# 4,4-Difluoroproline as a Unique <sup>19</sup>F NMR Probe of Proline Conformation

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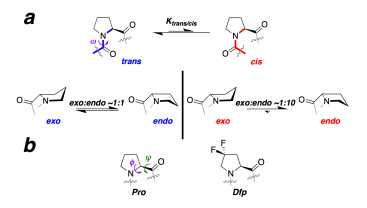
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#### **Abstract**

Despite the importance of proline conformational equilibria (trans versus cis amide, exo versus *endo* ring pucker) on protein structure and function, there is a lack of convenient ways to probe proline conformation. 4,4-Difluoroproline (Dfp) was identified to be a sensitive <sup>19</sup>F NMRbased probe of proline conformational biases and of cis-trans isomerism. Within model compounds and disordered peptides, the diastereotopic fluorines of Dfp exhibit similar chemical shifts ( $\Delta\delta_{FF} = 0$ –3 ppm) when a trans X–Dfp amide bond is present. In contrast, the diastereotopic fluorines exhibit a large ( $\Delta\delta_{FF} = 5-12$  ppm) difference in chemical shift in a cis X-Dfp prolyl amide bond. DFT calculations, X-ray crystallography, and solid-state NMR spectroscopy indicated that the  $\Delta\delta_{FF}$  directly reports on the relative preference of one proline ring pucker over the other: a fluorine which is pseudo-axial (i.e. the pro-4R-F in an exo ring pucker, or the pro-4S-F in an endo ring pucker) is downfield, while a fluorine which is pseudo-equatorial (i.e. pro-4S-F when exo, or pro-4R-F when endo) is upfield. Thus, when a proline is disordered (a mixture of exo and endo ring puckers, as at trans-Pro in peptides in water), it exhibits a small  $\Delta\delta$ . In contrast, when the Pro is ordered (i.e. when one ring pucker is strongly preferred, as in cis-Pro amide bonds, where the endo ring pucker is strongly favored), a large  $\Delta\delta$  is observed. Dfp can be used to identify inherent induced order in peptides and to quantify proline cis-trans isomerism. Using Dfp, we discovered that the stable polyproline II helix (PPII) formed in the denatured state (8 M urea) exhibits essentially equal populations of the exo and endo proline ring puckers. In addition, the data with Dfp suggested the specific stabilization of PPII by water over other polar solvents. These data strongly support the importance of carbonyl solvation and  $n \rightarrow \pi^*$ interactions for the stabilization of PPII. Dfp was also employed to quantify proline cis-trans isomerism as a function of phosphorylation and the R406W mutation in peptides derived from the intrinsically disordered protein tau. Dfp is minimally sterically disruptive and can be incorporated in expressed proteins, suggesting its broad application in understanding proline cistrans isomerization, protein folding, and local order in intrinsically disordered proteins.

## Introduction

Proline exhibits two key conformational equilibria: trans versus cis amide bond isomerism at the  $\omega$  torsion angle, and the exo versus endo pyrrolidine ring pucker (Figure 1). Proline ring pucker correlates with main chain conformation, with the exo ring pucker promoting more compact values of  $\phi$  and  $\psi$  and the endo ring pucker associated with more extended values. Proline cis-trans isomerization is often a rate-limiting step in protein folding, can function as a molecular timer, and has been implied to have a role in some protein misfolding diseases (e.g. the role of the phosphorylation-dependent prolyl isomerase Pin1 in tau aggregation in Alzheimer's disease). Thus, the development of new approaches to rapidly and site-specifically identify cis versus trans proline amide bonds in solution could have broad applications in understanding protein folding and structure.



**Figure 1**. (a) Summary of conformational equilibria and preferences of the proline (top) amide bond and (bottom) ring pucker. (b) Comparison of proline with 4,4-difluoroproline (Dfp).

Proline *cis-trans* isomerism is typically identified by the presence of an extra set of resonances in <sup>1</sup>H NMR spectroscopy (or sets of resonances, if multiple prolines are present). However, within globular proteins, only a single proline amide conformation is typically observed, so the number of resonances does not provide insights into the identity of the proline amide rotameric state. Residue-specific information on the identity of proline *cis* versus *trans* 

amide bond conformation, in peptides and proteins, can be provided by <sup>13</sup>C NMR chemical shifts ( $\delta$ ): in proline residues with *trans* amide bonds, the <sup>13</sup>C chemical shifts ( $\delta$ ) of C $\beta$  and C $\gamma$  are similar ( $\Delta\delta_{\beta\gamma}$  typically 1-4 ppm), whereas in cis proline amide bonds, these chemical shifts exhibit greater dispersion ( $\Delta\delta_{\beta\gamma}$  5-10 ppm).<sup>7-13</sup> Indeed, in the absence of X-ray crystallography, the  $\Delta\delta_{\beta\gamma}$  is the best-resolved indicator of cis versus trans amide bond rotameric state in larger proteins, and can be directly observed from <sup>1</sup>H-<sup>13</sup>C HSQC spectra. The alternative observation of interresidue NOEs between H $\alpha$  of the residue before proline and either Pro H $\alpha$  (in cis amide bonds) or Pro Hδ (in trans amide bonds) can become problematic due to spectral overlap with other proline resonances or with the water resonance, in addition to requiring assignment of all key resonances. The problem of spectral overlap is particularly severe in proline-rich sequences or in intrinsically disordered regions of proteins, where each residue typically exhibits similar chemical shifts (substantial spectral overlap/low chemical shift dispersion) and where standard sequential residue assignment (via NOEs between an amide hydrogen and the Hα of the prior residue, or between  $H\alpha$  and the amide hydrogen of the subsequent residue) breaks down. <sup>12,14-16</sup>

The challenge of identification of amide bond conformational state is not unique to NMR spectroscopy. Indeed, surveys of the PDB have found that cis amide bonds are commonly misidentified as trans amide bonds in lower resolution structures (especially > 2 Å), with the percentage of proline amides in a cis conformation observed to be greater in higher-resolution structures than in lower-resolution structures.<sup>17,18</sup>

We previously described a method (termed proline editing) for the site-specific modification of proline residues within peptides. Proline editing allows the modification within peptides of a 4R-hydroxyproline residue to any of a series of 4-substituted prolines. One

amino acid synthesized via proline editing was 4.4-difluoroproline (Dfp) (Figure 1b).<sup>22</sup> Moroder and others have demonstrated that 4,4-difluoroproline exhibits conformational preferences similar to those of proline (i.e. a minimal effect of Dfp compared to Pro on exolendo ring pucker or trans/cis amide conformational equilibria in peptides). 19,21,23-30 4,4-Difluoroproline can also be incorporated in expressed proteins via proline auxotrophic bacteria, with expression resulting in proteins with Dfp described for barstar, elastin-mimetic peptides, thioredoxin, and insulin. 23,31-34 Dfp may also be incorporated into proteins via native chemical ligation or expressed-protein ligation, as demonstrated in β2-microglobulin.<sup>35</sup> Dfp has also been incorporated by direct peptide synthesis in collagen-mimetic peptides.<sup>24</sup> In barstar and collagen-mimetic peptides, variants with Dfp had stabilities similar to those with proline. In thioredoxin and β2-microglobulin, decreased stability to denaturation was observed, potentially due to steric issues via observed interactions with proline. In both cases, a similar global protein structure (as determined by HSQC experiments on β2-microglobulin) or protein folding kinetics (thioredoxin) was present. There is also a structure of an insulin derivative with Dfp in the PDB (5uu3), although no data on the impacts of Dfp on insulin stability were described.<sup>33</sup>

In the course of our work on proline editing, we examined the <sup>19</sup>F NMR spectra of the peptides Ac-TYDfpN-NH<sub>2</sub> and Ac-TADfpN-NH<sub>2</sub> (Figure 12 of ref. <sup>21</sup>). In the *trans* amide conformation, both diastereotopic fluorines exhibited similar chemical shifts ( $\Delta\delta_{FF} = 0$  to 1.3 ppm); indeed, in the peptide Ac-TYDfpN-NH<sub>2</sub>, only a single peak was observed, due to  $\Delta\delta_{FF}$  (in Hz) <  $^2J_{FF}$ .<sup>36</sup> In contrast, in the *cis* amide conformation of both peptides, a large chemical shift dispersion of the fluorines was observed ( $\Delta\delta_{FF} \sim 8$ –10 ppm). Raines observed similar <sup>19</sup>F NMR data for Ac-Dfp-OMe in CDCl<sub>3</sub> and in D<sub>2</sub>O ( $\Delta\delta_{FF}$  (*trans*) 0.4–4.0 ppm,  $\Delta\delta_{FF}$  (*cis*) = 4.6–8.5 ppm).<sup>24</sup> These data suggested the possibility of using 4,4-difluoroproline as a probe of proline

cis-trans isomerization in peptides and proteins, analogous to using  $\Delta\delta_{\beta\gamma}$ , but with the greater sensitivity and simplified spectra of <sup>19</sup>F NMR compared to <sup>13</sup>C NMR. <sup>37-45</sup>

