Loss of stability and unfolding cooperativity in hPGK1 upon gradual structural perturbation of its N-terminal domain hydrophobic core

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Phosphoglycerate kinase has been a model for the stability, folding cooperativity and catalysis of a two-domain protein. The human isoform 1 (hPGK1) is associated with cancer development and rare genetic diseases that affect several of its features. To investigate how mutations affect hPGK1 folding landscape and interaction networks, we have introduced mutations at a buried site in the N-terminal domain (F25 mutants) that either created cavities (F25L, F25V, F25A), enhanced conformational entropy (F25G) or introduced structural strain (F25W) and evaluated their effects using biophysical experimental and theoretical methods. All F25 mutants folded well, but showed reduced unfolding cooperativity, kinetic stability and altered activation energetics according to the results from thermal and chemical denaturation analyses. These alterations correlated well with the structural perturbation caused by mutations in the N-terminal domain and the destabilization caused in the interdomain interface as revealed by H/D exchange under native conditions. Importantly, experimental and theoretical analyses showed that these effects are significant even when the perturbation is mild and local. Our approach will be useful to establish the molecular basis of hPGK1 genotype–phenotype correlations due to phosphorylation events and single amino acid substitutions associated with disease.
Phosphoglycerate kinase (PGK) is a glycolytic enzyme that catalyzes the reversible phosphotransfer reaction between ATP and 3-phosphoglycerate (3-PG) to form ADP and 1,3 bisphosphoglycerate (1,3-BPG) in the presence of Mg\(^{2+}\). Its main activity is to generate ATP in glycolysis, and this activity is highly conserved among all living organisms. Structurally, PGK is typically a monomeric protein of about 45 kDa with two well-defined domains: an N-terminal domain (NTD) that binds 3-PG/1,3-BPG (usually the first 200 amino acids), a C-terminal domain (CTD) (typically the last 200 amino acids) that binds the nucleotide and both domains are linked by a 10–15 amino acids long \(\alpha\)-helix (Fig. 1). Binding of Mg\(^{2+}\) triggers the transition between the inactive open conformation to a closed conformation that leads to catalysis. Importantly, PGKs from different organisms have been crystalized in different ligation states (only for the human isoform 1, hPGK1, we retrieved 16 different structures by February 2022 from the Protein Data Bank; Table S1), and metal binding may trigger long-range cooperative communication between domains (Fig. 1). These facts make PGK a highly suitable model to investigate allosteric communication upon ligand binding (or mutations). 

Phosphoglycerate kinase (PGK) is a glycolytic enzyme that catalyzes the reversible phosphotransfer reaction between ATP and 3-phosphoglycerate (3-PG) to form ADP and 1,3 bisphosphoglycerate (1,3-BPG) in the presence of Mg\(^{2+}\). Its main activity is to generate ATP in glycolysis, and this activity is highly conserved among all living organisms. Structurally, PGK is typically a monomeric protein of about 45 kDa with two well-defined domains: an N-terminal domain (NTD) that binds 3-PG/1,3-BPG (usually the first 200 amino acids), a C-terminal domain (CTD) (typically the last 200 amino acids) that binds the nucleotide and both domains are linked by a 10–15 amino acids long \(\alpha\)-helix (Fig. 1). Binding of Mg\(^{2+}\) triggers the transition between the inactive open conformation to a closed conformation that leads to catalysis. Importantly, PGKs from different organisms have been crystalized in different ligation states (only for the human isoform 1, hPGK1, we retrieved 16 different structures by February 2022 from the Protein Data Bank; Table S1), and metal binding may trigger long-range cooperative communication between domains (Fig. 1). These facts make PGK a highly suitable model to investigate allosteric communication upon ligand binding (or mutations). 

Particularly in the last decade, research on hPGK1 has attracted additional attention for several reasons: (1) hPGK1 is multifunctional (a so-called moonlighting protein). As we have previously mentioned, it is essential for glycolysis, but it also phosphorylates other proteins, is involved in DNA replication and repair, activates NAD+-nucleoside analogues used in anticancer and antiviral activities, and plays a key role in autophagy. (2) hPGK1 can operate in different cellular compartments. Obviously, due to its role in glycolysis, it is found at high levels in the cytosol. However, due to its multifunctionality, it can also localize in the nucleus, mitochondria and extracellularly. These findings suggest that non-native states in the energy landscape of hPGK1 may play important roles in the transport to different compartments. For instance, to be imported to mitochondria, proteins are unfolded by the mitochondrial import machinery or even presented to this machinery in a partially unfolded state by Hsp70 chaperones. In the particular case of hPGK1, the interaction with Hsc70 and Hsp70 chaperones inside the cell is likely, thus suggesting a role of partially (un)folded states of hPGK1 in mitochondrial import of this enzyme (this import is associated with cancer metabolism and tumorigenesis). (3) Alterations in hPGK1 activity, intracellular location and stability are associated with several diseases, including cancer, hemolytic anemia, myopathy and neurological dysfunction (possibly including amyotrophic lateral sclerosis). In most cases, a deficiency in hPGK1 associated with disease is concomitant with a modification of the protein sequence by mutations. These mutations often reduce catalytic efficiency, protein stability and unfolding cooperativity, and increase aggregation propensity. In several cases, a single mutation manifests this loss of function through several of these detrimental mechanisms, and mutations associated with cancer development or genetic hPGK1 deficiency (OMIM 300653) behave similarly regarding these mechanisms. Therefore, it is plausible that missense mutations may propagate their destabilizing effect across the structure of hPGK1 affecting different functional features such as activity, stability, or folding cooperativity. In this work, we analyzed the wild-type (WT) protein as well as five mutants aimed to progressively change the magnitude of the local destabilizing effect at a single site. To do so, we mutated a fully buried and large hydrophobic residue (F25; see Fig. 1 and Table S1) to the smaller residues (L, V, A and G) expected to create cavities of different size in the hydrophobic core of the NTD, increase conformational entropy (mutation to G) as well as a mutation that increases the size of the side-chain (mutation to W) expected to generate structural strain. These mutations are very suitable for this type of study (particularly cavity-creating mutations) because their effects on protein stability and energy landscapes have been extensively characterized by experiments and computation using typically smaller and simpler proteins. In addition, although F25 is not found mutated in the ClinVar (https://www.
ncbi.nlm.nih.gov/clinvar/) or COSMIC (https://cancer.sanger.ac.uk/cosmic) databases, a significant fraction of the mutations compiled in these databases can be classified as affecting buried positions and expected to create cavities, affecting glycine residues (i.e. altering the backbone flexibility) or introducing conformational strain (Fig. 1C and Table S2). Our approach uses a combination of experimental biophysical techniques including optical spectroscopy, calorimetry, thermal and chemical denaturation and hydrogen/deuterium exchange monitored by mass spectrometry (HDX-MS) with statistical mechanical calculations.

