# **Protein Data Bank Contents Guide:**

# **Atomic Coordinate Entry Format Description**

## Version 3.20

# Document Published by the wwPDB

This format complies with the PDB Exchange Dictionary (PDBx) http://mmcif.pdb.org/dictionaries/mmcif\_pdbx.dic/Index/index.html.

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## 1. Introduction

The Protein Data Bank (PDB) is an archive of experimentally determined three-dimensional structures of biological macromolecules that serves a global community of researchers, educators, and students. The data contained in the archive include atomic coordinates, crystallographic structure factors and NMR experimental data. Aside from coordinates, each deposition also includes the names of molecules, primary and secondary structure information, sequence database references, where appropriate, and ligand and biological assembly information, details about data collection and structure solution, and bibliographic citations.

This comprehensive guide describes the "PDB format" used by the members of the worldwide Protein Data Bank (wwPDB; Berman, H.M., Henrick, K. and Nakamura, H. Announcing the worldwide Protein Data Bank. *Nat Struct Biol* **10**, 980 (2003)). Questions should be sent to info@wwpdb.org

Information about file formats and data dictionaries can be found at http://wwpdb.org.

Version History:

Version 2.3: The format in which structures were released from 1998 to July 2007.

Version 3.0: Major update from Version 2.3; incorporates all of the revisions used by the wwPDB to integrate uniformity and remediation data into a single set of archival data files including IUPAC nomenclature. See <u>http://www.wwpdb.org/docs.html</u> for more details.

Version 3.1: Minor addenda to Version 3.0, introducing a small number of changes and extensions supporting the annotation practices adopted by the wwPDB beginning in August 2007 including chain ID standardization and biological assembly .

Version 3.15: Minor addenda to Version 3.20, introducing a small number of changes and extensions supporting the annotation practices adopted by the wwPDB beginning in October 2008 including DBREF, taxonomy and citation information.

Version 3.20: Current version, minor addenda to Version 3.1, introducing a small number of changes and extensions supporting the annotation practices adopted by the wwPDB beginning in December 2008 including DBREF, taxonomy and citation information.

September 15 2008, initial version 3.20.

November 15 2008, add examples for Refmac template and coordinate with alternate conformation.

December 24 2008, update REMARK 3 templates/examples, add Norine database in DBREF, update REMARK 500 on chiral center.

February 12 2009, update example in REMARK 210 and record format in NUMMDL July 6 2009, update description for REVDAT, DBREF2, MASTER and extend number of columns for AUTHOR, JRNL, CAVEAT, KEYWDS, etc.

December 22, 2009, update CAVEAT and REMARK 265.

April 21, 2010, update REMARK 5 and add BUSTER-TNT template in REMARK 3.

December 06, 2010, update maximum number of atoms for model. Update REMARK 3 with B value type for Refmac template.

March 30, 2011, correct description and examples for FORMUL and CONECT records. Change template in REMARK 630.

### **Basic Notions of the Format Description**

#### **Character Set**

Only non-control ASCII characters, as well as the space and end-of-line indicator, appear in a PDB coordinate entry file. Namely:

abcdefghijklmnopqrstuvwxyzABCDEFGHIJKLMNOPQRSTUVWXYZ

1234567890

` - = [ ] \ ; ' , . / ~ ! @ # \$ % ^ & \* ( ) + { } | : " < > ?

The use of punctuation characters in the place of alphanumeric characters is discouraged.

The space, and end-of-line:. The end-of-line indicator is system-specific character; some systems may use a carriage return followed by a line feed, others only a line-feed character.

#### **Special Characters**

Greek letters are spelled out, i.e., alpha, beta, gamma, etc.

Bullets are represented as (DOT).

Right arrow is represented as -->.

Left arrow is represented as <--.

If "=" is surrounded by at least one space on each side, then it is assumed to be an equal sign, e.g., 2 + 4 = 6.

Commas, colons, and semi-colons are used as list delimiters in records that have one of the following data types:

List

SList

**Specification List** 

Specification

If a comma, colon, or semi-colon is used in any context other than as a delimiting character, then the character must be escaped, i.e., immediately preceded by a backslash, "\".

Example - Use of "\" character:

COMPND COMPND COMPND COMPND COMPND COMPND	<pre>MOL_ID: 1; 2 MOLECULE: GLUTATHIONE SYNTHETASE; 3 CHAIN: A; 4 SYNONYM: GAMMA-L-GLUTAMYL-L-CYSTEINE\:GLYCINE LIGASE 5 (ADP-FORMING); 6 EC: 6.3.2.3; 7 ENGINEERED: YES</pre>
COMPND	<pre>MOL_ID: 1;</pre>
COMPND	2 MOLECULE: S-ADENOSYLMETHIONINE SYNTHETASE;
COMPND	3 CHAIN: A, B;
COMPND	<pre>4 SYNONYM: MAT, ATP\:L-METHIONINE S-ADENOSYLTRANSFERASE;</pre>
COMPND	5 EC: 2.5.1.6;
COMPND	6 ENGINEERED: YES;
COMPND	7 BIOLOGICAL_UNIT: TETRAMER;
COMPND	8 OTHER_DETAILS: TETRAGONAL MODIFICATION

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## **Record Format**

Every PDB file is presented in a number of lines. Each line in the PDB entry file consists of 80 columns. The last character in each PDB entry should be an end-of- line indicator.

Each line in the PDB file is self-identifying. The first six columns of every line contains a record name, that is left-justified and separated by a blank. The record name must be an exact match to one of the stated record names in this format guide.

The PDB file may also be viewed as a collection of record types. Each record type consists of one or more lines.

Each record type is further divided into fields.

Each record type is detailed in this document. The description of each record type includes the following sections:

- Overview
- Record Format
- Details
- Verification/Validation/Value Authority Control
- Relationship to Other Record Types
- Examples
- Known Problems

For records that are fully described in fixed column format, columns not assigned to fields must be left blank.

## **Types of Records**

It is possible to group records into categories based upon how often the record type appears in an entry.

<u>One time, single line:</u> There are records that may only appear one time and without continuations in a file. Listed alphabetically, these are:

RECORD TYPE	DESCRIPTION	
CRYST1	Unit cell parameters, space group, and Z.	
END	Last record in the file.	
HEADER	First line of the entry, contains PDB ID code, classification, and date of deposition.	
NUMMDL	Number of models.	
MASTER	Control record for bookkeeping.	
ORIGXn	Transformation from orthogonal coordinates to the submitted coordinates ( $n = 1, 2, or 3$ ).	
SCALEn	Transformation from orthogonal coordinates to fractional crystallographic coordinates (n = 1, 2, or 3).	

It is an error for a duplicate of any of these records to appear in an entry.

<u>One time, multiple lines</u>: There are records that conceptually exist only once in an entry, but the information content may exceed the number of columns available. These records are therefore continued on subsequent lines. Listed alphabetically, these are:

RECORD TYPE	DESCRIPTION	
AUTHOR	List of contributors.	
CAVEAT	Severe error indicator.	
COMPND	Description of macromolecular contents of the entry.	
EXPDTA	Experimental technique used for the structure determination.	
MDLTYP	Contains additional annotation pertinent to the coordinates presented in the entry.	
KEYWDS	List of keywords describing the macromolecule.	
OBSLTE	Statement that the entry has been removed from distribution and list of the ID code(s) which replaced it.	
SOURCE	Biological source of macromolecules in the entry.	
SPLIT	List of PDB entries that compose a larger macromolecular	

	complexes.	
SPRSDE	List of entries obsoleted from public release and replaced by current entry.	
TITLE	Description of the experiment represented in the entry.	

The second and subsequent lines contain a continuation field, which is a right-justified integer. This number increments by one for each additional line of the record, and is followed by a blank character.

<u>Multiple times, one line:</u> Most record types appear multiple times, often in groups where the information is not logically concatenated but is presented in the form of a list. Many of these record types have a custom serialization that may be used not only to order the records, but also to connect to other record types. Listed alphabetically, these are:

RECORD TYPE	DESCRIPTION
ANISOU	Anisotropic temperature factors.
АТОМ	Atomic coordinate records for standard groups.
CISPEP	Identification of peptide residues in cis conformation.
CONECT	Connectivity records.
DBREF	Reference to the entry in the sequence database(s).
HELIX	Identification of helical substructures.
HET	Identification of non-standard groups heterogens).
HETATM	Atomic coordinate records for heterogens.
LINK	Identification of inter-residue bonds.
MODRES	Identification of modifications to standard residues.
MTRIXn	Transformations expressing non-crystallographic symmetry (n = 1, 2, or 3). There may be multiple sets of these records.
REVDAT	Revision date and related information.
SEQADV	Identification of conflicts between PDB and the named sequence database.
SHEET	Identification of sheet substructures.
SSBOND	Identification of disulfide bonds.

