DDLB (d) and DHFR (g) are presented as a directed graph where the nodes represent metastable states, with representative structures shown in the boxes. The edges represent the transitions in the 80% most likely pathways. The nodes corresponding to the native state and the kinetically trapped states are framed in green and red, respectively. The transitions (arrows) that are observed in only the fast or slow variants are marked in orange and blue, respectively. b, e–h. Jensen–Shannon divergence of the co-translational microstate distributions (left; Supplementary Equation (21)) and post-translational metastable state distributions (right; Supplementary Equation (22)) of CAT-III (b), DDLB (e) and DHFR (h), comparing fast and slow variants, with a red line indicating zero divergence. aa, amino acids. c, i. Co-translational pathway probabilities (* → end state C*) and post-translational pathway probabilities from the co-translational end states to the native state (C* → native state) for CAT-III (c), DDLB (f) and DHFR (i). The pathway probabilities whose changes were >5% are highlighted in yellow. The post-translational pathway probabilities are normalized by the corresponding co-translational pathway probabilities that involve the particular co-translational end state.

Unlike CAT-III, DDLB co-translationally formed entangled structures (states C8 and C9) that persisted to form entangled post-translational states (states P1, P3, P4 and P8). In CAT-III, the divergence in co-translational folding pathways between fast and slow translation schedules monotonically increased with increasing chain length (Fig. 6b). However, the co-translational folding pathways of DDLB diverged starting at a nascent chain length 110 residues but reconverged around 190 residues, diverging again starting at 210 residues.
residues (Fig. 6e). The first divergence involved a transition to state C4 with the fast schedule that did not occur with the slow schedule (orange arrow in Fig. 6d). This state ultimately interconverted to state C3, which is an obligatory intermediate in both schedules. The second divergence started at state C7, in which 6% more nascent chains went to entangled state C8 with the slow schedule than with the fast schedule, and the rest went to state C10. Parallel co- and post-translational folding pathways then arose. In one pathway, transitions were observed from C10→C11→P2, while the other transitions from C8→C9→P1 involved entangled structures in all states (Fig. 6d; see also Supplementary Videos 3 and 4 illustrating the processes of entanglement and folding). The increased probability of this entangled pathway during slow translation decreased the probability that DDLB would post-translatationally reach the native state P10 (3.9 versus 11.7% in slow versus fast translation; Fig. 6f), while DDLB_in, partitioned 5.8% (95% CI = 3.2–8.3%) more protein (gross amount; that is, soluble plus insoluble) into the near-native kinetic trap P4 after post-translation. Thus, we again see that entangled states act as sinks with altered transition state barriers for the enzyme's reaction (Fig. 3f). These population shifts among these states and the folding pathways they take part in lead to long-lived changes in specific activity.

Discussion

A detailed understanding of the molecular mechanisms giving rise to the coupling of enzyme activity to synonymous mutations would provide biochemists, molecular biologists, evolutionary biologists and biomedical researchers with a framework with which to interpret experiments on the influence of codon usage on protein structure and function, cellular phenotype, disease and some of the selection pressures shaping mRNA sequence evolution. In this study, we developed a novel multiscale model that qualitatively recapitulates the experimental observations for the specific activity of CAT-III variants and correctly predicts the trends in specific activity changes of DDLB and DHFR variants, giving us confidence that the model is realistic. This provided us with the opportunity to study the structures, pathways and kinetics that give rise to this phenomenon.

Our key findings are: (1) changes in elongation kinetics induced by synonymous mutations can alter co-translational nascent chain structural ensembles and folding pathways; (2) for some enzymes, such as CAT-III and DDLB, these changes in the structural ensemble can persist long after the nascent chain has been released from the ribosome; (3) this persistence arises from conformational states that are long-lived kinetic traps; (4) these kinetic traps arise at the molecular level from deep entanglements that slowly disentangle because they require the unfolding of already folded protein segments; (5) these entanglements are non-covalent lasso topologies in which a closed loop is formed by a backbone segment connecting two residues that form a non-covalent native contact, and another segment threads through this loop; (6) many of these entangled structures have decreased catalytic efficiencies due to structural perturbation of their active sites; (7) some entangled structures are very similar to the native state, exposing similar extents of hydrophobic surface area and chaperone binding motifs; and (8) because of their near-native conformations, these entangled structures can bypass the chaperone and degradation machinery of the cell and do not exhibit an increased propensity to aggregate. This situation results in a soluble fraction of enzymes with decreased specific activities that can persist for long time periods in cells.

