

Cis–trans isomerization of omega dihedrals in proteins

Pierrick Craveur · Agnel Praveen Joseph ·
Pierre Poulain · Alexandre G. de Brevern ·
Joseph Rebehmed

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Abstract Peptide bonds in protein structures are mainly found in *trans* conformation with a torsion angle ω close to 180° . Only a very low proportion is observed in *cis* conformation with ω angle around 0° . *Cis–trans* isomerization leads to local conformation changes which play an important role in many biological processes. In this paper, we reviewed the recent discoveries and research achievements in this field. First, we presented some interesting cases of biological processes in which *cis–trans* isomerization is directly implicated. It is involved in protein folding and various aspect of protein function like dimerization interfaces, autoinhibition control, channel gating, membrane binding. Then we reviewed conservation studies of *cis* peptide bonds which emphasized evolution constraints in term of sequence and local conformation. Finally we made an overview of the numerous molecular dynamics studies and prediction methodologies already developed to

take into account this structural feature in the research area of protein modeling. Many *cis* peptide bonds have not been recognized as such due to the limited resolution of the data and to the refinement protocol used. *Cis–trans* proline isomerization reactions represents a vast and promising research area that still needs to be further explored for a better understanding of isomerization mechanism and improvement of *cis* peptide bond predictions.

Keywords Protein structures · Amino acids · Proline · Protein folds · Steric constraints · Secondary structures · Protein flexibility · Structural alphabet · Protein blocks · Protein data bank, backbone angle

Introduction

Proteins are major components of all cells. Composed of a sequence of amino acids, they are the support of critical

P. Craveur and A. Praveen Joseph contributed equally.
A. G. de Brevern and J. Rebehmed contributed equally.

P. Craveur · P. Poulain · A. G. de Brevern (✉) ·
J. Rebehmed (✉)
INSERM UMR-S 665, Dynamique des Structures et Interactions
des Macromolécules Biologiques (DSIMB), Université Denis
Diderot–Paris 7, INTS, 6, rue Alexandre Cabanel,
75739 Paris cedex 15, France
e-mail: alexandre.debrevern@univ-paris-diderot.fr

J. Rebehmed
e-mail: joseph.rebehmed@univ-paris-diderot.fr

P. Craveur · P. Poulain · A. G. de Brevern · J. Rebehmed
Univ. Paris Diderot, Sorbonne Paris Cité,
UMR-S 665, 75739 Paris, France

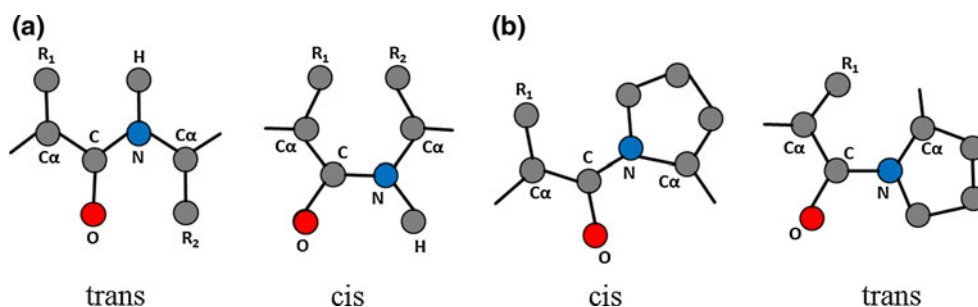
P. Craveur · P. Poulain · A. G. de Brevern · J. Rebehmed
Institut National de la Transfusion Sanguine (INTS),
75739 Paris, France

P. Craveur · P. Poulain · A. G. de Brevern · J. Rebehmed
Laboratoire d'Excellence GR-Ex, 75739 Paris, France

A. P. Joseph
National Centre for Biological Sciences, UAS-GKVK Campus,
Bellary Road, Bangalore 560 065, Karnataka, India

Present Address:
P. Poulain
Ets Poulain, Pointe-Noire, Congo

Fig. 1 **a** Peptide bonds can adopt two different conformations, *trans* and *cis*. In the *trans* isomer, the C=O and N–H groups point in the opposite directions, whereas in the *cis* isomer, they point in the same direction. **b** When the second residue of a peptide bond is a proline, the *cis*-form is more frequent



biological functions. Amino acids are composed of a succession of N, C α , C and O atoms defining the protein backbone, while more versatile side-chains give them specific physiochemical specificities. The backbone chain of proteins is constituted of three repeating dihedral angles: the phi-angle (φ) described the torsion angle around the N–C α bond, the psi-angle (ψ) around the C α –C bond, and the omega-angle (ω) around the C–N bond—peptide bond. The φ and ψ -angle define the structure of a protein backbone. Due to the delocalization of the carbonyl π electrons and the nitrogen lone pair, the ω angle tends to be planar and therefore is very rigid. From a theoretical point of view, the planar peptide bonds, between the carbonyl carbon atom and the amide nitrogen, in protein structures are more favorable in *trans* conformation as the amide hydrogen of *trans* peptide bonds offers less steric repulsion to the preceding C α atom than it does in the *cis* isomers (see Fig. 1).