In related work, Hilvert and coworkers described the use of a site-specifically incorporated Dfp residue to probe the mechanism of protein misfolding and amyloid formation in  $\beta$ 2-microglobulin. In this work, they focused on the use of Dfp to reduce the activation barrier for proline *cis-trans* isomerization.<sup>35</sup> However, by <sup>19</sup>F NMR spectroscopy, both in peptides and within the protein, in both the folded and the denatured states, they observed proline rotamer-dependent chemical shift differences between diastereotopic fluorines ( $\Delta\delta_{FF} = 4$ –14 ppm in *cis* amides,  $\Delta\delta_{FF} < 1$  ppm in *trans* amides). Collectively, these results suggested the ready distinction between *cis* and *trans* amide bonds via 4,4-difluoroproline. Therefore, we sought to more generally examine the utility of Dfp as a conformational probe, via the site-specific incorporation of 4,4-difluoroproline into a series of model compounds and peptides. This work is intended to determine the principles for the interpretation of the <sup>19</sup>F NMR spectra of Dfp, in order to enable subsequent applications to peptides and proteins containing Dfp. In addition, we apply Dfp within a proline-rich peptide context, in order to address outstanding questions on the roles of solvation in PPII structure and on the structure of proline in the urea-denatured state.

# **Experimental**

**Small molecule synthesis**. **3**, **6**, and **7** were synthesized from commercially available **1** using variations on previously described approaches. Synthesis, purification, and characterization details are in the Supporting Information.

X-ray crystallography. 3 crystallized via slow evaporation of a solution of 3 in CDCl<sub>3</sub>. Details of crystal structure determination are in the Supporting Information. The crystallographic

information files of **3** have been deposited with the Cambridge Crystallographic Data Centre under CCDC 1566681 (data collection at 200 K) and CCDC 2279640 (data collection at 100 K).

**Peptide synthesis**. Peptides were synthesized by standard Fmoc solid-phase peptide synthesis and purified to homogeneity. Synthesis, purification, and conformational analysis of the peptide Ac-TYDfpN-NH<sub>2</sub> were described previously.<sup>19,21</sup> Synthesis, purification, and characterization details of other peptides are in the Supporting Information.

<sup>19</sup>**F NMR spectroscopy**. Solution-state <sup>19</sup>F NMR spectroscopy was conducted on a Brüker 400 MHz NMR spectrometer with a BBFO probe or with a cryogenic QNP probe. Resonances were referenced internally to residual TFA ( $\delta = -76.55$  ppm). Solid-state magicangle spinning <sup>19</sup>F NMR spectroscopy was conducted on a Brüker AVIII 500 MHz NMR spectrometer with a 4 mm Brüker HX probe, using spinning rates of 13 kHz and 15 kHz. Solid-state <sup>19</sup>F NMR spectra were referenced externally with 5-F-tryptophan. Additional data and analysis, including <sup>1</sup>H NMR data on compounds, expanded views of <sup>19</sup>F NMR spectra, and compilations of spectral data, are in the Supporting Information.

**Circular dichroism**. CD spectra were recorded on a Jasco J-810 or a Jasco J-1500 spectropolarimeter. Data represent the average of at least three independent trials. Data were averaged but were not smoothed.

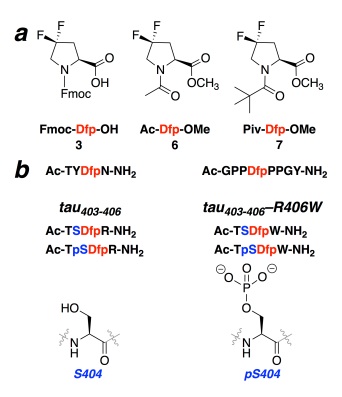
Computational chemistry. Calculations were conducted with Gaussian 09.<sup>46</sup> Geometry-optimized structures of Ac-Dfp-OMe, Piv-Dfp-OMe, Ac-Dfp-NMe<sub>2</sub>, and Piv-Dfp-NMe<sub>2</sub> were generated, with variation of main-chain conformation (α versus PPII), ring pucker (*exo* versus *endo*), proline amide rotamer (ω torsion angle, *trans* versus *cis*), and implicit solvent (H<sub>2</sub>O versus CHCl<sub>3</sub>). Final geometry optimization calculations were conducted using the M06-2X DFT functional and the 6-311++G(3d,3p) basis set with IEFPCM implicit solvation.<sup>47-49</sup> NMR

shielding parameters were then calculated using the GIAO method and the same combinations of functional, basis set, and solvation model. <sup>50,51</sup> <sup>19</sup>F chemical shifts in Dfp were referenced to those of TFA, based on the average calculated chemical shifts of the fluorines in TFA, which had been geometry optimized using the same methods as molecules with Dfp. Calculations on 3•CHCl<sub>3</sub> were conducted using the coordinates from the crystal structure file, with the positions of the heavy atoms fixed (optimization of H atoms only) based on those observed crystallographically, or alternatively with the positions of the heavy atoms other than fluorine fixed (optimization of H and F atoms). The minimal version of 3 (EtOCO-Dfp-OH) was generated via computational removal of the Fmoc aromatic carbon atoms and the addition of hydrogen atoms on the missing valences in the resulting ethyl group within GaussView, followed by restrained geometry optimization and calculation of the NMR shielding parameters. Additional details of calculations are in the Supporting Information.

#### **Results**

Synthesis of model compounds and peptides containing 4,4-difluoroproline. In order to test the applicability of Dfp as a conformational probe, a series of distinct model compounds and peptide contexts was considered (Figure 2). Dfp was examined within the simple model compounds Ac-Dfp-OMe and Piv-Dfp-OMe (Piv = pivaloyl). The pivaloyl group was employed based on our prior work to understand the role of  $n\rightarrow\pi^*$  interactions on conformation using X–Hyp(4-NO<sub>2</sub>-Bz)-OMe (X = acyl capping group) compounds. There, we found a significant increase in order using pivaloyl capping groups, including a stronger  $n\rightarrow\pi^*$  interaction and strong promotion of PPII and of the *trans*-proline amide conformation.<sup>52,53</sup> In addition, we also

examined peptides in which we have previously explored the impact of residue identity on proline *cis-trans* isomerism using <sup>1</sup>H NMR spectroscopy, in order to compare the results obtained via <sup>1</sup>H NMR spectroscopy versus via <sup>19</sup>F NMR spectroscopy with Dfp. Peptides examined included Ac-TY*Prox*N-NH<sub>2</sub> (*Prox* = 4-substituted proline) model peptides previously used to understand the roles of aromatic residues and of 4-substituted prolines on proline *cis-trans* isomerism. <sup>19,21,54,55</sup> Dfp was also placed within the Ac-GPPXPPGY-NH<sub>2</sub> host-guest peptide model system context used previously to characterize the PPII propensities of guest residues (including all 20 encoded amino acids) at position **X**. <sup>56-60</sup>



**Figure 2**. Summary of compounds and peptides examined herein. (a) Acyl capping groups were installed on Dfp to generate Ac-Dfp-OMe (6) and Piv-Dfp-OMe (7). The pivaloyl group strongly promotes intercarbonyl  $n \rightarrow \pi^*$  interactions and the *trans*-proline amide rotamer. Synthesis and characterization of 6 and 7 are described in the Supporting Information. (b) Dfp was incorporated into model peptides and into tetrapeptides derived from the *tau* C-terminal domain.

Finally, we examined the use of Dfp to quantify proline *cis-trans* isomerism in the context of a peptide derived from an intrinsically disordered protein. In the microtubule-binding protein *tau*, phosphorylation states (e.g. phosphorylation at Ser404) and mutations (R406W) in the *tau* C-terminal domain are associated with increased neurodegeneration in Alzheimer's disease and other neurodegenerative disorders. In peptides derived from *tau*, we recently demonstrated that both Ser404 phosphorylation and the R406W modification lead to an increased population of *cis*-proline at Pro405. Therefore, we examined Dfp at this position in peptides derived from residues 403-406 of *tau*, in order to understand the roles of both Ser404 phosphorylation and the R406W modification on proline *cis-trans* isomerism.

