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Proline/sidechain C–H/O interactions stabilize *cis*-proline

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Bioinformatics analysis was conducted on proteins in the PDB to identify local structures that can stabilize the *cis*-proline conformation. C–H/O interactions were observed between a sidechain oxygen and Pro C–H α in the *cis*-proline conformation at Glu–Pro, Asp–Pro, Gln–Pro, Asn–Pro, Ser–Pro, and Thr–Pro sequences. These C–H/O interactions are apparently most stabilizing at Glu–Pro sequences, which have a substantially higher than average frequency of *cis*-proline (7.1% of all Glu–Pro amide bonds in the PDB). DFT calculations were conducted to understand the bases and geometries of C–H/O interactions in these sequences. Computationally, these residues all exhibit close C–H/O interactions (substantially below the 2.72 Å sum of the van der Waals radii of H and O), with the closest C–H/O interactions observed with the anionic oxygens of Glu and Asp, and with closer interactions for the anionic residues than the neutral carboxamides Gln and Asn. DFT calculations revealed that C–H/O interactions also stabilize *cis*-proline at phosphoserine–proline and phosphothreonine–proline sequences, with closer C–H/O interactions in the dianionic forms of phosphorylated residues that predominate at physiological pH. These results also provide an explanation for the observed higher activation barrier for amide bond isomerism at phosphoserine–proline and phosphothreonine–proline sequences. Calculations suggested that C–H/O interactions mediated by these residues could also stabilize non-proline *cis* amide bonds, which are often functionally important when observed.

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Introduction

cis amide bonds exhibit substantial changes in structure compared to *trans* amide bonds (Fig. 1a), both in peptides and in globular proteins.^{1,2} In addition, *cis*–*trans* isomerism at proline amide bonds is important in the dynamics of proteins.^{3–6} In isolated peptide sequences, *cis* amide bonds are higher energy than *trans* amide bonds, due to both (1) the steric clash between the C α of adjacent residues that is present in *cis* amides and (2) $n \rightarrow \pi^*$ interactions that stabilize the *trans* amide conformation (Fig. 1).^{3,7} For proline residues, the C α /C α steric clash in *cis*-proline is partially counteracted by a C α /C δ steric clash in *trans*-proline. Due to these counterbalancing interactions, 5% of all proline residues in the PDB exhibit the *cis*-proline conformation, with higher frequencies of *cis*-proline often observed in simple peptides.^{8–10} In contrast, for non-proline residues, the absence of a similar steric clash in the *cis*-amide conformation leads to a much lower frequency of *cis* amide (~0.03% of all non-proline amide bonds).¹¹

Due to the geometric constraints imposed by the *cis* amide conformation, the vast majority of structures with *cis* amide bonds are β -turns (type VI), with C α /C α distances between the

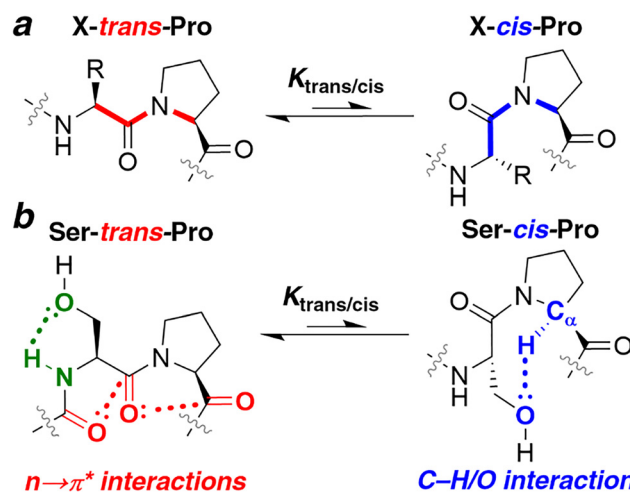


Fig. 1 Proline amide *cis*–*trans* isomerism and stabilizing interactions in each amide conformation. (a) Proline *cis*–*trans* amide isomerism. (b) The Ser sidechain hydroxyl can act as an electron donor for hydrogen bonds or can engage in other noncovalent interactions. When in the pre-proline ($i - 1$) position, the Ser hydroxyl can stabilize the *trans*-Pro conformation via the propagation of $n \rightarrow \pi^*$ interactions across proline. Alternatively, the Ser hydroxyl can also stabilize the *cis*-Pro conformation via a C–H/O interaction between Pro C–H α and a Ser sidechain oxygen lone pair. Numerous additional sidechain–main chain interactions are also possible in Ser–Pro sequences.

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i and $i + 3$ residues less than 7 Å.^{12–14} In addition, amide *cis-trans* isomerism is often the slow step in protein folding, with a timescale (seconds to minutes) that is much longer than that of other protein folding transitions (microseconds to milliseconds).^{4,15,16} At proline residues, the *cis-trans* amide interconversion is catalyzed by prolyl isomerases, including cyclophilins, FKBP, and parvulins.^{17,18} At non-proline residues, where the energy difference between *trans*-amide and *cis*-amide is higher in the unfolded state, and thus where a larger native barrier exists for the *trans*-to-*cis* conversion (leading to $t_{1/2} \sim 1000$ s), amide isomerism is mediated by molecular chaperones such as DnaK.^{19–22}

cis amide bonds are highly evolutionarily conserved, likely due to the substantial differences in structure and relative orientations of the protein chains on each side of the amide bond when comparing *trans* versus *cis* amides.²³ Proteins can exhibit switches in function as a result of proline *cis-trans* isomerism, with different activities or interaction partners in each amide conformation about a specific proline residue.^{5,6,24–27}

Despite the functional importance of proline *cis-trans* isomerism, there is still an incomplete understanding of the factors stabilizing or promoting the *cis*-proline conformation.^{9,28,29} Within globular proteins, the three-dimensional folded structure certainly plays a significant role in stabilizing the *cis*-amide conformation. Folding, however, plays a more minor role in intrinsically disordered regions of proteins (IDPs), where the dynamics of proline *cis-trans* isomerism are often critical.^{2,30} Moreover, even within globular proteins, there are substantial differences in the frequency of *cis*-proline depending on the adjacent residues. Proline-proline, aromatic-proline, and proline-aromatic sequences are substantially more likely to adopt the *cis*-proline conformation, both in globular proteins and within disordered peptides.^{31–37} The higher frequency of *cis*-proline in aromatic-proline and proline-aromatic sequences is primarily due to C-H/ π interactions, between the aromatic (π) ring and the proline ring and/or C-H α of the pre-proline residue, which stabilize the *cis*-proline conformation.³⁸

However, other bases for the stabilization of *cis*-proline are less well understood. For example, prior bioinformatics analyses have indicated that Glu-Pro sequences in the PDB also exhibit a significantly higher than typical frequency of *cis*-proline.^{9,28} Moreover, local sequences can dramatically impact the dynamics of proline *cis-trans* isomerism. Protein serine/threonine phosphorylation results in a ~ 1 kcal mol⁻¹ higher barrier for proline *cis-trans* isomerism. The substantially slower dynamics (~ 5 – 10 -fold lower rates of interconversion) are resolved *in vivo* by the phosphorylation-dependent prolyl isomerase Pin1.^{39–44} Pin1 overexpression is associated with cancers due to Pin1 promoting cell cycle progression.⁴⁵ In contrast, Pin1 is depleted in Alzheimer's disease, and suppression of Pin1 levels in disease models leads to more rapid *tau* aggregation and neurodegeneration.⁴⁶ Phosphorylation-dependent proline *cis-trans* isomerization in the RNA polymerase II C-terminal domain is also central to transcription.^{47–50} However, the bases for these observations of effects of phosphorylation on proline *cis-trans* isomerization rates have remained unexplained.

Recently, we demonstrated that *cis*-proline is stabilized in Ser-Pro sequences *via* a C-H/O interaction^{51–58} between the sidechain Ser oxygen and the Pro C-H α (Fig. 1b).^{59,60} This work was supported by data from small-molecule X-ray crystallography, solution-state NMR spectroscopy, bioinformatics analysis of the PDB, and DFT calculations on model peptides. In Ser-Pro, the *cis*-stabilizing effect of C-H/O interactions is counterbalanced by hydrogen-bonding interactions between the Ser hydroxyl and backbone amides that are present in *trans*-proline. Similarly, based on NMR data and molecular modeling, we recently proposed that Ser/Thr phosphorylation increases the activation barrier at pSer-Pro and pThr-Pro sequences in part *via* stabilization of *cis*-proline *via* C-H/O interactions between a phosphate oxygen and Pro C-H α .⁶¹

C-H/O interactions can be described electrostatically, as the interaction between the partial negative charge (δ^-) on an oxygen with the partial positive charge (δ^+) on the hydrogen of a polarized C-H bond. The H α of amino acids exhibit significant δ^+ due to bond polarization *via* the amide nitrogen and carbonyl carbon that are attached to C α .^{55,56} This reduced electron density on H α is reflected in the more downfield chemical shift of protein H α (~ 4 – 5 ppm) compared to other aliphatic hydrogens in proteins.⁶²

Importantly, however, C-H/O interactions have only a modest dependence on electrostatics,^{63–65} and are primarily stabilizing *via* stereoelectronic (molecular orbital-based) effects. Stabilization occurs due to electron delocalization between oxygen lone pair orbitals and the antibonding (σ^*) orbital of the C-H bond. Evidence for the predominantly stereoelectronic nature of C-H/O interactions comes both from experiment (due to only partial charges being involved, the underlying electrostatics energies are inherently weak in water, and thus the observation of these interactions being stabilizing in water is inconsistent with a primary electrostatics basis) and from calculations. For example, only minimal differences in calculated interaction energies of the model C-H/O interaction DMF-CHCl₃ are observed between $\epsilon = 9$ and $\epsilon = 1\,000\,000$.⁶⁵

X-ray crystallography is also consistent with a predominantly stereoelectronic nature for C-H/O interactions. H \cdots O distances significantly below the 2.72 Å sum of the van der Waals radii of H and O are inconsistent with a purely electrostatics-based interaction, but are consistent with electron delocalization (partial covalency) being important in the interaction.⁶⁵ C-H/O interactions are also dependent on the electron density on the oxygen, with a more electron-rich oxygen being a better electron donor.⁶⁶ Thus, calculations on model systems demonstrate a closer H \cdots O distance and stronger interaction with oxygens that are part of dianions than monoanions, and with anions compared to neutral oxygens.

