cis–trans-Amide isomerism of the 3,4-dehydroproline residue, the ‘unpuckered’ proline

Vladimir Kubyshkin* and Nediljko Budisa*

Full Research Paper

Address:
Institute of Chemistry, Technical University of Berlin,
Müller-Breslau-Str., 10, 10623, Berlin, Germany

Email:
Vladimir Kubyshkin* - kubyshkin@win.tu-berlin.de; Nediljko Budisa* - nediljko.budisa@tu-berlin.de

* Corresponding author

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Abstract

Proline (Pro) is an outstanding amino acid in various biochemical and physicochemical perspectives, especially when considering the cis–trans isomerism of the peptidyl-Pro amide bond. Elucidation of the roles of Pro in chemical or biological systems and engineering of its features can be addressed with various Pro analogues. Here we report an experimental work investigating the basic physicochemical properties of two Pro analogues which possess a 3,4-double bond: 3,4-dehydroproline and 4-trifluoromethyl-3,4-dehydroproline. Both indicate a flat pyrroline ring in their crystal structures, in agreement with previous theoretical calculations. In solution, the peptide mimics exhibit an almost unchanged equilibrium of the trans/cis ratios compared to that of Pro and 4-trifluoromethylproline derivatives. Finally we demonstrate that the 3,4-double bond in the investigated structures leads to an increase of the amide rotational barriers, presumably due to an interplay with the transition state.

Introduction

The sole genetically encoded secondary amino acid proline (Pro, 1) is known for its unique properties in biological systems. In particular, Pro residues are often found in the s-cis peptidyl-Pro conformation, due to the low energy difference between the s-trans and the s-cis conformational states (ca. 3–4 kJ/mol) [1,2]. In addition, the high energy barrier of the s-cis-s-trans isomerization (84–89 kJ/mol) stabilizes the amide conformers kinetically (Scheme 1) [3]. By comparison of the amide rotational rates of peptidyl-Pro with the ones of the closest Pro structural analogues, azetidine-2-carboxylic acid (norproline) and piperocil acid (homoproline) [4], it appears that the high isomerization barrier is a feature associated with the 5-membered pyrrolidine ring of Pro [5].

The pyrrolidine ring of Pro can be found in several conformations, designated as the exo- and endo-puckers, as well as in twisted forms [6,7]. Various ring substituents can significantly shift the equilibrium towards a high preference for one particu-
Table 1: pKₐ Values determined for the amino acids 1–4 and their N-acetyl derivatives.

| Xaa    | ammonium group pKₐ | carboxyl group pKₐ | ΔpKₐ |
|--------|---------------------|--------------------|-------|
|        | in Xaa s-trans      | in Ac-Xaa s-cis    |       |
| 1, Pro | 10.68               | 3.55               | 2.85  | 0.70 |
| 2, Dhp | 9.78                | 3.03               | 2.37  | 0.66 |
| 3, TfmPro | 8.46            | 3.21               | 2.57  | 0.64 |
| 4, TfmDhp | 7.60            | 2.65               | 1.99  | 0.66 |

ΔpKₐ = pKₐ(s-trans) − pKₐ(s-cis).

Results and Discussion

Firstly, we determined the pKₐ of the ammonium group in the free amino acids (Table 1). Overall the values demonstrate a 0.9 pKₐ reduction upon introduction of the 3,4-double bond, and a 2.2 pKₐ reduction for the 4-CF₃-group. Thus, both modifications lead to electron depletion of the ring system. In addition, the larger effect upon incorporation of the CF₃-group indicates that both analogues 3 and 4 possess a similar orientation of the side-chain substituent with respect to the amine group. Previously we speculated that the CF₃-group in 3 should adopt an equatorial conformation, and thus stabilize the exo-pucker, based upon considerations of the vicinal J-couplings (in Ac-TfmPro-OMe, 7) [35]. Considering that the CF₃-substituent should be located within the plane of the 3-pyrroline ring in 4, this, indeed, has an orientation close to the equatorial CF₃-group placement in 3, and is not axial.

Next, we determined the pKₐ of the carboxyl groups in N-acetyl amino acids for two rotameric forms separately (Table 1). All fluoromethyl-3,4-dehydroproline (TfmDhp, 4) in simple models. Data on Pro (1) and (4S)-trifluoromethylproline (3) is used for comparison.
four compounds exhibited similar $\Delta \Delta pK_a$ values [36,37]. Though, absolute acidity was depressed with both the 3,4-double bond and 4-CF$_3$ substitutions, the former had a stronger impact. Thus it is evident that the 3,4-double bond significantly increases the electrophilicity of the carbonyl group of the amino acid residue.

The effect on the $s$-trans/$s$-cis equilibrium was revealed upon NMR investigations of the conventional methyl esters of the N-acetyl amino acids (Ac-Xaa-OMe) [38]. The equilibrium $K_{s$-trans/$s$-cis} constants in the model compounds were found to be: 5 – $4.97 \pm 0.07$, 6 – $5.45 \pm 0.09$, 7 – $4.31 \pm 0.05$ and 8 – $4.82 \pm 0.03$ (50 mM, D$_2$O, 296 K). In terms of the free energy the 3,4-double bond increased the relative stability of the $s$-trans conformer by 0.2–0.3 kJ/mol, whereas the 4-CF$_3$-group demonstrated an opposite effect of about 0.3 kJ/mol (standard error $\pm 0.1$ kJ/mol). Despite both effects being rather marginal, this indicates that the increase of the electrophilicity of the terminal carbonyl groups (as seen previously in Ac-Xaa acidity) does not have a significant impact on the intramolecular interaction between the two carbonyl groups (as seen from $K_{s$-trans/$s$-cis} values). Similarly, Jenkins et al. reported on bicyclic proline analogues and demonstrated that axially oriented electron withdrawing substituents (4-fluoro and 4-hydroxy groups) maintained the $K_{s$-trans/$s$-cis} equilibrium values of the parent unsubstituted structure [39].