X-ray crystallography of Fmoc-Dfp-OH. In order to incorporate Dfp into peptides by solid-phase peptide synthesis, commercially available Boc-4,4-difluoroproline **1** was converted to Fmoc-4,4-difluoroproline **3** as previously described (Figure 3a).<sup>22</sup> **3** crystallized from CDCl<sub>3</sub>. The crystal structure of **3** (Figure 3b) exhibited a compact value of  $\phi$  but an extended value of  $\psi$  ( $\phi = -59^{\circ}$ ,  $\psi = +175^{\circ}$ ), a trans amide bond ( $\omega = +171^{\circ}$ ), and a  $C\gamma$ -exo ring pucker ( $\chi_2 = +31^{\circ}$ ). The value of  $\psi$  observed is notably larger than is typical for a proline exo ring pucker.<sup>70,71</sup> In the crystal assembly, a close intermolecular hydrogen bond was observed between the carboxylic acid hydrogen and the carbamate carbonyl oxygen of an adjacent molecule (O•••H distance 1.71 Å) (Figure 3c). Only a weak  $n \rightarrow \pi^*$  interaction, with poor interaction geometry, was observed between consecutive carbonyls (O•••C=O distance 3.02 Å,  $\angle_{OCO} = 77^{\circ}$  (ideal 109°)). Interestingly, the difluoroproline carbonyl exhibited a close C–H/O interaction (2.60 Å) between one ProHβ and the carbamate carbonyl (Figure 3d). These observations are consistent

with our recent report that C–H/O interactions are a general mode for structure stabilization and molecular assembly at proline residues.<sup>77</sup>

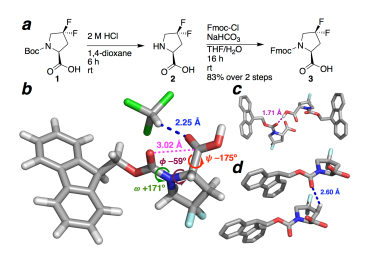
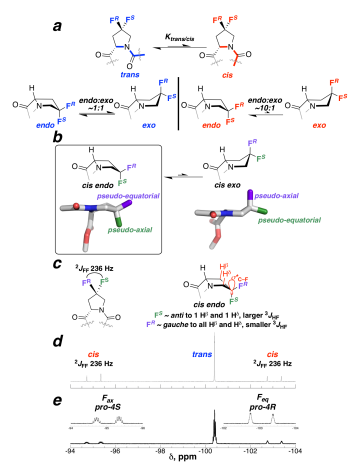


Figure 3. (a) Synthesis of Fmoc-Dfp-OH (3) from commercially available Boc-Dfp-OH (1). (b) Crystal structure (100 K data collection) of Fmoc-Dfp-OH•CHCl<sub>3</sub>. The structure exhibited a close C–H/O interaction (blue) between the carboxylic acid carbonyl and the C–H of chloroform. (c) Crystal assembly of Fmoc-Dfp-OH is mediated in part via hydrogen bonding between the carboxylic acid hydrogen and the carbamate carbonyl of an adjacent molecule. (d) Intermolecular C–H/O interaction between Pro C–Hβ and an adjacent carbonyl. Bond lengths to hydrogen were normalized in Mercury.

<sup>19</sup>F NMR spectroscopy of 4,4-difluoroproline in model compounds and model peptides. The previously reported <sup>1</sup>H-coupled <sup>19</sup>F NMR spectrum of Ac-TYDfpN-NH<sub>2</sub> in water (5 mM phosphate buffer, 25 mM NaCl, pH 4) revealed that, in the *cis* amide conformation, the diastereotopic fluorines exhibited highly divergent <sup>19</sup>F chemical shifts (Figure 4).<sup>21</sup> Each diastereotopic fluorine in the *cis*-proline conformation appears as a doublet due to coupling between the geminal fluorines ( $^2J_{FF} = 236$  Hz) (Figure 4d). In contrast, for *trans*-proline, because

the chemical shift difference (in Hz) between the fluorines is less than their coupling constant  $(\Delta\delta \text{ (Hz)} < {}^2J_{FF})$ , the fluorines were isochronous and appear as a singlet in the  ${}^{19}\text{F}$  NMR spectrum.  ${}^{36}$ 



**Figure 4.** (a) Summary of the conformational equilibria and preferences of the Dfp amide bond and ring pucker. (b) The *pro-4S* fluorine is pseudo-axial in an *endo* ring pucker and pseudo-equatorial in an *exo* ring pucker, while the opposite is observed for the *pro-4R* fluorine. The structures shown are for a model with *cis*-proline, but similar relationships of the diastereotopic fluorines are present for *trans*-proline. Pseudo-axial fluorines are approximately *anti*-periplanar to 1 Hβ and 1 Hγ and thus exhibit larger  $^3J_{\rm HF}$  coupling constants with these hydrogens, while pseudo-equatorial fluorines are *gauche* to both Hβ and to both Hδ hydrogens, and thus exhibit only smaller coupling constants with these hydrogens. (c) The diastereotopic fluorines exhibit strong  $^2J_{\rm FF}$  coupling, which causes each fluorine resonance to appear as a doublet, as is observed here in the *cis* conformation. The larger  $^3J_{\rm HF}$  observed for the pseudo-axial fluorine are due to orbital overlap between  $\sigma^*_{\rm C-F}$  and the  $\sigma_{\rm C-H}$ , which also leads to longer C–F bonds for the pseudo-axial fluorines. (d)  $^1H$ -decoupled and (e)  $^1H$ -coupled  $^{19}F$  NMR spectra of the peptide Ac-TYDfpN-NH<sub>2</sub> in 90% H<sub>2</sub>O/10% D<sub>2</sub>O with 5 mM phosphate buffer pH 4.0 and 25 mM NaCl. The *cis* peaks are shown in the inset, expanded to emphasize the differences in  $^1H$ - $^{19}F$  coupling in the

predominantly pseudo-axial fluorine (downfield) versus the predominantly pseudo-equatorial fluorine (upfield).

In order to understand the basis for the observed <sup>19</sup>F chemical shifts, we examined the <sup>1</sup>H-coupled <sup>19</sup>F NMR spectra (Figure 4e). The <sup>19</sup>F resonances in *cis*-proline exhibited distinct  ${}^3J_{\rm HF}$  coupling patterns of the fluorines to the  $\beta$  and  $\delta$  hydrogens. When a fluorine in Dfp is pseudo-axial (i.e. the *pro-4S*-fluorine in an *endo* ring pucker, or the *pro-4R* fluorine in an *exo* ring pucker), the fluorine is nearly *anti*-periplanar to one H $\beta$  and to one H $\delta$  (Figure 4b), and thus should exhibit larger coupling constants with these two vicinal hydrogens. In contrast, when a fluorine in Dfp is pseudo-equatorial (i.e. the *pro-4R*-fluorine in an *endo* ring pucker, or the *pro-4S* fluorine in an *exo* ring pucker), the fluorine is approximately *gauche* to all vicinal hydrogens, and thus should exhibit only smaller  ${}^3J_{\rm HF}$  coupling constants.<sup>78</sup>

The *endo* and *exo* proline ring puckers are in rapid equilibrium due to a small activation barrier ( $\Delta G^{\ddagger} \sim 2 \text{ kcal mol}^{-1}$ ) for interconversion, with each diastereotopic fluorine being pseudo-axial in one ring pucker and pseudo-equatorial in the other ring pucker. Therefore, in a typical disordered conformation, with similar populations of *endo* and *exo* ring puckers,  $^{70,79-82}$  both diastereotopic fluorines should exhibit similar patterns of  $^3J_{\rm HF}$  coupling, due to the observed  $^3J_{\rm HF}$  representing the population-weighted average of the two ring puckers. Indeed, here for *trans*-proline, the coupling pattern appears as a quintet, indicating similar  $^3J_{\rm HF}$  of each fluorine with each of the 4 vicinal H $\beta$  and H $\delta$ .