Two concepts central to our explanation—intramolecular entanglement and subpopulations of kinetically trapped states—are not without precedent. The material properties of entangled synthetic polymers have long been studied and modelled in the field of polymer physics. Also, there has been a large amount of research focused on knotted proteins that often contain disulfide bonds or topologies in which when the protein’s ends are pulled in opposite directions the protein does not fully extend. These characteristics, however, were not present in our entanglements. The entanglements we observed formed a closed loop due to a non-covalent native contact, and if both ends of the protein were pulled, disentanglement would occur. Recently, this type of entanglement has been detected in almost one-third of the protein structures deposited in the Protein Data Bank. Thus, the potential for protein segments to form non-native, non-covalent lassos is plausible. Kinetically trapped states of proteins have been observed in single-molecule experiments probing the functioning of flavoenzymes. In one study, heterogeneous populations of the protein cholesterol oxidase were observed to stochastically switch very slowly between active and nonactive states. However, in that study, the influence of protein synthesis on the distribution of conformational states was not probed. Thus, a novel aspect of our discovery is that it combines the phenomena of entanglement and kinetic trapping as essential to the mechanism by which synonymous mutations affect co- and post-translational protein structure and function.

In contrast with CAT-III and DDLB, the changes in DHFR's co-translational structures and folding pathways did not persist post-translationality. This is consistent with previous studies showing that DHFR has fast folding kinetics and an absence of off-pathway intermediates and kinetic traps. Two reasons for this are that DHFR only negligibly populates an entangled conformation and that the entanglement it forms is shallow; its disentanglement requires only five residues to slide out of the closed loop and there is no requirement to unfold any other portions of the protein to do so. Thus, this shallow entanglement is rapidly disentangled without large structural rearrangements. In contrast, CAT-III and DDLB entanglements involve many more residues, requiring large structural rearrangements to become disentangled, leading to longer-lived entangled structures. Thus, the presence of entanglement is not sufficient to guarantee a kinetic trap. The nature of the entanglement and the native structure surrounding it is critical.

Long-lived entangled states have the potential to be probed using a variety of experimental techniques. Ensemble-level experiments, such as NMR and X-ray crystallography, often require appreciable populations to detect a substrate, which might make entanglement detection difficult. Many biophysical techniques used in protein folding (such as fluorescence resonance energy transfer, tryptophan fluorescence and circular dichroism) are sensitive to subpopulations, but tend to have low structural resolution, which would limit their capacity to distinguish near-native conformations. Therefore, hydrogen–deuterium exchange mass spectrometry and LiP-MS, which can localize conformational differences within a protein, combined with molecular modelling, seem promising in detecting signatures of entanglements. Cryogenic electron microscopy, with its ability to build structural classes from heterogeneous populations, might also be effective for systems of suitable size and resolution, although the increased flexibility of entangled regions may prove a challenge.

It is reasonable to expect that the enzymatic activity of initially misfolded proteins should increase with time as the protein relaxes to its native state. Based on the disentangling timescale of 10^4 s (see the section 'Deep entanglements are usually long lived'), we estimate that it would take more than 3 h for the specific activities between fast and slow variants to converge. This is consistent with what was found for CAT-III, where the specific activity of synonymous mutant CAT-III did not converge to that of the wild type within 20 min. For DDLB, because the LiP-MS results confirmed that the misfolded states can persist for longer than 2 h, it is reasonable to anticipate that the specific activities between fast and slow DDLB variants will take more than 2 h to converge, which is also consistent with our prediction. Further experiments, such as a time-dependent activity assay coupled with pulse–chase protein expression and production, could be applied to measure such timescales.

This study provides a plausible explanation of how synonymous mutations can alter enzyme activity in cells. Synonymous mutations

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alter translation elongation speeds and change the population of nascent chain conformations in entangled states that are near native but have lower catalytic efficiencies than that of the native state. Hence, the specific activity—a quantity averaged over the populations of proteins in different conformational states—can increase or decrease due to synonymous mutations. The experimental search for these entangled structures and their roles in influencing protein structure, function and phenotypes in cells is likely to be a fruitful area of research in the future.