<u>Multiple times, multiple lines</u>: There are records that conceptually exist multiple times in an entry, but the information content may exceed the number of columns available. These records are therefore continued on subsequent lines. Listed alphabetically, these are:

RECORD TYPE	DESCRIPTION	
FORMUL	Chemical formula of non-standard groups.	
HETNAM	Compound name of the heterogens.	
HETSYN	Synonymous compound names for heterogens.	
SEQRES	Primary sequence of backbone residues.	
SITE	Identification of groups comprising important entity sites.	

The second and subsequent lines contain a continuation field which is a right-justified integer. This number increments by one for each additional line of the record, and is followed by a blank character.

<u>Grouping:</u> There are three record types used to group other records. Listed alphabetically, these are:

RECORD TYPE	DESCRIPTION
ENDMDL	End-of-model record for multiple structures in a single coordinate entry.
MODEL	Specification of model number for multiple structures in a single coordinate entry.
TER	Chain terminator.

The MODEL/ENDMDL records surround groups of ATOM, HETATM, ANISOU, and TER records. TER records indicate the end of a chain.

<u>Other:</u> The remaining record types have a detailed inner structure. Listed alphabetically, these are:

RECORD TYPE	DESCRIPTION
JRNL	Literature citation that defines the coordinate set.
REMARK	General remarks; they can be structured or free form.

The wwPDB will use the following protocol in making changes to the way PDB coordinate entries are represented and archived. The purpose of the policy is to allow ample time for everyone to understand these changes and to assess their impact on existing programs. PDB format modifications are necessary to address the changing needs of PDB users as well as the changing nature of the data that is archived.

- 1. Comments and suggestions will be solicited from the community on specific problems and data representation issues as they arise.
- 2. Proposed format changes will be disseminated through <u>pdb-l@rcsb.org</u> and wwpdb.org.
- 3. A 60-day discussion period will follow the announcement of proposed changes. Comments and suggestions must be received within this time period. Major changes that are not upwardly compatible will be allotted up to twice the standard amount of discussion time.
- 4. The wwPDB will then work in consultation with the wwPDB Advisory Committee and the equivalent partner Scientific Advisory Committees to evaluate and reconcile all suggestions. The final decision will be officially announced via <u>pdb-l@rcsb.org</u> and wwpdb.org.
- 5. Implementation will follow official announcement of the format change. Major changes will not appear in PDB files earlier than 60 days after the announcement, allowing sufficient time to modify files and programs.

### **Order of Records**

All records in a PDB coordinate entry must appear in a defined order. Mandatory record types are present in all entries. When mandatory data are not provided, the record name must appear in the entry with a NULL indicator. Optional items become mandatory when certain conditions exist. Record order and existence are described in the following table:

RECORD TYPE	EXISTENCE	CONDITIONS IF OPTIONAL
HEADER	Mandatory	
OBSLTE	Optional	Mandatory in entries that have been replaced by a newer entry.
TITLE	Mandatory	
SPLIT	Optional	Mandatory when large macromolecular complexes are split into multiple PDB entries.
CAVEAT	Optional	Mandatory when there are outstanding errors such as chirality.
COMPND	Mandatory	
SOURCE	Mandatory	
KEYWDS	Mandatory	
EXPDTA	Mandatory	
NUMMDL	Optional	Mandatory for NMR ensemble entries.
MDLTYP	Optional	Mandatory for NMR minimized average Structures or when the entire polymer chain contains C alpha or P atoms only.
AUTHOR	Mandatory	
REVDAT	Mandatory	
SPRSDE	Optional	Mandatory for a replacement entry.
JRNL	Optional	Mandatory for a publication describes the experiment.
REMARK 0	Optional	Mandatory for a re-refined structure
REMARK 1	Optional	
REMARK 2	Mandatory	
REMARK 3	Mandatory	
REMARK N	Optional	Mandatory under certain conditions.

DBREF	Optional	Mandatory for all polymers.
DBREF1/DBREF2	Optional	Mandatory when certain sequence database accession and/or sequence numbering does not fit preceding DBREF format.
SEQADV	Optional	Mandatory if sequence conflict exists.
SEQRES	Mandatory	Mandatory if ATOM records exist.
MODRES	Optional	Mandatory if modified group exists in the coordinates.
НЕТ	Optional	Mandatory if a non-standard group other than water appears in the coordinates.
HETNAM	Optional	Mandatory if a non-standard group other than water appears in the coordinates.
HETSYN	Optional	
FORMUL	Optional	Mandatory if a non-standard group or water appears in the coordinates.
HELIX	Optional	
SHEET	Optional	
SSBOND	Optional	Mandatory if a disulfide bond is present.
LINK	Optional	Mandatory if non-standard residues appear in a polymer
CISPEP	Optional	
SITE	Optional	
CRYST1	Mandatory	
ORIGX1 ORIGX2 ORIGX3	Mandatory	
SCALE1 SCALE2 SCALE3	Mandatory	
MTRIX1 MTRIX2 MTRIX3	Optional	Mandatory if the complete asymmetric unit must be generated from the given coordinates using non-crystallographic symmetry.
MODEL	Optional	Mandatory if more than one model is present in the entry.
АТОМ	Optional	Mandatory if standard residues exist.
ANISOU	Optional	
TER	Optional	Mandatory if ATOM records exist.
HETATM	Optional	Mandatory if non-standard group exists.

ENDMDL	Optional	Mandatory if MODEL appears.
CONECT	Optional	Mandatory if non-standard group appears and if LINK or SSBOND records exist.
MASTER	Mandatory	
END	Mandatory	

## Sections of an Entry

The following table lists the various sections of a PDB entry (version 3.2) and the records within it:

SECTION	DESCRIPTION	RECORD TYPE
Title	Summary descriptive remarks	HEADER, OBSLTE, TITLE, SPLIT, CAVEAT, COMPND, SOURCE, KEYWDS,EXPDTA, NUMMDL, MDLTYP, AUTHOR, REVDAT, SPRSDE, JRNL
Remark	Various comments about entry annotations in more depth than standard records	REMARKs 0-999
Primary structure	Peptide and/or nucleotide sequence and the relationship between the PDB sequence and that found in the sequence database(s)	DBREF, SEQADV, SEQRES MODRES
Heterogen	Description of non-standard groups	HET, HETNAM, HETSYN, FORMUL
Secondary structure	Description of secondary structure	HELIX, SHEET
Connectivity annotation	Chemical connectivity	SSBOND, LINK, CISPEP
Miscellaneous features	Features within the macromolecule	SITE
Crystallographic	Description of the crystallographic cell	CRYST1
Coordinate transformation	Coordinate transformation operators	ORIGXn, SCALEn, MTRIXn,
Coordinate	Atomic coordinate data	MODEL, ATOM, ANISOU, TER, HETATM, ENDMDL
Connectivity	Chemical connectivity	CONECT

Bookkeeping

Summary information, end-of-file marker

MASTER, END

### **Field Formats and Data Types**

Each record type is presented in a table which contains the division of the records into fields by column number, defined data type, field name or a quoted string which must appear in the field, and field definition. Any column not specified must be left blank.

Each field contains an identified data type that can be validated by a program. These are:

DATA TYPE	DESCRIPTION
AChar	An alphabetic character (A-Z, a-z).
Atom	Atom name.
Character	Any non-control character in the ASCII character set or a space.
Continuation	A two-character field that is either blank (for the first record of a set) or contains a two digit number right-justified and blank-filled which counts continuation records starting with 2. The continuation number must be followed by a blank.
Date	A 9 character string in the form DD-MMM-YY where DD is the day of the month, zero-filled on the left (e.g., 04); MMM is the common English 3-letter abbreviation of the month; and YY is the last two digits of the year. This must represent a valid date.
IDcode	A PDB identification code which consists of 4 characters, the first of which is a digit in the range 0 - 9; the remaining 3 are alpha-numeric, and letters are upper case only. Entries with a 0 as the first character do not contain coordinate data.
Integer	Right-justified blank-filled integer value.
Token	A sequence of non-space characters followed by a colon and a space.
List	A String that is composed of text separated with commas.
LString	A literal string of characters. All spacing is significant and must be preserved.
LString(n)	An LString with exactly n characters.
Real(n,m)	Real (floating point) number in the FORTRAN format Fn.m.
Record name	The name of the record: 6 characters, left-justified and blank-filled.
Residue name	One of the standard amino acid or nucleic acids, as listed below, or the non-standard group designation as defined in

	the HET dictionary. Field is right-justified.
SList	A String that is composed of text separated with semi-colons.
Specification	A String composed of a token and its associated value separated by a colon.
Specification List	A sequence of Specifications, separated by semi-colons.
String	A sequence of characters. These characters may have arbitrary spacing, but should be interpreted as directed below.
String(n)	A String with exactly n characters.
SymOP	An integer field of from 4 to 6 digits, right-justified, of the form nnnMMM where nnn is the symmetry operator number and MMM is the translation vector.

To interpret a String, concatenate the contents of all continued fields together, collapse all sequences of multiple blanks to a single blank, and remove any leading and trailing blanks. This permits very long strings to be properly reconstructed.