Collectively, these results suggested the possibility that protein sidechain oxygens might more generally interact with Pro C-H α to stabilize the *cis*-proline conformation *via* C-H/O interactions. We examine this hypothesis herein, *via* a combination of bioinformatics analysis of the PDB and DFT calculations. The potential for local proline/sidechain C-H/O interactions will be examined *via* bioinformatics analysis of structures with



cis-proline. The structures identified *via* bioinformatics will then be analyzed *via* DFT calculations to determine the potential bases for main chain/sidechain C–H/O interactions to stabilize the *cis*-proline conformation.

Methods

Bioinformatics

On May 9, 2024, a search of the Protein Data Bank (PDB) was conducted for structures with resolution ≤ 2.5 Å and with sequence similarities less than 30% using the PISCES server.⁶⁷ The search yielded 15 599 total structures. Perl scripts were written to extract lists of structures containing Glu–Pro, Gln–Pro, Asp–Pro, Asn–Pro, Ser–Pro, and Thr–Pro sequences. Additional perl scripts were written to individually analyze the lists of X–Pro structures and calculate dihedral angles, specified interatomic distances, amide hydrogen positions, and H α positions for the examined residues. The data sets were manually refined to exclude structures with nearby broken backbone bonds and structures that contained Pro with positive ϕ dihedral angles. The data sets used for analyses contained 4453 EP structures (4138 with *trans*-proline and 315 with *cis*-proline), 3519 QP structures (3354 with *trans*-proline and 165 with *cis*-proline), 5322 DP structures (5155 with *trans*-proline and 167 with *cis*-proline), 4648 NP structures (4440 with *trans*-proline and 208 with *cis*-proline), 5188 SP structures (4895 with *trans*-proline and 293 with *cis*-proline), and 5392 TP structures (5188 with *trans*-proline and 204 with *cis*-proline). Additional details are in the SI.

Computational chemistry

Calculations were conducted with Gaussian 09.⁶⁸ Initial structures derived from the PDB were subjected to iterative geometry optimization. For all molecules, final optimization was conducted using the M06-2X DFT functional with the 6-311++G(d,p) basis set (including diffuse and polarization functions).^{69,70} Calculations were conducted in implicit water (IEFPCM) unless stated otherwise.⁷¹

Rotamer analysis to estimate interaction energies⁷² (Fig. S15) was conducted on molecules with Glu[−], Glu⁰, or Gln *via* rotation of the χ_2 torsion angle to *t* (180°) within GaussView, followed by full geometry optimization from this initial structure. The calculated interaction energies were determined *via* comparison of the calculated electronic energies of the interacting *versus* non-interacting rotamer. Interaction energies as a function of solvent were determined similarly, *via* full geometry optimization of the interacting and non-interacting rotamers in the indicated solvent (IEFPCM), vacuum ($\epsilon = 1$), or condition with artificial dielectric constant ($\epsilon = 1000$; 10 000, 100 000; or 1 000 000), with the calculated relative electronic energies of the interacting *versus* non-interacting rotamer compared (Table S13). C–H/O interaction distances as a function of solvent and residue identity were also determined (Tables S14 and S15).

Structures with phosphoserine or phosphothreonine were conducted *via* bond rotations on models described previously,⁷³

followed by bond rotations within GaussView and subsequent full geometry optimization. Structures at non-proline *cis* amide bonds were generated *via* modification of the relevant structures with Pro, followed by full geometry optimization.

Additional geometric details of all computational models, as well as their relative energies and the coordinates for all models, are in the SI.

NBO analysis was conducted using NBO6 as implemented within Gaussian09.^{74,75} Visualization was conducted in GaussView 5 with isovalues of 0.02. Atoms in Molecules (AIM) analysis was conducted using Multiwfn.^{76–78}

Results

Bioinformatics analysis of X–Pro sequences in the PDB

In order to examine the possibility of C–H/O interactions stabilizing *cis*-proline, we examined high-resolution structures of X–Pro sequences in the PDB for all amino acids with an oxygen in an aliphatic sidechain (X = Asp, Asn, Glu, Gln, Ser, Thr).⁶⁷ Post-translational modifications can also add oxygens to protein sidechains (*e.g.* Cys sulfenic acid (Cys–SOH or Cys–SO[−]), sulfinate (Cys–SO₂[−]), and sulfonate (Cys–SO₃[−]); phosphoserine or phosphothreonine). However, there is no example of an X-ray crystal structure of an isolated X–*cis*-proline in the PDB with X = one of these post-translationally modified amino acids. All structures were analyzed for the conformation at the X–Pro amide bond (*trans*-proline, $\omega \sim 180^\circ$; or *cis*-proline, $\omega \sim 0^\circ$) and the conformations at the main chain (ϕ , ψ) and sidechain (χ_1 , χ_2 , and/or χ_3 , as appropriate) for both residues. In addition, the distance between any sidechain oxygen atom and Pro H α was measured for all structures. The position of H α was calculated geometrically from the positions of the heavy atoms attached to Pro C α (Fig. S1), as the position of H α is typically not included in PDB files.

Glu–Pro sequences exhibited the highest frequency of *cis*-Pro (7.1% of Glu–Pro sequences) (Table 1), and were observed at greater than the typical 5.3% frequency of *cis*-Pro at all X–Pro sequences. Ser–Pro sequences also had an elevated frequency of *cis*-Pro, as we had observed previously.⁵⁹ In contrast, Gln–Pro and Asn–Pro had modestly lower than average frequencies of *cis*-Pro, and Thr–Pro and Asp–Pro had substantially lower than average frequencies of *cis*-Pro (Table 1). These frequencies of

Table 1 Frequencies of proline amide conformations observed at X–Pro sequences in the PDB. Structures in the PDB containing X–Pro sequences, where X is a residue with an oxygen-containing atom in the sidechain that is capable of engaging in a C α –H/O interaction, were analyzed and separated by proline amide isomer

X	Number of structures			% of structures	
	Total	X– <i>trans</i> -Pro	X– <i>cis</i> -Pro	X– <i>trans</i> -Pro	X– <i>cis</i> -Pro
Glu	4453	4138	315	92.9	7.1
Ser	5188	4893	295	94.3	5.7
Gln	3519	3354	165	95.3	4.7
Asn	4648	4440	208	95.5	4.5
Thr	5392	5188	204	96.2	3.8
Asp	5322	5155	167	96.9	3.1



cis-proline for each amino acid are consistent with prior analyses that were conducted on fewer structures and with less stringent resolution limits than were used in the current analysis.^{8,9}

Distances for potential C–H/O interactions were examined for all amino acids in both *trans*-Pro and *cis*-Pro (Fig. 2, Table 2, Tables S2 and S3). These data indicated that C–H/O interactions were present between a sidechain oxygen and Pro C–H α for a significant number of structures with *cis*-Pro, with potential C–H/O interactions defined initially as O \cdots H α distances < 3.00 Å. Notably, in *cis*-Pro for all amino acids, a bimodal or trimodal distribution of O \cdots H α distances was observed (Fig. 2), with one distribution of distances distinguished by O \cdots H α distances < 3.00 Å. In contrast, essentially no structures with *trans*-Pro had O \cdots H α distances < 4 Å (Table S3), indicating that Pro C–H α /O interactions are not significant for *trans*-Pro.

Analysis as a function of distance and amino acid type indicated substantial differences in frequencies of potential C–H/O interactions. The highest frequency of close O \cdots H α distances was observed for Glu-*cis*-Pro and Asp-*cis*-Pro sequences: ~50% of these structures had an O \cdots H α distance \leq 3.0 Å, and 37% of these structures had an O \cdots H α distance \leq 2.72 Å. Notably, 20% of these structures had an O \cdots H α distance \leq 2.50 Å,

Table 2 Frequencies of C α –H/O interactions at X-*cis*-Pro sequences in the PDB as a function of H α \cdots O distance. The frequencies of X-*cis*-Pro structures with H α \cdots O distances \leq 2.30 Å, 2.50 Å, 2.75 Å, and 3.00 Å are indicated

X- <i>cis</i> -Pro	% of X- <i>cis</i> -Pro with C α –H \cdots O distances \leq			
	2.30 Å	2.50 Å	2.75 Å	3.00 Å
Glu	5.7	20.3	38.1	47.3
Ser	3.1	16.0	31.4	38.6
Gln	0.6	12.7	26.1	34.5
Asn	3.8	17.8	36.1	46.6
Thr	4.9	12.7	16.7	18.1
Asp	7.2	21.0	39.5	52.1

which is well below the sum of the van der Waals radii of H and O, and a distance that would be associated with a particularly favorable C–H/O interaction despite the superficial steric clash at these distances. In contrast, Gln-*cis*-Pro and Thr-*cis*-Pro sequences had the lowest frequencies of O \cdots H α distances that would be consistent with a C–H/O interaction. Notably, however, Thr-*trans*-Pro sequences had the third highest frequency of very close (\leq 2.30 Å) O \cdots H α distances, suggesting that, while Thr-*cis*-Pro are less likely to exhibit a C–H/O interaction, they were still capable of exhibiting close C–H/O interactions.