Results and discussion

F25 mutations do not impair protein expression of hPGK1 as a folded protein. Upon expression in E. coli and purification with two chromatographic steps, IMAC (Immolized-Metal Affinity Chromatography) and SEC (Size-Exclusion Chromatography) (Fig. 2A) we obtained hPGK1 proteins of a very high purity and also confirmed that all variants showed similar hydrodynamic behavior to that of the WT protein (i.e. monomeric; the elution volume was very similar to that of ovalbumin, with a MW of 44 kDa). Since hPGK1 is often purified with certain small molecules bound to the protein (possibly nucleotides or small polynucleotides, based on the UV–visible absorption and Circular Dichroism (CD) spectra and consistent with the ability of the enzyme to interact with such type of molecules, e.g. ADP and ATP), we therefore included a step of precipitation with streptomycin between the IMAC and the SEC. After this step, protein samples contained virtually no additional signals in the near-UV range (Fig. 2B).

Further analyses were carried out using CD to test the effect of mutations on the conformation of the protein. Far-UV CD spectra revealed the typical spectra for a protein rich in α-helix such as hPGK1 (Fig. 2C) and closely resembled results from previous studies. We must note that the far-UV CD intensity was slightly lower for the mutant F25G, suggesting that this mutation could slightly distort the overall secondary structure of the protein. Mutants F25G and F25W also reduced the strongest signal in the near-UV CD spectra (centered at about 290 nm; consistent with the dichroic signal from Trp residues) suggesting local distortions of the tertiary structure (Fig. 2D).

These analyses support that the F25 mutants largely maintained the overall structure of the WT protein.

Thermal stability of hPGK1 variants. Thermal denaturation of hPGK1 has been described well by using a simple two-state irreversible denaturation model. The origin of this irreversibility is likely protein aggregation, as indicated by the similar profiles obtained by DSC and light scattering measurements. To initially assess the effect of mutations on hPGK1 thermal stability, we acquired the CD signal varying the temperature from 20 to 75 °C. Both far- and near-UV CD monitored thermal denaturation experiments provided similar results (Fig. 3). The apparent $T_m$ decreased according to the size of cavity (from 2–3 °C in F25L to 6–8 °C in F25G,
compared with those values of WT), whereas F25W (that may introduce certain conformational strain) showed an apparent \( T_m \) very similar to that of F25A.

To provide deeper knowledge of the effects of F25 mutants on the kinetic stability (i.e. the rate constant for irreversible denaturation, \( k \) and activation energy, \( E_a \)) of hPGK1, we applied DSC using previously described protocols for this protein\(^3\). The results from these analyses are summarized in Fig. 4 and Tables 1 and 2. In agreement with thermal denaturation monitored by CD, we observed a progressive decrease in \( T_m \) in the cavity-making
mutants (F25G decreases the \( T_m \) by \( \sim 7 \) °C). The mutant F25W showed again a similar behavior to that of F25A, with a decrease of \( \sim 4.5 \) °C. The destabilization induced in the thermal denaturation process by each individual mutation using different probes (DSC, far-UV and near-UV CD spectroscopies) are very close at similar scan rates (Figs. 3 and S1–S2).

It is also interesting to note that the decrease in thermal stability correlates strongly with the \( E_a \) values for irreversible denaturation and the \( T_m \) values as determined by DSC (Fig. 4B–D and Tables 1 and 2). An overall linear fitting of \( E_a \) versus \( T_m \) provides a value for the change in activation heat capacity (\( \Delta C_p^{≠} \)) of 10.6 ± 0.5 kcal mol\(^{-1}\) in a very good agreement with our previous analyses that yielded a value of 9.1 ± 0.8 kcal mol\(^{-1}\) (Fig. 4D). Notably, the mutation F25G decreased the \( E_a \) value by 75 kcal mol\(^{-1}\) (basically to a half of that for the WT protein).

**Figure 4.** Thermal stability of hPGK1 variants by DSC. (A) Normalized DSC scans (upon subtraction of chemical baselines) for all variants at 10 µM protein concentration and 4 °C min\(^{-1}\). Lines are best-fits to a two-state irreversible model; (B) Arrhenius plot for all variants, derived from experiments at three different scan rates (1, 2.5 and 4 °C min\(^{-1}\)). The \( k \) values were determined using Eq. (4). (C) Average ± s.d. for activation energies (\( E_a \) values) for the irreversible denaturation of hPGK1 variants. These values are derived from Arrhenius plots and are very consistent with those obtained from individual fittings (see Table 1). (D) \( E_a \) values derived from individual fittings at different scan rates (black) and from Arrhenius plots (red) are very consistent and linearly depend on the average \( T_m \) from individual fittings (x-axis). Errors for fittings are S.D. for different experiments whereas those for Arrhenius are those from linear fittings (see panel B). Experiments were carried out in 20 mM HEPES–NaOH 200 mM NaCl, pH 7.4. All the analysis were carried out following 29,41.

| Variant | Parameter | \( E_a \) (kcal·mol\(^{-1}\)) | \( \Delta E_a \) (kcal·mol\(^{-1}\)) | \( k_{37°C} \) (min\(^{-1}\)) | \( \Delta \Delta G^{≠} \) \( 37°C \) (var-WT) (kcal·mol\(^{-1}\)) |
|---------|-----------|----------------------------|-----------------------------|----------------------------|---------------------------------|
| WT      | 168.0 ± 3.3 | 0.0 ± 4.7                  | 1.8 ± 0.4·10\(^{-4}\)       | 0.00 ± 0.19                |
| F25L    | 149.3 ± 3.1 | −18.7 ± 4.4               | 2.9 ± 0.6·10\(^{-3}\)       | −1.71 ± 0.19               |
| F25V    | 136.9 ± 2.7 | −31.1 ± 3.8               | 1.5 ± 0.2·10\(^{-4}\)       | −2.73 ± 0.16               |
| F25A    | 114.3 ± 2.1 | −53.7 ± 3.0               | 1.3 ± 0.1·10\(^{-3}\)       | −4.06 ± 0.14               |
| F25G    | 92.9 ± 1.2  | −75.1 ± 3.5               | 1.1 ± 0.1·10\(^{-3}\)       | −5.37 ± 0.15               |
| F25W    | 117.8 ± 2.1 | −50.2 ± 3.0               | 9.2 ± 1.0·10\(^{-3}\)       | −3.84 ± 0.15               |

Table 1. Kinetic parameters derived from Arrhenius plots calculated from DSC data in the absence of urea. \( k_{37°C} \) was obtained using 37°C (310.15 K) as reference temperature \((T^*)\) in Eq. (2).
These results have remarkable implications for the kinetic stability of hPGK1 variants in vitro, since the decrease in $T_m$ values correlated with the decrease in $E_a$ values, and thus, the extrapolated values of the kinetic rate constants at 37 °C (and therefore, the associated half-lives for irreversible denaturation) vary strongly among variants. We observed a gradual decrease of the half-life for irreversible denaturation correlating with the size of the cavity, from a small ~15-fold in F25L to a large ~6000-fold in F25G. Again, F25A and F25W showed similarly low kinetic stabilities (about 600-fold lower than that of the WT protein; extrapolated to 37 °C from Arrhenius plots) (Table 1).

It is worth noting that previous studies have shown that mutations associated with hPGK1 deficiency (a rare X-linked genetic disorder3,30) also showed different effects on kinetic stability when extrapolated to 37 °C, up to 5 orders of magnitude lower than that of the WT protein; extrapolated to 37 °C from Arrhenius plots (Table 1).