The first protein structures obtained by X-ray crystallography clearly agree with this point (Ramachandran and Sasisekharan 1968). The occurrence of a peptide bond in the *cis* conformation was occasional. Only a specific case had been underlined with the Xaa-Pro peptide bond (Xaa represents any amino acids), where both *cis* and *trans* isomers experience similar steric clashes with the neighboring and are nearly equivalent energetically. Therefore, it was considered that the vast majority of *cis* peptides are observed with a proline. The *cis* conformation for such bond becomes feasible due to a steric interaction between the C δ -atom of the proline side chain and the backbone atoms destabilizing the *trans* conformer (Stewart et al. 1990; Weiss et al. 1998a). Nonetheless non-proline *cis* bonds are also found in proteins, although they occur much less frequently (Jabs et al. 1999). In a nonredundant set of 571 proteins, 5.2 % of Xaa–Pro and 0.03 % of Xaa–non-Pro peptide bonds were found in *cis* conformation. In random coil polypeptides, the proportion of Xaa-Pro *cis* bonds varies between 5 and 30 %. The size of this population is influenced mainly by the identity of neighboring residues (Brandts et al. 1975; Grathwohl and Wuthrich 1976; Raleigh et al. 1992). Figure 2 presents the *cis* and *trans* conformational isomers of a cysteine–proline peptide

bond in the Bovine NPC2 (Nieman-Pick C2) protein (PDB code 1NEP) (Friedland et al. 2003).

Even if they represent a very limited number of residues, the *cis* conformers have biological implications. It was shown for the first time (35 years ago) by Schmid et al. that the *cis*–*trans* isomerization of peptide bonds on the N-terminal side of a proline plays an important role in the folding process of the protein (Schmid and Baldwin 1978). With the increasing of three-dimensional structures of proteins available today, the importance of *cis* peptide bonds started to emerge. Later on, the importance of the *cis*–*trans* isomerization process of the peptide bond in protein folding has grown bigger (Levitt 1981; Kiefhaber et al. 1990; Wedemeyer et al. 2002). Additionally, the isomerization reaction is also used as a molecular timer in a number of biological processes, including cell signaling (Sarkar et al. 2007; Wulf et al. 2005), ion channel gating (Lummiss et al. 2005) and gene expression (Nelson et al. 2006). Its deregulation is related to pathological conditions, such as cancer (Suizu et al. 2006), amyloid formation (Eakin et al. 2006; Pastorino et al. 2006) and Alzheimer's disease (Lu et al. 2007).

In this review, we will underline some characteristic biological examples of *cis*–*trans* isomerization of omega dihedrals in proteins. Then, we will highlight the interest of structural analyses to conclude with the different *in silico* approaches applied on this subject.

I. Importance in biological processes

The *cis*–*trans* isomerization of omega angles consists of a local conformation change that is often compensated by local variations of backbone angles of the residues flanking the *cis* peptide, thereby avoiding an important global change. Nonetheless, these small changes are sometimes biologically crucial. Some interesting cases are presented below.

Active sites and dimerization interface

Many *cis* peptide bonds were found to be located at the active sites of proteins and dimerization interfaces. For

Fig. 2 Representation of the *cis* and *trans* conformations in a cysteine–proline (Cys–Pro) peptide bond. The displayed structure is the Bovine NPC2 (Niemann–Pick C2) protein (PDB code 1NEP) (Friedland et al. 2003) obtained from the Protein Data Bank. In the *cis* conformation, the alpha carbons are locked on the same side of the peptide bond, whereas in the *trans* conformation, they lie on opposite sides of the peptide bond