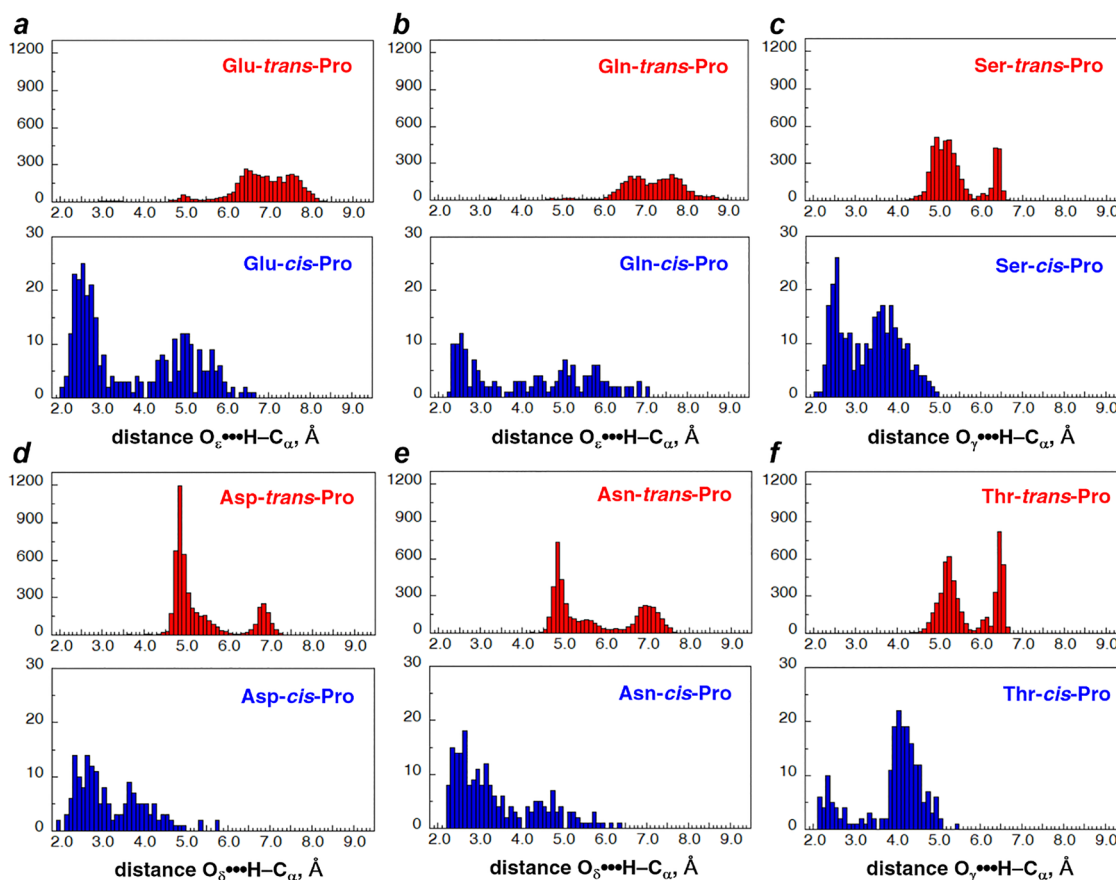


Fig. 2 C α –H \cdots O distances at X-Pro sequences in the PDB. Minimum H α \cdots O distances at (a) Glu-Pro, (b) Gln-Pro, (c) Ser-Pro, (d) Asp-Pro, (e) Asn-Pro, and (f) Thr-Pro sequences as a function of Pro amide conformation (for each, (top) *trans* (red) and (bottom) *cis* (blue)).



Protein sidechain conformations exhibiting C–H/O interactions in *cis*-proline

The distributions of sidechain rotamers were examined for each amino acid.⁷⁹ Comparison of the rotamer distributions in *trans*-Pro versus in *cis*-Pro (Table 3) indicated several trends. Glu and Gln residues exhibited a significantly higher frequency of the g^- χ_1 rotamer in Glx-*cis*-Pro, while Gln (but not Glu) exhibited a significantly higher frequency of the g^+ χ_2 rotamer. For Asp and Asn, the major χ_1 rotamer was *t*, for both *trans*-Pro and *cis*-Pro. Most notably, Ser and Thr exhibited a substantially higher fraction of the *t* χ_1 rotamer for *cis*-Pro than *trans*-Pro; the *t* rotamer was previously⁵⁹ identified to be associated with C–H/O interactions in Ser-*cis*-Pro.

In order to identify the sidechain conformations associated with potential C–H/O interactions, all structures with O \cdots H α distances ≤ 2.72 Å were analyzed separately (Fig. 3, Fig. S3–S6). These results indicated strong conformational preferences when a C–H/O interaction was present. Glu and Gln exhibited C–H/O interactions predominantly with the g^- χ_1 rotamer, with the *t* χ_1 rotamer a minor conformation for C–H/O interactions with Glu (Fig. 3a). In contrast, for Asp and Asn, C–H/O interactions were associated predominantly with the *t* χ_1 rotamer. Similarly, for Ser and Thr, C–H/O interactions were observed almost exclusively with the *t* χ_1 rotamer.

Glu and Gln also showed a strong preference for the g^- χ_2 rotamer for C–H/O interactions, with Glu also exhibiting a small population with the g^+ χ_2 rotamer (Fig. 3b, Fig. S4). The Asp and Asn χ_2 rotamers (Fig. 3b, Fig. S5), and the equivalent Glu and Gln χ_3 rotamers (Fig. 3c, Fig. S4), showed a broader distribution between -60° and $+60^\circ$.

Individual structures from the PDB were analyzed for all of these identified combinations of conformations with potential C–H/O interactions (Fig. 4). All structures exhibited clear evidence of C–H/O interactions, including good interaction geometries. Analysis of these structures indicated two different interaction modes for Glu and Gln: a more common one where the Glx χ_1 and χ_2 rotamers were both g^- , and a less common one where the χ_1 rotamer was *t* and the χ_2 rotamer was g^+ . These results are consistent with analysis of sidechain torsion angle distributions in C–H/O interactions (Fig. 3). Similarly, Asp and

Asn each had two different combinations of χ_1 and χ_2 rotamers, (*t*, $0 \pm 90^\circ$) and ($\sim -100^\circ$, $\sim -30^\circ$). Ser and Thr exhibited C–H/O interactions through the *t* χ_1 rotamer. Close C–H/O interactions were observed in all type VI β -turn subtypes^{12–14} (VIa1 [PcisD], VIa2 [BcisD], and VIb [PcisP and BcisP]) (Fig. 5, Fig. S8, Tables S10 and S11). For Glu, Gln, and Ser, C–H/O interactions were relatively overrepresented in type VIa β -turns (Pro in the δ conformation), while for Asp, Asn, and Thr, C–H/O interactions were substantially more likely in BcisP type VIb β -turns (type VIb: Pro in the PPII/ β conformations) (seen by comparing conformation frequencies in Tables S9 versus S11).

All residues examined exhibited evidence of frequent and close C–H/O interactions with *cis*-Pro (Table 2 and Fig. 4), but substantially different frequencies of *cis*-Pro (Table 1). For example, Glu and Asp have essentially identical likelihoods of C–H/O interactions in *cis*-Pro (found in $\sim 40\%$ – 50% of all *cis*-Pro structures), but very different frequencies of *cis*-Pro. Asp has been previously shown to frequently interact *via* hydrogen bonds with the local protein backbone.⁸⁰ Indeed, numerous examples were found of Asp-*trans*-Pro structures with hydrogen bonds to the pre-Asp (*i* – 1), Asp (*i*), and post-proline (*i* + 2 and *i* + 3) amide hydrogens (Fig. 6). These sidechain–main chain hydrogen bonds inherently compete with other conformations, and could relatively promote the *trans*-Pro conformation in Asp-Pro due to the greater strength of hydrogen bonds, despite favorable C–H/O interactions that are observed with Asp-*cis*-Pro.

DFT calculations on C–H/O interactions in Ac–X-*cis*-Pro–NHMe peptides

In order to better understand the nature of C–H/O interactions in stabilizing specific conformations in *cis*-Pro, DFT calculations were conducted on model Ac–X-*cis*-Pro–NHMe peptides, X = Glu, Gln, Asp, Asn, Ser, Thr. Quantum mechanics-based calculations were used instead of force field-based calculations due to the inherently quantum mechanical nature of C–H/O interactions, and the absence of full parametrization of these interactions in common force fields. Indeed, by typical molecular mechanics-based approaches, the close approach of the Pro H α to the sidechain oxygen would be treated as a repulsive interaction, due to H \cdots O distances that are below the sum of the van der Waals radii of these atoms. These systematic biases against close approaches of atoms in protein structure determination software could also impact the refinement of protein crystal structures, and thus the apparent distances observed in the PDB.⁵³

While C–H/O interactions were observed in all type VI β -turn subtypes (Fig. 5, Table S11), calculations were conducted on those structures in type VIa1 (PcisD) β -turns (Tables S5 and S6). Unlike other type VI β -turn subtypes, type VIa1 β -turns exhibit an *i*/*i* + 3 C=O \cdots H \cdots N main-chain/main-chain hydrogen bond, which restrains the geometry and provides interaction partners for these main-chain hydrogen-bonding groups, simplifying the calculations and analysis.^{12–14} Geometry optimization calculations were conducted using structures from the PDB (Fig. 4) as initial models, examining all different

Table 3 Sidechain χ rotamer populations of the pre-proline (X) residue at X-*trans*-Pro and X-*cis*-Pro sequences in the PDB. Populations of each sidechain χ rotamer at X-Pro sequences are represented as the percent of structures in each Pro amide conformation. Conformations that were significantly overrepresented in the *cis*-proline conformation relative to the *trans*-proline conformation for a given amino acid are indicated in italics