To further investigate the correlation found between the changes in $T_m$ and $E_a$ values, we have carried out DSC experiments in the presence of urea (Table 2 and Figs. S1–S2). These analyses allow to extract the kinetic $m^*$ value for irreversible denaturation from detailed DSC analyses at different scan rates and urea concentrations29,31,41. Comparing this value with the theoretical $m^*$ value derived from the protein size (i.e. in number of amino acids), we can estimate the degree of the native hPGK1 structure which is denatured in the transition state for irreversible thermal denaturation (Table 3). Actually, these analyses revealed that, in contrast to hPGK1-deficiency causing mutants characterized so far29, only the F25G showed a small decrease in value for the $m^*$ (Table 3). Thus, the Hammond effect (i.e. that the kinetic stability decreases as the transition state becomes more native-like) proposed for the hPGK1-deficiency causing mutants previously characterized29 does not seem to follow the same behavior that those of the F25 mutants characterized in the present work (Table 3).

| Variant | Urea (M) | $T_m$ (°C) | $\Delta H$ (kcal·mol$^{-1}$) | $E_a$ (kcal·mol$^{-1}$) |
|---------|---------|------------|----------------------------|----------------------------|
| WT      | 0       | 53.5 ± 0.7 | 162 ± 10                   | 168 ± 15                   |
|         | 0.5     | 51.2 ± 0.7 | 142 ± 12                   | 140 ± 7                    |
|         | 1       | 48.7 ± 0.9 | 136 ± 13                   | 115 ± 4                    |
| F25L    | 0       | 51.6 ± 0.6 | 156 ± 6                    | 150 ± 12                   |
|         | 0.5     | 49.2 ± 0.7 | 133 ± 6                    | 129 ± 4                    |
|         | 1       | 46.4 ± 0.9 | 126 ± 16                   | 105 ± 1                    |
| F25V    | 0       | 50.4 ± 0.7 | 153 ± 11                   | 140 ± 12                   |
|         | 0.5     | 47.5 ± 0.9 | 133 ± 14                   | 106 ± 5                    |
|         | 1.1     | 44.3 ± 1.3 | 114 ± 22                   | 80 ± 2                     |
| F25V    | 0       | 48.9 ± 0.8 | 153 ± 14                   | 119 ± 9                    |
|         | 0.5     | 45.8 ± 1.1 | 124 ± 16                   | 89 ± 3                     |
|         | 1       | 42.8 ± 1.7 | 109 ± 18                   | 68 ± 3                     |
| F25G    | 0       | 46.7 ± 0.9 | 155 ± 15                   | 94 ± 3                     |
|         | 0.5     | 43.5 ± 1.6 | 123 ± 11                   | 71 ± 4                     |
|         | 1       | 41.1 ± 1.9 | 108 ± 19                   | 59 ± 5                     |
| F25W    | 0       | 49.2 ± 0.7 | 140 ± 17                   | 120 ± 6                    |
|         | 0.5     | 46.3 ± 0.9 | 120 ± 18                   | 93 ± 2                     |
|         | 1       | 43.0 ± 1.5 | 96 ± 16                    | 72 ± 5                     |

Table 2. Energetic parameters for the denaturation of hPGK1 variants determined from DSC data in the absence or the presence of urea. Data are the mean ± s.d. for three different scan rates (4, 2.5 and 1 °C min$^{-1}$) at 7–15 µM protein concentration and using a two-state irreversible model.

| Variant | $m^*$ (kcal·mol$^{-1}$·M$^{-1}$) | $m^*/m^*_{\text{unf}}$ |
|---------|---------------------------------|------------------------|
| WT      | 2.69 ± 0.36                     | 0.59 ± 0.08            |
| F25L    | 2.47 ± 0.34                     | 0.54 ± 0.07            |
| F25V    | 2.74 ± 0.51                     | 0.60 ± 0.11            |
| F25A    | 2.60 ± 0.60                     | 0.57 ± 0.13            |
| F25G    | 1.93 ± 0.42                     | 0.43 ± 0.09            |
| F25W    | 2.63 ± 0.48                     | 0.58 ± 0.11            |

Table 3. Degree of structural unfolding in the transition state for the irreversible denaturation of hPGK1 variants determined by DSC. $m^*$ values were determined from DSC experiments and Eq. (5). $m^*_{\text{unf}}$ is the theoretical value (4.54 kcal mol$^{-1}$ M$^{-1}$) for the full unfolding of the protein39 based on a protein size of 417 residues42.

These results have remarkable implications for the kinetic stability of hPGK1 variants in vitro, since the decrease in $T_m$ values correlated with the decrease in $E_a$ values, and thus, the extrapolated values of the kinetic rate constants at 37 °C (and therefore, the associated half-lives for irreversible denaturation) vary strongly among variants. We observed a gradual decrease of the half-life for irreversible denaturation correlating with the size of the cavity, from a small ~15-fold in F25L to a large ~6000-fold in F25G. Again, F25A and F25W showed similarly low kinetic stabilities (about 600-fold lower than that of the WT protein; extrapolated to 37 °C from Arrhenius plots) (Table 1).

It is worth noting that previous studies have shown that mutations associated with hPGK1 deficiency (a rare X-linked genetic disorder3,30) also showed different effects on kinetic stability when extrapolated to 37 °C, up to 5 orders of magnitude lower than that of the WT protein in some cases. Interestingly, mutations found in COSMIC database also showed reduced stability against thermal and chemical denaturation39.

To further investigate the correlation found between the changes in $T_m$ and $E_a$ values, we have carried out DSC experiments in the presence of urea (Table 2 and Figs. S1–S2). These analyses allow to extract the kinetic $m^*$ value for irreversible denaturation from detailed DSC analyses at different scan rates and urea concentrations29,31,41. Comparing this value with the theoretical $m^*$ value derived from the protein size (i.e. in number of amino acids), we can estimate the degree of the native hPGK1 structure which is denatured in the transition state for irreversible thermal denaturation (Table 3). Actually, these analyses revealed that, in contrast to hPGK1-deficiency causing mutants characterized so far29, only the F25G showed a small decrease in value for the $m^*$ (Table 3). Thus, the Hammond effect (i.e. that the kinetic stability decreases as the transition state becomes more native-like) proposed for the hPGK1-deficiency causing mutants previously characterized29 does not seem to follow the same behavior that those of the F25 mutants characterized in the present work (Table 3).
Figure 5. Urea-induced denaturation of hPGK1 variants at 25 °C. 5 µM of hPGK1 variants were incubated for at least 2 h in the presence of increasing urea concentrations in HEPES–NaOH 20 mM NaCl 200 mM 1 mM β-ME, pH 7.4. Then the far-UV CD spectra were acquired and appropriate blanks subtracted. Data are from two independent experiments, and in each experiment the sample without urea was triplicated and averaged. Lines are best-fits to an apparent two-state equilibrium denaturation model (Eq. 6).