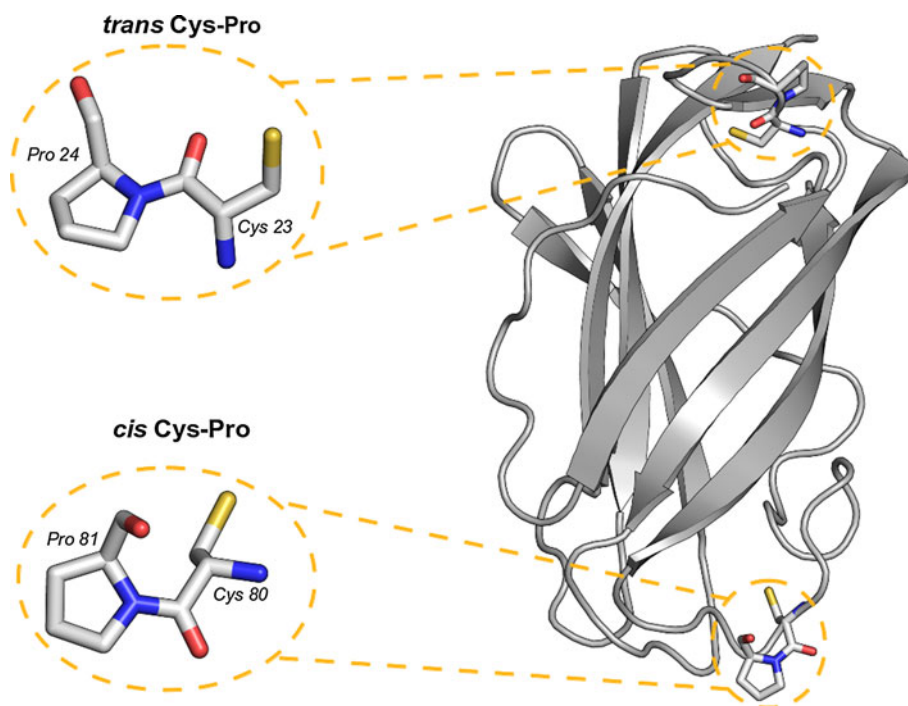
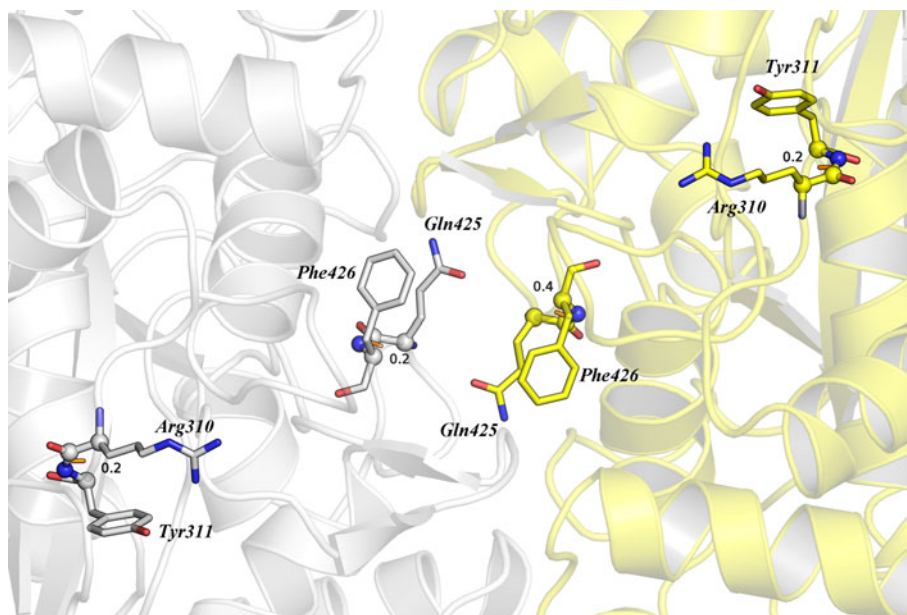


Fig. 3 Structure of the factor XIII dimer [PDB code 1F13, (Weiss et al. 1998b)]. One subunit is colored in *silver* and the second in *yellow*. Two non-proline *cis* peptide bonds are shown in each subunit with partner residues in sticks. One is between Arg310 and Tyr311 close to the active site and the other is between Gln425 and Phe426 at the dimerization interface. The value of the ω angle for the different *cis* peptide bonds is labeled on the figure and marked by an *orange bar*. The atoms composing the ω angle are represented in *spheres* (color figure online)



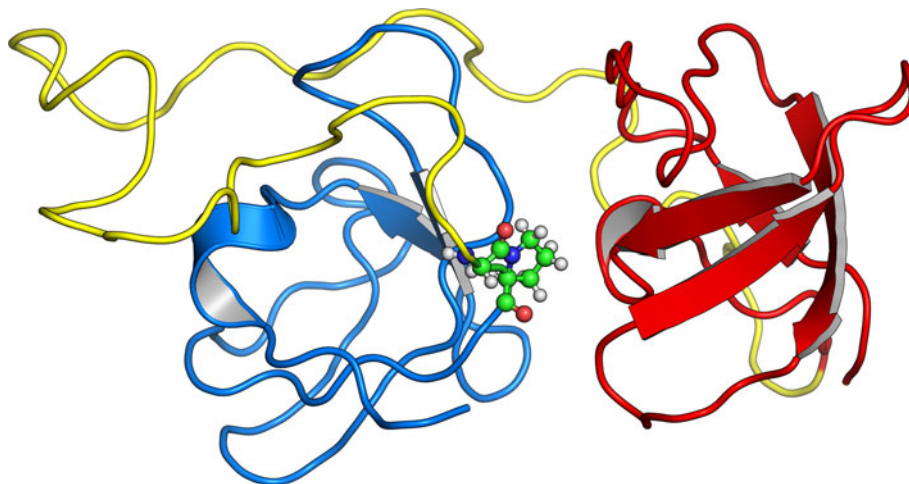
instance, the recombinant human cellular factor XIII zymogen (Weiss et al. 1998b), which catalyzes the enzymatic cross-linking of fibrin monomers into stable polymers and protects polymers from plasmatic and nonspecific degradation, has two non-proline *cis*-peptide bonds in its crystal structure. One is between Arg310 and Tyr311 close to the active site and the other is between Gln425 and Phe426 at the dimerization interface (see Fig. 3). In the same way, the structure of Chitinase B, an hydrolytic enzyme that breaks down glycosidic bonds in chitin, holds

a non-proline *cis* bond between Glu144 and Tyr145 in the active site; these two residues are involved in direct contact with the substrate and are also in the vicinity of the dimer interface (van Aalten et al. 2001).