Rotamer	Glu		Gln		Asp	Asn	Ser	Thr		
	χ_1	χ_2	χ_1	χ_2	χ_1	χ_1	χ_1	χ_1		
<i>trans</i>	g^-	–60	62.4	53.6	65.9	35.7	19.1	33.0	23.6	40.5
	g^+	+60	9.2	31.5	9.6	13.4	6.8	7.6	39.3	52.4
	<i>t</i>	± 180	28.4	14.8	24.5	50.8	74.0	59.4	37.1	7.1
<i>cis</i>	g^-	–60	71.7	41.9	75.2	10.9	29.3	32.7	28.7	33.8
	g^+	+60	5.7	18.1	5.5	45.5	7.8	3.4	15.7	45.1
	<i>t</i>	± 180	22.5	40.0	19.4	43.6	62.9	63.9	55.6	21.1



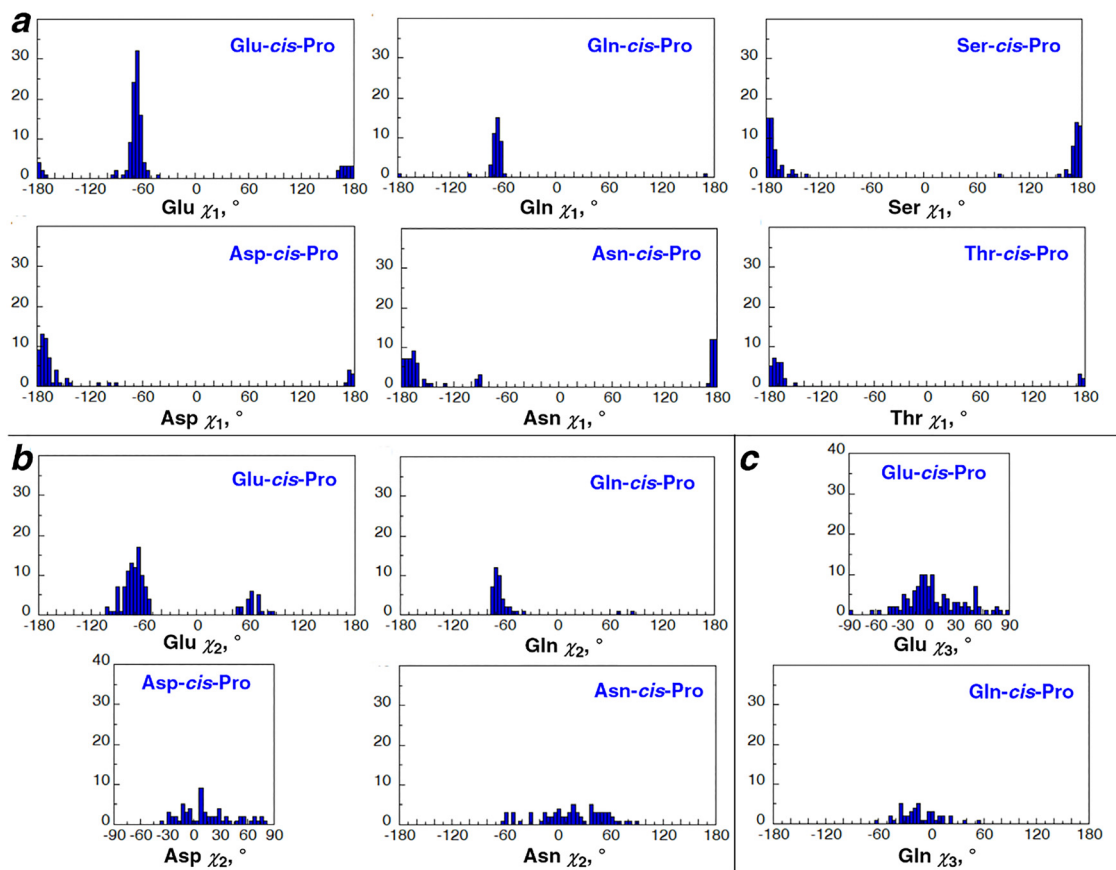


Fig. 3 Sidechain χ rotamers at pre-proline (X) residues in X-*cis*-Pro sequences in the PDB exhibiting C-H/O interactions with Pro C-H α . In structures with X-*cis*-Pro exhibiting C-H/O interactions (C_{α} -H \cdots O distances ≤ 2.72 Å), pre-proline residues were analyzed for sidechain χ dihedral angles at (a) χ_1 , (b) χ_2 , (c) χ_3 . C-H/O interactions at Asp-*cis*-Pro, Asn-*cis*-Pro, Ser-*cis*-Pro, and Thr-*cis*-Pro are mediated predominantly via the t χ_1 rotamer (t , $\chi \sim \pm 180^\circ$). A small number of C-H/O interactions at Asn-*cis*-Pro and Asp-*cis*-Pro sequences are observed with a χ_1 dihedral angle around -90° (7% and 5%, respectively). C-H/O interactions at Gln-*cis*-Pro are mediated predominantly via the g^- χ_1 rotamer (g^- , $\chi \sim -60^\circ$) and the g^- χ_2 rotamer, with a small number of C-H/O interactions at Gln-*cis*-Pro sequences mediated by the t χ_1 rotamer and the g^+ χ_2 rotamer (g^+ , $\chi \sim +60^\circ$) (5%). In contrast, C-H/O interactions at Glu-*cis*-Pro are mediated by either the combination of the g^- χ_1 rotamer and the g^- χ_2 rotamer (82%) or via the combination of the t χ_1 rotamer and the g^+ χ_2 rotamer (18%). Due to the symmetrical oxygens of carboxylates, the Glu χ_3 and Asp χ_2 dihedral angles were standardized to a range of -90° to $+90^\circ$.

combinations of sidechain torsion angles that exhibited C-H/O interactions.^{68–71,81}

As was seen in the PDB analysis, Glu and Gln exhibited two distinct combinations of conformations with C-H/O interactions, with $(\chi_1, \chi_2) = (g^-, g^-)$ or (t, g^+) (Fig. 7). Closer H \cdots O interactions were observed in the (g^-, g^-) sidechain rotamer pair for both residues, and these (g^-, g^-) structures were also lower in energy than the (t, g^+) structures by 1.3–1.5 kcal mol $^{-1}$ in these calculations. The (g^-, g^-) structures exhibited a more classical oxygen-based type of C-H/O interaction, with hydrogen bond-like geometries.

In contrast, the (t, g^+) structures exhibited longer O \cdots H distances, but had a geometry that is more typical of cation/ π or C-H/ π interactions,^{82–84} with the Pro H α interacting closely with the π face of the carboxylate or carboxamide (a geometry also observed in the PDB, Table S7). Indeed, these structures had very close $C_{=O}\cdots H\alpha$ distances (2.35–2.40 Å), well below the 2.90 Å sum of the van der Waals radii of C and H. C-H/O interactions with carbonyls may be mediated via either of the

oxygen lone pairs (O_s or O_p) or via the π molecular orbital serving as the electron donor that interacts with the C-H σ^* orbital.⁶⁵ The calculations indicate that these distinct interaction modes may be alternatively observed in different conformations of the Glu or Gln sidechains.

Notably, closer C-H/O interactions were observed with Glu than with Gln. These results are consistent with bioinformatics data (Table 2), which indicated a substantially greater likelihood of close C-H/O interactions for Glu than Gln. The closer interactions are likely due to the greater electron density on the anionic carboxylate oxygen of Glu than on the formally neutral carboxamide oxygen of Gln, along with the higher energy of the filled orbitals of the more electron-rich carboxylate than the less electron-rich carboxamide. Consistent with this interpretation, geometry optimization calculations on the neutral (acidic) form of Glu (Glu 0) demonstrated a significantly longer O \cdots H α distance for the neutral acid compared to the anionic carboxylate (Fig. 8). The neutral Glu 0 also exhibited longer C-H/O interaction distances than the Gln carboxamide (2.52 Å versus



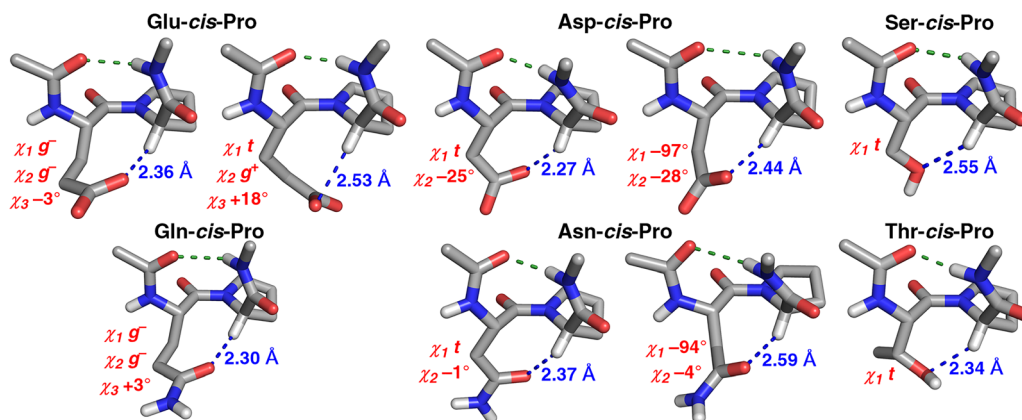


Fig. 4 Interaction modes of C–H/O interactions in type VIa1 β -turns in the PDB. Type VIa1 β -turns (PPII-*cis*- δ , PcisD) were analyzed at Glu-*cis*-Pro [pdb 7f82 (left) and 4yzo (right)], Asp-*cis*-Pro [pdb 4y7s (left) and 3ksx (right)], Ser-*cis*-Pro (pdb 4d0q), Gln-*cis*-Pro (pdb 3l8a), Asn-*cis*-Pro [pdb 4i79 (left) and 3oyv (right)], and Thr-*cis*-Pro (pdb 7u9u) sequences in the PDB. A structure of Gln in the type VIa1 β -turn conformation with the combination of $\chi_1 = t$, $\chi_2 = g^+$ was not observed, although interactions with this rotamer pair were observed in other type VI β -turn subtypes (VIb [BcisP] pdb 2jks, VIb [PcisP] pdb 3oyv). Additional geometric information is in Fig. S9.