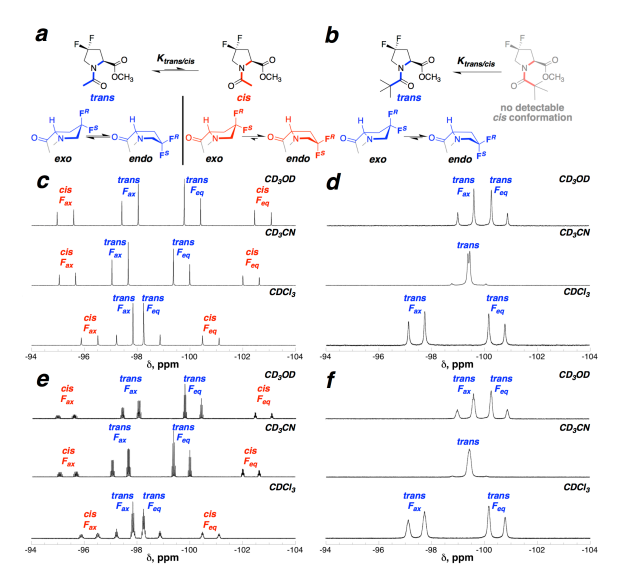
In contrast, if one ring pucker is significantly preferred, then the fluorines should exhibit distinct chemical shifts and patterns of coupling constants, corresponding to those of the major ring pucker present in solution. Proline predominantly adopts an *endo* ring pucker when in the *cis* amide conformation.<sup>70,71,80</sup> Based on the different patterns of the coupling constants in the

fluorines (larger apparent  ${}^3J_{\rm HF}$  coupling constants observed at the downfield fluorine, but all smaller apparent  ${}^3J_{\rm HF}$  values at the upfield fluorine), the downfield fluorine resonance ( $\delta = -95.0$  ppm) in this peptide in water was assigned as the *pro-4S* fluorine. In an *endo* conformation (Figure 4b), the *pro-4S* fluorine is nearly *anti*-periplanar to one H $\beta$  and to one H $\delta$ , and thus should exhibit larger coupling constants ( ${}^3J_{\rm HF}$  over 30 Hz has been observed for strong hyperconjugative interactions (Figure 4c). In contrast, the *pro-4R* fluorine in the *endo* ring pucker would be approximately *gauche* to all H $\beta$  and H $\delta$ , and thus should exhibit smaller overall coupling constants. A similar pattern was previously seen in the  ${}^1H$ -coupled  ${}^1P$ F NMR spectrum of Ac-Dfp-OMe. These results suggested that the downfield  ${}^1P$ F resonance might be associated with a [predominantly] pseudo-axial fluorine, while the upfield  ${}^1P$ F resonance might be associated with a [predominantly] pseudo-equatorial fluorine.

We emphasize that proline cis and trans amide conformations are in slow exchange on the NMR timescale ( $\Delta G^{\ddagger} \sim 20 \text{ kcal mol}^{-1}$ ), and thus the resonances associated with the trans-proline and cis-proline conformations are distinct. In contrast, the interconversion between the endo and exo ring puckers ( $\Delta G^{\ddagger} \sim 2 \text{ kcal mol}^{-1}$ ) is rapid (conformers in fast exchange) on the NMR timescale, and thus the proline ring puckers are not observed as distinct resonances by NMR spectroscopy. Instead, the NMR spectrum represents the population-weighted average of the  $^{19}F$  chemical shifts and coupling constants of a fluorine in both the endo and exo ring puckers.

The model compounds Ac-Dfp-OMe and Piv-Dfp-OMe were similarly examined by <sup>19</sup>F NMR spectroscopy as a function of solvent, in deuterated methanol, acetonitrile, and chloroform (Figure 5). In all solvents, Ac-Dfp-OMe exhibited an equilibrium between the *trans*-proline and *cis*-proline rotamers, with a higher population of *trans*-proline in all solvents (Figure 5c).<sup>23,24</sup> The

trans-proline peaks also exhibited strong second-order effects, in which the doublet of each fluorine is not symmetric, with the magnitude of the second-order effects dependent on the chemical shift difference between the fluorines (smaller  $\Delta\delta$  = more pronounced second-order effects).<sup>36</sup>



**Figure 5**. Solvent-dependent <sup>19</sup>F NMR spectra of (left) Ac-Dfp-OMe and (right) Piv-Dfp-OMe. (a) Ac-Dfp-OMe conformational equilibria: *cis-trans* amide isomerization and ring pucker interconversion. (b) Piv-Dfp-OMe conformational equilibria: only the *trans* amide conformation was observed. (c,d) Solvent dependent <sup>1</sup>H-decoupled <sup>19</sup>F NMR spectra of (c) Ac-Dfp-OMe and (d) Piv-Dfp-OMe, (e,f) Solvent-dependent <sup>1</sup>H-coupled <sup>19</sup>F NMR spectra of (e) Ac-Dfp-OMe and (f) Piv-Dfp-OMe. Data were collected at 298 K. For spectra with Piv-Dfp-OMe, broadening is observed due to proline *cis-trans* isomerization that is in intermediate exchange on the NMR

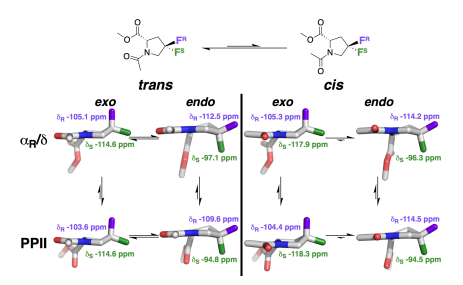
timescale at this temperature. The assignment of the *endo* ring pucker as the main ring pucker for Ac-cis-Dfp-OMe is further supported by the H $\alpha$  resonances of the <sup>1</sup>H NMR spectra, which exhibit simplified coupling patters compared to the H $\alpha$  resonances for *trans*-Dfp, with the cis-Dfp coupling pattern consistent with that expected for an *endo* ring pucker as the dominant species (Figures S1, S4, S7, S10).  $F_{ax}$  indicates the diastereotopic fluorine which is pseudo-axial in the major proline ring pucker, while  $F_{eq}$  indicates the diastereotopic fluorine which is pseudo-equatorial in the major proline ring pucker in the indicated proline amide conformation under the described conditions.

In all cases, by <sup>19</sup>F NMR, the *cis*-proline conformation exhibited a larger chemical shift difference between the diastereotopic fluorines than was observed for *trans*-proline. For Piv-Dfp-OMe, only the *trans*-proline rotamer was observed (Figure 5d), as expected based on prior results. <sup>52,53</sup> The patterns of chemical shift differences between the diastereotopic fluorines in Piv-Dfp-OMe ( $\Delta\delta_{FF} = 0$ –3.1 ppm) were similar to those observed for *trans*-proline in Ac-Dfp-OMe ( $\Delta\delta_{FF} = 1.1$ –2.4 ppm), and significantly smaller than those of Ac-Dfp-OMe with the *cis*-proline conformation ( $\Delta\delta_{FF} = 4.6$ –7.5 ppm).

The  $^{1}$ H-coupled  $^{19}$ F NMR spectra of Ac-Dfp-OMe (Figure 5e) exhibited similar patterns of  $^{3}J_{HF}$  coupling constants as were observed for Ac-TYDfpN-NH<sub>2</sub>: the downfield fluorine exhibited overall larger apparent  $^{3}J_{HF}$ , while the upfield fluorine exhibited smaller apparent coupling constants. These results suggested that the fluorine with a downfield  $^{19}$ F resonance was predominantly pseudo-axial, while the upfield  $^{19}$ F resonance was predominantly pseudo-equatorial, with a large chemical shift difference between the fluorines indicating a strong bias toward one ring pucker (as is present for *cis*-proline and the *endo* ring pucker), while a small chemical shift difference indicates a weak preference for one ring pucker (as is observed for *trans*-proline in water).  $^{70,80,81}$ 

Calculations on 4,4-difluoroproline in model compounds. In order to further understand the relationships between proline ring pucker, proline main-chain  $(\phi,\psi)$  conformation, proline

amide ( $\omega$ ) rotamer, and <sup>19</sup>F chemical shift, we conducted DFT calculations on Ac-Dfp-OMe and Ac-Piv-OMe (Figure 6).<sup>83,84</sup> All conformers were subjected to full geometry optimization, followed by analysis of the calculated chemical shifts as a function of conformation. These results were unequivocal: in all conformations in both molecules, the pseudo-axial fluorine (*pro-4R* in an *exo* ring pucker, *pro-4S* in an *exo* ring pucker) was downfield, and the pseudo-equatorial fluorine (*pro-4S* in an *exo* ring pucker, *pro-4R* in an *endo* ring pucker) was upfield. The chemical shift differences between the diastereotopic fluorines in any individual conformation of Ac-Dfp-OMe were 9.5–20 ppm, depending on the main-chain conformation. Similar results were observed in implicit water and in implicit chloroform, indicating that the effect of conformation on <sup>19</sup>F chemical shift has minimal solvent dependence, and thus should be general in different contexts (e.g. a solvent-exposed proline versus a proline buried in a protein hydrophobic core).