## 2. Title Section

This section contains records used to describe the experiment and the biological macromolecules present in the entry: HEADER, OBSLTE, TITLE, SPLIT, CAVEAT, COMPND, SOURCE, KEYWDS, EXPDTA, AUTHOR, REVDAT, SPRSDE, JRNL, and REMARK records.

### HEADER

#### Overview

The HEADER record uniquely identifies a PDB entry through the idCode field. This record also provides a classification for the entry. Finally, it contains the date when the coordinates were deposited to the PDB archive.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"HEADER"	
11 - 50	String(40)	classification	Classifies the molecule(s).
51 - 59	Date	depDate	Deposition date. This is the date the coordinates were received at the PDB.
63 - 66	IDcode	idCode	This identifier is unique within the PDB.

#### Details

\* The classification string is left-justified and exactly matches one of a collection of strings. A class list is available from the current wwPDB Annotation Documentation Appendices (http://www.wwpdb.org/docs.html). In the case of macromolecular complexes, the classification field must present a class for each macromolecule present. Due to the limited length of the classification field, strings must sometimes be abbreviated. In these cases, the full terms are given in KEYWDS.

\* Classification may be based on function, metabolic role, molecule type, cellular location, etc. This record can describe dual functions of a molecules, and when applicable, separated by a comma ",". Entries with multiple molecules in a complex will list the classifications of each macromolecule separated by slash "/".

#### Verification/Validation/Value Authority Control

The verification program checks that the deposition date is a legitimate date and that the ID code is well-formed.

PDB coordinate entry ID codes do not begin with 0. "No coordinates", or NOC files, given as 0xxx

codes, contained no structural information and were bibliographic only. These entries were subsequently removed from PDB archive.

The classification found in HEADER also appears in KEYWDS, unabbreviated and in no strict order.

#### Example

1 2 3 4 5 7 6 8 HEADER PHOTOSYNTHESIS 28-MAR-07 2UXK HEADER TRANSFERASE/TRANSFERASE INHIBITOR 17-SEP-04 1XH6HEADER MEMBRANE PROTEIN, TRANSPORT PROTEIN 20-JUL-06 2HRT

### OBSLTE

#### **Overview**

OBSLTE appears in entries that have been removed from public distribution.

This record acts as a flag in an entry that has been removed ("obsoleted") from the PDB's full release. It indicates which, if any, new entries have replaced the entry that was obsoleted. The format allows for the case of multiple new entries replacing one existing entry.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"OBSLTE"	
9 - 10	Continuation	continuation	Allows concatenation of multiple records
12 - 20	Date	repDate	Date that this entry was replaced.
22 - 25	IDcode	idCode	ID code of this entry.
$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode	rIdCode rIdCode rIdCode rIdCode rIdCode rIdCode rIdCode rIdCode rIdCode	ID code of entry that replaced this one. ID code of entry that replaced this one.

#### Details

\* It is PDB policy that only the principal investigator and/or the primary author who submitted an entry has the authority to obsolete it. All OBSLTE entries are available from the PDB archive (<u>ftp://ftp.wwpdb.org/pub/pdb/data/structures/obsolete</u>).

\* Though the obsolete entry is removed from the public archive, the initial citation that reported the structure is carried over to the superseding entry.

#### Verification/Validation/Value Authority Control

wwPDB staff adds this record at the time an entry is removed from release.

#### **Relationships to Other Record Types**

None.

Example

	1 2	3	4	5	6	7	8
123456789	01234567890	1234567890	1234567890	L23456789012	345678901234	156789012345	567890
OBSLTE	31-JAN-94	1MBP	2MBP				

## TITLE

#### Overview

The TITLE record contains a title for the experiment or analysis that is represented in the entry. It should identify an entry in the same way that a citation title identifies a publication.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"TITLE "	
9 - 10	Continuation	continuation	Allows concatenation of multiple records.
11 - 80	String	title	Title of the experiment.

#### Details

\* The title of the entry is free text and should describe the contents of the entry and any procedures or conditions that distinguish this entry from similar entries. It presents an opportunity for the depositor to emphasize the underlying purpose of this particular experiment.

\* Some items that may be included in TITLE are:

- Experiment type.
- Description of the mutation.
- The fact that only alpha carbon coordinates have been provided in the entry.

#### Verification/Validation/Value Authority Control

This record is free text so no verification of format is required. The title is supplied by the depositor, but staff may exercise editorial judgment in consultation with depositors in assigning the title.

#### **Relationships to Other Record Types**

COMPND, SOURCE, EXPDTA, and REMARKs provide information that may also be found in TITLE. You may think of the title as describing the experiment, and the compound record as describing the molecule(s).

#### **Examples**

1234567812345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890TITLERHIZOPUSPEPSIN COMPLEXED WITH REDUCED PEPTIDE INHIBITORTITLESTRUCTURE OF THE TRANSFORMED MONOCLINIC LYSOZYME BYTITLE2 CONTROLLED DEHYDRATION

TITLENMR STUDY OF OXIDIZED THIOREDOXIN MUTANT (C62A,C69A,C73A)TITLE2 MINIMIZED AVERAGE STRUCTURE

### SPLIT (added)

#### Overview

The SPLIT record is used in instances where a specific entry composes part of a large macromolecular complex. It will identify the PDB entries that are required to reconstitute a complete complex.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"SPLIT "	
9 - 10	Continuation	continuation	Allows concatenation of multiple records.
$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode	idCode idCode idCode idCode idCode idCode idCode idCode idCode idCode idCode idCode	ID code of related entry. ID code of related entry.
72 - 75 77 - 80	IDcode IDcode	idCode idCode	ID code of related entry. ID code of related entry.

#### Details

\* The SPLIT record can be continued on multiple lines, so that all related PDB entries are cataloged.

#### Verification/Validation/Value Authority Control

This record will be generated at the time of processing the component PDB files of the large macromolecular complex when all complex constituents are deposited.

#### **Relationships to Other Record Types**

REMARK 350 will contain an amended statement to reflect the entire complex.

#### Examples

```
        1
        2
        3
        4
        5
        6
        7
        8

        123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890
        SPLIT
        1VOQ
        1VOV
        <
```

### CAVEAT

#### **Overview**

CAVEAT warns of errors and unresolved issues in the entry. Use caution when using an entry containing this record.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"CAVEAT"	
9 - 10	Continuation	continuation	Allows concatenation of multiple records.
12 - 15	IDcode	idCode	PDB ID code of this entry.
20 - 79	String	comment	Free text giving the reason for the CAVEAT.

#### Details

\* The CAVEAT will also be included in cases where the wwPDB is unable to verify the transformation of the coordinates back to the crystallographic cell. In these cases, the molecular structure may still be correct.

#### Verification/Validation/Value Authority Control

CAVEAT will be added to entries known to be incorrect.

## **COMPND** (updated)

#### **Overview**

The COMPND record describes the macromolecular contents of an entry. Some cases where the entry contains a standalone drug or inhibitor, the name of the non-polymeric molecule will appear in this record. Each macromolecule found in the entry is described by a set of token: value pairs, and is referred to as a COMPND record component. Since the concept of a molecule is difficult to specify exactly, staff may exercise editorial judgment in consultation with depositors in assigning these names.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"COMPND"	
8 - 10	Continuation	continuation	Allows concatenation of multiple records.
11 - 80	Specification list	compound	Description of the molecular components.

#### Details

\* The compound record is a Specification list. The specifications, or tokens, that may be used are listed below:

TOKEN	VALUE DEFINITION
MOL_ID	Numbers each component; also used in SOURCE to associate the information.
MOLECULE	Name of the macromolecule.
CHAIN	Comma-separated list of chain identifier(s).
FRAGMENT	Specifies a domain or region of the molecule.
SYNONYM	Comma-separated list of synonyms for the MOLECULE.
EC	The Enzyme Commission number associated with the molecule. If there is more than one EC number, they are presented as a comma-separated list.
ENGINEERED	Indicates that the molecule was produced using recombinant technology or by purely chemical synthesis.
MUTATION	Indicates if there is a mutation.
OTHER_DETAILS	Additional comments.

\* In the case of synthetic molecules, the depositor will provide the description.

\* For chimeric proteins, the protein name is comma-separated and may refer to the presence of a linker (protein\_1, linker, protein\_2).

\* Asterisks in nucleic acid names (in MOLECULE) are for ease of reading.

\* No specific rules apply to the ordering of the tokens, except that the occurrence of MOL\_ID or FRAGMENT indicates that the subsequent tokens are related to that specific molecule or fragment of the molecule.

\* When insertion codes are given as part of the residue name, they must be given within square brackets, i.e., H57[A]N. This might occur when listing residues in FRAGMENT or OTHER\_DETAILS.

\* For multi-chain molecules, e.g., the hemoglobin tetramer, a comma-separated list of CHAIN identifiers is used.

#### Verification/Validation/Value Authority Control

CHAIN must match the chain identifiers(s) of the molecule(s). EC numbers are also checked.

#### **Relationships to Other Record Types**

In the case of mutations, the SEQADV records will present differences from the reference molecule. REMARK records may further describe the contents of the entry. Also see verification above.

