# LSM2241 Lecture 7 supplement and exercises: $\beta$ turns

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#### 1 Introduction

This supplement discusses a few concepts from the structural biology material we cover in the structure lecture, and in the associated practical. I've included a few exercises for you to work through. I will provide solutions to the exercises after I receive the first five (5) attempted solutions from students, sent to me via email.

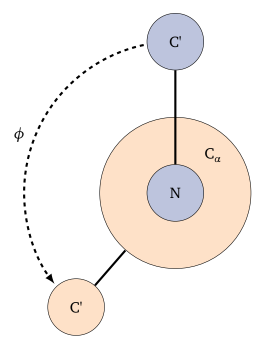
#### **Exercise on canonical secondary structures**

The diagram of fig:newman is called a Newman projection. The  $\phi$  angle for a typical anti-parallel beta sheet is shown, as it is in the lecture. Draw the Newman projection for the typical  $\phi$  and  $\psi$  angles of a *parallel* beta sheet, and the typical  $\phi$  and  $\psi$  angles of an alpha helix. Given the handedness of amino acids in proteins, we know that the alpha carbon  $C_{\alpha}$  is asymmetric for all amino acids except glycine. As a consequence, where will the  $H_{\alpha}$  and R groups appear on the projections?

Consider the angles you just drew for the  $\alpha$  helix. Which direction does it appear to turn? Is an  $\alpha$  helix left handed or right handed? Does this make sense? What if anything is missing for it to make sense?

#### 2 What defines a $\beta$ turn?

A  $\beta$  turn is a tight turn that involves four residues (*three* peptide bonds) between two close amino acids in adjacent elements of secondary structure. (A  $\beta$  turn is unrelated to  $\beta$  sheets;  $\beta$  turns can occur between sheets or helices.) We usually refer to the amino acids in the turn by number: n, n+1, n+2, and n+3. The n and n+3 residues must have  $C_{\alpha}$  atoms within 7Å of each other to be



**Figure 1.** Diagram for a  $\phi$  angle calculation. Note that the angle is measured from the atom in the front to the atom in the back. This diagram shows a  $\phi$  angle of -139, corresponding to a canonical antiparallel  $\beta$  sheet

considered a beta turn. The different types of  $\beta$  turns are characterized by different backbone angles and propensities of different amino acids.

Since the amino acids n and n+3 are within 7Å, the major distinctions between types of  $\beta$  turns are in the backbone  $\phi$  and  $\psi$  angles for the amino acids n+1 and n+2. However, analysis of turns in the PDB shows that some amino acids are more likely to occur than others at particular types of turns. This propensity can be used to predict the occurrence of a turn.

## 3 Type I $\beta$ turn

The most common  $\beta$  turn is a type I turn. It is defined by idealized  $\phi$  and  $\psi$  angles at n+1 and n+2 positions, but of course the real angles may deviate from the ideal. Hutchinson and Thornton looked at a large number of turns in the PDB, and found average backbone dihedral angles for Type I turns corresponding to the values in tab:typeI (Hutchinson and Thornton 1994).

Type I turns very often have a proline at either position n or n + 1, and very rarely have a proline at position n + 2. Hydrogen bond acceptors (histidine, apsaragine, aspartate, serine, and cystine) are common at position n, so they can accept a hydrogen bond from the backbone nitrogen of position n + 3. The propensities for different amino acids at different turn types can be found at this page.

An example type I turn can be seen in residues 39-42 of the structure of subtilisin Carlsberg (PDB entry 1CSE) which has the sequence HPDL. A close up stereo diagram is show in fig:typeI. In the practical you may give yourself a headache trying to see three dimensional figures from stereo diagrams on the screen, but you can do the same on the printed page without the headache. Try it!

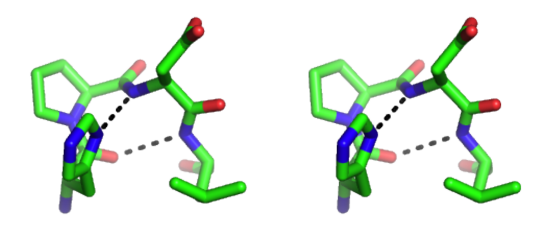
Type I  $\beta$  turns commonly show hydrogen bonding between the carbonyl carbon of position n and the amino group of position n + 3. This is highlighted as a dashed line in the figure. There is

position	$\phi$	$\psi$
n+1	-64(-60)	-27(-30)
n+2	-90(-90)	-7(0)

**Table 1.** Average  $\phi$  and  $\psi$  angles at the key positions in Type I  $\beta$  turns in the PDB, from Hutchinson and Thornton 1994. Values in parentheses correspond to idealized turns, rather than actual turns.

an additional hydrogen bond, also shown, in this structure from the backbone amino nitrogen of position n + 2 to the histidine ring nitrogen at position n.