Modulation of signaling: autoinhibition control

Autoinhibition is a natural control of protein activities and is typically regulated by extrinsic factors. For instance, the Kit kinase is autoinhibited through an intramolecular

Fig. 4 A NMR model of Crk adaptor [PDB code 2L3S, (Sarkar et al. 2010)]. The presence of a heterogeneous proline residue undergoing *cis*–*trans* isomerization provides a potential binding site for PPIase enzymes. The Gly237–Pro238 bond (in green) is located on the linker (colored in yellow) between the two SH3 domains (colored in red and blue) (color figure online)



interaction with the juxtamembrane (JM) domain. Kit JM behaves as an autonomously folding domain that binds directly to the amino-terminal lobe of the Kit kinase and prolongs its induction time to activation (Chan et al. 2003).

Another example is the adaptor molecule Crk that binds to several tyrosine-phosphorylated proteins. It is involved in several signaling pathways, recruiting cytoplasmic proteins in the vicinity of tyrosine kinase through src-homology domains 2-phosphotyrosine interaction. Recently, Sarkar and collaborators (Sarkar et al. 2007, 2010) showed that autoinhibition of Crk can be also controlled by an intrinsic intramolecular switch afforded by a *cis*–*trans* isomerization. A proline of the linker tethering the two SH3 domains of the Crk adaptor protein interconverts between the *cis* and *trans* conformation (see Fig. 4). In the *cis* conformation, the two domains interact intramolecularly, leading to autoinhibition. The isomerization facilitates the interactions between domains of the same protein. Conversely, in the *trans* conformation, Crk exists in an extended, uninhibited conformation that serves to activate the protein upon ligand binding. This interconversion between the *cis* and *trans* conformations can be accelerated by the action of peptidyl isomerases. The authors concluded that proline isomerization make an ideal switch that can regulate the kinetics of activation, thereby modulating the dynamics of signal response.

Channel gating

The 5-hydroxytryptamine type 3 receptor (5-HT₃R) is a transmembrane, G-protein-coupled protein. It belongs to the superfamily of ligand-gated ion channels and is closely related to the nicotinic acetylcholine receptor. Composed of five subunits arranged around a central ion conducting pore, it is permeable to potassium and calcium ions and is well known to bind serotonin leading to an excitatory response in neurons.

Lumms and co-workers (Lumms et al. 2005; Melis et al. 2009) suggest that *cis*–*trans* isomerization of a single proline provides the switch that interconverts the open and closed states of the channel. A model has been proposed for 5-HT₃R gating whereby *trans*–*cis* isomerization of a specific proline (Pro308) at the apex of the M2–M3 loop would induce a conformational change in M2, resulting in a displacement of the hydrophobic zone that occludes the pore and opening the channel. A series of proline analogues was incorporated at the position 308 by mutagenesis. Proline analogues that strongly favor the *trans* conformation produced nonfunctional channels. Molecular rearrangement, after ligand binding, of the protein backbone at Pro308 from the *trans* conformation to the *cis* conformation can be the structural mechanism that opens the receptor pore. Figure 5 underlines this particular position.

Proline has unique structural and conformational properties, and it is found with anomalously high frequency in the transmembrane regions of ion channels and transporters, suggesting a key role in structural changes associated with transmembrane signaling (Brandl and Deber 1986; Sansom and Weinstein 2000).

Membrane binding

The winged helix gene Brain factor-1 (BF1 or Forkhead box protein G1, FOXG1) has a pleiotropic role in the development of the cerebral hemispheres of the brain. Mice lacking BF1 have defects in the morphogenesis of the structures of the dorsal and the ventral telencephalon with important consequences, e.g., for development of the vertebrate olfactory system. This 489-residue long protein, is known to bind membrane. In the process of phospholipid binding, Pro22 of BF1 may undergo, in the presence of calcium ions, isomerization from the *trans* to *cis* conformation in a step essential for membrane binding (Evans and Nelsestuen 1996). Perera and co-workers used molecular dynamics

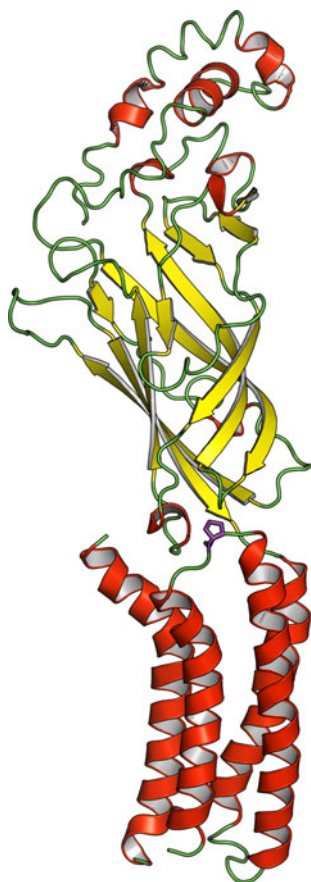


Fig. 5 Model of the 5-hydroxytryptamine type 3 receptor. Pro308, in its classical *trans* form, is shown as *sticks* and colored in *purple*. This model was generated by SPARKS (Yang et al. 2011) (color figure online)

(MD) simulations to investigate the relative importance of the two conformations of Pro22 to the structural and dynamical properties of BF1 (Perera et al. 1998). They underlined that the overall structural changes associated with the *trans*–*cis* isomerization is negligible; the calculated electrostatic potential energy surfaces is very similar and cannot explain the binding specificities. Only minor modifications to the hydrogen bond network take place.