#### Examples

```
3
                                 4
                                          5
                                                   6
                                                           7
                                                                    8
                2
       1
COMPND
       MOL ID: 1;
COMPND
       2 MOLECULE: HEMOGLOBIN ALPHA CHAIN;
COMPND 3 CHAIN: A, C;
COMPND 4 SYNONYM: DEOXYHEMOGLOBIN ALPHA CHAIN;
COMPND 5 ENGINEERED: YES;
      6 MUTATION: YES;
COMPND
      7 MOL_ID: 2;
COMPND
COMPND 8 MOLECULE: HEMOGLOBIN BETA CHAIN;
COMPND 9 CHAIN: B, D;
COMPND 10 SYNONYM: DEOXYHEMOGLOBIN BETA CHAIN;
COMPND 11 ENGINEERED: YES;
COMPND 12 MUTATION: YES
COMPND
      MOL ID: 1;
COMPND 2 MOLECULE: COWPEA CHLOROTIC MOTTLE VIRUS;
      3 CHAIN: A, B, C;
COMPND
      4 SYNONYM: CCMV;
COMPND
COMPND
      5 MOL ID: 2;
COMPND
      6 MOLECULE: RNA (5'-(*AP*UP*AP*U)-3');
COMPND
      7 CHAIN: D, F;
COMPND
      8 ENGINEERED: YES;
COMPND
      9 MOL_ID: 3;
```

COMPND 10 MOLECULE: RNA (5'-(\*AP\*U)-3'); COMPND 11 CHAIN: E; COMPND 12 ENGINEERED: YES COMPND MOL\_ID: 1; COMPND 2 MOLECULE: HEVAMINE A; COMPND 3 CHAIN: A; COMPND 4 EC: 3.2.1.14, 3.2.1.17; COMPND 5 OTHER\_DETAILS: PLANT ENDOCHITINASE/LYSOZYME

### SOURCE (updated)

#### **Overview**

The SOURCE record specifies the biological and/or chemical source of each biological molecule in the entry. Some cases where the entry contains a standalone drug or inhibitor, the source information of this molecule will appear in this record. Sources are described by both the common name and the scientific name, e.g., genus and species. Strain and/or cell-line for immortalized cells are given when they help to uniquely identify the biological entity studied.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"SOURCE"	
8 - 10	Continuation	continuation	Allows concatenation of multiple records.
11 - 79	Specification List	srcName	Identifies the source of the macromolecule in a token: value format.
Details			
TOKEN			JE DEFINITION
MOL_ID			pers each molecule. Same as appears in COMPND.
SYNTHETIC		Ind	icates a chemically-synthesized source.
FRAGMENT			omain or fragment of the molecule may be cified.
ORGANISM_SCI	ENTIFIC	Scie	entific name of the organism.
ORGANISM_COM	MON	Com	mon name of the organism.
ORGANISM_TAX	ID	NCB	I Taxonomy ID number of the organism.
STRAIN		Ide	ntifies the strain.

VARIANT	Identifies the variant.
CELL_LINE	The specific line of cells used in the experiment.
ATCC	American Type Culture Collection tissue culture number.
ORGAN	Organized group of tissues that carries on a specialized function.
TISSUE	Organized group of cells with a common function and structure.
CELL	Identifies the particular cell type.
ORGANELLE	Organized structure within a cell.
SECRETION	Identifies the secretion, such as saliva, urine, or venom, from which the molecule was isolated.
CELLULAR_LOCATION	Identifies the location inside/outside the cell.
PLASMID	Identifies the plasmid containing the gene.
GENE	Identifies the gene.
EXPRESSION_SYSTEM	Scientific name of the organism in which the molecule was expressed.
EXPRESSION_SYSTEM_COMMON	Common name of the organism in which the molecule was expressed.
EXPRESSION_SYSTEM_TAXID	NCBI Taxonomy ID of the organism used as the expression system.
EXPRESSION_SYSTEM_STRAIN	Strain of the organism in which the molecule was expressed.
EXPRESSION_SYSTEM_VARIANT	Variant of the organism used as the expression system.
EXPRESSION_SYSTEM_CELL_LINE	The specific line of cells used as the expression system.
EXPRESSION_SYSTEM_ATCC_NUMBER	Identifies the ATCC number of the expression System.
EXPRESSION_SYSTEM_ORGAN	Specific organ which expressed the molecule.
EXPRESSION_SYSTEM_TISSUE	Specific tissue which expressed the molecule.
EXPRESSION_SYSTEM_CELL	Specific cell type which expressed the molecule.
EXPRESSION_SYSTEM_ORGANELLE	Specific organelle which expressed the molecule.
EXPRESSION_SYSTEM_CELLULAR_LOCATION	Identifies the location inside or outside the cell which expressed the molecule.

EXPRESSION_SYSTEM_VECTOR_TYPE	Identifies the type of vector used, i.e., plasmid, virus, or cosmid.
EXPRESSION_SYSTEM_VECTOR	Identifies the vector used.
EXPRESSION_SYSTEM_PLASMID	Plasmid used in the recombinant experiment.
EXPRESSION_SYSTEM_GENE	Name of the gene used in recombinant experiment.
OTHER_DETAILS	Used to present information on the source which is not given elsewhere.

\* The srcName is a list of tokens: value pairs describing each biological component of the entry.

\* As in COMPND, the order is not specified except that MOL\_ID or FRAGMENT indicates subsequent specifications are related to that molecule or fragment of the molecule.

\* Only the relevant tokens need to appear in an entry.

\* Molecules prepared by purely chemical synthetic methods are described by the specification SYNTHETIC followed by "YES" or an optional value, such as NON-BIOLOGICAL SOURCE or BASED ON THE NATURAL SEQUENCE. ENGINEERED must appear in the COMPND record.

\* In the case of a chemically synthesized molecule using a biologically functional sequence (nucleic or amino acid), SOURCE reflects the biological origin of the sequence and COMPND reflects its synthetic nature by inclusion of the token ENGINEERED. The token SYNTHETIC appears in SOURCE.

\* If made from a synthetic gene, ENGINEERED appears in COMPND and the expression system is described in SOURCE (SYNTHETIC does NOT appear in SOURCE).

\* If the molecule was made using recombinant techniques, ENGINEERED appears in COMPND and the system is described in SOURCE.

\* When multiple macromolecules appear in the entry, each MOL\_ID, as given in the COMPND record, must be repeated in the SOURCE record along with the source information for the corresponding molecule.

\* Hybrid molecules prepared by fusion of genes are treated as multi-molecular systems for the purpose of specifying the source. The token FRAGMENT is used to associate the source with its corresponding fragment.

- When necessary to fully describe hybrid molecules, tokens may appear more than once for a given MOL\_ID.
- All relevant token: value pairs that taken together fully describe each fragment are grouped following the appropriate FRAGMENT.

• Descriptors relative to the full system appear before the FRAGMENT (see third example below).

\* ORGANISM\_SCIENTIFIC provides the Latin genus and species. Virus names are listed as the scientific name.

\* Cellular origin is described by giving cellular compartment, organelle, cell, tissue, organ, or body part from which the molecule was isolated.

\* CELLULAR\_LOCATION may be used to indicate where in the organism the compound was found. Examples are: extracellular, periplasmic, cytosol.

\* Entries containing molecules prepared by recombinant techniques are described as follows:

- The expression system is described.
- The organism and cell location given are for the source of the gene used in the cloning experiment.

\* Transgenic organisms, such as mouse producing human proteins, are treated as expression systems.

\* New tokens may be added by the wwPDB.

#### Verification/Validation/Value Authority Control

The biological source is compared to that found in the sequence databases. The Tax ID is identified and the corresponding scientific and common names for the organism is matched to a standard taxonomy database (such as NCBI).

#### **Relationships to Other Record Types**

Each macromolecule listed in COMPND must have a corresponding source.

#### Examples

3 5 6 7 1 2 4 MOL ID: 1; SOURCE 2 ORGANISM SCIENTIFIC: AVIAN SARCOMA VIRUS; SOURCE SOURCE 3 ORGANISM TAXID: 11876 SOURCE 4 STRAIN: SCHMIDT-RUPPIN B; SOURCE 5 EXPRESSION\_SYSTEM: ESCHERICHIA COLI; 6 EXPRESSION SYSTEM TAXID: 562 SOURCE 7 EXPRESSION\_SYSTEM\_PLASMID: PRC23IN SOURCE

For a Chimera protein:

SOURCE SOURCE	<pre>MOL_ID: 1; 2 ORGANISM SCIENTIFIC: MUS MUSCULUS, HOMO SAPIENS;</pre>
SOURCE	3 ORGANISM_COMMON: MOUSE, HUMAN;
SOURCE SOURCE	3 ORGANISM_TAXID: 10090, 9606 5 EXPRESSION SYSTEM: ESCHERICHIA COLI;
SOURCE	6 EXPRESSION_SYSTEM_TAXID: 344601
SOURCE	6 EXPRESSION_SYSTEM_STRAIN: B171;
SOURCE SOURCE	7 EXPRESSION_SYSTEM_VECTOR_TYPE: PLASMID; 8 EXPRESSION SYSTEM PLASMID: P4XH-M13;
DODICE	• EXTRESSION_SISTEM_TEROMID. TAXI-MIS,

### KEYWDS

#### **Overview**

The KEYWDS record contains a set of terms relevant to the entry. Terms in the KEYWDS record provide a simple means of categorizing entries and may be used to generate index files. This record addresses some of the limitations found in the classification field of the HEADER record. It provides the opportunity to add further annotation to the entry in a concise and computer-searchable fashion.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"KEYWDS"	
9 - 10	Continuation	continuation	Allows concatenation of records if necessary.
11 - 79	List	keywds	Comma-separated list of keywords relevant to the entry.