**Figure 6**. Geometry-optimized structures of Ac-Dfp-OMe in implicit water in the *trans* and *cis* amide conformations, each with the *exo* and *endo* ring puckers in the indicated regions of Ramachandran space. The calculated <sup>19</sup>F chemical shifts for the *pro-4R* (purple) and *pro-4S* (green) fluorines are indicated in each conformation. Equivalent <sup>19</sup>F NMR analysis for Piv-Dfp-OMe, Ac-Dfp-NMe<sub>2</sub>, and Piv-Dfp-NMe<sub>2</sub> as a function of conformation are in the Supporting Information (Tables S24-S25, S28-S29). <sup>19</sup>F NMR chemical shifts were calculated using the GIAO method with the M06-2X functional and the 6-311++G(3d,3p) basis set in implicit water. Additional details on the conformations, including their relative energies (Tables S22-S23, S26-S27), as well as coordinates for all structures in implicit H<sub>2</sub>O and implicit CHCl<sub>3</sub>, are in the Supporting Information.

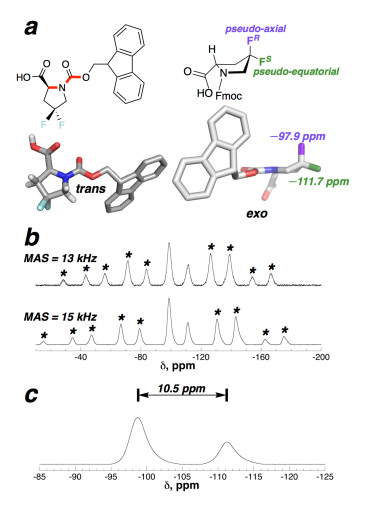
The same general results were also observed for Piv-Dfp-OMe. In addition, the *endo* ring pucker was also observed with relatively more downfield chemical shifts for the pseudo-axial fluorine, although these effects were smaller than the effect of being pseudo-axial versus pseudo-equatorial. Similar results were also observed for the model compounds with C-terminal amides Ac-Dfp-NMe<sub>2</sub> and Piv-Dfp-NMe<sub>2</sub>; see the Supporting Information for details (Tables S24-S25, S28-S29). The experimentally observed chemical shifts inherently represent a population-weighted average of the chemical shifts across the full conformational ensemble present in solution. In addition, these calculations only considered the conformations that represent local energy minima, while in solution molecules are dynamic and significantly populate conformations that are near but not identical to the energy minima (e.g., as observed in a

Ramachandran plot). Overall, in the context of prior work on <sup>19</sup>F chemical shift calculations <sup>83,84</sup> and the required referencing of all chemical shifts to the calculated TFA <sup>19</sup>F chemical shift, the correlation between calculated and observed <sup>19</sup>F chemical shifts was reasonable when one major conformation is expected to predominate, e.g. for *cis*-proline, where the *endo* ring pucker is most populated; see also temperature-dependent NMR data below.

The calculations suggested that the basis for the chemical shift difference of the fluorines is likely directly related to the inherent stereoelectronic effects that stabilize fluorines when pseudo-axial. These effects are analogous to the stereoelectronic effects that stabilize alcohols in the axial position ( $\alpha$  anomer) at the anomeric carbon in carbohydrates. When a fluorine is pseudo-axial, it is stabilized by hyperconjugative interactions between the electron-accepting C-F antibonding orbital ( $\sigma^*_{C-F}$ ) and an electron-donating C-H bonding orbital ( $\sigma^*_{C-H}$ ) and to which it is approximately *anti*-periplanar. Stabilization occurs via hyperconjugation of  $\sigma^*_{C-F}$  with both C-H $\delta$  and C-H $\delta$  (which results in a *syn*-periplanar [overlapping orbitals] relationship of  $\sigma^*_{C-F}$  with each  $\sigma^*_{C-H}$ ) (Figure 4c). Electron delocalization between these C-H  $\sigma$  bonding orbitals and the C-F  $\sigma^*$  antibonding orbital (overlap of the  $\sigma^*_{C-H}$  with the  $\sigma^*_{C-F}$ , a  $\sigma^*$  interaction) is stabilizing. This electron delocalization also results in a longer C-F bond for the pseudo-axial C-F than the pseudo-equatorial C-F, as was seen computationally and is observed more generally for hyperconjugation.

Solid-state NMR spectroscopy of crystalline Fmoc-Dfp-OH. These computational results suggested that, in any molecule with a strong bias for one proline ring pucker, Dfp substitution for Pro should result in a large chemical shift difference between the diastereotopic fluorines. In the X-ray crystal structure of Fmoc-Dfp-OH (Figure 3), Dfp was observed to adopt an *exo* ring pucker, and therefore should exhibit a large difference between the chemical shifts of its

fluorines. Therefore, we examined the crystals of Fmoc-Dfp-OH by solid-state <sup>19</sup>F NMR spectroscopy, as a function of spinning rate by magic angle spinning (MAS).<sup>89-91</sup> The solid-state <sup>19</sup>F NMR spectrum (Figure 7) indicated two central-band resonances, associated with the two diastereotopic fluorines. The observed 12.5 ppm chemical shift difference between the fluorines was very similar to the calculated 11.0 ppm chemical shift difference between the diastereotopic fluorines for Ac-Dfp-OMe in the crystallographically observed conformation (exo ring pucker and PPII conformation). In addition, calculations on Fmoc-Dfp-OH•CHCl<sub>3</sub> and the computationally simpler molecule EtOC(O)-Dfp-OH•CHCl<sub>3</sub>, with the geometry of each restrained to that observed crystallographically, were consistent with the solid-state NMR data, with the pseudo-axial pro-4R fluorine downfield and the pseudo-equatorial pro-4S fluorine upfield, and a 12-14 ppm chemical shift difference between those fluorines, similar to that observed by solid-state MAS NMR; see the Supporting Information for details (Tables S30-S31). In this molecule with only a single conformation, the calculated <sup>19</sup>F chemical shifts were within 1 ppm of those observed experimentally, consistent with typical errors in <sup>19</sup>F chemical shift calculations.83,84



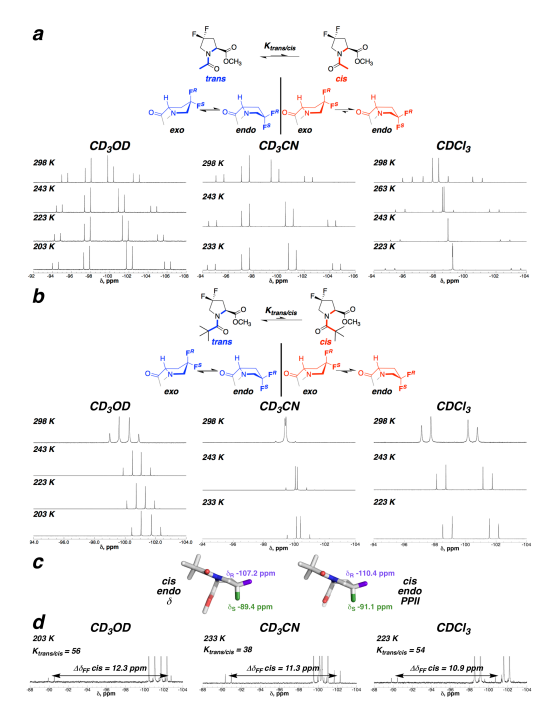
**Figure 7**. Solid-state <sup>19</sup>F NMR spectroscopy of Fmoc-Dfp-OH in the crystalline form. (a) Crystal structure of Fmoc-Dfp-OH in the *trans*-proline conformation, which exhibits an *exo* ring pucker, with the *pro-4R* (purple) and *pro-4S* (green) fluorines indicated. The electron density map of Fmoc-Dfp-OH shows no evidence of an alternative conformation present in the unit cell. (b) Solid-state <sup>19</sup>F NMR spectra of crystalline Fmoc-Dfp-OH. The spectra were obtained using magic-angle spinning (MAS) with spinning frequencies of 13 kHz and 15 kHz. The central bands of the spectra, with chemical shifts of –98.7 ppm and –111.2 ppm, correspond to the *pro-4R* and *pro-4S* fluorines, respectively. All other signals present in the spectra are spinning side bands (indicated with asterisks). The spectra were referenced with external polycrystalline 5F-Trp at 87.0 ppm, as a secondary reference to CCl<sub>3</sub>F at 0 ppm. (c) Expanded view of the central bands in the MAS <sup>19</sup>F NMR spectra of crystalline Fmoc-Dfp-OH, revealing a chemical shift difference (Δδ) of 10.5 ppm between the fluorines assigned as the *pro-4R* (downfield) and *pro-4S* (upfield) fluorines. Additional details are in the Supporting Information (Figures S25-S26).