Protein stability

It was also observed that *cis*–*trans* isomerization of the peptide bond can impact proteins stability. Truckses and collaborators encountered an example with the *Staphylococcus aureus* nuclease. This enzyme is an endo-exonuclease that preferentially digests single-stranded nucleic acids. They observed a mixture of *cis* and *trans* configuration of the Lys116–Pro117 peptide bond. In terms of free energy, they measured that approximately 17 % of the increase in protein stability manifests itself as stabilization of the *cis* configuration at Lys116–Pro117 (Truckses et al. 1996).

II. Analyses and conservation of *cis*–*trans* peptide bonds in proteins

Analyses

As *cis* conformations are occasional, they are expected to occur for specific biological and physical reasons; they would also be more conserved. MacArthur and Thornton calculated the ω peptide angle distribution in the Cambridge Structural Database of small molecules (Allen et al. 1979). They found only 54 on 1,712 peptides with *cis* conformation. They extended their approach to a nonredundant set of proteins from the Brookhaven protein database (Bernstein et al. 1977) and they found out that peptide bonds in the *cis* conformation represented less than 0.1 % of the peptide bonds in their dataset (MacArthur and Thornton 1996).

Two years later Weiss and co-workers analyzed specifically the *cis*-peptide bonds (Weiss et al. 1998a). They used a nonredundant set of 571 X-ray protein structures taken from the Brookhaven protein database (Bernstein et al. 1977) with a resolution of 3.5 Å or better. *Cis* conformation represented 0.28 % of all omega angles. They underlined that most of the peptide bonds in *cis* conformation occur where the peptide bond is an imide (Xaa–Pro) rather than an amide bond (Xaa–non-Pro). The authors also discussed the discrepancy between the fraction of *cis* peptide bonds observed and what can be predicted from free enthalpy values. One of the main results of this study was the significant correlation between the resolution of the structure solved and the number of *cis* peptides detected. High-resolution structures (resolution <2.0 Å) contain almost twice the number of Xaa–Pro bonds than medium- and low-resolution structures (resolution ≥ 2.5 Å) and almost four times the number of Xaa–non-Pro bonds.

This conclusion brought our attention to the fact that many *cis* peptide bonds may have passed unnoticed due to an automatic usage of refinement programs (Huber and Steigemann 1974). Moreover, Gunasekaran and co-workers showed that half of disallowed regions residues (often due to *cis*-conformation) in the Ramachandran plot are linked to biological functions (Gunasekaran et al. 1996). These unnoticed *cis* peptide bonds could impact the understanding of many protein function mechanisms.

Evolution

Due to their structural and functional importance, *cis* peptide bonds in proteins are expected to be conserved during evolution. The first conservation study was made on a set of nonhomologous and high-resolution structures (sequence identity <25 % and resolution ≤ 2.0 Å) (Lorenzen et al. 2005). The set contained 1,729 proteins

corresponding to 742 *cis* and 14,502 *trans* prolyl residues. They show, as previously reported (Pal and Chakrabarti 1999), that position preceding *cis*-proline is strongly occupied by the aromatic amino acids tryptophan and tyrosine, and by small residue glycine, as well as proline itself. Phenylalanine and tyrosine, but not tryptophan, are the most present in the succeeding position of the prolyl residue. This predominance for aromatic and small residues at neighboring positions of *cis*-proline could be respectively explained by π interactions between the aromatic and the proline rings, and by the lack of steric hindrance (Wu and Raleigh 1998). The authors analyzed the impact of local and global sequence homology and showed that (i) *cis* prolyl residues are more often conserved than *trans* prolyl residues, (ii) *cis* and *trans* prolyl residues are more conserved than the surrounding amino acids, and (iii) neighboring amino acids have a direct influence on the probability of forming a *cis* prolyl residue.

The residue conservation in term of structural homology is also an important feature. It was firstly analyzed to illustrate the usefulness of structure alignments database to improve comparative protein modeling (Sali and Overington 1994). The underlying idea is that restraints of *cis/trans* isomerism in a modeled sequence could be provided by homologous structures. Despite the small number of example in the used dataset (238 *cis*-prolines), the authors estimated that the knowledge of equivalent conformations increases the prediction success from 0 to 82.9 % for *cis*-proline and from 93.3 to 96.2 % for *trans*-proline, compared with the only overall stereochemical preference of proline taken into account. The same conclusion was also obtained by structurally aligning proteins from same SCOP (Murzin et al. 1995) superfamily (Lorenzen et al. 2005). The best decision about the presence of a *cis* prolyl residue can be made by modeling according to the highest resolution structure with more than 20 % sequence identity to the query protein.

