

## 3. CIF DATA DEFINITION AND CLASSIFICATION

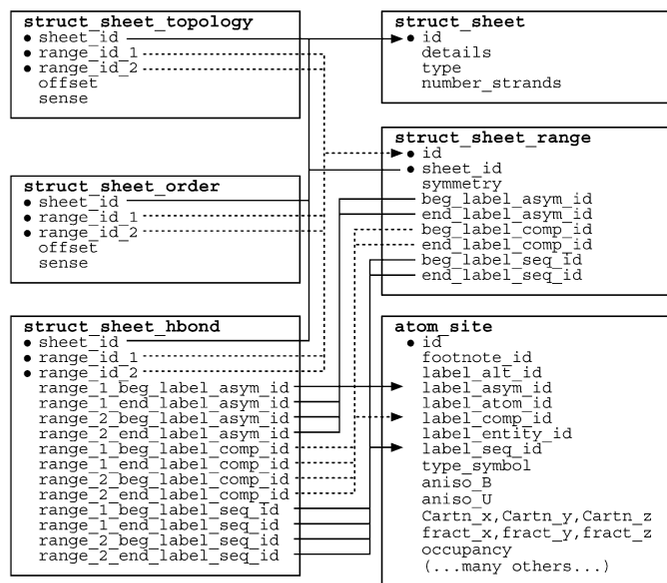


Fig. 3.6.7.14. The family of categories used to describe  $\beta$ -sheets. Boxes surround categories of related data items. Data items that serve as category keys are preceded by a bullet ( $\bullet$ ). Lines show relationships between linked data items in different categories with arrows pointing at the parent data items.

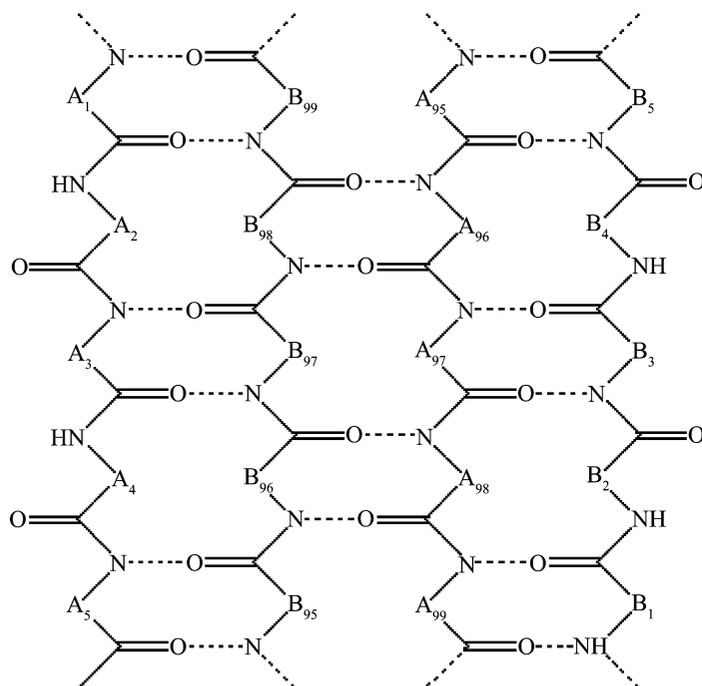


Fig. 3.6.7.15. A hypothetical  $\beta$ -sheet to be described with data items in the STRUCT\_SHEET, STRUCT\_SHEET\_ORDER, STRUCT\_SHEET\_RANGE and STRUCT\_SHEET\_HBOND categories. Note that the strands come from two different polypeptides, labelled A and B.

In the more detailed and more general method for describing  $\beta$ -sheets, data items in the STRUCT\_SHEET\_RANGE category specify the range of residues that form strands in the sheet, data items in the STRUCT\_SHEET\_ORDER category specify the relative pairwise orientation of strands and data items in the STRUCT\_SHEET\_HBOND category provide details of specific hydrogen-bonding interactions between strands (see Fig. 3.6.7.15 and Example 3.6.7.14). Note that the specifiers for the strand ranges include the amino acid ( $*_{comp\_id}$  and  $*_{seq\_id}$ ), the chain ( $*_{asym\_id}$ ) and a symmetry code ( $_{struct\_sheet\_range.symmetry}$ ). Thus sheets that are composed of strands from more than one polypeptide chain

Example 3.6.7.14. A hypothetical  $\beta$ -sheet described with data items in the STRUCT\_SHEET, STRUCT\_SHEET\_ORDER, STRUCT\_SHEET\_RANGE and STRUCT\_SHEET\_HBOND categories.

```

loop_
  _struct_sheet.id
  _struct_sheet.number_strands
  S1 4

loop_
  _struct_sheet_order.sheet_id
  _struct_sheet_order.range_id 1
  _struct_sheet_order.range_id 2
  _struct_sheet_order.sense
  S1 1 2 anti-parallel
  S1 2 3 anti-parallel
  S1 3 4 anti-parallel
  S2 1 2 anti-parallel

loop_
  _struct_sheet_range.sheet_id
  _struct_sheet_range.id
  _struct_sheet_range.beg_label_comp_id
  _struct_sheet_range.beg_label_asym_id
  _struct_sheet_range.beg_label_seq_id
  _struct_sheet_range.end_label_comp_id
  _struct_sheet_range.end_label_asym_id
  _struct_sheet_range.end_label_seq_id
  S1 1 PRO A 1 LEU A 5
  S1 2 CYS B 95 PHE B 99
  S1 3 CYS A 95 PHE A 99
  S1 4 PRO B 1 LEU B 5

loop_
  _struct_sheet_hbond.sheet_id
  _struct_sheet_hbond.range_id 1
  _struct_sheet_hbond.range_id 2
  _struct_sheet_hbond.range_1_beg_label_atom_id
  _struct_sheet_hbond.range_1_beg_label_seq_id
  _struct_sheet_hbond.range_2_beg_label_atom_id
  _struct_sheet_hbond.range_2_beg_label_seq_id
  S1 1 2 A 3 O 97
  S1 2 3 B 98 O 96
  S1 3 4 A 97 O 3
  
```

or from polypeptides in more than one asymmetric unit can be described.

It is conventional to assign the number 1 to an outermost strand. The choice of which outermost strand to number as 1 is arbitrary, but would usually be the strand encountered first in the amino-acid sequence. The remaining strands are then numbered sequentially across the sheet.

In some simple cases, the complete hydrogen bonding of the sheet could be inferred from the strand-range pairings and the relationship between the strands (parallel or antiparallel). However, in most cases it is necessary to specify at least one hydrogen bond between adjacent strands in order to establish the registration. The data items in the STRUCT\_SHEET\_HBOND category can be used to do this. Hydrogen bonds also need to be specified precisely when a sheet contains a nonstandard feature such as a  $\beta$ -bulge. This is a case where it is sufficient to specify a single hydrogen-bonding interaction to establish the registration; here only the  $*_{beg}$  or  $*_{end}$  data items need to be used to reference the atom-label components. However, it is preferable, wherever possible, to specify the initial and final atoms of the two ranges participating in the hydrogen bonding.

## 3.6.7.5.8. Molecular sites

The data items in these categories are as follows:

- (a) STRUCT\_SITE
- $_{struct\_site.id}$
  - $_{struct\_site.details}$