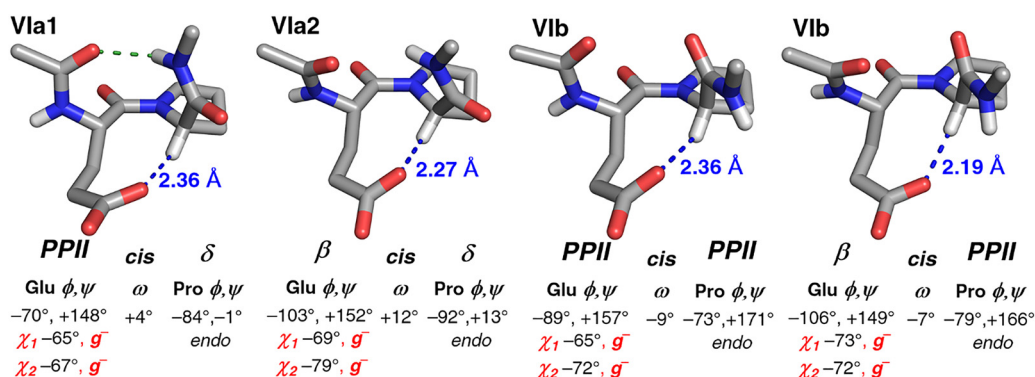


Fig. 5 Different type VI β -turns with C–H/O interactions at Glu-*cis*-Pro in the PDB. Protein structures with C–H/O interactions (blue) in different type VI β -turn subtypes at E-*cis*-P sequences (from left: PDB 7f82: type VIa1, PcisD; PDB 4lr2: type VIa2, BcisD; PDB 5deq: type VIb, PcisP; pdb 3wvn: type VIb, BcisP).

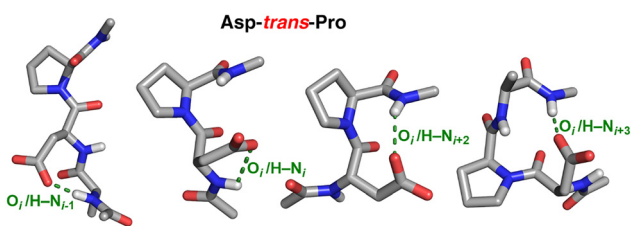


Fig. 6 Asp-*trans*-Pro structures in the PDB stabilized *via* local hydrogen bonds between main-chain amide hydrogens and the Asp carboxylate. (left to right) pdb 5ik4, pdb 5w2f, pdb 6a9w, and pdb 3cov.

2.31 Å), consistent with the more electron-rich nature (and higher-energy π molecular orbitals) of the amide compared to the acid.

Two interaction geometries were identified as stable energy minima for Asp, $(\chi_1, \chi_2) = (t, -10^\circ)$ or $(t, +80^\circ)$, while one was identified for Asn $(\chi_1, \chi_2) = (t, -75^\circ)$ (Fig. 7). The $(t, -10^\circ)$ geometry for Asp exhibited the closest $O \cdots H\alpha$ distance of any structure in these calculations, as well as a classical hydrogen

bond-like geometry, consistent with a very favorable C–H/O interaction. However, notably, this structure was also quite strained in the ω torsion angle ($\omega = +20^\circ$), which imposes an inherent torsional energy cost that would result in destabilization due to reduced electron delocalization in that amide bond. The Asp $(\chi_1, \chi_2) = (t, +80^\circ)$ structure and the Asn $(\chi_1, \chi_2) = (t, -75^\circ)$ structure had significantly longer $O \cdots H\alpha$ distances, but both exhibited a favorable C–H/ π interaction geometry, with close $C \cdots H\alpha$ distances, similar to that observed in the (t, g^+) rotamer of Glu and Gln. As was the case with Glu *versus* Gln, in structures with comparable geometries, closer interaction distances were observed for Asp than Asn, consistent with closer C–H/O interactions with more electron-rich anionic carboxylate electron donors than with neutral amide carbonyls.

C–H/O interactions in the Ser and Thr structures were mediated *via* the *t* rotamer, with $O \cdots H\alpha$ distances that were substantially below 2.72 Å (Fig. 7). Interestingly, a closer interaction was observed with Thr than Ser. These results are consistent with bioinformatics data indicating that, while Thr had the lowest overall frequency of C–H/O interactions in



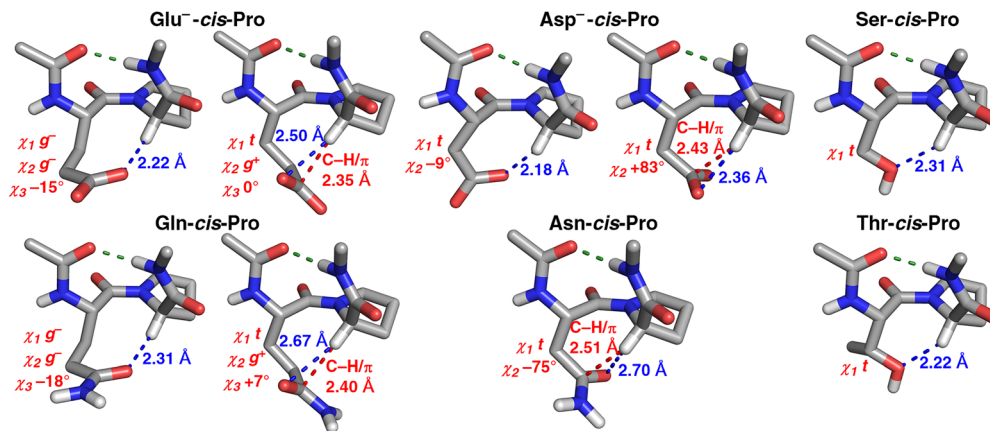


Fig. 7 C–H/O interactions with Pro C–H_α in geometry-optimized structures with X–*cis*-Pro amide bonds in type VIa1 β-turns. Structures from the PDB (Fig. 4) were modified to Ac–X–*cis*-Pro–NHMe and then subjected to geometry optimization to examine the conformations at pre-proline residues which can stabilize the *cis*-Pro conformation *via* C–H/O interactions (H_α···O distances, blue). Glu, Gln, Asp, and Asn all have one interaction mode where the nearest interaction is a C–H/π interaction between the amide or carboxylate π orbitals and Pro C–H_α (red). C–H/π interactions were defined as C_{C=O}···H distances at or below the sum of the van der Waals radii of C and H (≤2.90 Å). Conformations at Asp–Pro with Asp χ₁ = –100°, χ₂ = –30° and Asn–Pro with Asn χ₁ = –90°, χ₂ = –30° were observed with C–H/O interactions with Pro C–H_α in the PDB but optimized to alternative structures with intrasidic C–H/O interactions, and thus were not included. Additional geometric information is in Fig. S10.

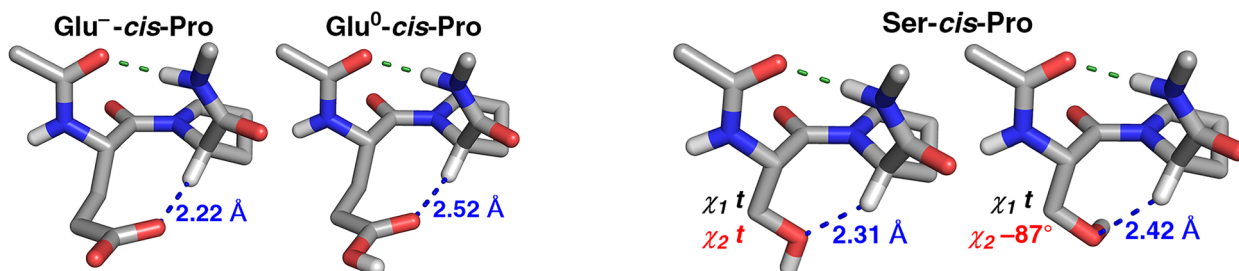


Fig. 8 Effect of Glu ionization state on C–H/O interactions in geometry-optimized structures of Glu–*cis*-Pro in a type VIa1 β-turn conformation. Structures of Ac–Glu–*cis*-Pro–NHMe in a type VIa1 β-turn conformation with either the (left) anionic or (right) neutral sidechain ionization state. Additional geometric information is in Fig. S12.

Fig. 9 Effects of Ser χ₂ rotamer on C–H/O interaction distance in Ser–*cis*-Pro. Structures of Ac–Ser–*cis*-Pro–NHMe in a type VIa1 β-turn conformation were geometry-optimized as a function of the χ₂ dihedral angle of Ser. The sidechain hydroxyl of both Ser and Thr residues does not have a reliably determined hydrogen position in structures in the PDB. When Ser χ₂ adopts the *t* rotamer, the C–H/O interaction with Pro C–H_α is significantly closer than when it adopts a χ₂ of –87°. Additional geometric information is in Fig. S11. Geometry optimization calculations on anionic Ser–*cis*-Pro show that the Ser oxyanion exhibits a closer C–H/O interaction than either structure with neutral Ser (2.23 Å, Fig. S13).

cis-Pro (<20% of Thr–*cis*-Pro had a C–H/O interaction, Table 2), Thr–*cis*-Pro C–H/O interactions that were present were particularly likely to be close (≤2.30 Å).