#### Details

\* The KEYWDS record contains a list of terms relevant to the entry, similar to that found in journal articles. A phrase may be used if it presents a single concept (e.g., reaction center). Terms provided in this record may include those that describe the following:

- Functional classification.
- Metabolic role.
- Known biological or chemical activity.
- Structural classification.

\*Other classifying terms may be used. No particular ordering is required. A number of PDB entries contain complexes of macromolecules. In these cases, all terms applicable to each molecule should be provided separated by a comma.

\*Note that the terms in the KEYWDS record duplicate those found in the classification field of the HEADER record. Terms abbreviated in the HEADER record are unabbreviated in KEYWDS.

#### Verification/Validation/Value Authority Control

Terms used in the KEYWDS record are subject to scientific and editorial review. A list of terms, definitions, and synonyms will be maintained by the wwPDB. Every attempt will be made to provide some level of consistency with keywords used in other biological databases.

#### **Relationships to Other Record Types**

HEADER records contain a classification term which must also appear in KEYWDS. Scientific judgment will dictate when terms used in one entry to describe a molecule should be included in other entries with the same or similar molecules.

#### Example

 1
 2
 3
 4
 5
 6
 7
 8

 123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890
 KEYWDS
 LYASE, TRICARBOXYLIC ACID CYCLE, MITOCHONDRION, OXIDATIVE

 KEYWDS
 2
 METABOLISM
 KEYWDS
 2
 METABOLISM
 KEYWDS
 1
 KEYWDS
 1

## **EXPDTA** (updated)

#### **Overview**

The EXPDTA record presents information about the experiment.

The EXPDTA record identifies the experimental technique used. This may refer to the type of radiation and sample, or include the spectroscopic or modeling technique. Permitted values include:

X-RAY DIFFRACTION FIBER DIFFRACTION NEUTRON DIFFRACTION ELECTRON CRYSTALLOGRAPHY ELECTRON MICROSCOPY SOLID-STATE NMR SOLUTION NMR SOLUTION SCATTERING

\*Note:Since October 15, 2006, theoretical models are no longer accepted for deposition. Any theoretical models deposited prior to this date are archived at <a href="http://ftp.wwpdb.org/pub/pdb/data/structures/models">ftp://ftp.wwpdb.org/pub/pdb/data/structures/models</a>. Please see the documentation from previous versions for the related file format description.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"EXPDTA"	
9 - 10	Continuation	continuation	Allows concatenation of multiple records.
11 - 79	SList	technique	The experimental technique(s) with optional comment describing the sample or experiment.

### Details

\* EXPDTA is mandatory and appears in all entries. The technique must match one of the permitted values. See above.

\* If more than one technique was used for the structure determination and is being represented in the entry, EXPDTA presents the techniques as a semi-colon separated list.

### Verification/Validation/Value Authority Control

The verification program checks that the EXPDTA record appears in the entry and that the technique matches one of the allowed values. It also checks that the relevant standard REMARK is added, as in

the cases of NMR or electron microscopy studies, that the appropriate CRYST1 and SCALE values are used.

### **Relationships to Other Record Types**

If the experiment is an NMR or electron microscopy study, this may be stated in the TITLE, and the appropriate EXPDTA and REMARK records should appear. Specific details of the data collection and experiment appear in the REMARKs.

In the case of a polycrystalline fiber diffraction study, CRYST1 and SCALE contain the normal unit cell data.

### Examples

5 7 2 3 4 6 8 1 EXPDTA X-RAY DIFFRACTION EXPDTA NEUTRON DIFFRACTION; X-RAY DIFFRACTION EXPDTA SOLUTION NMR EXPDTA ELECTRON MICROSCOPY

## NUMMDL (added)

### Overview

The NUMMDL record indicates total number of models in a PDB entry.

Record Forma	t		
COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"NUMMDL"	
11 - 14	Integer	modelNumber	Number of models.

### Details

\* The modelNumber field lists total number of models in a PDB entry and is left justified.

\* If more than one model appears in the entry, the number of models included must be stated.

\* NUMMDL is mandatory if a PDB entry contains more than one models.

### Verification/Validation/Value Authority Control

The verification program checks that the modelNumber field is correctly formatted.

### Example

## **MDLTYP** (added)

### Overview

The MDLTYP record contains additional annotation pertinent to the coordinates presented in the entry.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"MDLTYP"	
9 - 10	Continuation	continuation	Allows concatenation of multiple records.
11 - 80	SList	comment	Free Text providing additional structural annotation.

### Details

\* The MDLTYP record will be used by the wwPDB to highlight certain features of the deposited coordinates as described below.

\* For entries that are determined by NMR methods and the coordinates deposited are either a minimized average or regularized mean structure, this record will contain the tag "MINIMIZED AVERAGE" to highlight the nature of the deposited coordinates in the entry.

\* Where the entry contains entire polymer chains that have only either C-alpha (for proteins) or P atoms (for nucleotides), the MDLTYP record will be used to describe the contents of such chains along with the chain identifier. For these polymeric chains, REMARK 470 (Missing Atoms) will be omitted.

\* If multiple features need to be described in this record, they will be separated by a ";" delineator.

\* Where an entry has multiple features requiring description in this record including MINIMIZED AVERAGE, the MINIMIZED AVERAGE value will precede all other annotation.

\* New descriptors may be added by the wwPDB.

### Verification/Validation/Value Authority Control

The chain\_identifiers described in this record must be present in the COMPND, SEQRES and the coordinate section of the entry.

### Example

1	2	3	4	5	6	7	8

123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 MDLTYP MINIMIZED AVERAGE

MDLTYP CA ATOMS ONLY, CHAIN A, B, C, D, E, F, G, H, I, J, K ; P ATOMS ONLY, MDLTYP 2 CHAIN X, Y, Z

MDLTYP MINIMIZED AVERAGE; CA ATOMS ONLY, CHAIN A, B

## AUTHOR

### **Overview**

The AUTHOR record contains the names of the people responsible for the contents of the entry.

#### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"AUTHOR"	
9 - 10	Continuation	continuation	Allows concatenation of multiple records.
11 - 79	List	authorList	List of the author names, separated by commas.

### Details

\* The authorList field lists author names separated by commas with no subsequent spaces.

\* Representation of personal names:

- First and middle names are indicated by initials, each followed by a period, and precede the surname.
- Only the surname (family or last name) of the author is given in full.
- Hyphens can be used if they are part of the author's name.
- Apostrophes are allowed in surnames.
- Umlauts and other character modifiers are not given.

\* Structure of personal names:

- There is no space after any initial and its following period.
- Blank spaces are used in a name only if properly part of the surname (e.g., J.VAN DORN), or between surname and Jr., II, or III

Abbreviations that are part of a surname, such as Jr., St. or Ste., are followed by a period and a space before the next part of the surname.

\* Representation of corporate, organization or university names:

• Group names used for one or all of the authors should be spelled out in full.

• The name of the larger group comes before the name of a subdivision, e.g., University of Somewhere, Department of Chemistry.

\* Structure of list:

- Line breaks between multiple lines in the authorList occur only after a comma.
- Personal names are not split across two lines.

\* Special cases:

• Names are given in English if there is an accepted English version; otherwise in the native language, transliterated if necessary.

#### Verification/Validation/Value Authority Control

The verification program checks that the authorList field is correctly formatted. It does not perform any spelling checks or name verification.

#### **Relationships to Other Record Types**

The format of the names in the AUTHOR record is the same as in JRNL and REMARK 1 references.

#### Example

1 2 3 4 5 6 7 8 12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 AUTHOR M.B.BERRY, B.MEADOR, T.BILDERBACK, P.LIANG, M.GLASER, AUTHOR 2 G.N.PHILLIPS JR., T.L.ST. STEVENS

## **REVDAT** (updated)

### **Overview**

REVDAT records contain a history of the modifications made to an entry since its release.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"REVDAT"	
8 - 10	Integer	modNum	Modification number.
11 - 12	Continuation	continuation	Allows concatenation of multiple records.
14 - 22	Date	modDate	Date of modification (or release for new entries) in DD-MMM-YY format. This is not repeated on continued lines.
24 - 27	IDCode	modId	ID code of this entry. This is not repeated on continuation lines.
32	Integer	modType	An integer identifying the type of modification. For all revisions, the modification type is listed as 1
40 - 45 47 - 52 54 - 59 61 - 66	LString(6) LString(6) LString(6) LString(6)	record record record record	Modification detail. Modification detail. Modification detail. Modification detail.