Table 4. Apparent denaturation equilibrium parameters with urea at 25 °C determined by Far-UV CD. $C_m$ and $m_{\text{urea}}$ values were retrieved from fittings of experiments shown in Fig. 5. For F25G, we could only obtained reasonable fittings when $C_m$ or $m_{\text{urea}}$ values were fixed, and thus some gross estimation was only possible.
Urea denaturation of F25 mutants reveals effects on both unfolding cooperativity and resistance to the urea-induced unfolding. An interesting feature of PGKs is that their chemical denaturation is generally reversible (e.g. using urea or guanidium chloride). Therefore, this type of analysis is amenable for comparing the effects of mutations on thermodynamic stability and unfolding cooperativity. Urea-denaturation of WT hPGK1 resembles quite well a two-state folder although the equilibrium value is somewhat lower than the expected for a protein of this size. Importantly, urea-induced denaturation of 14 mutants, including those found in cancer cell lines and hPGK1-deficient patients, shows reduced unfolding cooperativity (in some cases the $m_{\text{urea}}$ value was half of that of the WT protein). Interestingly, only in a few cases changes in the $C_m$ values were observed.

To compare the behavior of F25 mutants versus those of naturally-occurring and disease-associated, we carried out similar urea-induced unfolding experiments (Fig. 5). As previously described, denaturation of hPGK1 provides a value for the unfolding free energy of about 8 kcal mol$^{-1}$, based on the linear extrapolation method (Table 4). The F25 mutants showed changes in urea-induced unfolding compared to the WT protein. The mutants F25L and F25V, as well as the strain-inducing F25W, caused a significant decrease in the unfolding cooperativity and $C_m$ values (note that in this case F25W resembles F25V). Some mutants decreased the $m_{\text{urea}}$ value by ≤ 50% of the value for the WT protein (Table 4). The most destabilizing variants were F25A (for which we could obtain reasonable and physically meaningful values) and F25G (that showed a denaturation profile similar to that of F25A but we could not get reliable estimations of $C_m$ or $m_{\text{urea}}$, see Table 4). Therefore, for F25G and F25A, the

**Figure 6.** Conformational stability of WT and F25 mutants as determined by proteolysis with thermolysin. (A) Representative SDS-PAGE gels of hPGK1 variants proteolysis kinetics; (B) Densitometric analysis of results shown in panel A with fittings to a single exponential function (left panel) and the corresponding half-lives (right panel). Thermolysin concentration was 1 µM. Experiments were carried out in 20 mM HEPES–NaOH 200 mM NaCl, pH 7.4 at 25 °C.
unfolding cooperativity was severely reduced. Overall, these results suggest that increasing the cavity size or introducing conformational strain, at least at the F25 site in hPGK1, increase the population of non-native state conformations, possibly even under native conditions. We further tested this hypothesis by carrying out additional experimental and computational analyses under native conditions.

**Proteolysis suggests a mild increase in the population of partially folded states in the hPGK1 mutants under native conditions.** Proteolysis has been shown to provide information on the popula-

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**Figure 7.** Kinetics of deuterium (D) incorporation to hPGK1 WT. (A) % of D incorporation at different time points for the entire hPGK1 protein. Note that segments 13–17, 265–269 and 345–347 were not detected. (B) Selected segments displaying complex kinetics. Lines generally show best-fits to a double-exponential function including a burst phases. The best-fit parameters are compiled in Table 5. Experiments were carried out in 20 mM HEPES–NaOH, 200 mM NaCl, 0.5 mM TCEP, pH 7.4 at 25 °C.
particularly relevant in the domain-domain interface (Figs. 8 and S10, 11). F25W showed the strongest effects in stability and these propagated to regions further from the mutated sites, close to F25, while these effects were more extensive due to the mutation F25A (Fig. 8). Mutants F25G and certain regions in the protein (Figs. S5–S9). Mutants F25L and F25V showed mild stability decreases in regions that different conformational substates are populated in the native state ensemble of hPGK1.

be, at least, partially explained by the structural destabilization of the NTD: CTD interface.

F25V begins to perturb this interface, and mutants F25G and F25W have the largest effect, affecting 30–40% of F25 mutations on this interface from HDX-MS data (Fig. S11). These analyses showed that even the mutation C-terminal domains (NTD and CTD, respectively) (see Figs. S10 and S11). We next analyzed the effects of the by isothermal urea denaturation; Table 4) are likely associated with the interaction surface between the N- and protein segments is highly different (Fig. 7A). Second, the exchange kinetics are complex, and phenomenologically described in many cases by a double exponential function with a burst-phase. N. app.- not applicable.

Table 5. Best-fit parameters for selected segments of WT hPGK1 determined by HDX-MS. Good fittings were obtained using a two-exponential function (A∞ is the total amplitude of %D uptake at t = ∞, A1 and k1 are the amplitude and rate constant for the fast phase, while A2 and k2 are those for the slow phase). *The fitting to a double-exponential function did not converge. The values reported corresponded to a fitting using a one-exponential function with a burst-phase. N. app.- not applicable.

Perturbations in the native ensemble of F25 mutants monitored by HDX-MS. To provide experimental information on the effects of F25 mutations on the local stability of the native state ensemble, we carried out HDX-MS experiments. Analysis of the WT protein in a ligand-free conformation (Fig. 7) provided some interesting results. First, the apparent local stability based on the overall exchange rate for diverse protein segments is highly different (Fig. 7A). Second, the exchange kinetics are complex, and phenomenologically described in many cases by a double exponential function with a burst-phase (Fig. 7B and Table 5). This suggests that different conformational substates are populated in the native state ensemble of hPGK1.

Similar experiments were carried out with the F25 mutants. F25 mutations selectively affected the stability of certain regions in the protein (Figs. S5–S9). Mutants F25L and F25V showed mild stability decreases in regions close to F25, while these effects were more extensive due to the mutation F25A (Fig. 8). Mutants F25G and F25W showed the strongest effects in stability and these propagated to regions further from the mutated sites, particularly relevant in the domain-domain interface (Figs. 8 and S10, 11).

The kinetic stability (as determined by DSC; Tables 1, 2 and 3) and unfolding cooperativity (as determined by isothermal urea denaturation; Table 4) are likely associated with the interaction surface between the N- and C-terminal domains (NTD and CTD, respectively) (see Figs. S10 and S11). We next analyzed the effects of the F25 mutations on this interface from HDX-MS data (Fig. S11). These analyses showed that even the mutation F25V begins to perturb this interface, and mutants F25G and F25W have the largest effect, affecting 30–40% of the residues forming this interface (Fig. S11A) and these effects are of larger magnitude (Fig. S11B). These results strongly suggest that the effects of F25 mutations on conformational stability and unfolding cooperativity can be, at least, partially explained by the structural destabilization of the NTD: CTD interface.