Calculations were conducted on Ser–*cis*-Pro with two different χ₂ torsion angles, which represent two different positions of the serine hydroxyl hydrogen and are not directly determinable in the PDB. Notably, the O···H_α distance depended on the χ₂ torsion angle (Fig. 9), and also depended on the Ser ionization state (Fig. S13), with a closer distance for the atypical anionic Ser. While anionic Ser is strongly thermodynamically disfavored, these results suggest that hydrogen bonding to the sidechain hydroxyl might modulate electron density on oxygen and impact the strength and geometry of C–H/O interactions in *cis*-Pro.

Calculations on the energetics and inherent bases of C–H/O interactions stabilizing *cis*-Pro

In order to further understand the underlying basis or bases for C–H/O interactions that stabilize *cis*-Pro, natural bond orbital (NBO) calculations^{74,75} were conducted on models derived from

all X–*cis*-Pro sequences. NBO calculations convert the wavefunction into localized molecular orbitals, which can be used to approximate the component molecular orbitals that contribute to stabilization *via* electron delocalization.

In Glu–*cis*-Pro and Gln–*cis*-Pro structures with (χ₁, χ₂) = (g[–], g[–]), NBO calculations indicated substantial stabilization due to interaction of the Pro C–H antibonding (σ*) orbital with two oxygen lone pairs of the carboxylate of Glu or the carbonyl of Gln (Fig. 10a). In contrast, in Glu–*cis*-Pro and Gln–*cis*-Pro structures with (χ₁, χ₂) = (t, g⁺), interactions with the oxygen lone pairs were greatly reduced (Fig. 10b). However, in this case, NBO calculations demonstrated greater interaction of the Pro C–H σ* orbital with the π molecular orbitals, consistent with the hypothesis above that these structures could be interpreted as being stabilized primarily *via* a C–H/π interaction mode.



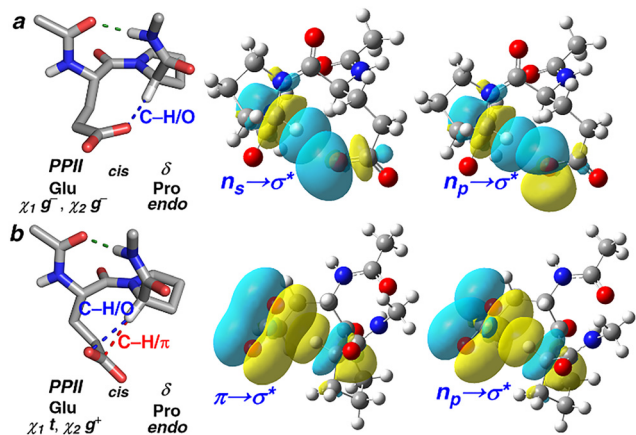


Fig. 10 Natural bond orbital (NBO) calculations on structures with C-H/O and C-H/ π interactions stabilizing Glu-*cis*-Pro. NBO calculations were performed on Glu-*cis*-Pro structures in type VIa1 β -turns with Glu (a) $\chi_1 = g^-$, $\chi_2 = g^-$ or (b) $\chi_1 = t$, $\chi_2 = g^+$, in order to understand the component molecular orbitals that contribute to stabilization of a C-H/O interaction. When Glu $\chi_1 = g^-$, $\chi_2 = g^-$, both the s-like (n_s , left) and p-like (n_p , right) lone pairs on the interacting Glu sidechain oxygen exhibit substantial orbital overlap with Pro C-H α σ^* , with the extent of orbital overlap correlating with the amount of electron delocalization. In contrast, when Glu $\chi_1 = t$, $\chi_2 = g^+$, the overlap between Pro C-H α σ^* and the Glu lone pairs is reduced (right), while the overlap with the 3-atom carboxylate π system is more significant (left). Depictions of global molecular orbitals derived from full molecular orbital calculations are in the SI (Fig. S14).

Full molecular orbital calculations, which properly address the global (rather than localized) nature of molecular orbitals, clearly demonstrate electron delocalization between the Glx carboxylate or carboxamide and the Pro C-H α , with through-space electron delocalization stabilizing these conformations (Fig. S14).

The wavefunctions from these calculated structures were also examined *via* Atoms in Molecules (AIM) analysis.^{76–78} All structures exhibited bond critical points (BCPs) between sidechain atoms and Pro H α (Fig. S16). These results are consistent with NBO calculations indicating through-space electron delocalization and partial covalency in these C-H/O interactions. The highest electron densities at the BCPs were for Glu⁻ in the (g^- , g^-) conformation, for Thr, and for Asp in the (t , -10°) conformation. These results are consistent with these conformations exhibiting the closest C-H/O interactions (Fig. 7). Notably, these electron densities were also similar to those observed in the model acetylene-water C-H/O interaction (Fig. S16).⁸⁵

In most structures, the BCP was located on an approximately linear path between the sidechain oxygen and Pro H α . However, for either Glu⁻ or Gln in the (t , g^+) conformation, and for Asn in the (t , -75°) conformation, the BCP was on paths between the Pro H α and the carbonyl carbon (Fig. S16). For Asp in the (t , $+80^\circ$), a curved path was observed. The results for these conformations are consistent with our analysis above, that these C-H/O interaction modes are more properly described as C-H/ π interactions, with electron delocalization between the conjugated π systems and σ^* of Pro C-H α .

In order to understand the energetics and the underlying primary bases for C-H/O interactions that stabilize *cis*-Pro, calculations were conducted comparing the structures identified above with structures in which one sidechain torsion angle was rotated away from the C-H/O interaction, with subsequent full geometry optimization (Fig. S15).⁷² For structures with Glu and Gln, the structures resulting from rotating the χ_2 torsion angle to t (180°) and subsequent geometry optimization placed the sidechain away from the backbone, allowing an estimation of the C-H/O interaction energy *via* the energy differences between the interacting and non-interacting rotamers. In contrast, in structures with Asp, Asn, Ser, and Thr, the resultant rotated structures exhibited a new hydrogen bond with the backbone, consistent with the high frequency of sidechain-backbone interactions observed for these amino acids, and precluding application of this approach.

For Glu, the (χ_1 , χ_2) = (g^- , g^-) rotamer exhibited an interaction energy of -3.0 kcal mol⁻¹ in implicit water using this approach of comparing energies of different rotamers (Table S12). In contrast, the Glu (χ_1 , χ_2) = (t , g^+) rotamer had an interaction energy of -1.8 kcal mol⁻¹, with this less favorable interaction energy consistent with its less frequent observation in the PDB. For Gln, the comparable energies were less favorable than those with Glu ($[g^-, g^-] = -2.4$ kcal mol⁻¹, $[t, g^+] = -1.4$ kcal mol⁻¹), consistent with stronger C-H/O interactions with Glu than with Gln. Finally, the (g^- , g^-) rotamer exhibited the weakest interaction energy (-2.0 kcal mol⁻¹) for neutral Glu⁰, consistent with the less electron-rich nature of a carboxylic acid (Glu⁰) compared to a carboxamide (Gln) and the longer O \cdots H α distances observed in geometry optimization calculations (Fig. 8). Two important caveats to these rotamer-based interaction energies are (1) that they do not address inherent differences in the energies of the conformations, though these are expected to be relatively small at χ_2 ; and (2) more importantly, that they do not fully address differences in solvation energies (*e.g.*, differences in competitive hydrogen bonds at the oxygen or between water molecules) and other solvation (*e.g.*, organized solvation at H α by water).

The interacting and non-interacting conformer pairs of Glu, Gln, and Glu⁰ were used to explore the roles of electrostatics *versus* stereoelectronic effects in the energetics of C-H/O interactions. CM5 calculations⁸⁶ of charge on the interacting oxygens and Pro H α atoms in these structures indicate relatively modest partial charges (Glu -0.49 , Gln -0.39 , Glu⁰ carbonyl O -0.34 ; Pro H α $+0.10$ to $+0.12$). These partial charges would be expected to be associated with minimal favorable energetics *via* electrostatics in water for these structures with C-H/O interactions. Moreover, considering the very close interactions of the sidechain carbonyl carbon with Pro H α in the (t , g^+) rotamers (C \cdots H α 2.35–2.40 Å, well below the 2.90 Å sum of van der Waals radii of these atoms), the CM5 charges on the carbonyl carbon ($+0.16$ in Glu and $+0.26$ in Gln) and Pro H α ($+0.10$ to $+0.12$) suggest that the very close approach of these atoms in a C-H/ π interaction mode should be unfavorable due to electrostatic repulsion between positively charged C and H atoms.



Geometry optimization calculations were conducted on both the interacting and non-interacting structures as a function of solvent dielectric constant (ϵ), examining the interaction distance and the energy difference between interacting and non-interacting rotamers in vacuum (gas phase) ($\epsilon = 1$), hexane ($\epsilon = 1.9$), CHCl_3 ($\epsilon = 4.7$), CH_2Cl_2 ($\epsilon = 8.9$), acetone ($\epsilon = 20.5$), acetonitrile ($\epsilon = 36$), DMSO ($\epsilon = 47$), H_2O ($\epsilon = 78$), and artificial solvent conditions with $\epsilon = 1000$, $\epsilon = 10\,000$, $\epsilon = 100\,000$, and $\epsilon = 1\,000\,000$. These latter high-dielectric-constant conditions will effectively fully screen out purely electrostatics-based interactions, since electrostatics interaction energy ($E_{\text{electrostatics}}$) scales as $E_{\text{electrostatics}} \sim 1/\epsilon$.