**Online content**

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41557-022-01091-z.

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Methods

Coarse-grained simulation model
To model transitions between the native state and the unfolded state, we utilized a Gō-based coarse-grained model for all of the proteins studied. Briefly, this coarse-grained model represents each residue as a single interaction site centred on the Cα position and uses a structure-based potential energy function (see Supplementary Equation (1)). The force field parameters were optimized by training the parameters to reproduce the folding stability free energies of 18 small single-domain proteins, followed by assignment of the minimum values that can reproduce the structural stability for a given protein. To map the simulation timescale to an experimental timescale, we used the scaling factor α, which is the ratio of bulk experimental folding times to simulated folding times (see Supplementary Table 4). The high-resolution crystal structure PDB 4V9D was used to coarse-grain the E. coli ribosome, with the A- and P-site transfer RNA (tRNA) molecules modelled on the PDB structure 5JTE. The entire SOS subunit, including the A- and P-site tRNAs, was coarse-grained using the three-/four-point model of ribosomal RNA and the Cα model for ribosomal protein and then truncated to include only those coarse-grained interaction sites within 30 Å of the centre line of the exit tunnel and within 20 Å of the peptidyl transferase centre (identified as A2602 in the 23S ribosomal RNA of the E. coli ribosome), as well as the interaction sites at the ribosome surface near the exit tunnel opening. This resulted in 4,577 interaction sites for the cropped coarse-grained E. coli ribosome.

Protein synthesis on the ribosome was modelled using a continuous synthesis protocol that describes A-site tRNA binding, peptidyl transfer, tRNA translocation and ribosome trafficking at each nascent chain length using codon translation rates obtained from the model of Fluitt at al. (see Supplementary Equations (11) and (13)). Post-translational dynamics were modelled by simulating the nascent protein in the absence of the ribosome. Simulations for both co- and post-translational folding were performed via Langevin dynamics with a collision frequency of 0.05 ps⁻¹ and a time step of 15 fs using the relevant databases EzCatDB, UniProt and RCSB PDB. To identify candidate enzymes that may have kinetic traps, we created a dataset of well-characterized monomeric E. coli enzymes by searching the relevant databases EzCatDB, UniProt and RCSB PDB. The wild-type enzymes were then parameterized and their synthesis and post-translational dynamics were simulated with a 14-d wall time. The candidates were selected using the scoring function

\[
\text{Score} = \frac{1}{2} \times \left[ \left( 1 - \left( 1 + \frac{t_{\text{post}}}{t_{\text{eq}}} \right)^{-0.2} \right) + \Theta_{\text{binding}} \right] \times 100. \tag{1}
\]

where \(t_{\text{post}}\) is the mean post-translational folding time calculated using the double-pathway kinetics scheme of Supplementary Equation (6) with no delay time (that is, \(t_1 = t_2 = 0\)) and \(\Theta_{\text{binding}}\) is an indicator of whether (equals 1) or not (equals 0) misfolding occurs at or near the substrate-binding site and persists to the end of the simulation. We assigned equal weights for these two terms because they are considered equally important in identifying long-lived kinetic traps that have perturbed enzymatic functions. A higher score indicates a higher possibility for the enzyme to have long-lived kinetic traps that have misfolding in the substrate-binding pocket during its post-translational folding dynamics. Further details of the screening can be found in Supplementary Methods Section 9.

Characterizing post-translational misfolded structures
The misfolded structures of the post-translational folding process were characterized using the metrics \(G\) and \(Q_{\text{act}}\). \(G\) is an order parameter that measures the extent to which there is a change of entanglement in a given structure compared with the native structure and is calculated as

\[
G = \frac{1}{N} \sum_{i,j} \Theta((i,j) \in \text{nc} \cap g(i,j) \neq g^{\text{native}}(i,j)), \tag{2}
\]

where \((i,j)\) is one of the native contacts in the native crystal structure, \(\text{nc}\) is the set of native contacts formed in the current structure, \(g(i,j)\) and \(g^{\text{native}}(i,j)\) are, respectively, the total linking number of the native contact \((i,j)\) in the current and native structures estimated using Supplementary Equation (16) (see Supplementary Methods Section 11 for details). \(N\) is the total number of native contacts within the native structure, and the selection function \(\Theta\) equals 1 when the condition is true and 0 when it is false. The larger \(G\), the larger the number of residues that have changed their entanglement status relative to the native state. That is, \(G\) reports on the presence of non-native entanglements in structures.