A very recent study consisted of analyzing the preferences of amino acid types and local backbone conformations associated with *cis* peptide bonds. In this study, a larger set of high-quality protein structures solved by x-ray crystallography with a resolution better than 1.6 Å was used (Joseph et al. 2012). The authors emphasized the variations between *cis* and *trans* conformers in structurally similar proteins. They described the local conformations associated with these peptides using a Structural alphabet named Protein Blocks (PBs) (de Brevern et al. 2000; Etchebest et al. 2005; Joseph et al. 2010). PBs are local approximation of protein structures and are useful to analyze local protein conformations [see for instance (Jallu et al. 2012)]. Considering the Xaa-Proline *cis* bonds, 34 % are not conserved in structural homologous proteins; and proline has a high tendency to get replaced by another

amino acid in the *trans* conformer (87 %). The authors showed that the change between *cis* and *trans* conformations are efficiently utilized for the emergence of new function among similar protein folds. Many of the non-proline *cis* peptides are more prone to such changes than the Xaa-Pro peptides (Joseph et al. 2012). This study also underline the fact that aligned structural regions can easily have *cis* and *trans* conformation in front of each other despite the huge change of omega angles; in most cases, the other angles compensate the local change in the backbone directions.

III. In silico

Prediction of cis-trans isomerization in proteins

In regards to other related research areas, the number of studies aiming to predict the peptide bond conformation between amino acids is limited. The main limitation of these approaches is that the fraction of peptide bonds adopting a *trans* conformation is 350 times more important than the *cis* conformation, and even 20 times more for the prolyl peptide bond. Therefore, the prediction tools of *cis-trans* isomerization focus exclusively on Xaa-Pro *cis* peptide bonds.

The beginning of the 1990s witnessed the first attempt to predict peptide bond conformation of prolines in proteins. The methodology consisted of determining sequence patterns flanking a centered proline (Frommel and Preissner 1990). The authors considered windows up to the sixth residues around the proline, and their physicochemical properties. They identified six different patterns to discriminate between the *cis* and *trans* conformations. They correctly predicted about 73 % of *cis* prolyl residues and did not predict any proline in *trans*-conformation as *cis* conformation. Nonetheless, the dataset used was too restricted, with only 242 Xaa-Pro cases to make reliable prediction on larger dataset.

COPS algorithm (Pahlke et al. 2005a, b) was the first to predict the peptide bond also for Xaa-non-Pro bonds. The algorithm is based on an extension of the Chou-Fasman parameters (Chou and Fasman 1974). It predicts conformation of the peptide bond by taking into account only the secondary structure of amino acid triplets. Despite a large dataset (8,584 proteins yielded 25,663 amino acids in *cis* conformation and ~11 million amino acids in *trans* conformation), a sensitivity of 35 % and an accuracy of 66 % underlined the difficulty of the approach.

Many methods are based on Support Vector Machine (SVM) (Vapnik 1999). The first study of prolyl *cis/trans* isomerization took into account amino acid properties such as secondary structure prediction, solvent accessibility prediction and the physicochemical properties of the

surrounding amino acids (Wang et al. 2004). They used larger dataset than previous studies (2,193 structures) and reported a better success rate. However, no software or web server had been available, making it difficult to assess its real performance as the validation steps seem biased.

More interestingly, Song and co-workers proposed another method based on SVM in 2006 (Song et al. 2006). It used the amino acid compositions of local sequence flanking centered proline residues (different window sizes were tested), the position specific scoring matrix (PSSMs) extracted from PSI-BLAST (Altschul et al. 1997) and the predicted secondary structure extracted from PSIPRED (Jones 1999). During this study, different kernel functions and parameters were tested for the SVM training and testing to unbiased the prediction. Their method yielded a prediction accuracy of 71.5 % and a Matthews Correlation Coefficient (MCC) (Matthews 1975) of 0.43. The authors also compared their newly developed method with those published (Frommel and Preissner 1990; Pahlke et al. 2005b) (see Table 6 of (Song et al. 2006)). The authors pointed out the fact that the method accuracy depends on the training set size, and mentioned that they have also developed a web server named CISPEPpred (<http://sunflower.kuicr.kyoto-u.ac.jp/~sjn/cispep/>).

Exarchos and co-workers presented another method to predict all types of *cis* peptide bonds. It is related to previous studies using evolutionary profiles, secondary structure information, solvent accessibility predictions for each amino acid and the physicochemical properties of the surrounding residues with SVMs. The methodology was more complex than the previous ones for the selection of features. They also used a large dataset of high quality proteins (3,050 structures) with resolution <2.0 Å and sequence identity <25 %. They made distinction of the four peptide bonds *cis*–Pro, *cis*–non-Pro, *trans*–Pro and *trans*–non-Pro and also developed the PBOND web server (that it currently no more available) (Exarchos et al. 2009a). They reported a sensitivity up to 77 % and an accuracy up 74 %. The extracted patterns were also compared against the PROSITE database, in order to gain insight into the implications of *cis* prolyl bonds in protein functions (Exarchos et al. 2009b).

Molecular dynamics simulations

The usefulness of molecular dynamics simulation techniques for the study of *cis*–*trans* isomerization of proline has been demonstrated in many studies. A first interest in computational approach was given by Levitt in the beginning of the 1980s (Levitt 1981). He performed conformational energy calculations to study the role of the proline residues in the folding of the bovine pancreatic trypsin inhibitor (BPTI) by forcing the *trans*–*cis* isomerization of

four prolines. It underlined the existence of a potential alternative folded native-like conformation for either isomeric form when the *cis* form is calculated to destabilize slightly the folded structure.