Statistical mechanical analysis of the effects of F25 mutations. To further understand the extent to which mutations affect the different regions of the protein, we introduced mutations in silico (using PyMol) and studied three representative F25 mutants (F25V, F25A and F25G) via the WSME model. In each of the cases, no further modulation of structure was introduced as these are primarily truncating mutations. The mutated structures were fed into the WSME model and the free-energy profiles were predicted without changing any model parameters. The resulting ranking of destabilization matched the expectations from experiments, i.e. F25G > F25A > F25V (Fig. 9A) and also indicated that the folded state of the NTD and CTD is uncoupled to some extent in the native state ensemble of hPGK1 (intermediates I1 and I2, Fig. 9B). In addition, we calculated the positive coupling free energies (∆G+) for these variants, a parameter that reports on the degree of relative thermodynamic coupling between residues that are folded44. The effect of each of the mutations (mut) versus the WT protein results in the mean difference in coupling free energies (∆∆G+) or perturbation, defined as

| Parameter | Segment | A∞ (%D) | A1 (%D) | k1 (s⁻¹) | A2 (%D) | k2 (s⁻¹) | r² |
|-----------|---------|---------|---------|---------|---------|---------|----|
| 12        | 100.5 ± 1.8 | 45.1 ± 2.7 | 7.2 ± 1.1·10⁻³ | 48.9 ± 2.7 | 1.3 ± 0.2·10⁻⁴ | 0.9986 |
| 19–21     | 21.1 ± 0.7 | 19.6 ± 1.2 | 2.0 ± 0.4·10⁻⁴ | 4.2 ± 0.9 | 1.2 ± 0.7·10⁻⁴ | 0.9956 |
| 36        | 96.5 ± 0.3 | 16.9 ± 0.4 | 1.3 ± 0.1·10⁻² | 17.0 ± 0.3 | 1.2 ± 0.1·10⁻⁴ | 0.9998 |
| 89–92     | 96.6 ± 3.7 | 26.4 ± 4.7 | 6.1 ± 2.9·10⁻³ | 31.7 ± 5.0 | 7.5 ± 3.2·10⁻³ | 0.9858 |
| 142–145   | 98.0 ± 1.0 | 28.1 ± 2.0 | 1.4 ± 0.3·10⁻² | 9.2 ± 1.9 | 3.4 ± 1.7·10⁻⁴ | 0.9957 |
| 192–194*  | 97.8 ± 1.4 | 98.5 ± 1.5 | 6.5 ± 0.3·10⁻³ | N.app | N.app | 0.9990 |
| 213–216   | 76.4 ± 1.4 | 32.4 ± 1.9 | 5.8 ± 0.9·10⁻³ | 12.9 ± 2.0 | 9.3 ± 3.8·10⁻⁵ | 0.9968 |
| 244       | 32.2 ± 3.1 | 10.3 ± 0.5 | 8.1 ± 1.1·10⁻³ | 16.0 ± 3.0 | 1.8 ± 0.5·10⁻³ | 0.9986 |
| 296       | 94.9 ± 3.2 | 46.6 ± 4.7 | 5.7 ± 1.4·10⁻³ | 22.3 ± 4.8 | 1.2 ± 0.7·10⁻⁴ | 0.9931 |
| 348–353   | 48.7 ± 0.6 | 17.4 ± 1.1 | 1.0 ± 0.1·10⁻³ | 27.9 ± 1.0 | 4.8 ± 0.6·10⁻⁵ | 0.9997 |
| 381*      | 97.8 ± 0.8 | 41.3 ± 3.6 | 2.4 ± 0.5·10⁻² | N.app | N.app | 0.9836 |
| 404–405   | 96.5 ± 3.3 | 30.2 ± 8.0 | 3.6 ± 1.8·10⁻² | 57.8 ± 3.7 | 7.7 ± 1.4·10⁻³ | 0.9934 |

Statistical mechanical analysis of the effects of F25 mutations. To further understand the extent to which mutations affect the different regions of the protein, we introduced mutations in silico (using PyMol) and studied three representative F25 mutants (F25V, F25A and F25G) via the WSME model. In each of the cases, no further modulation of structure was introduced as these are primarily truncating mutations. The mutated structures were fed into the WSME model and the free-energy profiles were predicted without changing any model parameters. The resulting ranking of destabilization matched the expectations from experiments, i.e. F25G > F25A > F25V (Fig. 9A) and also indicated that the folded state of the NTD and CTD is uncoupled to some extent in the native state ensemble of hPGK1 (intermediates I1 and I2, Fig. 9B). In addition, we calculated the positive coupling free energies (∆G+) for these variants, a parameter that reports on the degree of relative thermodynamic coupling between residues that are folded44. The effect of each of the mutations (mut) versus the WT protein results in the mean difference in coupling free energies (∆∆G+) or perturbation, defined as
ΔG_{mut} = ΔG_{WT}, (plotted as a function of the residue number; Fig. 9C). These results show a similar ranking of the mutations (F25G > F25A > F25V) according to their negative impact on thermodynamic coupling of folded regions in the native state ensemble.

As expected from the decoupled nature of the two domains, the perturbations are mostly localized to the NTD mirroring experimental observations. The mean perturbation magnitude (μ), i.e. the average of the values in Fig. 9C corresponding to the first 200 residues (N-terminal domain), follows experimental HDX patterns with the F25G being the most perturbed (Fig. 9D, top panel). The corresponding standard deviation in perturbation (σ), that measures the extent of distribution, followed a similar trend (Fig. 9D, bottom panel), highlighting that the origins of the complex HDX-MS kinetics are a manifestation of population redistribution in the conformational landscape. The perturbation, when mapped onto the structure, vividly reveals the extent to which a single mutation affects neighbouring residues and the extent of coupling between them (Fig. 9E).

It is interesting to note that local stability measurements using HDX-MS (Fig. 8) and statistical mechanical model (Fig. 9) show that destabilization caused by F25 mutations are mostly restricted to the NTD and increases with the predicted size of the cavity (F → L < F → V < F → A) and conformational entropy (F → A < F → G). These analyses indicate that gradual local destabilization (due to alterations of the interaction networks) is connected to the effects on the global stability (i.e. reversible by urea or kinetically-controlled by temperature) and unfolding cooperativity (Figs. 4 and 5 and Tables 1, 2 and 4). Since hPGK1 WT resembles a two-state (un)-folder by urea, our results support that it displays certain heterogeneity of substates populated in the native state ensemble. It is likely that F25 mutations enhance such heterogeneity shifting the equilibrium towards partially unfolded substates, thus accelerating HDX and reducing the apparent unfolding cooperativity. This link between local
destabilization of the NTD and global denaturation effects (i.e. thermal and chemical denaturation) could be also partially explained by the effects of the F25 mutations on the NTD: CTD interface (Fig. S11).

Conclusions

Understanding how mutations affect the native state ensemble is essential to obtain insight into mutational effects on protein stability and function as well as their potential pathological consequences45,46. This is a challenging task since proteins display multiple features (stability, enzymatic function, regulation, biomacromolecular interactions, different subcellular locations, etc.) and a large genetic diversity exists in human population45–47. In this work, we have investigated how mutations targeting a fully buried Phe residue propagate their effects across the structure of the model two-domain hPGK1 enzyme, associated with disease3,28,30. In our strategy, we progressively decreased the size of the side-chain (F→L, F→V, F→A), additionally increased the entropy of the side chain (F→G) or potentially introduced conformational strain (F→W). A combination of biophysical studies (thermal and chemical denaturation, proteolysis), HDX-MS and statistical mechanical calculations showed that even small local perturbations (e.g. F→L or F→V) have remarkable effects on the unfolding cooperativity and energetics. Intriguingly, the mutation aimed to introduce conformational strain (F→W) has severe effects on thermal/kinetic stability, denaturation energetics, unfolding cooperativity and local stability of the NTD. Although mutations creating a protein cavity or introducing conformational strain could show similar effects on protein stability32, in our case a mutation that would increase by ~20% the size of the side-chain (F25W) shows comparable effects that one reducing this size by ~50% (F25A) (both based on the average volumes reported by48). The molecular origin of this apparently stronger effect of conformational strain is unclear.