\[Q_{\text{act}} = \frac{\sum_{i,j} (\Theta_{\text{Current}}(i,j) - \Theta_{\text{Native}}(i,j))}{\sum_{i,j} \Theta_{\text{Current}}(i,j)}\]  \tag{3}

where \(i\) and \(j\) are the residue indices and satisfy \(j > i + 3\), \(I\) is the intersection set of residues within secondary structure elements (α helical or β strands) and the substrate-binding pocket, \(J\) is the set of all residues within secondary structure elements and \(\Theta_{\text{Current}}(i,j)\) and \(\Theta_{\text{Native}}(i,j)\) are step functions that equal 1 when residues \(i\) and \(j\) have native contact and 0 when \(i\) and \(j\) do not have native contact in the current structure and native structure, respectively. Native contacts are considered formed when the distance between the Cα atoms of residues \(i\) and \(j\) does not exceed 1.2 times their native distance and the native distance does not exceed 8 Å. In the case of CAT-III, residue set \(I\) also includes native contacts in the trimer interface region (residues 25–33 and 150–157) to monitor the folding of the trimer interface as well. Note that the native contacts used in calculating \(Q_{\text{act}}\) are restricted to those within secondary structure elements, while the entire set of native contacts is used to calculate \(G\). Details of the assessment procedure can be found in Supplementary Methods Section 12.

Specific activity estimation
For each metastable state \(i\), identified using the Markov state modelling procedure reported in Supplementary Methods Section 12, we randomly selected five conformations from all of the microstates (based on the probability distribution of the microstates within the metastable state) and back-mapped them to all-atom structures (see Supplementary Methods Section 13). We then used QM/MM simulations (see Supplementary Methods Section 14) to calculate the transition state barrier for each of the five conformations, and from these, we determined the median activation barrier height \(\Delta G^\ddagger\). Only the metastable states that formed a near-native active site (\(Q_{\text{act}} \geq 0.6\)), as well as the native state, were taken to estimate \(\Delta G^\ddagger\), while the others were considered to have an infinite barrier height (zero reaction rate). Assuming the rate constants of each metastable state have similar pre-exponential factors that can be treated as a constant, the specific activity for a protein can be estimated using the probability distribution of metastable states (state probability \(p\)) and the activation free energy barrier height \(\Delta G^\ddagger\) of each state as

\[\text{Specific activity} = \frac{A}{M} \sum_{i=1}^{N} \zeta_i \times e^{-\frac{\Delta G^\ddagger_i}{RT}}, \tag{4}\]

where \(m\) is the molecular weight of the enzyme, \(A\) is the pre-exponential factor allowing us to convert \(\Delta G^\ddagger\) to the reaction rate constant, \(N\) is the
total number of metastable states, \( R \) is the gas constant and \( T \) is the temperature. In many codon usage studies,\(^1\)\(^2\), the relative specific activity is often used to compare the enzymatic activities of a protein produced by synonymous mRNA variants. In this study, the maximum specific activity among the fast and slow synonymous mRNA variants of an enzyme was used for normalization:

\[
\begin{align*}
SA_{fast}^* &= \frac{SA_{fast}}{\max(SA_{fast},SA_{slow})} \\
SA_{slow}^* &= \frac{SA_{slow}}{\max(SA_{fast},SA_{slow})}
\end{align*}
\]

where \( SA^\ast \) is the relative specific activity under saturating conditions. Note that the pre-exponential factor \( A \) and the molecular weight \( m \) cancel out when the specific activity is normalized; therefore, they do not need to be estimated. The details of estimating the state probability \( p_i \) (accounting for the soluble fraction only) and the activation free energy barrier height (\( \Delta G^\circ \)) are presented in Supplementary Methods Section 14. The materials and experimental methods for measuring the specific activities of the DDLB and DHFR variants are presented in Supplementary Methods Sections 19 and 20, respectively.