### Details

\* Each time revisions are made to the entry, a modification number is assigned in increasing (by 1) numerical order. REVDAT records appear in descending order (most recent modification appears first). New entries have a REVDAT record with modNum equal to 1 and modType equal to 0. Allowed modTypes are:

- 0 Initial released entry.
- 1 Other modification.

\* Each revision may have more than one REVDAT record, and each revision has a separate continuation field.

\* Modification details are typically PDB record names such as JRNL, SOURCE, TITLE, or COMPND. A special modification detail VERSN indicates that the file has undergone a change in version. The current version will be specified in REMARK 4.

### Verification/Validation/Value Authority Control

The modType must be one of the defined types, and the given record type must be valid. If modType is 0, the modId must match the entry's ID code in the HEADER record.

### **Relationships to Other Record Types**

In the case of a version revision, the current will be specified in REMARK 4.

### Template

7 2 5 1 3 4 6 8 15-OCT-99 1ABC REMARK REVDAT 2 1 REVDAT 1 09-JAN-89 1ABC 0 2 3 4 5 6 7 8 1 REVDAT 11-MAR-08 2ABC JRNL 2 1 VERSN REVDAT 09-DEC-03 2ABC 0 1

## SPRSDE

### Overview

The SPRSDE records contain a list of the ID codes of entries that were made obsolete by the given coordinate entry and removed from the PDB release set. One entry may replace many.

It is wwPDB policy that only the principal investigator of a structure has the authority to obsolete it.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"SPRSDE"	
9 - 10	Continuation	continuation	Allows for multiple ID codes.
12 - 20	Date	sprsdeDate	Date this entry superseded the listed entries. This field is not copied on continuations.
22 – 25	IDcode	idCode	ID code of this entry. This field is not copied on continuations.
$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode IDcode	sIdCode sIdCode sIdCode sIdCode sIdCode sIdCode sIdCode sIdCode sIdCode	ID code of a superseded entry. ID code of a superseded entry.

### Details

\* The ID code list is terminated by the first blank sldCode field.

### Verification/Validation/Value Authority Control

wwPDB checks that the superseded entries have actually been removed from release.

### **Relationships to Other Record Types**

The sprsdeDate is usually the date the entry is released, and therefore matches the date in the REVDAT 1 record. The ID code found in the idCode field must be the same as one found in the idCode field of the HEADER record.

### Example

	1 2	3		4	5	6	7	8
123456789	01234567890	12345678903	12345	678901234	56789012345	6789012345	56789012345	67890
SPRSDE	17-JUL-84	4HHB	1HHB					
SPRSDE	27-FEB-95	1GDJ	1LH4	2LH4				

## JRNL (updated)

#### Overview

The JRNL record contains the primary literature citation that describes the experiment which resulted in the deposited coordinate set. There is at most one JRNL reference per entry. If there is no primary reference, then there is no JRNL reference. Other references are given in REMARK 1.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"JRNL "	
13 - 79	LString	text	See Details below.

### Details

\* The following tables are used to describe the sub-record types of the JRNL record.

\* The AUTH sub-record is mandatory in JRNL. This is followed by TITL, EDIT, REF, PUBL, REFN, PMID and DOI sub- record types. REF and REFN are also mandatory in JRNL. EDIT and PUBL may appear only if the reference is to a non-journal.

### <u>1. AUTH</u>

\* AUTH contains the list of authors associated with the cited article or contribution to a larger work (i.e., AUTH is not used for the editor of a book).

\* The author list is formatted similarly to the AUTHOR record. It is a comma-separated list of names. Spaces at the end of a sub-record are not significant; all other spaces are significant. See the AUTHOR record for full details.

\* The authorList field of continuation sub-records in JRNL differs from that in AUTHOR by leaving no leading blank in column 20 of any continuation lines.

\* One author's name, consisting of the initials and family name, cannot be split across two lines. If there are continuation sub-records, then all but the last sub-record must end in a comma.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(4)	"AUTH"	Appears on all continuation records.
17 - 18	Continuation	continuation	Allows a long list of authors.

20 - 79 List authorList List of the authors.

### 2. TITL

\* TITL specifies the title of the reference. This is used for the title of a journal article, chapter, or part of a book. The TITL line is omitted if the author(s) listed in authorList wrote the entire book (or other work) listed in REF and no section of the book is being cited.

\* If an article is in a language other than English and is printed with an alternate title in English, the English language title is given, followed by a space and then the name of the language (in its English form, in square brackets) in which the article is written.

\* If the title of an article is in a non-Roman alphabet the title is transliterated.

\* The actual title cited is reconstructed in a manner identical to other continued records, i.e., trailing blanks are discarded and the continuation line is concatenated with a space inserted.

\* A line cannot end with a hyphen. A compound term (two elements connected by a hyphen) or chemical names which include a hyphen must appear on a single line, unless they are too long to fit on one line, in which case the split is made at a normally-occurring hyphen. An individual word cannot be hyphenated at the end of a line and put on two lines. An exception is when there is a repeating compound term where the second element is omitted, e.g., "DOUBLE- AND TRIPLE-RESONANCE". In such a case the non-completed word "DOUBLE-" could end a line and not alter reconstruction of the title.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(4)	"TITL"	Appears on all continuation records.
17 - 18	Continuation	continuation	Permits long titles.
20 - 79	LString	title	Title of the article.

### <u>3. EDIT</u>

\* EDIT appears if editors are associated with a non-journal reference. The editor list is formatted and concatenated in the same way that author lists are.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6 10	Record name LString(1)	"REMARK" "1"	
13 - 16	LString(4)	"TITL"	Appears on all continuation records.
17 - 18	Continuation	continuation	Permits long titles.

20 - 79 LString title Title of the article.

### <u>4. REF</u>

\* REF is a group of fields that contain either the publication status or the name of the publication (and any supplement and/or report information), volume, page, and year. There are two forms of this sub-record group, depending upon the citation's publication status.

4a. If the reference has not been published yet, the sub-record type group has the form:

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"JRNL "	
13 - 16	LString(3)	"REF"	
20 - 34	LString(15)	"TO BE PUBLISHE	D"

\* Publication name (first item in pubName field):

If the publication is a serial (i.e., a journal, an annual, or other non-book or non-monographic item issued in parts and intended to be continued indefinitely), use the abbreviated name of the publication as listed in PubMed with periods.

If the publication is a book, monograph, or other non-serial item, use its full name according to the Anglo-American Cataloguing Rules, 2nd Revised Edition; (AACR2R). (Non-serial items include theses, videos, computer programs, and anything that is complete in one or a finite number of parts.) If there is a sub-title, verifiable in an online catalog, it will be included using the same punctuation as in the source of verification. Preference will be given to verification using cataloging of the Library of Congress, the National Library of Medicine, and the British Library, in that order.

If a book is part of a monographic series: the full name of the book (according to the AACR2R) is listed first, followed by the name of the series in which it was published. The series information is given within parentheses and the series name is preceded by "IN:" and a space. The series name should be listed in full unless the series has an accepted ISO abbreviation. If applicable, the series name should be followed, after a comma and a space, by a volume (V.) and/or number (NO.) and/or part (PT.) indicator and its number and/or letter in the series.

\* Supplement (follows publication name in pubName field):

If a reference is in a supplement to the volume listed, or if information about a "part" is needed to distinguish multiple parts with the same page numbering, such information should be put in the REF sub-record.

A supplement indication should follow the name of the publication and should be preceded by a comma and a space. Supplement should be abbreviated as "SUPPL." If there is a supplement number or letter, it should follow "SUPPL." without an intervening space. A part indication should also

follow the name of the publication and be preceded by a comma and a space. A part should be abbreviated as "PT.", and the number or letter should follow without an intervening space.

If there is both a supplement and a part, their order should reflect the order printed on the work itself.

\* Report (follows publication name and any supplement or part information in pubName field):

If a book has a report designation, the report information should follow the title and precede series information. The name and number of the report is given in parentheses, and the name is preceded by "REPORT:" and a space.

\* Reconstruction of publication name:

The name of the publication is reconstructed by removing any trailing blanks in the pubName field, and concatenating all of the pubName fields from the continuation lines with an intervening space. There are two conditions where no intervening space is added between lines: when the pubName field on a line ends with a hyphen or a period, or when the line ends with a hyphen (-). When the line ends with a period (.), add a space if this is the only period in the entire pubName field; do not add a space if there are two or more periods throughout the pubName field, excluding any periods after the designations "SUPPL", "V", "NO", or "PT".

\* Volume, page, and year (volume, first page, year fields respectively):

The REF sub-record type group also contains information about volume, page, and year when applicable.

In the case of a monograph with multiple volumes which is also in a numbered series, the number in the volume field represents the number of the book, not the series. (The volume number of the series is in parentheses with the name of the series, as described above under publication name.)