Temperature-dependent <sup>19</sup>F NMR spectra of model compounds. Overall, the calculations indicated that observed chemical shift differences between the diastereotopic fluorines of Dfp reflect the inherent preferences at a given residue for one proline ring pucker over the other. As

the energy differences between the proline ring puckers are small in most contexts, the effects of these energy differences on the populations of the *exo* versus *endo* ring puckers should be manifested as a function of temperature. At lower temperatures, a larger population of the lower-energy ring pucker versus the higher-energy ring pucker should be observed (as  $\Delta G = -RT$  ln  $K_{\text{exo/endo}}$ ), which based on the calculations would be manifested as a larger chemical shift difference between the fluorines at lower temperature. This analysis assumes only small enthalpic effects on the stability of one ring pucker versus the other. Therefore, we examined the <sup>19</sup>F NMR spectra of Ac-Dfp-OMe and Piv-Dfp-OMe as a function of temperature (Figure 8, Table 1).

**Table 1**. Summary of temperature-dependent <sup>19</sup>F NMR data for Ac-Dfp-OMe and Piv-Dfp-OMe as a function of solvent.

				cis		cis	cis trans		
				downfield	upfield		downfield	upfield	
	Solvent	Temperature, K	K <sub>trans/cis</sub>	δ, ppm	δ, ppm	Δδ <sub>FF</sub> , ppm	δ, ppm	δ, ppm	Δδ <sub>FF</sub> , ppm
Ac- Dfp- OMe	CD₃OD	298	2.4	-95.3	-102.8	7.5	-97.7	-100.1	2.4
		243	2.6	-94.7	-104.6	9.9	-97.7	-101.2	3.5
		223	2.6	-94.5	-105.3	10.8	-97.6	-101.6	4.0
		203	2.5	-94.3	-106.1	11.8	<b>-</b> 97.6	-102.1	4.5
	CD₃CN	298	2.3	-95.4	-102.3	6.9	-97.4	-99.7	2.3
		243	2.7	-94.8	-104.1	9.3	-97.4	-100.8	3.4
		233	2.8	-94.7	-104.5	9.8	-97.4	-101.0	3.6
	CDCl₃	298	3.1	-96.2	-100.8	4.6	<b>–</b> 97.5	-98.6	1.1
		263	3.7	-95.7	-101.9	6.2	-98.2	-98.9	0.7
		243	4.0	-95.4	-102.6	7.2	-98.5	-99.2	0.7
		223	4.1	-95.1	-103.3	8.2	-98.8	-99.5	0.7
Piv- Dfp- OMe	CD₃OD	298	n.d.	_	_	_	-99.3	-100.6	1.3
		243	30	-91.0	-101.6	10.6	-100.2	-101.2	1.0
		223	37	-90.6	-102.0	11.4	-100.5	-101.8	1.3
		203	56	-90.2	-102.5	12.3	-100.8	-102.0	1.2
	CD₃CN	298	n.d.	_	_	_	-99.1	-99.8	0.7
		243	32	-90.9	-101.7	10.8	-99.8	-100.5	0.7
		233	38	-90.7	-102.0	11.3	-99.8	-100.7	0.9
	CDCl₃	298	n.d.	_	_	_	-97.4	-100.5	3.1
		243	37	-90.6	-100.5	9.9	-98.4	-101.5	3.1
		223	54	-90.1	-101.0	10.9	-98.8	-101.9	3.1



**Figure 8**. Temperature-dependent  $^{1}$ H-decoupled  $^{19}$ F NMR spectra of (a) Ac-Dfp-OMe and (b) Piv-Dfp-OMe in CD<sub>3</sub>OD, CD<sub>3</sub>CN, and CDCl<sub>3</sub>. (c) Calculated optimized structures and  $^{19}$ F chemical shifts of Piv-Dfp-OMe in CHCl<sub>3</sub> with *cis*-proline and an *endo* ring pucker. Calculations indicated a  $\geq 2$  kcal mol<sup>-1</sup> preference for the *endo* versus the *exo* ring pucker in Piv-*cis*-Dfp-OMe. Structures were subjected to NMR calculations using the GIAO method with the M06-2X functional and the 6-311++G(3d,3p) basis set. (d) Low-temperature  $^{19}$ F NMR spectra of Piv-Dfp-OMe, with expansion of the chemical shift range and truncation of the *trans*-proline peaks in order to emphasize the minor *cis*-proline peaks, in CD<sub>3</sub>OD, CD<sub>3</sub>CN, and CDCl<sub>3</sub>. The peak broadening observed for Piv-Dfp-OMe at 298 K is likely due to its lower barrier for the *cis*-to-*trans* isomerization compared to Ac-Dfp-OMe, which causes its spectrum to represent intermediate exchange (broadening) rather than slow exchange on the NMR timescale.

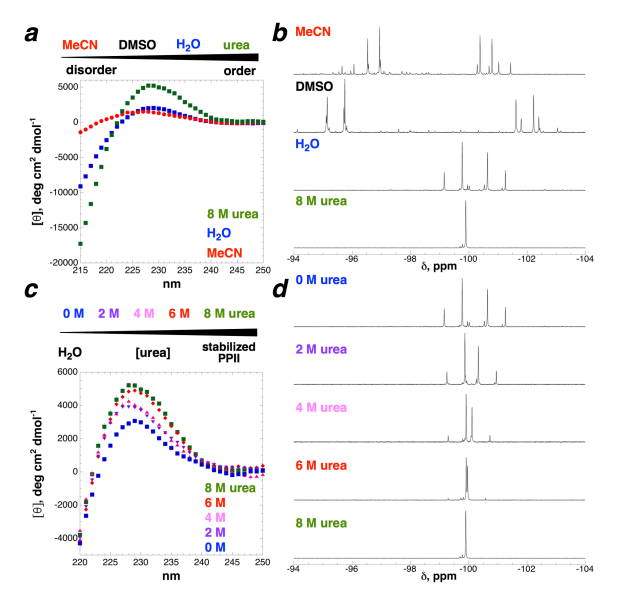
For Ac-Dfp-OMe, in methanol and in acetonitrile (Figure 8a, left and middle panels; Table 1), larger chemical shift differences between the diastereotopic fluorines were observed at lower temperatures. This increase in chemical shift dispersion between the fluorines in a given amide conformation was observed in both the *cis*-proline and the *trans*-proline conformations. These results are consistent with higher populations of the lowest-energy ring pucker being present at lower temperatures, as is expected by  $\Delta G = -RT \ln K_{\rm exo/endo}$ . In chloroform, this increased dispersion of the diastereotopic fluorines at lower temperatures was also observed for *cis*-proline (Figure 8a, right; Table 1). In contrast, for *trans*-proline, the fluorine chemical shifts became more similar at lower temperature, including the observation of only a single peak for *trans*-Dfp in CDCl<sub>3</sub> at  $\leq 243$  K. These data indicated that the *endo* and *exo* ring puckers became more similar in energy at lower temperatures in these conditions, suggesting an additional enthalpic component impacting ring pucker populations for Ac-*trans*-Dfp-OMe in CDCl<sub>3</sub>.

For Piv-Dfp-OMe (Figure 8b), the temperature-dependent <sup>19</sup>F NMR data indicated the possibility of a change in the main-chain conformation, as suggested by the more upfield <sup>19</sup>F chemical shifts at lower temperature. Notably, at lower temperatures, the *cis*-proline conformation of Piv-Dfp-OMe was also observed as a very minor species ( $\sim 2-3\%$  of the total population) by <sup>19</sup>F NMR (Figure 8cd). The *pro-4S* fluorine  $\delta$  was particularly downfield, with a large  $\Delta\delta$  observed for *cis*-proline in all solvents. Calculations on Piv-Dfp-OMe indicated that the *exo* ring pucker is particularly disfavored for *cis*-Dfp, with the pivaloyl acyl group  $\geq$  2 kcal mol<sup>-1</sup> higher energy for either *cis–exo* conformer than either *cis–endo* conformer. In addition, the calculated <sup>19</sup>F chemical shifts for Piv-*cis*-Dfp-OMe with an *endo* ring pucker were the most downfield chemical shifts identified computationally, as was also observed experimentally. Finally, the <sup>19</sup>F NMR spectra permitted the identification and quantification of this very low

population of *cis*-proline, which could not be definitively identified in the equivalent <sup>1</sup>H NMR spectra. These data emphasize the advantages of the simplified <sup>19</sup>F spectra in the identification of minor *cis*-proline species in solution.

Understanding the effects of solvation and urea on PPII. The polyproline II helix (PPII) conformation has been proposed to be stabilized by solvation with water, although other bases for PPII stability have alternatively been proposed. PPII stability have alternatively been proposed. In order to better understand the role of solvation in stabilization of PPII, the peptide Ac-GPPDfpPPGY-NH<sub>2</sub> was examined by circular dichroism in water and in acetonitrile (Figure 9). In water, this peptide exhibited a somewhat reduced magnitude of mean residue ellipticity at 228 nm compared to the peptide with proline at the central (guest) residue. These data indicate that Dfp strongly supports PPII, although it modestly destabilizes PPII relative to proline. These results are consistent with the somewhat greater likelihood of Dfp to induce cis amide bonds compared to proline in the model peptides Ac-Dfp-OMe or Ac-TADfpN-NH<sub>2</sub>, as well as with a somewhat weaker  $n \rightarrow \pi^*$  interaction for Dfp compared to proline. PPII CD signature.