**Reporting summary**
Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

**Data availability**
Data supporting the main findings of this study are available within the Article and its Supplementary Information and source data files. We cannot feasibly provide all ~5.3 TB of molecular dynamics trajectory data, but we provide the input data that were used to perform the simulations in this study in the repository subdirectory https://github.com/obrien-lab/cg_simtk_protein_folding/blob/master/example/input_data.tar.xz. All of the data that support the findings of this study, as well as the biological materials that were used to test the enzymatic activity of the DDLB and DHFR variants and for the Lip-MS experiments, are available from the corresponding author upon reasonable request. The raw mass spectrometry data for DDLB and DHFR have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD031425. A website (https://obrien-lab.github.io/visualize_entanglements/) was created to provide interactive visualization of the key misfolded, entangled structures predicted in this study. Source data are provided with this paper.

**Code availability**
All of the computer code developed in this work is available in the GitHub repositories https://github.com/obrien-lab/cg_simtk_protein_folding and https://github.com/obrien-lab/Activation-Energy-Estimation-Workflow under the MIT License. Detailed instructions on code usage, basic theory and examples of the input/output are available in the wiki pages of the above repositories.

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**Author contributions**
E.P.O. designed the research. Y.J. developed the computational methods with contributions from E.P.O. Y.J. wrote the computer code and carried out the simulations and computations. S.S.N. and S.J.B. designed the experimental validation for the DDLB variants. S.S.N., I.S., E.P.O. and S.J.B. designed the experimental validation for the DHFR variants. S.S.N., I.S. and P.P. performed the specific activity experiments. S.D.F. designed the Lip-MS experiments for DDLB and DHFR. P.T. and Y.X. performed the Lip-MS experiments. All of the authors analyzed the data and wrote the manuscript.

**Competing interests**
The authors declare no competing interests.
Additional information
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Software and code

Policy information about availability of computer code

Data collection

Simulation and modeling: All computer code developed in this work is available in the GitHub repositories https://github.com/obrien-lab/cg_simtk_protein_folding and https://github.com/obrien-lab/Activation-Energy-Estimation-Workflow, under the MIT license. MD simulations were performed via OpenMM v7.4.1 and Amber 17.

Enzyme activity measurements: UV-visible spectra were recorded on a Cary 60 spectrometer from Varian (Agilent Technologies, Santa Clara, CA) using the WinUV software package to control the instrument.

Limited Proteolysis experiments: No software was used in data collection. A Thermo Q-Exactive HF-x Orbitrap mass spectrometer was used to analyze protein digests.

Data analysis

Simulation and modeling: Computer code developed in this work for analyzing the simulation results is available in the GitHub repositories https://github.com/obrien-lab/cg_simtk_protein_folding and https://github.com/obrien-lab/Activation-Energy-Estimation-Workflow, under the MIT license. Visual Molecular Dynamics v1.9.1 was used for molecular/trajecotry visualization and image/movie generation. Python v3.7, PyEMMA v2.5.7, ParmEd v3.4.1, Scipy, Numpy, Stride, Mfitral v1.9.7, KrotPul v0.4.1, WHAM v2.0.11, and ChaperonIS v1.0 were also used for analyzing simulation data.

Enzyme activity measurements: Data were analyzed by using Python v3.7.

Limited Proteolysis experiments: Proteome Discoverer (PD) Software Suite v2.4, Thermo Fisher) and the Minora Algorithm were used to analyze mass spectra and perform Label Free Quantification (LFQ) of detected peptides. PD output files were outputted in a three-level hierarchy (protein > peptide group > consensus feature) and were further processed utilizing custom Python analyzer scripts that are available on GitHub: https://github.com/FriedLabHU/Refoldibility-Tools/.