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"JRNL "	
13 - 16	LString(3)	"REF "	
17 - 18	Continuation	continuation	Allows long publication names.
20 - 47	LString	pubName	Name of the publication including section or series designation. This is the only field of this sub-record which may be continued on successive sub-records.
50 - 51	LString(2)	"V."	Appears in the first sub-record only, and only if column 55 is non-blank.
52 - 55	String	volume	Right-justified blank-filled volume information; appears in the first sub-record only.
57 - 61	String	page	First page of the article; appears in

the first sub-record only.

63 - 66 Integer year Year of publication; first sub-record only.

#### <u>5. PUBL</u>

\* PUBL contains the name of the publisher and place of publication if the reference is to a book or other non-journal publication. If the non-journal has not yet been published or released, this sub-record is absent.

\* The place of publication is listed first, followed by a space, a colon, another space, and then the name of the publisher/issuer. This arrangement is based on the ISBD(M) International Standard Bibliographic Description for Monographic Publications (Rev.Ed., 1987) and the AACR2R, and is used in public online catalogs in libraries. Details on the contents of PUBL are given below.

\* Place of publication:

Give the place of publication. If the name of the country, state, province, etc. is considered necessary to distinguish the place of publication from others of the same name, or for identification, then follow the city with a comma, a space, and the name of the larger geographic area.

If there is more than one place of publication, only the first listed will be used. If an online catalog record is used to verify the item, the first place listed there will be used, omitting any brackets. Preference will be given to the cataloging done by the Library of Congress, the National Library of Medicine, and the British Library, in that order.

\* Publisher's name (or name of other issuing entity):

Give the name of the publisher in the shortest form in which it can be understood and identified internationally, according to AACR2R rule 1.4D.

If there is more than one publisher listed in the publication, only the first will be used in the PDB file. If an online catalog record is used to verify the item, the first place listed there will be used for the name of the publisher. Preference will be given to the cataloging of the Library of Congress, the National Library of Medicine, and the British Library, in that order.

\* Ph.D. and other theses:

Theses are presented in the PUBL record if the degree has been granted and the thesis made available for public consultation by the degree-granting institution. The name of the degree-granting institution (the issuing agency) is followed by a space and "(THESIS)".

\* Reconstruction of place and publisher:

The PUBL sub-record type can be reconstructed by removing all trailing blanks in the pub field and concatenating all of the pub fields from the continuation lines with an intervening space.

### Continued lines do not begin with a space.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"JRNL "	
13 - 16	LString(4)	"PUBL"	
17 - 18	Continuation	continuation	Allows long publisher and place names.
20 - 70	LString	pub	City of publication and name of the publisher/institution.

### 6. REFN (changed)

\* REFN is a group of fields that contain encoded references to the citation. No continuation lines are possible. Each piece of coded information has a designated field.

\* There are two forms of this sub-record type group, depending upon the publication status.

6a. This form of the REFN sub-record type group is used if the citation has not been published.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"JRNL "	
13 - 16	LString(4)	"REFN"	

6b. This form of the REFN sub-record type group is used if the citation has been published.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"JRNL "	
13 - 16	LString(4)	"REFN"	
36 - 39	LString(4)	"ISSN" or "ESSN"	International Standard Serial Number or Electronic Standard Serial Number.
41 - 65	LString	issn	ISSN number (final digit may be a letter and may contain one or more dashes).

### 7. PMID (added)

\* PMID lists the PubMed unique accession number of the publication related to the entry.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	 "JRNL "	

13 - 16	LString(4)	"PMID"	
20 - 79	Integer	continuation	unique PubMed identifier number assigned to the publication describing the experiment. Allows for a long PubMed ID number.

#### 8. DOI (added)

\* DOI is the Digital Object Identifier for the related electronic publication ("e-pub"), if applicable.

\* Every DOI consists of a publisher prefix, a fore-slash ("/"), and then a suffix which can be any length and may include a combination of numbers and alphabets. For example: 10.1073/PNAS.0712393105

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"JRNL "	
13 - 16	LString(4)	"DOI "	
20 - 79	LString	continuation	Unique DOI assigned to the publication describing the experiment. Allows for a long DOI string.

### Verification/Validation/Value Authority Control

wwPDB verifies that this record is correctly formatted.

Citations appearing in JRNL may not also appear in REMARK 1.

### **Relationships to Other Record Types**

The publication cited as the JRNL record may not be repeated in REMARK 1.

### Example

3 5 7 1 2 4 6 AUTH G.FERMI, M.F.PERUTZ, B.SHAANAN, R.FOURME JRNL TITLTHE CRYSTAL STRUCTURE OF HUMAN DEOXYHAEMOGLOBIN AT JRNL JRNL TITL 2 1.74 A RESOLUTION JRNLREF J.MOL.BIOL. V. 175 159 1984 JRNLREFN ISSN 0022-2836 JRNL PMID 6726807 10.1016/0022-2836(84)90472-8 JRNL DOI

### **Known Problems**

\* Interchange of bibliographic information and linking with other databases is hampered by the lack of labels or specific locations for certain types of information or by more than one type of information

being in a particular location. This is most likely to occur with books, series, and reports. Some of the points below provide details about the variations and/or blending of information.

\* Titles of the publications that require more than 28 characters on the REF line must be continued on subsequent lines. There is some awkwardness due to volume, page, and year appearing on the first REF line, thereby splitting up the title.

\* Information about a supplement and its number/letter is presented in the publication's title field (on the REF lines in columns 20 - 47).

\* When series information for a book is presented, it is added to the REF line. The number of REF lines can become large in some cases because of the 28-column limit for title information in REF.

\* Books that are issued in more than one series are not accommodated.

\* Pagination is limited to the beginning page.

## REMARK

### **Overview**

REMARK records present experimental details, annotations, comments, and information not included in other records. In a number of cases, REMARKs are used to expand the contents of other record types. A new level of structure is being used for some REMARK records. This is expected to facilitate searching and will assist in the conversion to a relational database.

The very first line of every set of REMARK records is used as a spacer to aid in reading.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"REMARK"	
8 - 10	Integer	remarkNum	Remark number. It is not an error for remark n to exist in an entry when remark n-1 does not.
12 - 79	LString	empty	Left as white space in first line of each new remark.

### **REMARKs 0-5**

REMARK 0, 1, 2, 3, 4 and 5 detailed below, are specific for re-refinement, references, resolution, final refinement, PDB File Format version and obsolete statement, respectively.

## **REMARK 0** (added), Re-refinement notice

REMARK 0 identifies entries in which a re-refinement has been performed using the data from an existing entry.

### Template

5 2 3 4 6 7 8 1 REMARK 0 0 THIS ENTRY YYYY REFLECTS AN ALTERNATIVE MODELING OF THE REMARK 0 ORIGINAL STRUCTURAL DATA (RXXXXSF or XXXX.MR) DETERMINED BY REMARK 0 AUTHORS OF THE PDB ENTRY XXXX: REMARK REMARK 0 AUTHOR INITIALS, AUTHOR LAST NAME

Note: In entries where REMARK 0 is included as described above, remarks REMARK 1 and REMARK 900 will also reflect the reuse of existing experimental data.

REMARK 200 REMARK: AUTHOR USED THE SF(MR) DATA FROM ENTRY XXXX.

NOTE: the rest of REMARKs 200 and 280 are blank, since the re-refinement author did not collect original data.

## **REMARK 1 (updated), Related publications**

REMARK 1 lists important publications related to the structure presented in the entry. These citations are chosen by the depositor. They are listed in reverse-chronological order. Citations are not repeated from the JRNL records. After the first blank record and the REFERENCE sub-record, the sub-record types for REMARK 1 are the same as in the JRNL sub-record types. For details, see the JRNL section.

### **Record Format and Details**

As with all other remarks, the first line is empty and is used as a spacer.

The following tables are used to describe the sub-record types of REMARK 1.

### 1. REFERENCE

Each reference is preceded by a line indicating the reference number in the entry.

COLUMNS	DATA TYPE	FIELD	DEFINITION	
1 – 6	Record name	"REMARK"		
10	LString(1)	"1"		
12 - 20	LString(9)	"REFERENCE"		
22 - 70	Integer	refNum	Reference number. Starts with 1 and increments by 1.	

### <u>2. AUTH</u>

AUTH contains the list of authors of the reference.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(4)	"AUTH"	Appears on all continuation records.
17 - 18	Continuation	continuation	Allows a long list of authors.
20 - 79	List	authorList	List of the authors.

See JRNL AUTH for details.

### <u>3. TITL</u>

TITL specifies the title of the reference.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(4)	"TITL"	Appears on all continuation records.
17 - 18	Continuation	continuation	Permits long titles.
20 - 79	LString	title	Title of the article.

See JRNL TITL for details.

### <u>4. EDIT</u>

EDIT appears if editors are associated with a non-journal reference.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(4)	"TITL"	Appears on all continuation records.
17 - 18	Continuation	continuation	Permits long list of editors.
20 - 79	LString	editorList	List of the editors.