For Glu-*cis*-Pro structures with $(\chi_1, \chi_2) = (g^-, g^-)$, the closest $\text{O} \cdots \text{H}_\alpha$ distance (2.05 Å) and most favorable interaction energy ($-5.7 \text{ kcal mol}^{-1}$) were observed in vacuum, consistent with electrostatics being an important component of C-H/O interactions in vacuum, as well as the inherent destabilization of charge without counterbalancing opposite charges. In contrast, in all solvent conditions with $\epsilon \geq 4.7$ (CHCl_3 and all more polar conditions), the interaction distances ($2.22 \text{ Å} \pm 0.02 \text{ Å}$) and interaction energies (-2.7 to $-3.1 \text{ kcal mol}^{-1}$) were similar (Tables S13 and S14). Notably, among these conditions, the most favorable interaction energies were actually with $\epsilon = 1\,000\,000$, conditions that functionally screen out all electrostatics interactions. These results indicate that C-H/O interactions are highly favorable even under conditions with large dielectric constants, suggesting that outside of the most non-polar conditions (vacuum or hexanes), electrostatics play a minimal role in the strength of these C-H/O interactions. Moreover, calculations with neutral oxygen sources (Gln or Glu⁰), in the $(\chi_1, \chi_2) = (g^-, g^-)$ rotamer, indicate that the $\text{O} \cdots \text{H}_\alpha$ interaction distances ($2.305 \text{ Å} \pm 0.01 \text{ Å}$ for Gln, $2.38 \text{ Å} \pm 0.01 \text{ Å}$ for Glu⁰) and interaction energies ($-2.3 \pm 0.2 \text{ kcal mol}^{-1}$ for Gln, $-1.96 \pm 0.13 \text{ kcal mol}^{-1}$ for Glu⁰) have minimal solvent dependence. For example, the $\text{O} \cdots \text{H}_\alpha$ distance for Gln was

nearly identical in vacuum ($\epsilon = 1$) (2.313 Å) and in $\epsilon = 1\,000\,000$ (2.315 Å), as was also the case for Glu⁰ (2.382 Å versus 2.387 Å).

In structures of Glu⁻ or Gln with $(\chi_1, \chi_2) = (t, g^+)$, the interaction energies and geometries similarly showed minimal dependence on solvent dielectric constant in all conditions other than calculations in vacuum or hexane for Glu⁻. The close $\text{C}_\pi \cdots \text{H}_\alpha$ interaction distances ($2.36 \text{ Å} \pm 0.03 \text{ Å}$ for Glu⁻, $2.40 \text{ Å} \pm 0.01 \text{ Å}$ for Gln; both of these are $\sim 0.5 \text{ Å}$ below the 2.90 Å sum of the van der Waals radii of C and H) also exhibit minimal dependence on the solvent dielectric constant. These results provide further evidence that C-H/O interactions and C-H/ π interactions are fundamentally stereoelectronic in nature, and that the molecular orbital basis of these interactions renders them functionally solvent-independent for their inherent interaction energies.

C-H/O interactions in pSer-*cis*-Pro and pThr-*cis*-Pro

Ser/Thr phosphorylation leads to an increased barrier for proline *cis-trans* isomerization at pSer-Pro and pThr-Pro sequences.³⁹⁻⁴¹ Recently, we proposed the possibility that this increased barrier to isomerization was due to the combination of (a) stabilization of the ground state of *trans*-Pro via a phosphate-amide hydrogen bond⁷³ and (b) stabilization of the ground state of *cis*-Pro via a phosphate-Pro C-H α C-H/O interaction,^{60,61} with neither of these interactions present and/or impacting the energy of the transition state of isomerization. In order to explore this hypothesis, we conducted calculations on pSer-*cis*-Pro and pThr-*cis*-Pro structures, in order to examine potential C-H/O interactions with the phosphate that could stabilize *cis*-Pro (Fig. 11). Calculations were conducted in the physiologically most important dianionic ionization state, in the monoanionic state that can be a significant population in solution, and in the non-physiologically relevant neutral form (Fig. 11).

We identified two different pairs of sidechain conformations that exhibit phosphate-proline C-H/O interactions,

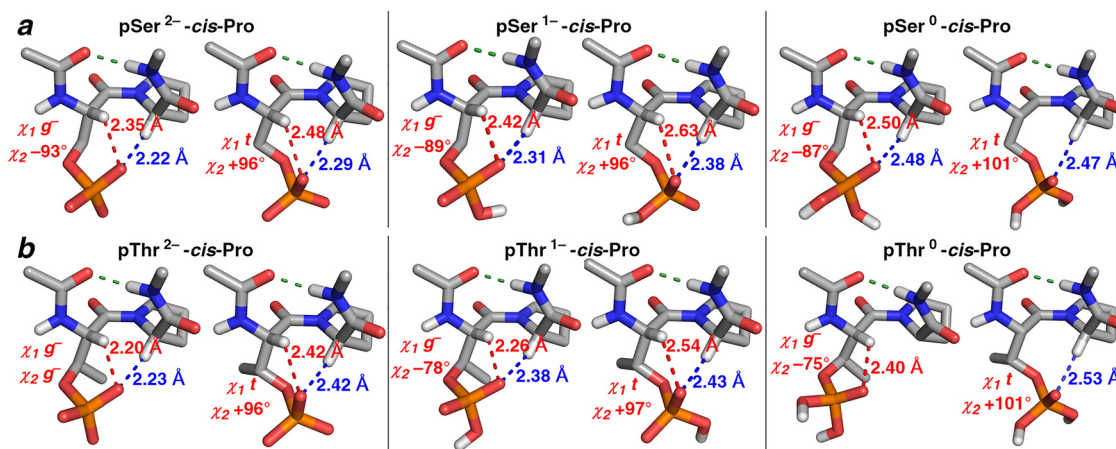


Fig. 11 C-H/O interactions in pSer-*cis*-Pro and pThr-*cis*-Pro as a function of phosphate ionization state. Geometry-optimized structures of (a) Ac-pSer-*cis*-Pro-NHMe and (b) Ac-pThr-*cis*-Pro-NHMe in a type VIa1 β -turn conformation in different ionization states. pSer-*cis*-Pro tends to optimize to include a C-H/O interaction with Pro C-H α (blue). Longer bidentate interactions with both Pro C-H α (blue) and Ser C-H α (red) are also observed in some structures. Conversely, pThr-*cis*-Pro tends to include bidentate C-H/O interactions with both Thr C-H α (red) and Pro C-H α (blue) that have similar $\text{H}_\alpha \cdots \text{O}$ distances. Additional geometric information is in Fig. S17.



$(\chi_1, \chi_2) = (g^-, -90^\circ)$ or $(t, +95^\circ)$. For both pSer and pThr, closer $H\alpha \cdots O$ distances were observed in the $(g^-, -90^\circ)$ rotamer pair. Closer $H\alpha \cdots O$ distances were also observed in the dianionic form than in the monoanionic form, with both closer than with the neutral phosphate. Due to multiple Lewis basic oxygens in the phosphate, C–H/O interactions were observed to both the Pro C–H α and to the pSer/pThr C–H α . Notably, even in structures with the neutral phosphate, close C–H/O interactions were observed, at Pro C–H α and/or at the pSer/pThr H α . In addition, AIM analysis demonstrated BCPs between a phosphate oxygen and Pro H α in all of these structures (Fig. S18). These results are consistent with the hypothesis that C–H/O interactions can stabilize *cis*-proline when preceded by pSer or pThr, and that *cis*-proline-stabilizing C–H/O interactions contribute to the higher activation barrier for proline *cis*–*trans* isomerization at these sites in proteins.

C–H/O interactions in *cis*-non-proline amide bonds

0.03% of non-proline amide bonds are in the *cis*-amide conformation.¹¹ Because *cis*-non-proline amide bonds are inherently ~ 5 kcal mol^{−1} higher in energy than those of *trans*-non-proline,^{3,20} their observation in proteins requires stabilization of the *cis* amide by local and/or global structures. The low frequencies of *cis*-non-proline amide bonds preclude bioinformatics analysis as conducted above.⁸⁷ However, human phospholipase D exhibits a particularly close C–H/O interaction between Gly C–H α and the Asp carboxylate, which we propose to stabilize the *cis*-Gly conformation (Fig. 12).⁸⁸ We had previously observed⁵⁹ that C–H/O interactions stabilize a Ser–*cis*-Arg conformation that is widely conserved in bacterial xanthine–guanine phosphoribosyl-transferases.^{89,90} While not supported by the power of statistics, these structures suggest that C–H/O interactions can be one component stabilizing *cis*-non-proline conformations.

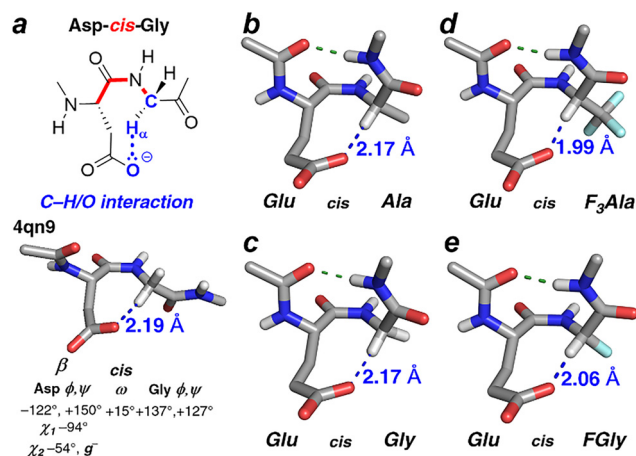


Fig. 12 Stabilization of non-proline *cis*-amide bonds via C–H/O interactions. (a) X-ray crystal structure of pdb 4qn9 containing an Asp–*cis*-Gly sequence with a close C–H/O interaction ($H\alpha \cdots O$ distance 2.19 Å, blue). (b)–(e) Results of geometry-optimization calculations on the X–*cis*-nonPro sequences (b) Glu–*cis*-Ala, (c) Glu–*cis*-Gly, (d) Glu–*cis*-F₃Ala, and (e) Glu–*cis*-FGly in the type VIa1 β -turn conformation. Geometry-optimized structures of Glu–*cis*-Val, Glu–*cis*-Ser, Glu–*cis*-Tle, Glu–*cis*-FAla, and Glu–*cis*-F₂Ala are in the SI (Fig. S21 and S22).