Another example was presented later by Ikura and co-workers (Ikura et al. 1997). Using the *Staphylococcus aureus* nuclease (SNase), they studied the urea-induced unfolding transition and five proline mutants, coupling experimental methods (circular dichroism, absorption spectroscopy, refolding–unfolding kinetics of the proteins by stopped-flow circular dichroism and stopped-flow absorption techniques) and *in silico* approaches (molecular dynamics). Two phases were observed in the unfolding of the wild-type and mutant proteins (that contained a specific Pro117); a fast phase corresponding to the unfolding of the *trans* isomer and a slow phase corresponding to that of the *cis* isomer. On the basis of these results, they also discussed the folding scheme of SNase.

Hodel and co-workers obtained thermodynamic and structural description of the conformational equilibrium of a particular protein loop. The authors used an exhaustive conformational search that identified several substrates followed by free energy simulations between the substrates (Hodel et al. 1995). These simulations correctly predicted a small free-energy difference between the *cis* and *trans* forms composed of larger, compensating differences in enthalpy and entropy. The structural predictions of these simulations were qualitatively consistent with known X-ray structures of nuclease variants.

Another interesting approach was developed by Darve and co-workers with an adaptive biasing force molecular dynamics simulation (Darve et al. 2008). This approach was used to force *cis*–*trans* isomerization at Pro32 of β 2-microglobulin and to calculate the relative free energy in the folded and unfolded state (Fogolari et al. 2011). The authors carried clockwise and anticlockwise rotations to sample the whole $-180:180$ interval of the ω angle between His31 and Pro32 to generate the energy profile. They underlined clearly that *cis* state is favored in the native structure.

In our laboratory, we have also observed the effect of *cis* and *trans* conformation on protein dynamics of the platelet integrin α IIb β 3. Composed of two subunits, it is implicated into fibrillar aggregation and responsible of alloimmune thrombocytopenia or Glanzmann's thrombasthenia. Through *in silico* approaches, we showed the impact of punctual mutation (Jallu et al. 2010, 2012). We also studied the mutation S163P of the β 3 subunit (see Fig. 6). As no observation is known about the local conformation of the Pro163 residue, we tested both *cis* and *trans* conformations with long MD simulations [see Jallu et al. (2012) for technical methodology and results]. We observed that both isomers are highly rigid; they do not

Fig. 6 β I domain of the β 3 subunit of the platelet integrin α IIb β 3. Structure colored in purple and silver contained the *trans* and *cis* Ser162-Pro163 peptide bond, respectively (color figure online)

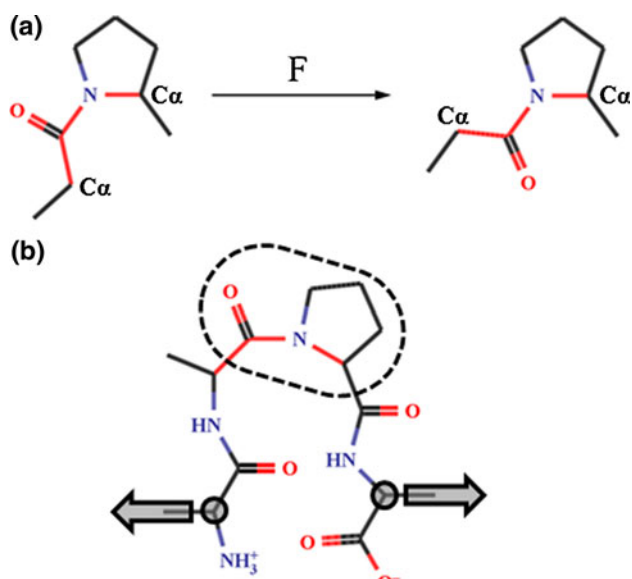
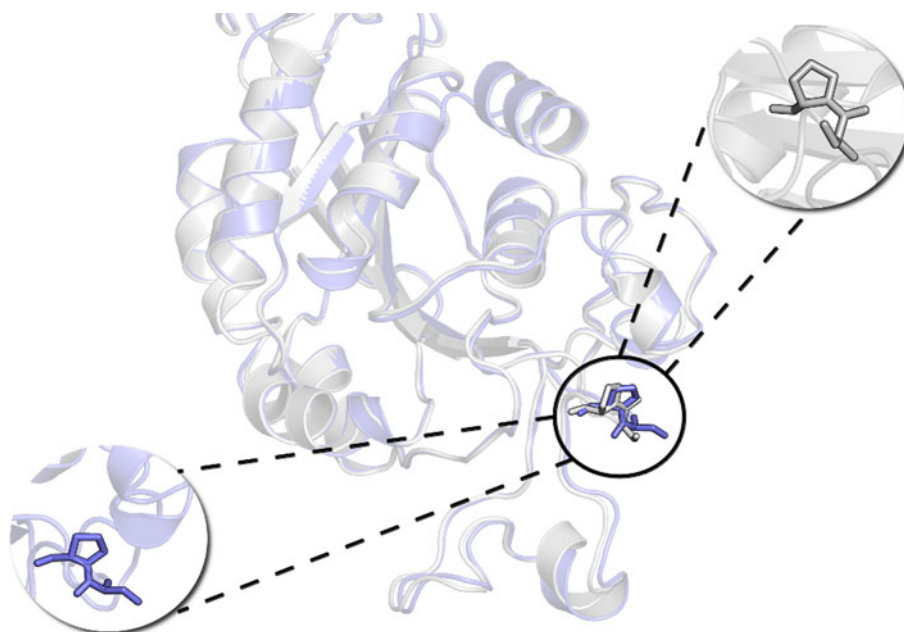