We also analyzed X-ray crystal structures of the Ac-Xaa-OMe models. All four compounds crystallized in the $s$-trans conformation: 5 and 7 as racemates [40], 6 and 8 as single enantiomers (Figure 2). In addition, rac-Boc-TfmDhp (9) crystals were subjected for analysis as well. The latter crystal structure demonstrated a pseudo $s$-cis conformation due to the helical hydrogen bond structure established between the free carboxylic and the carbamate groups as C(=O)–O–H···O=C(–O$\,^t$-Bu)–N. The same has been recently reported in particular in the cases of N-Boc-2-methylproline [41] and N-Boc 4,5-difluoromethanoprolpine [42].

In the crystal structure the ring conformations were found to be: twisted $endo$-pucker for 5, $exo$-pucker for 7, and compounds 6, 8 and 9 exhibited reasonably flat pyrroline rings, that is in agreement with previous computational works. (Alternative ‘flattened’ proline analogues – 4,5-methanoprolines have also been characterized, see [43,44].) For 6 and 8 the $\varphi$-angles were found to be $-69$ and $-67^\circ$ respectively, which are typical values for a Pro residue. Importantly, both structures did not indicate any pyramidalization around the amide nitrogen atom. The N–C=O→C=O(OMe) angle was 94–97°, which is below the optimal Bürgi–Dunitz trajectory angle [45,46]. The $\varphi$-angles in 6 and 8 found in the crystals and the $K_{s$-trans/$s$-cis} values found in solution both indicate close conformational similarities between Dhp and Pro fragments.

Finally, we identified the amide rotation rates and corresponding activation barriers by EXSY NMR (D$_2$O, 310 K). In order to establish a proper reference system we correlated the observed activation energies with the $pK_a$ of the ammonium groups in the corresponding free amino acids (Scheme 2). The resulting correlation indicates a remarkable offset in the activation energy in the 3,4-double bond containing residues (Figure 3). This effect was not only observed in water, but also persisted in organic solvents (MeOD, DMSO, CDCl$_3$, see Supporting Information File 1). The activation energies are thus generally increased by the presence of the double bond by ca. 2.7 kJ/mol, that corresponds to about a factor of 1/3 of the rotational rates.

Considering that no nitrogen pyramidalization has been observed in the crystal structures, this would indicate a destabilization of the transition state by the 3,4-double bond in Dhp.
Scheme 2: The relationship between the $pK_a$ of the ammonium function in the amino acid and the amide rotational barrier in proline analogues. The substituents that impose a $pK_a$ depression effect should also decrease the content of the resonance structure with the separate charges in the ground state of the corresponding amide, that leads to a lowering of the rotational barrier.

Figure 3: The double bond between C$_3$ and C$_4$ atoms in 3,4-dehydroproline residues induces an increase in the amide rotational barriers in Ac-Xaa-OMe.

5-CF$_3$-group impose electron-withdrawing effects on the functional groups of the amino acids. Though, the carboxyl function is influenced more strongly by the double bond, whereas the amino group is more affected by the structurally proximal 4-CF$_3$-substituent. Conversely, the backbone conformational properties and the s-trans/s-cis energy differences remain nearly non-affected in both cases. Finally, the 3,4-double bond was found to increase the barrier of the amide rotation presumably due to the repulsive effect between the amide oxygen and the double bond in the syn/exo transition state. Thus 3,4-dehydroproline can be considered as a potential structural ‘freezer’ for polypeptide structures.

Supporting Information

The crystal structures are deposited in Cambridge Structural Database under the following IDs: 5-CCDC1443104, 6- CCDC1443105, 8- CCDC1443103, 9- CCDC1443102. The crystal structure of 7 have already been discussed in [35] and the deposit number was CCDC1042476. The structure files can be retrieved free of charge at http://www.ccdc.cam.ac.uk.

Supporting Information File 1

Experimental procedures, values for the amide rotational barriers in different solvents, copies of the NMR spectra and ellipsoid diagrams of the X-ray crystal structures. [http://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-12-57-S1.pdf]

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and TfmDhp. Indeed, it is well known, that the peptidyl-Pro amide bond rotation proceeds via the syn/exo transition state, where the oxygen atom of the amide group moves under the pyrrolidine ring, approaching to C$_3$ and C$_4$ atoms [47]. Repulsion between the oxygen lone pairs and the double bond in the Dhp residue could cause the experimentally evident increase in the rotation barriers.

Conclusion

In summary, we performed the experimental characterization of proline analogues with a 3,4-double bond: Dhp and TfmDhp. Our results confirmed ‘flattening’ of the proline ring by the double bond, in agreement with what has been previously suggested by theoretical studies. Both, the 3,4-double bond and the