Urea strongly promotes PPII, which has led to proposals that PPII is a major conformation in the denatured states of proteins. <sup>92,96,98-100</sup> The CD spectrum of this peptide in 8 M urea exhibited a much stronger CD signature of PPII, consistent with prior data that urea promotes PPII, including in the Ac-GPPXPPGY-NH<sub>2</sub> peptide host-guest context (Figure S8 of reference <sup>101</sup>).



**Figure 9**. <sup>19</sup>F NMR and circular dichroism (CD) spectroscopy of the model peptide Ac-GPPDfpPPGY-NH<sub>2</sub> as a function of solvation. (a) CD spectra in buffered H<sub>2</sub>O (with 5 mM phosphate pH 4 and 25 mM KF), acetonitrile, and 8 M urea. Experiments were conducted at 25 °C. Data represent the average of at least three independent trials. (b) <sup>1</sup>H-decoupled <sup>19</sup>F NMR spectra as a function of solvent. (c) CD spectra as a function of urea concentration. (d) <sup>1</sup>H-decoupled <sup>19</sup>F NMR spectra as a function of urea concentration. Experiments in organic solvents and experiments in (c) and (d) were conducted without added buffer or salt. <sup>1</sup>H NMR spectra of the peptide in buffered H<sub>2</sub>O, in DMSO, and in acetonitrile, as well as temperature-dependent <sup>1</sup>H and <sup>19</sup>F NMR spectra in buffered H<sub>2</sub>O, are in Supporting Information (Figures S29-S31, S35-S36).

The peptide Ac-GPPDfpPPGY-NH<sub>2</sub> has the possibility of 32 (=2<sup>5</sup>) species in slow exchange, due to *cis-trans* isomerism at 5 X–Pro amide bonds. However, this peptide exhibited only one major species by <sup>19</sup>F NMR in water, with only very small amounts of species with *cis*-Pro observed, consistent with CD data that PPII was adopted in the model peptide (Figure 9).<sup>102</sup> In contrast, much higher populations of species with *cis*-proline amide bonds were observed both in acetonitrile and in DMSO, which is observable in the large number of minor peaks present in the <sup>19</sup>F NMR spectra in these solvents.<sup>103</sup> Interestingly, the major species in both organic solvents also had a larger difference in <sup>19</sup>F chemical shifts of the fluorines, suggesting a stronger preference for one ring pucker in these solvents compared to in water. In contrast, the <sup>19</sup>F NMR spectrum in 8 M urea exhibited one major species, with just a single major peak in the <sup>19</sup>F NMR spectrum, that is, that the chemical shifts of the diastereotopic fluorines were almost identical. These <sup>19</sup>F NMR data, in combination with the data and calculations above, demonstrate that similar populations of the *endo* and *exo* ring puckers were present in this peptide in 8 M urea.

These results were confirmed by CD and NMR data on Ac-GPPDfpPPGY-NH<sub>2</sub> as a function of concentration of urea (Figures 9cd). As expected, higher concentrations of urea resulted in higher populations of PPII by CD. However, increased PPII by CD also correlated with smaller chemical shift differences between the diastereotopic fluorines. These results indicate that PPII is most favorable when the proline ring adopts *both* ring puckers in similar populations. This result stands in stark contrast to the implications of extensive data with 4-substituted prolines, which has suggested that the *exo* ring pucker (favored by 4*R*-substituted hydroxyprolines and 4*R*-fluoroproline) promotes PPII, while the *endo* ring pucker (favored by 4*S*-substituted hydroxyprolines and 4*S*-fluoroproline) relatively disfavors PPII, compared to proline. <sup>59,104-106</sup> Herein, using Dfp as a probe of proline ring pucker, we identify that the highest

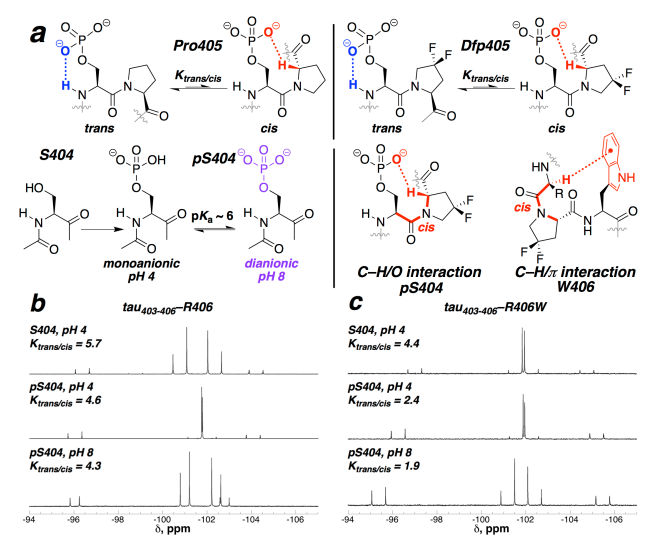
population of PPII is present when *both* proline ring puckers are present in similar populations, based on the observed similar <sup>19</sup>F chemical shifts of the diastereotopic fluorines. This result is logical from both entropic and enthalpic perspectives: selection of one ring pucker over the other would be inherently energetically unfavorable. The results herein suggest that urea promotes PPII in part by making the *endo* and *exo* proline ring puckers equienergetic, and thus equally populated, resulting in neither an entropic cost nor an enthalpic cost for adopting PPII. These data are also consistent with urea strongly promoting PPII via solvation of the acceptor carbonyl for  $n\rightarrow\pi^*$  interactions, as we recently proposed.<sup>96</sup>

Quantifying proline cis-trans isomerization and identifying induced order in peptides from an intrinsically disordered protein. Finally, we examined the use of Dfp to quantify proline cis-trans isomerism in model peptides consisting of residues 403-406 from the C-terminal domain of the tau protein, whose misfolding and aggregation are associated with neurodegeneration in Alzheimer's disease. 63,67,107-109 We previously observed that both Ser404 phosphorylation and the R406W modification lead to increased population of cis-Pro405, both in the shorter  $tau_{403-406}$  peptides Ac-TSPR-NH<sub>2</sub> and Ac-TSPW-NH<sub>2</sub> and in larger  $tau_{395-411}$  peptides.<sup>69</sup> The data in shorter peptides recapitulated the data in larger tau peptides, whose analysis was complicated by the intrinsically disordered nature of these sequences, which results in poor <sup>1</sup>H chemical shift dispersion. Using Dfp, we observed similar effects of phosphorylation and the R406W modification on increasing the population of *cis*-proline as was observed by <sup>1</sup>H NMR on peptides with proline, but here with much easier data interpretation due to the simplified nature of <sup>19</sup>F NMR spectra (Figure 10, Table 2). In addition, in our prior work we proposed that the combination of both Ser-404 phosphorylation and the R406W modification (Figure 10c, bottom) would lead to increased order, in both trans-proline and cis-proline, due to the promotion of turn

conformations. The <sup>19</sup>F NMR data are consistent with increased ordering in *both* proline amide isomers, based on increased dispersion of the <sup>19</sup>F chemical shifts, which indicates a greater ordering of the proline ring pucker. These results indicate that Dfp can be employed both to quantify proline *cis-trans* isomerization in complex peptides and to identify increased order in peptides, independent of proline amide isomerization state. These results should be broadly applicable to probe protein folding, protein misfolding, and changes in protein structure due to post-translational modifications, protein mutations, and protein-protein interactions.