Although this set of F25 mutations has not direct relationship with disease (i.e. not found in ClinVar or COSMIC databases), it represents a large fraction of naturally-occurring missense variants (Fig. 1C and Table S2). It is also worth noting that COSMIC mutations in hPGK1 (e.g. V216F and F241S)29 as well as several mutations analyzed from inherited hPGK1 deficiency leading to changes in the size-chain or the conformational entropy of the protein backbone (e.g. L89P and S320N)30 show effects on hPGK1 conformational landscape consistent with those reported here for the F25 mutants. We propose that our approach, combining biophysical analyses, HDX-MS studies and statistical mechanical calculations may allow to obtain high-resolution conformational information on the effects of cancer-associated mutations36, mutations associated with hPGK1 deficiency30 and cancer-associated post-translational modifications49. In addition, alterations in protein dynamics (in the ns-ms time scale) might be evaluated by molecular dynamics simulations50–52. We expect that similar analyses on mutations found in COSMIC (https://cancer.sanger.ac.uk/cosmic/gene/analysis?ln=PGK1_ENST00000644362),
Material and methods

Protein expression and purification. hPGK1 WT cDNA was chemically synthesized by GenScript (Leiden, The Netherlands) and cloned into the pET15b plasmid. The cloning strategy introduced in the N-terminal Meti of the hPGK1 WT the following sequence: MGSSHHHHHSGGLVPRGSH1. Residues underlined constitute the His6-tag used to purify the proteins by immobilized-metal affinity chromatography (IMAC) and the sequence in bold is the thrombin cleavage site. Mutants were generated by standard site-directed mutagenesis (GenScript, Leiden, The Netherlands). pET15b-hPGK1 plasmids were used to transform BL21(DE3) E. coli cells (Agilent Technologies, Santa Clara, CA, USA) for protein expression. For each variant, 40 mL of Luria–Bertani (LB) medium containing 0.1 mg mL⁻¹ ampicillin (Canvax Biotech, Córdoba, Spain) were inoculated with transformed cells and grown for 16 h at 37 °C. These cultures were diluted into 800 mL of LB containing 0.1 mg mL⁻¹ ampicillin, grown at 37 °C for 3 h to reach an optical density of about 1.0 and then these were transferred to 25 °C and induced with 0.5 mM IPTG (Isopropyl β-d-1-thiogalactopyranoside, Canvax Biotech, Córdoba, Spain). After 6 h, bacteria were harvested by centrifugation and frozen at −80 °C overnight. Cells were resuspended in binding buffer, BB (20 mM Na-phosphate, 300 mM NaCl, 50 mM imidazole, pH 7.4) containing 1 mM PMSF (phenylmethylsulfonyl fluoride, Sigma-Aldrich) and sonicated in an ice bath. These extracts were centrifuged (20,000 g, 30 min, 4 °C) and the supernatants were loaded into IMAC (Ni-Sepharose immobilized-metal affinity chromatography, from Cytiva, Barcelona, Spain) columns, washed with 40 bed volumes of BB and eluted in BB containing a final imidazole concentration of 500 mM. These eluates were treated with streptomyycin 10% W/V to remove (poly)-nucleotides that may be bound to the hPGK1 protein. After 30 min of centrifugation at 20,000 g and 4 °C, supernatants were loaded into PD-10 columns (Cytiva, Barcelona, Spain) and buffer exchanged to the storage buffer (20 mM HEPES–NaOH, 200 mM NaCl, pH 7.4). These samples were flash-frozen in liquid N₂ and stored at −80 °C.

To remove protein aggregates and increase protein purity, proteins were subjected to size exclusion chromatography on a Superdex 200 16/60 column (GE Healthcare) using 20 mM HEPES–NaOH, 200 mM NaCl, stored at −80 °C.

Material and methods

Thermal denaturation by CD spectroscopy. hPGK1 WT cDNA was chemically synthesized by GenScript (Leiden, The Netherlands) and cloned into the pET15b plasmid. The cloning strategy introduced in the N-terminal Meti of the hPGK1 WT the following sequence: MGSSHHHHHSGGLVPRGSH1. RESIDUES UNDERLINED CONSTITUTE THE HIS6-TAG USED TO PURIFY THE PROTEINS BY IMMobilIZED-METAL AFFINITY CHROMATOGRAPHY (IMAC) AND THE SEQUENCE IN BOLD IS THE THROMBIN CLEAVAGE SITE. MUTANTS WERE GENERATED BY STANDARD SITE-DIRECTED MUTAGENESIS (GENSCRIPT, LEIDEN, THE NETHERLANDS). pET15B-HPGK1 PLASMIDS WERE USED TO TRANSFORM BL21(DE3) E. COLI CELLS (AGILENT TECHNOLOGIES, SANTA CLARA, CA, USA) FOR PROTEIN EXPRESSION. FOR EACH VARIANT, 40 mL OF LURIA–BERTANI (LB) MEDIUM CONTAINING 0.1 MG mL⁻¹ AMPICILLIN (CANVAX BIOTECH, CÓRDOBA, SPAIN) WERE INOCULATED WITH TRANSFORMED CELLS AND GROWN FOR 16 H AT 37 °C. THESE CULTURES WERE DILUTED INTO 800 mL OF LB CONTAINING 0.1 MG mL⁻¹ AMPICILLIN, GROWN AT 37 °C FOR 3 H TO REACH AN OPTICAL DENSITY OF ABOUT 1.0 AND THEN THESE WERE TRANSFERRED TO 25 °C AND INDUCED WITH 0.5 mM IPTG (ISOPROPYL β-D-1-THIOGALACTOPYRANOside, CANVAX BIOTECH, CÓRDOBA, SPAIN). AFTER 6 H, BACTERIA WERE HARVESTED BY CENTRIFUGATION AND FROZEN AT −80 °C OVERNIGHT. CELLS WERE RESUSPENDED IN BINDING BUFFER, BB (20 mM Na-phosphate, 300 mM NaCl, 50 mM imidazole, pH 7.4) CONTAINING 1 mM PMSF (PHENYLmethylsULFONYL FLuorIDE, SIGMA-ALDRICH) AND SONICATED IN AN ICE BATH. THESE EXTRACTS WERE CENTRIFUGED (20,000 g, 30 min, 4 °C) AND THE SUPERNATANTS WERE LOADED INTO IMAC (Ni-Sepharose immobilized-metal affinity chromatography, FROM CYTIVA, BARCELONA, SPAIN) COLUMNS, WASHED WITH 40 BED VOLUMES OF BB AND ELUTED IN BB CONTAINING A FINAL IMIDAZOLE CONCENTRATION OF 500 mM. THESE ELUATES WERE TREATED WITH STREPTOMYcIN 10% W/V TO REMOve (POLY)-nucleotides THAT MAY BE BOUND TO THE hPGK1 PROTEIN. AFTER 30 MIN OF CENTRIFUGATION AT 20,000 g AND 4 °C, SUPERNATANTS WERE LOADED INTO PD-10 COLUMNS (CYTIVA, BARCELONA, SPAIN) AND BUFFER EXCHANGED TO THE STORAGE BUFFER (20 mM HEPES–NaOH, 200 mM NaCl, pH 7.4). THESE SAMPLES WERE FLASH-FROZEN IN LIQUID N₂ AND STORED AT −80 °C.