In order to further explore this possibility, we conducted DFT calculations on Ac–X–*cis*-Ala–NHMe structures, X = Glu, Gln, Asp, Asn, Ser, Thr, pSer, and pThr. All residues exhibited C–H/O interactions between a sidechain oxygen and Ala C–H α , with $H\alpha \cdots O$ distances and interaction geometries similar to those observed with Pro (Fig. 12b and Fig. S19, S20). Calculations were also conducted on Ac–Glu–*cis*-Z–NHMe structures, Z = Ala, Gly, Ser, Val, *tert*-leucine (Tle), in order to examine the dependence of the interaction on the residue identity at the *cis* amide. All structures exhibited close C–H/O interactions, with the shortest $H\alpha \cdots O$ distances for Ser and the longest for Val (Fig. 12b, c, Fig. S21, Table S16).

In order to test the ability to electronically tune the strength of C–H/O interactions *via* inductive effects, Ac–Glu–*cis*-Z–NHMe structures with Z = fluorinated amino acids β -F-Ala, β, β -F₂-Ala, β, β, β -F₃-Ala, and α -F-Gly were also examined computationally. Consistent with structure being driven by favorable C–H/O interactions, introduction of fluorines led to closer $H\alpha \cdots O$ distances,^{64,91} with the structure with trifluoroalanine exhibiting a 1.99 Å $H\alpha \cdots O$ distance, comparable to that of typical hydrogen bonds (Fig. 12d, e, Fig. S22, Table S17).

Discussion

cis-Proline amide bonds are associated with substantial changes in protein structure, dynamics, and function.^{1–6} However, there is an incomplete understanding of local structural effects that stabilize *cis*-proline. We previously identified that *cis*-proline in Ser–Pro sequences is stabilized by a C–H/O interaction between the sidechain Ser oxygen and the proline C–H α .⁵⁹

Herein, we have more broadly examined the possibility of C–H/O interactions stabilizing *cis*-proline amide bonds *via* sidechain oxygen lone pairs interacting with Pro C–H α bonds. Bioinformatics analysis of *cis*-proline amide bonds at Glu–Pro, Gln–Pro, Asp–Pro, Asn–Pro, Ser–Pro, and Thr–Pro sequences indicate high frequencies of structures with $O \cdots H\alpha$ distances at or below the 2.72 Å sum of the van der Waals radii of O and H, with a significant number of structures with $O \cdots H\alpha$ distances < 2.30 Å.

Analysis of structures with C–H/O interactions, combined with DFT calculations on these structures, identified two interaction modes of the sidechain with *cis*-proline for Glu, Gln, Asp, and Asn: one with an oxygen lone pair-directed C–H/O interaction ($[\chi_1, \chi_2] = [g^-, g^-]$ for Glu and Gln, $[\chi_1, \chi_2] = [t, \sim 0^\circ]$ for Asp, Asn), and one with a C–H/ π -type interaction geometry of the Pro C–H α with the C=O π molecular orbitals ($[\chi_1, \chi_2] = [t, g^+]$ for Glu and Gln, $[\chi_1, \chi_2] = [t, \sim -80^\circ]$ for Asp, and $[\chi_1, \chi_2] = [t, \sim +80^\circ]$ for Asn). C–H/O interactions with Ser and Thr are mediated *via* the t χ_1 rotamer.

These C–H/O interactions were further investigated computationally, in order to understand the bases for these interactions. In water, the primary basis of interaction strength is not an electrostatics interaction between the partial negative charge on the oxygen and the partial positive charge on the polarized hydrogen. Instead, C–H/O interactions are



fundamentally driven by through-space electron delocalization, a molecular orbitals-based effect that leads to orbital mixing between electron-rich oxygen lone pair orbitals (n_s or n_p) on the sidechain oxygen or the π orbitals on the carbonyl and the proline C–H antibonding orbital (σ^*), as $n_O \rightarrow \sigma^*_{C-H}$ or $\pi_{C=O} \rightarrow \sigma^*_{C-H}$ interactions. These quantum-mechanical effects on structure are currently not well addressed by typical molecular mechanics/force field-based approaches.

More broadly, because the interaction strengths of stereo-electronic effects are inherently less dependent on solvent than are electrostatics interactions, they have the potential to impact protein structures and interactions in diverse contexts and in ways that might be underappreciated.^{64,65} Considered only in electrostatics terms, C–H/O interactions should be insignificant in water, given the very modest partial charges present on (for example) protein H α ($\sim +0.1$). Moreover, at the sidechain oxygen atoms, a C–H/O interaction is weaker than a hydrogen bond of those oxygen atoms to water.⁹² However, the C–H/O interaction also obviates the need for water solvation at Pro H α , allowing the released water molecules to engage in more favorable water–water hydrogen bonds. C–H/O interactions also convert hydrogen bonds of water with the sidechain oxygen to water–water hydrogen bonds that are likely similar in strength. In addition, by releasing water molecules at both sites, the C–H/O interaction also has potential advantages in translational entropy. Notably, in membrane environments, C–H/O interactions can occur without a water desolvation energy cost.^{93,94} Globally, these questions can be properly addressed in future QM/MM simulations that explicitly include the quantum mechanical nature both of C–H/O interactions and of hydrogen bonds to water.^{95,96}

Glu–Pro sequences have previously been identified to have a significantly higher than average frequency of the *cis*-Pro conformation.^{8,9} However, no basis for this higher frequency of *cis*-Pro has been identified. The data herein suggest that C–H/O interactions between the Glu carboxylate and Pro H α specifically stabilize the *cis*-Pro conformation relative to the *trans*-Pro conformation, leading to an increased population of *cis*-Pro at Glu–Pro sequences. C–H/O interactions also impact the conformational ensemble present in structures with *cis*-Pro. More broadly, C–H/O interactions are one of a series of competing sidechain–mainchain interactions that impact conformations in both *trans*-Pro and *cis*-Pro.

Intrinsically disordered proteins (IDPs) and unfolded states of proteins exhibit mixtures of *trans*-Pro and *cis*-Pro at proline sites.^{2,4,10,16} In contrast, in folded proteins, typically a single amide conformation is present at each proline, due to the constraints of tertiary structure and the differences in local structure in *trans*-Pro versus *cis*-Pro.²³ The work herein provides insights into transient structures that can be present when the *cis*-Pro conformation is present in IDPs, as well as local sequence elements that can promote *cis*-Pro.

DFT calculations also demonstrated that *cis*-proline in phosphoserine–proline and phosphothreonine–proline sequences can be stabilized by favorable C–H/O interactions mediated by the electron-rich phosphate group. This stabilization of the

cis-proline conformation by C–H/O interactions (relative to the transition state for isomerization, where these interactions are not present) provides an explanation for the higher barrier and the significantly slower rate of proline *cis*–*trans* isomerization at these sites in proteins.⁴⁰

The strength of the C–H/O interactions at pSer–Pro and pThr–Pro sites, as well as at Glu–Pro and Asp–Pro sites, is impacted by ionization state (charge).^{66,97} Phosphorylated amino acids typically exist primarily in the dianionic forms, but significant populations of the monoanionic forms can also be present or predominant.^{73,98,99} Notably, local environment impacts the pK_a values of ionizable groups.^{100,101} The results herein provide a context by which changes in local environment could impact proline *cis*–*trans* isomerism at these sites *via* changes in sidechain ionization state.

Slower proline *cis*–*trans* isomerization at pSer–Pro and pThr–Pro sites is a central element both in transcription (*via cis*–*trans* isomerization in the Pol II CTD) and in cell cycle progression and cell division (*via* numerous proteins).^{39,47–49} It has also been implicated in misfolding of the *tau* protein in Alzheimer's disease. The phosphorylation-dependent prolyl isomerase Pin1/Ess1 is critical in these processes.^{41,42,44,45} The identified relevant Ser/Thr phosphorylation sites in these proteins are all in IDPs and/or in intrinsically disordered regions (IDRs) of proteins. In addition, other prolyl isomerases have been identified to impact the phase separation behavior and aggregation of IDPs and IDRs.^{44,102–104} The results herein suggest that phosphate–Pro H α C–H/O interactions are important to the dynamics of these proline *cis*–*trans* isomerization events. Moreover, these results suggest that transient protein structures or protein–protein interactions that engage with sidechain phosphates or with other sidechain oxygens could impact the structures and dynamics of the conformational ensemble, and thereby change protein function.

Finally, all encoded amino acids have a C–H α equivalent to that in proline. We demonstrate computationally that sidechain–main chain C–H/O interactions are also capable of stabilizing *cis* amide bonds at non-proline residues. Collectively, these results suggest that C–H/O interactions between a sidechain oxygen and the main-chain C–H α at the subsequent residue can stabilize a *cis* amide bond both at proline and at non-proline residues, and that the interaction strength is dependent on the identity of both residues.

Conflicts of interest

There are no conflicts to declare.

Data availability

Summaries of key data are included in the main manuscript. Additional data are in the supplementary information (SI), including additional bioinformatics analysis and the coordinates for all structures determined computationally.



Supplementary information is available. See DOI: <https://doi.org/10.1039/d5cp03423j>.

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