Fig. 7 **a** Schematic presentation of the *cis* to *trans* isomerization. Peptide bonds are highlighted in red. **b** The simulated AAPA peptide with QM region separated from the MM region by a dashed line. In the Force-clamp molecular dynamics, the $C\alpha$ atoms (gray spheres) were subjected to a constant force in opposite directions (color figure online)

locally move during the different simulations, but due to the steric constraints they also keep distinct local conformations, i.e., distinct PBs series (Poulain, Jallu and de Brevern, unpublished).

QM/MM studies

QM/MM methods were applied to study the electronic forces that facilitate the peptidyl *cis* to *trans*

interconversion, because conventional force fields based on classical mechanics do not take into account the effect of electronic state changes. The general mechanism of peptide bond isomerization involves a change of the bond order from a partial double-bond to a single bond, followed by a return to partial double-bond character. This pyrrolidine N changing from a planar sp^2 state to a pyramidal sp^3 state allows the rotation (Yonezawa et al. 2009; Kang 2006). By cons, QM cannot be applied to whole proteins systems because the computational cost exceeds current resources.

Umbrella sampling methods were used to explore the free-energy landscape during the isomerization of a dipeptide (Yonezawa et al. 2009). It showed that the *trans* state is more stable than the *cis* state by 4 kcal/mol and the high energy barrier separating the two states is about 20 kcal/mol.

Force-clamp molecular dynamics (FCMD) simulations were used to accelerate the interconversion from *cis* to *trans* conformation of a tetrameric peptide (AAPA) (Chen et al. 2012) (Fig. 7). Its termini $C\alpha$ atoms were subjected to a constant force in opposite directions. The authors observed the variation of different order parameters related to the isomerization of the peptide bond in respect to the simulation time: torsion angle of the peptide bond, distance between the $C\alpha$ atoms, and length of the peptide bond. Force releases the partial double bond and converts it to a single bond. Rotation around the peptide bond can then occur and afterward the partial double bond reforms in the *trans* conformation. Their study demonstrated that *cis* to *trans* isomerization can be triggered by mechanical force.

QM/MM MD simulations in combination with mean reaction force (MRF) (Vohringer-Martinez and Toro-Labbe 2011) were applied to Pin1, an enzyme that specifically

catalyzes the isomerization of peptide bonds between phosphorylated threonine or serine residues and proline. The authors wanted to discern the catalysis influence of Pin1 enzyme in the isomerization (Vohringer-Martinez et al. 2012). They observed the influence of the enzyme and identified structural and electronic contributions to activation barriers and reaction free energies for the isomerization. They carried out simulations of the peptide in two different environments: in solution and in the enzyme. The enzyme induces a rotation of the ψ angle of the preceding residue to avoid unfavorable interactions with the rotating part of the peptide.

Conclusion

The *cis–trans* isomerization and conformations are well known from researchers involved in protein structures analyses. Nonetheless, as they are quite nonfrequent, they are not often considered. This review emphasizes two points: first, the biological importance of the *cis* conformation of peptide bonds in protein structures and functions; and second, the possibility that many *cis* peptide bonds have not been recognized as such due to the limited resolution of the data and to the refinement protocol used.

Even if the number of Xaa–Pro peptide bonds in the *cis* and *trans* conformations are quite unbalanced, it is possible to observe some general preferences concerning the preceding amino acid types, with aromatic and small residues favoring the *cis* isomer slightly for example. In the same way, pertinent methodology for *cis* peptide bonds prediction seems to give quite correct prediction rate in regards to the difficulty. Accurate prediction of *cis–trans* isomerization in proteins would have many important applications towards the understanding of protein structure and function. Nonetheless, an effort is still for an accurate prediction of the Xaa–non-Pro *cis* peptide bonds.

An interesting observation was done on structural homologues that are well superimposed, but share different *cis/trans* conformation. In an important number of cases, no significant difference can be observed in terms of structural local conformations (Joseph et al. 2012). The difference of ω angles is compensated by the neighboring dihedrals.

It has been shown, that this local similarity could be associated with biological specificity. For instance, glycylamide ribonucleotide (GAR) transformylases and GAR synthetase have the same fold and bind similar substrates. Both enzymes bind ATP (or its derivatives) and GAR, but the conformation of bound ATP (or ADP) is different in the two crystal structures with *cis* and *trans* isomers. They also have similar binding sites, but the catalysis mechanisms are different. Hence, the *cis–trans* peptide changes can be

implicated in the emergence of new function among similar protein folds during evolution [see Joseph et al. (2012)].

Structure-based in silico approaches provide further details for the explanation of protein functions and specific mechanisms. Due to the computation time, most of the QM/MM studies were made on small peptides and proposed explanations may not be correct on a real protein where extra constraints can be applied on the peptide bond. *Cis–trans* proline isomerization reaction represents a vast and promising research area that still needs to be further studied.

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Conflict of interest The authors declare that they have no conflict of interest.

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