The actual dihedral angles for this structure are show in tab:1cse.



**Figure 2.** Stereo diagram of type I  $\beta$  turn of residues 39-42 of subtilisin from PDB entry 1CSE. Hydrogen bonding within the turn is shown as dashed lines. If you have learned how to see stereo diagrams in the practical, you should be able to see this turn in three dimensions.

position	$\phi$	$\psi$
40	-62	-22
41	-90	13.8

**Table 2.** Actual dihedral angles for the key positions of the turn shown in fig:typel. These angles were measured in PyMol.

#### **Exercise**

Draw the  $\phi$  and  $\psi$  angles in a Newman projection for the n+1 and n+2 residues of a type I  $\beta$  turn. Given the L handedness of amino acids, where on the projection would the  $H_{\alpha}$  and the side chain (the R group) appear?

Imagine a proline at position n + 1, and draw the side chain connected to the backbone of the protein. Can you see why proline is preferred at this position?

## 4 Type II $\beta$ turns

Type II  $\beta$  turns are distinguished from type I by the  $\phi$  and  $\psi$  angles at positions n+1 and n+2. Type II turns also characteristically have a hydrogen bond from the carbonyl carbon of position n to the amino nitrogen of position n+3, but the  $\psi$  angle at position n+1 and the  $\phi$  angle at position n+2 are almost completely opposite the type I turn case. The average and ideal values for a large number of Type II  $\beta$  turns in the PDB, identified by Hutchinson and Thornton 1994, are shown in tab:typeII.

Like type I turns, proline is favored at position n + 1. Note the similarity of the  $\phi$  angle for this position in both types of turns.

Because of the angular requirements at position n + 2 of a type II turn, this amino acid is a glycine in about 90% of cases, but may also be an asparagine or aspartate.

position	$\phi$	$\psi$
n+1	-60(-60)	131(120)
n+2	84(80)	1(0)

**Table 3.** Average  $\phi$  and  $\psi$  angles at the key positions in Type II  $\beta$  turns in the PDB, from Hutchinson and Thornton 1994. Values in parentheses correspond to idealized turns, rather than actual turns.

An example of a type II  $\beta$  turn is residues 75-78 of yeast iso-1-cytochrome c (PDB entry 1YCC), which has the sequence IPGT and is illustrated in fig:typeII. This shows the characteristic type II turn propensity for a glycine at position n + 2.

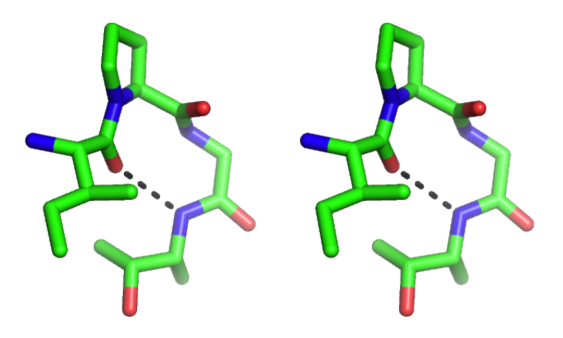


Figure 3. Type II  $\beta$  turn of residues 75-78 of yeast yeast iso-1-cytochrome c (PDB entry 1YCC). The hydrogen bond along the backbone between residues n and n+3 is shown with a dashed line. If you can see this in stereo, the difference from a Type I turn will be clear.

Again, the stereo diagram should give you a sense of how these torsion are different from those of type I turns, but to be specific, in this case the actual torsion angles are in tab:1ycc.

position	$\phi$	$\psi$
76	-66.3	128.8
77	99.7	-1.5

Table 4. Actual dihedral angles for the key positions of the turn shown in fig:typeII

#### **Exercise**

Draw the  $\phi$  and  $\psi$  angles in a Newman projection for the n+1 and n+2 residues of a type II  $\beta$  turn. Given the L handedness of amino acids, where on the projection would the  $H_{\alpha}$  and the side chain (the R group) appear?

Can you see from your Newman projections why a glycine is so strongly favored at position n + 2?

## 5 Other turn types

The other  $\beta$  turn types are also defined by characteristic torsion angles, and empirical amino acid propensities that make sense from structural considerations. The Type I' and Type II' turns are roughly the mirror images of the Type I and Type II turns, respectively, with the signs of all dihedral angles reversed.

#### References

Hutchinson, E G and J M Thornton (1994). "A revised set of potentials for beta-turn formation in proteins." In: *Protein science : a publication of the Protein Society* 3.12, pp. 2207–16. DOI: 10.1002/pro.5560031206 (cit. on pp. 2–4).