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- A list of figures that have associated raw data
- A description of any restrictions on data availability

Raw data for Figures 1 to 6 are provided. We cannot feasibly provide all ~5.3 TB of molecular dynamics trajectory data, but we provide the input data that was used to perform the simulations in the repository subdirectory https://github.com/obrien-lab/gg_simtk_protein_folding/blob/master/example/input_data.tar.gz. All the data that support the findings of this study, as well as the biological materials that were used for testing the enzymatic activity of DDLB and DHFR variants and for UPL-MS experiments, are available from the corresponding author upon reasonable request. The raw mass spectrometry data of DDLB and DHFR have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD014252. The PDB structures 3CLA, 4CSC, 1AQ2, 2ZCV, 1AJK, 1VHL, 3KDL, 3LFB, 2FMT, 1C2T, 1FDR, 3CV7, 20FP, 4KJK, 1PDA, 1RYK, 1LMB, 2ABD, 1CIE, 1MQ, 256B, 1L08, 1SHF, 1CO9, 1MUC, 1TE, 1WIT, 1POH, 2QPL, 1SPR, 1E05, 3CHY, 2RNZ, 5NYV, 5JTE, 6IDY, 3I8U, 4L8Y, 6ENJ, 6ENJ, 5LUB, 4V9D were used in this study. The databases ExCatDB (https://excatdb.cbr.cpn.ast.go.jp/exCatDB/), UnProt (https://www.uniprot.org/), RCSB PDB (https://www.rcsb.org/) and CATH (http://www.cathdb.info/) were used in this study. A website [https://obrien-lab.github.io/visualize_entanglements/] was created to provide interactive visualization of the key misfolded entangled structures predicted in this study.

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

- [x] Life sciences
- [ ] Behavioural & social sciences
- [ ] Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-flat.pdf

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size
No sample size calculation was conducted prior. The specific activity measurements of DDLB and DHFR were performed on five biological replicates, respectively. Each Limited Proteolysis experiment was conducted as biological triplicates to allow for statistical analysis. The numbers of simulation trajectories were chosen due to the limitations of our computational resources.

Data exclusions
No data were excluded.

Replication
The experiments for DDLB have been done for five biological replicates and all of them have the same trend of the enzymatic activity, which indicates the robustness and reproducibility of our findings. The experiments for DHFR have been done for five biological replicates as well and exhibit no statistical difference in the specific activities of fast and slow variants. Limited Proteolysis experiments were conducted on two separate experiments. Experimental findings were consistent in both experiments. Simulations were replicated 50 (DHFR) and 100 (DDLB and CAT-III) times for co-translational folding, 500 (DHFR) and 1,000 (DDLB and CAT-III) times for post-translational folding, 30 times for disentangling timescale estimation and 100 times for DDLB refolding simulations, with different random seeds to allow for the calculation of statistics. Due to different random seeds, behavior is different between trajectories. All trajectories ran to completion. All attempts at replication were successful.

Randomization
E. coli lysates were divided into either being either a native or a refolded sample depending on if they were unfolded and refolded or not. Covariates are not relevant in our study as all bacteria are cultured simultaneously under identical growth conditions and simultaneously prepared under identical methods. Random allocation is not relevant to the other experiments/simulations because they were not grouped to get treatments.

Blinding
Investigators know which E. coli lysates are native samples and which ones are refolded samples as limited proteolysis experiments are conducted after different refolding time points. Our analysis compares the limited proteolysis peptide profile of our refolded samples to our native samples, so blinding is not possible. Blinding is also not possible to the other experiments/simulations for the same reason.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.
### Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ☒ | Antibodies |
| ☒ | Eukaryotic cell lines |
| ☒ | Palaeontology and archaeology |
| ☒ | Animals and other organisms |
| ☒ | Human research participants |
| ☒ | Clinical data |
| ☒ | Dual use research of concern |

### Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ☒ | ChiP-seq |
| ☒ | Flow cytometry |
| ☒ | MRI-based neuroimaging |

### Antibodies

**Antibodies used**

Antibodies used in the Time-course analysis of DDLB expression:
- Primary antibody: Mouse anti-His6 IgG, Clone: His HB, Company: Invitrogen, catalog number: MA1-21315
- Secondary antibody: Goat anti-mouse-AP, Company: Millipore, Catalog number: 69266-3

**Validation**

[https://assets.thermofisher.com/IFS-Assets/LSG/flyers/commitment-antibody-performance-flyer.pdf](https://assets.thermofisher.com/IFS-Assets/LSG/flyers/commitment-antibody-performance-flyer.pdf)