To remove protein aggregates and increase protein purity, proteins were subjected to size exclusion chromatography on a Superdex 200 16/60 column (GE Healthcare) using 20 mM HEPES–NaOH, 200 mM NaCl, pH 7.4 as mobile phase at 1.5 mL min⁻¹ flow rate. Monomeric fractions were concentrated and stored at −80 °C after flash-freezing in liquid N₂.

UV-visible absorption spectra were recorded at 10–20 μM protein concentration in a Cary Eclipse 50 spectrophotometer using 1 cm pathlength cuvettes, using an extinction coefficient of ε280 = 27,960 M⁻¹ cm⁻¹ except for F25W for which we used ε280 = 33,460 M⁻¹ cm⁻¹.

Circular dichroism (CD) spectroscopy. CD spectra were registered in a Jasco J-710 spectropolarimeter with a Peltier element for temperature control. Before the acquisition of the far-UV and near-UV/visible CD spectra, protein samples were buffer exchanged to 20 mM K-phosphate pH 7.4 using PD-10 columns (Cytiva, Barcelona, Spain). Spectra in the far-UV range were acquired using 6–7 μM of protein in a 1 mm path-length quartz cuvette. These spectra were registered in the 200–260 nm range, at 25 °C, with a scan rate of 100 nm min⁻¹, a bandwidth of 1 nm and a time response of 2 s. Six spectra of each sample were acquired and averaged, and the corresponding blank (containing the same buffer but no protein) was acquired under the same conditions at the beginning and the end of each experimental series to assess for baseline drifts during the day, were averaged and subtracted from the spectra of protein samples. Similar conditions were used to acquire near-UV/visible CD spectra with some modifications. Protein concentration for these experiments was 15 μM and 10 spectra were acquired and averaged, the wavelength range was 270–330 nm and the path-length of the cuvette was 5 mm. To calculate mean molar residue ellipticities [Θ]MRW, Eq. (1) was used:

\[
[\text{Θ}]_{\text{MRW}} = \frac{\text{MRW} \cdot \text{Θ}}{10 \cdot I \cdot c}
\]

where MRW is the molecular weight of each hPGK1 variant divided by 416 (the residues of native hPGK1 minus 1), Θ is the observed ellipticity of the sample in mdeg, I is the pathlength of the cuvette in cm and c the protein concentration in mg·mL⁻¹.

Thermal denaturation by CD spectroscopy. For thermal denaturation experiments by CD spectroscopy, we used exactly the same samples used to acquire CD wavelength-dependent spectra. Samples were thermostatted at 20 °C for 10 min and then temperature was raised to 75 °C at a 2 °C min⁻¹ scan rate. Signals were acquired at 222 nm (for UV-CD in 1 mm path-length cuvettes) or 290 nm (for Near-UV CD in 5 mm path-length), using a bandwidth of 2 nm and a time response of 16 s. Raw data were normalized for linear pre- and post-transition baselines and the apparent Tm was determined as the temperature at which half of the unfolded fraction was observed by linear extrapolation. Thermal denaturation leads to irreversible protein aggregation (i.e. formation of protein aggregates of large size).30

Differential scanning calorimetry (DSC). DSC analyses were carried out using a VP-DSC differential scanning calorimeter (Malvern Panalytical, Malvern, UK) with a cell volume of 137 μL. Experiments were performed in 20 mM HEPES–NaOH, 200 mM NaCl pH 7.4 and using 7–15 μM protein. Temperature range was 5–75 °C and scan rates were 1, 2.5 or 4 °C min⁻¹. To investigate the effect of urea on hPGK1 denaturation, stock solutions of urea 5 M in 20 mM HEPES–NaOH, 200 mM NaCl pH 7.4 were diluted to the desired final concentration. All urea concentrations were checked by refractive index measurements.
The DSC traces were analysed using a two-state irreversible model\textsuperscript{29,41} (model 1), that describes very well the irreversible denaturation of hPGK1\textsuperscript{29}. In this model, denaturation is considered as a kinetically-controlled conversion of the native (N) to a final state (F) that cannot fold back:

\[ N \xrightarrow{k} F \]  
(Model 1)

This process is characterized by a strongly temperature-dependent first-order rate constant \( k \). We also assume that the temperature dependence of \( k \) follows a slightly modified version of the Arrhenius equation (Eq. 2):

\[
\ln k = \ln k_{T*} - \frac{E_a}{R} \left( \frac{1}{T} - \frac{1}{T*} \right) \tag{2}
\]

where \( T \) is the absolute temperature, \( T* \) is a reference absolute temperature, \( k_{T*} \) is the rate constant at \( T* \) and \( E_a \) is the activation energy and \( R \) is the ideal gas constant (1.987 cal mol\(^{-1}\) K\(^{-1}\)).

The fitting function used to extract relevant energetic parameters: \( k, E_a \), the enthalpy changes between N and F (\( \Delta H \)) and the \( T_m \) (the maximum of the excess heat capacity versus \( T \)); usually close to the temperature at which \( k = 1 \text{ min}^{-1} \) uses two linear pre- and post-transition temperature-dependent baselines smoothly connecting N and F signals. This procedure allow to isolate the calorimetric peak defined by the term \(- \Delta H \cdot \frac{dT \cdot \Delta C_p}{dT} \). This peak shows the dependence of the excess heat capacity \( C_{p,Exc} \) from \( \frac{dT \cdot \Delta C_p}{dT} \) which is described by Eq. (3)\textsuperscript{29,41,53}:

\[
\frac{dT_m}{dT} = \frac{E_a}{RT_m^2} \exp \left( \frac{E_a \cdot \Delta T}{RT_m^2} \right) \exp \left[ -\exp \left( \frac{E_a \cdot \Delta T}{RT_m^2} \right) \right] \tag{3}
\]

From \( C_{p,Exc} \) versus \( T \) profiles (i.e. after removing the chemical baselines), the value of the temperature-dependent rate constant \( k \) can be determined using Eq. (4)\textsuperscript{44}:

\[
k = \frac{v \cdot C_{p,Exc}}{\Delta H - < H >} \tag{4}
\]

where \( v \) is the scan rate in °C min\(^{-1}\), \( C_{p,Exc} \) is the excess heat capacity at a given \( T \), \( \Delta H < H > \) is enthalpy change of the transition and \( < H > \) is the excess enthalpy at a given \( T \). This equation allows to determine the \( T \)-dependence of \( k \) and thus yields the corresponding Arrhenius plots (Eq. 2).

To determine kinetic \( m \) values (\( m^* \)) from DSC studies we followed the procedure described by\textsuperscript{45}. Briefly, we carried out DSC experiments under the same conditions as described above but adding a final urea concentration in the range of 0.5–1.1 M. Then, \( m^* \) values were determined using Eq. (5):

\[
m^* = -\frac{E_a}{T_m} \left( \frac{dT_m}{d[\text{Urea}]} \right) - R \cdot T_m \cdot \left( \frac{d \ln \left( \frac{E_a}{RT_m^2} \right)}{dT_m} \right) \tag{5}
\]

where \( E_a, T_m, dT_m/d[\text{Urea}] \) can be readily calculated from fittings at different urea concentrations. Thus, for each protein variant, three different values of \( m^* \), one for each scan rate were obtained, and the data reported are the mean ± s.d. of these three values.