**Table 2**. Summary of  $^{19}$ F NMR data of  $tau_{403-406}$  peptides as a function of phosphorylation, residue 406 identity, and pH.

			cis		cis	trans		trans
peptide	рН	K <sub>trans/cis</sub>	δ <sub>downfield</sub> , ppm	δ <sub>upfield</sub> , ppm	Δδ <sub>FF</sub> , ppm	δ <sub>downfield</sub> , ppm	δ <sub>upfield</sub> , ppm	Δδ <sub>FF</sub> , ppm
TSDfpR	4.0	5.7	-96.3	-104.2	7.9	-100.8	-102.3	1.5
	4.0	4.6	-96.1	-104.0	7.9	-101.4	-102.1	0.7
TnCDfnD	6.5	3.9	-96.1	-102.9	6.8	-101.1	-102.3	1.2
TpSDfpR	7.2	4.1	-96.0	-102.8	6.8	-101.0	-102.4	1.4
	8.0	4.3	-96.0	-102.8	6.8	-101.0	-102.4	1.4
TSDfpW	4.0	4.4	-97.0	-104.8	7.8	-101.5	-102.3	0.8
	4.0	2.4	-96.3	-105.2	8.9	-101.6	-102.3	0.7
TnCDfnW	6.5	1.9	-95.6	-105.4	9.8	-101.3	-102.4	1.1
TpSDfpW	7.2	2.0	-95.4	-105.5	10.1	-101.2	-102.4	1.2
	8.0	1.9	-95.4	-105.5	10.1	-101.2	-102.4	1.2



**Figure 10**. (a) Phosphorylation of serine stabilizes the *trans*-proline and *cis*-proline conformations via an intraresidue phosphate-amide hydrogen bond and a phosphate-ProHα C-H/O interaction, respectively. This effect is dependent on the ionization state of the phosphate, with the strongest effects observed in the dianionic form of phosphoserine. The R406W modification induces C-H/π interactions which further stabilize the *cis*-proline conformation, but also promote turn conformations in *trans*-proline. (b) <sup>1</sup>H-decoupled <sup>19</sup>F NMR spectra of *tau*<sub>403-406</sub> R406 peptides as a function of phosphorylation and ionization state of the phosphate. The phosphorylated peptide has an increased *cis* population relative to the non-phosphorylated peptide at pH 4.0, and this population increases when the phosphate is dianionic. (c) <sup>1</sup>H-decoupled <sup>19</sup>F NMR spectra of *tau*<sub>403-406</sub> R406W peptides as a function of phosphorylation and ionization state of the phosphate. The non-phosphorylated R406W peptide has an increased *cis*-proline population in comparison to the native R406 peptide, and this effect further increases upon phosphorylation. <sup>1</sup>H-coupled <sup>19</sup>F NMR spectra, <sup>1</sup>H NMR spectra, TOCSY spectra, and summaries of the NMR data for each peptide are in the Supporting Information (Figures S39-S90, Tables S9-S21).

## **Discussion**

Proline *cis–trans* isomerism is a relatively slow conformational equilibrium (typical half-lives of seconds to minutes) due to the partial double-bond character of the amide bond restricting bond rotation.<sup>1-3,18,70,110</sup> In addition, proline *cis* and *trans* amide bonds are structurally distinct in their relative positioning of the N-terminal and C-terminal protein chains.<sup>2,111,112</sup> Based on these kinetic and structural effects, proline *cis–trans* isomerism can be a rate-determining step in protein folding, and can result in structural and functional switches within proteins.<sup>1,111-113</sup> Despite this fundamental importance of proline conformation on protein structure and dynamics, there is an absence of convenient, broadly accessible tools to rapidly interrogate proline *cis–trans* isomerism.

In 2001, Moroder and coworkers demonstrated that 4,4-difluoroproline (Dfp) has potentially unique properties.<sup>23</sup> Dfp is sterically similar to proline, allowing its substitution within proteins and at protein-protein interfaces.<sup>25</sup> Indeed, proline can be replaced by Dfp in expressed proteins using proline-auxotrophic bacteria and incubation with Dfp, as demonstrated both in globular proteins and in elastin.<sup>23,31</sup> Moreover, in contrast to the monosubstituted 4R- and 4S-fluoroprolines, Dfp has a minimal effect on proline conformational preferences.<sup>19,21,23,25,27,29,30</sup> Thus, as either simple acetylated amino acids or within peptides, Dfp and Pro exhibit similar relative populations of *cis* and *trans* amide bonds, as well as similar overall conformational preferences. Consistent with the NMR data on model compounds, both the globular protein barstar and collagen-mimetic peptides have similar  $T_{\rm m}$  values in their native forms with proline or with Pro substituted by Dfp.<sup>23,24,34</sup>

One notable distinction between proline and Dfp is dynamic: Dfp exhibits a modestly (~1 kcal/mol) reduced activation barrier for *cis-trans* isomerism compared to proline.<sup>23</sup> Thus, Dfp

could serve to accelerate protein folding or misfolding when proline cis–trans isomerism is rate limiting. In a dramatic example, Hilvert and coworkers demonstrated that  $\beta$ 2-microglobulin exhibits dynamic cis–trans isomerization at Pro32, with the cis conformation favoring the native state, but the trans conformation favoring protein misfolding and aggregation.<sup>35</sup> This misfolding is enhanced by low pH, at which histidine is cationic and electronically disfavors an aromatic-proline C–H/ $\pi$  interaction that stabilizes the cis conformation.<sup>57,114</sup> Hilvert further demonstrated that replacement of Pro32 with Dfp substantially enhanced  $\beta$ 2-microglobulin aggregation and amyloid formation, via the reduced cis-trans isomerization activation barrier of Dfp, which allowed the protein to more rapidly sample the aggregation-prone trans conformation.

Curiously, neither Moroder nor Conticello nor Rubini reported the  $^{19}F$  NMR spectra of proteins containing Dfp. $^{23,31,34}$  In our work on Dfp within model peptides, $^{21}$  we found the  $^{19}F$  NMR spectrum of the peptide Ac-TYDfpN-NH<sub>2</sub> to be similarly curious, with only a singlet observed for the *trans* conformation (Figure 4a), despite two diastereotopic fluorines that should furthermore exhibit strong  $^2J_{FF}$  coupling. In the *trans* conformation, the chemical shifts of the diastereotopic fluorines were similar (rendering them isochronous), whereas these chemical shifts were highly dispersed in the *cis* amide conformation. Hilvert subsequently made similar observations within  $\beta$ 2-microglobulin. $^{35}$  These data suggested the possibility that Dfp might be used as a sensitive  $^{19}F$  NMR-based probe of proline *cis-trans* isomerism. In contrast to methods based on differences in  $^{13}C$  chemical shift of proline  $C\beta$  and  $C\gamma$ , $^{7-10}$  which require either complete NMR resonance assignment, high peptide or protein concentrations, and/or the use of expensive  $^{13}C$ -labeled proline or complete  $^{13}C'$ 15N labeling of the protein and 3-D NMR spectroscopy, $^{12,13,15,16}$  Dfp provides a highly practical approach to interrogate proline conformational state. The rapidity of this approach, due to the sensitivity of  $^{19}F$  NMR

spectroscopy and the absence of additional resonances in the spectrum, provides an additional advantage in its greater temporal resolution (faster spectral acquisition allowing more frequent analysis of kinetics) compared to <sup>13</sup>C NMR-based methods. Due to its commercial availability (both as Boc-Dfp-OH and Fmoc-Dfp-OH) and its ability to be incorporated in proteins genetically via proline-auxotrophic bacteria, <sup>23,31</sup> Dfp might represent a unique tool for the rapid determination of the *cis-trans* isomerization state of prolines, within proteins and for the investigation of mechanisms of protein folding and assembly.

Dfp can be site-specifically incorporated at the site of a single proline, via standard solidphase peptide synthesis in peptides, or via the combination of synthetic peptides and/or expressed proteins using native chemical ligation and/or expressed proteins using expressed protein ligation/protein semisynthesis. 115,116 Using these technologies, Dfp can be employed to rapidly and exquisitely probe protein conformation at proline, via simple 1-D 19F NMR spectroscopy, with the absence of complications from other residues, from H<sub>2</sub>O, or from urea or other small molecules in solution. Indeed, <sup>19</sup>F NMR is extensively used to understand protein folding and other biological processes in complex media, including whole cells.<sup>38-40,117-119</sup> Herein, Dfp was applied both to quantify proline cis-trans isomerism, and to identify order versus disorder, via the difference in the chemical shifts of the diastereotopic fluorines. The observation of similar <sup>19</sup>F chemical shifts of the fluorines indicates inherent disorder, that is, structures in which both the *endo* and *exo* proline ring puckers are present in similar populations. In contrast, the observation of distinct chemical shifts of the diastereotopic fluorines is associated with local or global order, as it indicates a significant preference for one proline ring pucker over the other. Intrinsically disordered proteins (IDPs) are critical in cellular function, and changes in protein structure, including proline cis-trans isomerism, in IDPs have been implicated in disease. 120-122

More broadly, proline is specially situated as a locus for order and for intermolecular interactions, both within IDPs and within globular proteins. However, identification of these structural changes, especially at proline, poses significant challenges. The results herein suggest that Dfp might be uniquely useful for the practical identification of order, disorder, and proline *cis-trans* isomerism in proteins.

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## **Supporting Information Available**

<sup>1</sup>H, <sup>13</sup>C, and <sup>19</sup>F NMR spectra for synthesized small molecules, CD spectra, and full synthesis and characterization data of all peptides, as well as additional <sup>19</sup>F NMR data. This material is available free of charge via the Internet at the journal web site.

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