**Isothermal urea denaturation monitored by CD spectroscopy.** Urea was prepared at ~6 M in 20 mM HEPES–NaOH, 200 mM NaCl, 1 mM 2-mercaptoethanol (β-ME), pH 7.4 and its actual concentration was determined by refractive index measurements. Urea stock solutions were mixed with hPGK1 solutions in 20 mM HEPES–NaOH, 200 mM NaCl, 1 mM 2-mercaptoethanol (β-ME), pH 7.4 to a final concentration of 5 μM (in protein) and 0–5 M (in urea). Samples were incubated for at least 2 h and denaturation experiments were monitored by Far-UV CD spectroscopy similarly to those described in Section "Circular dichroism (CD) spectroscopy" but restricting the acquisition range to 215–260 nm.

Data at 222 nm were used for fittings assuming a two-state model (apparently valid for WT hPGK1 using different conformational probes)\textsuperscript{3,28–30} to provide the values of experimental equilibrium \( m \) (\( m_{\text{urea}} \)) and \( C_m \) (concentration of urea at which half-denaturation occurs) using Eq. (6):

\[
S = S_N + m_N \cdot [\text{Urea}] + S_U + m_U \cdot \left( \exp \left( \frac{\text{molar mass} \cdot (\text{Slope}_{\text{Native}} - \text{Slope}_{\text{Unfolded}})}{RT_m} \right) \right) / \left[ 1 + \exp \left( \frac{\text{molar mass} \cdot (\text{Slope}_{\text{Native}} - \text{Slope}_{\text{Unfolded}})}{RT_m} \right) \right] \tag{6}
\]

where \( S \) is the experimental CD signal as a function of the [Urea], \( S_N \) and \( S_U \) are the fitted CD signals for the Native and Unfolded state baselines at 0 M urea, respectively, \( m_N \) and \( m_U \) are the slopes of the native and unfolded state baselines, \( m_{\text{urea}} \) describes the unfolding cooperativity, \( R \) is the ideal gas constant and \( T \) is the experimental temperature (298.15 K). Although this model is likely an oversimplification, it provides a proxy to compare the cooperativity of irreversible chemical unfolding of hPGK1 variants.

**Proteolysis by thermolysin.** Thermolysin from *Geobacillus stearothermophilus* (Sigma-Aldrich, Madrid, Spain) was prepared in 20 mM HEPES–NaOH, 200 mM NaCl, 100 mM CaCl\(_2\), pH 7.4 and its concentration was determined spectrophotometrically using \( \varepsilon_{280} = 66,086 \text{ M}^{-1} \cdot \text{cm}^{-1} \) based on the protein primary sequence. Proteolysis was initiated after 5 min incubation at 25°C by mixing hPGK1 variants and thermolysin to a final concentrations of 10 μM and 0.5 or 1 μM, respectively. When the initial reaction was started contained 10 mM CaCl\(_2\). Aliquots were withdrawn at different times (2 min–96 h), mixed with 25 mM EDTA (final concentration),
pH 8 and denatured with Laemlli’s buffer (1:1 in V/V) at 95 °C for 5 min. Samples were resolved in SDS-PAGE gels (12% acrylamide) and analysed by densitometry using the software ImageJ (https://imagej.nih.gov/ij/).

The decay of the full-length protein (I) versus time (t) was fitted using the following single exponential function to provide the first-order kinetic constant k_{obs}:

$$I = I_0 \cdot \exp^{-k_{obs} \cdot t}$$

The half-life ($t_{1/2}$) for proteolysis was determined using: $t_{1/2} = \ln 2 / k_{obs}$. Errors corresponded to those from the fittings.

**Hydrogen/deuterium eXchange mass spectrometry (HDX-MS).** Amide hydrogen/deuterium exchange (HDX) of hPGK1 was studied for the WT and mutant variants F25A, F25G, F25L, F25V, F25W as described previously with some modifications. Briefly, 20 μM of protein solution was 10× diluted with a D,O-based 20 mM HEPEs–NaOH, 200 mM NaCl, 0.5 mM TCEP (tris(2-carboxyethyl)phosphine), pH 7.4, to start the exchange reaction. The exchange was terminated after 10 s, 30 s, 2 min, 5 min, 20 min, 1 h, 3 h, 20 min and 20 h by mixing (1:1) with 0.5 M Glycine–HCl, pH 2.3 and the samples were flash frozen in liquid N₂. Time points 30 s, 5 min and 3 h 20 min were replicated. Custom-made nepenthesin-2 column was used for online proteolysis and the peptides were trapped and desalted on a SecurityGuard™ pre-column (ULTRA Cartridges UK) against a database containing hPGK1 sequences, nepenthesin-2 and cRAP contaminant sequences. Decoy MS run performed using the identical LC settings but connected to ESI-timsTOF Pro with PASEF. LC–MS/MS data were exported using DataAnalysis 5.3 and searched by in-house developed program Deutex. Peptide identification was done in a separate data-dependent LC–MS/MS run performed using the identical LC settings but connected to ESI-timsTOF Pro with PASEF. LC–MS/MS data were exported to mgf files using DataAnalysis 5.3 and searched by MASCOT (Matrix Science, London, UK) against a database containing hPGK1 sequences, nepenthesin-2 and cRAP contaminant sequences. Decoy search was enabled with a false-discovery ratio <1% and an Ion Score cut-off 15.

**Computational analyses of the F25 mutational effects.** The Wako-Saitó-Muñoz-Eaton (WSME) model based perturbation analysis of hPGK1 was performed as discussed before for mutants of the protein NQO1. Briefly, an ensemble of 8,398,795 microstates is constructed employing the block–version of the WSME model—bWSME with a most probable block length of 5 consecutive residues—and the PDB structure 2XE7. Computations of the microstate energetics include stabilizing contributions from van der Waals interactions (heavy-atom interactions identified with a 5 Å cut-off with a mean-field interaction energy of -78 J mol⁻¹ per heavy atom contact), all-to-all electrostatics at pH 7 protonation state, and simplified solvation free energy (defined by the heat capacity change of −0.36 J mol⁻¹ K⁻¹ per native contact). In addition, an entropic penalty of −14.5 J mol⁻¹ K⁻¹ per residue is introduced when fixing an unfolded residue to a folded conformation. The balance between the stabilization (free-)energy terms and entropic penalty determines the free energy of every microstate, the statistical weight and hence the associated probability.

**Data availability**
The datasets generated and/or analyzed during the current study are available from the corresponding author on reasonable request (A.L.P.).

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A.L.P ., A.N.N. and M.C-M. analyzed and interpreted data in structural terms. A.L.P . wrote the initial

P .V . and P .M. analyzed mass spectrometry experiments. A.N.N. carried out and analyzed statistical mechanical

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Author contributions

A.L.P . conceived and supervised the project. A.L.P . and A.N.N. acquired funding; J.L.P-G. and A.L.P . expressed

and purified hPGK1 proteins, performed and analyzed spectroscopic and calorimetric experiments. J.L.P-G. car-

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Competing interests

The authors declare no competing interests.

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