See JRNL EDIT for details.

### <u>5. REF</u>

REF is a group of fields which contains the name of the publication.

5a. If it has not been published yet, the REF sub-record type has the form:

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(3)	"REF"	
20 - 34	LString(15)	"TO BE PUBLISHE	2D "

At the present time, there is no formal mechanism in place for monitoring the subsequent publication

of referenced papers. wwPDB relies upon the depositor to provide reference update information since preliminary information can change by the time of actual publication.

5b. If the reference has been published, then the REF sub-record type group contains information about the name of the publication, supplement, report, volume, page, and year, in the appropriate fields.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(3)	"REF"	
17 - 18	Continuation	continuation	Permits long publication names.
20 - 47	LString	pubName	Name of the publication including section or series designation. This is the only field of this record which may be continued on successive records.
50 - 51	LString(2)	"V."	Appears in the first record only, and only if column 55 is filled in.
52 - 55	String	volume	Right-justified blank-filled volume information; appears in the first sub-record only.
57 - 61	String	page	First page of the article; appears in the first sub-record only.
63 - 66	Integer	year	First record year of publication.

See JRNL REF for details.

#### <u>6. PUBL</u>

PUBL contains the name of the publisher and place of publication if the reference is to a book or other non-journal publication. If the reference has not yet been published or released, this sub-record is absent.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(4)	"PUBL"	
17 - 18	Continuation	continuation	Permits long publisher and city information.
20 - 70	LString	pub	Name of the publisher and city of

#### publication.

See JRNL PUBL for details.

#### 7. REFN (changed)

REFN is a group of fields which contains encoded references to the citation.

7a. If the citation has not been published, this form of the REFN sub-record type group is used.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(4)	"REFN"	

7b. If the citation has been published, this form of the REFN sub-record type group is used.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"1"	
13 - 16	LString(4)	"REFN"	
36 - 39	LString(4)	"ISSN" or "ESSN"	International Standard Serial Number or Electronic Standard Serial Number.
41 - 65	LString	issn	ISSN number.

See JRNL REFN for details.

#### 8. PMID (added)

\* PMID lists the PubMed unique accession number of the publication related to the entry.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"JRNL "	
13 - 16	LString(4)	"PMID"	
20 - 79	Integer	continuation	unique PubMed identifier number assigned to the publication describing the experiment. Allows for a long pubmed id number.

### 9. DOI (added)

\* DOI is the Digital Object Identifier for the related electronic publication ("e-pub"), if applicable.

\* Every DOI consists of a publisher prefix, a fore-slash ("/"), and then a suffix which can be any length and may include a combination of numbers and alphabets. For example: 10.1073/PNAS.0712393105

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"JRNL "	
13 - 16	LString(4)	"DOI "	
20 - 79	LString	continuation	Unique DOI assigned to the Publication describing the experiment. Allows for a long DOI string.

### Verification/Validation/Value Authority Control

wwPDB verifies that this record is correctly formatted.

### **Relationships to Other Record Types**

Citations appearing in REMARK 1 may not appear in JRNL.

### Examples

	1	2 3	4	5	6	7	8
12345678	901234567	89012345678901234	56789012345	6789012345	5789012345	678901234	567890
REMARK	1						
REMARK	1 REFERE	NCE 1					
REMARK	1 AUTH	J.N.BREG,J.H.J.	VAN OPHEUSD	EN,M.J.M.BU	JRGERING,		
REMARK	1 AUTH 2	2 R.BOELENS, R.KAP	TEIN				
REMARK	1 TITL	STRUCTURE OF AR	C REPRESSOR	IN SOLUTIO	ON: EVIDEN	ICE	
REMARK	1 TITL 2	2 FOR A FAMILY OF	B-SHEET DN	A-BINDING 1	PROTEIN		
REMARK	1 REF	NATURE		V. 346	586 199	0	
REMARK	1 REFN		ISSN 0028-	0836			
REMARK	1 PMID	2377232					
REMARK	1 DOI	10.1038/346586a	.0				
REMARK	1 REFERE	NCE 2					
REMARK	1 AUTH	J.N.BREG,R.BOEL	ENS,A.V.E.G	EORGE, R.KAI	PTEIN		
REMARK	1 TITL	SEQUENCE-SPECIF	IC 1H NMR A	SSIGNMENT A	AND SECONE	DARY	
REMARK	1 TITL 2	2 STRUCTURE OF TH	E ARC REPRE	SSOR OF BAG	CTERIOPHAG	Έ	
REMARK	1 TITL 3	3 P22 AS DETERMIN	ED BY 2D 1H	NMR SPECTI	ROSCOPY		
REMARK	1 REF	BIOCHEMISTRY		V. 28	9826 198	39	
REMARK	1 REFN		ISSN 0006-	2960			
REMARK	1 PMID	2611268					
REMARK	1						
REMARK	1 REFERE						
REMARK	1 AUTH	J.MAREK, J.VEVOD	•	•	•		
REMARK		2 L.A.SVENSSON,J.		,			
REMARK	1 TITL	CRYSTAL STRUCTU	RE OF THE H	ALOALKANE I	DEHALOGENA	<b>SE</b>	
REMARK		2 FROM SPHINGOMON	AS PAUCIMOB				
REMARK	1 REF	BIOCHEMISTRY			14082 200	0	
REMARK	1 REFN		ISSN 0006-	2960			

REMARK 1 PMID 11087355 REMARK 1 DOI 10.1021/bi001539c

### **Known Problems**

See JRNL for a listing of problems associated with references.

## **REMARK 2 (updated), Resolution**

REMARK 2 states the highest resolution, in Angstroms, that was used in building the model. As with all the remarks, the first REMARK 2 record is empty and is used as a spacer.

### **Record Format and Details**

\* The second REMARK 2 record has one of two formats. The first is used for diffraction studies, the second for other types of experiments in which resolution is not relevant, e.g., NMR.

\* For diffraction experiments,

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"REMARK"	
10	LString(1)	"2"	
12 - 22	LString(11)	"RESOLUTION."	
24 - 30	Real(7.2)	resolution	Resolution.
32 - 41	LString(10)	"ANGSTROMS."	

\* REMARK 2 when not a diffraction experiment:

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"REMARK"	
10	LString(1)	"2"	
12 - 38	LString(28)	"RESOLUTION. N	IOT APPLICABLE."
41 - 70	String	comment	Comment in free text.

\* Additional explanatory text may be included starting with the third line of the REMARK 2 record. For example, depositors may wish to qualify the resolution value provided due to unusual experimental conditions.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"REMARK"	
10	LString(1)	"2"	
12 - 22	LString(11)	"RESOLUTION."	
24 - 70	String	comment	Comment in free text.

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## **REMARK 3** (updated), Final refinement information

### Overview

REMARK 3 presents information on refinement program(s) used and related statistics. For nondiffraction studies, REMARK 3 is used to describe any refinement done, but its format is mostly free text.

### Details

\* The value "NULL" is given when there is no data available for a particular token.

\* If more than one refinement package was used, they may be named in "OTHER REFINEMENT REMARKS". However, REMARK 3 statistics are given for the final refinement run.

\* Entries which were refined using Refmac program, B VALUE TYPE may appear in REMARK 3 for "LIKELY RESIDUAL" or "UNVERIFIED" for entries which likely contain residual B value in ATOM record or cannot be verified.

The format of this remark changes with the evolution of refinement software. Selected representative templates or examples are provided here.

# Refinement using X-PLOR

## Template/example

3

REMARK

REMARK3PROGRAM: X-PLOR 3.851REMARK3AUTHORS: BRUNGERREMARK3DATA USED IN REFINEMENT.REMARK3RESOLUTION RANGE HIGH (ANGSTROMS) : 2.47REMARK3RESOLUTION RANGE LOW (ANGSTROMS) : 34.50REMARK3DATA CUTOFF (GIGMA(F)) : 0.000REMARK3DATA CUTOFF HIGH (ABS(F)) : NULLREMARK3DATA CUTOFF HIGH (ABS(F)) : NULLREMARK3DATA CUTOFF LOW (ABS(F)) : NULLREMARK3DATA CUTOFF ELECTIONSREMARK3COMPLETENESS (WORKING+TEST) (%) : NULLREMARK3FREE R VALUE TEST SET SELECTION : RANDOMREMARK3FREE R VALUE TEST SET SIZE (%) : NULLREMARK3FREE R VALUE TEST SET COUNT : 2819REMARK3FREE R VALUE TEST SET COUNT : 2819REMARK3FIT IN THE HIGHEST RESOLUTION BIN.REMARK3FIT IN THE HIGHEST RESOLUTION BIN.REMARK3FIT IN THE HIGHEST RESOLUTION BIN.REMARK3BIN RESOLUTION RANGE LOW (A) : NULLREMARK3BIN RESOLUTION RANGE LOW (A) : NULLREMARKBIN FREE R VALUE TEST SET SIZE (%) : NULLREMARKBIN FREE R VALUE TEST SET SIZE (%) : NULLREMARKBIN FREE R VALUE TEST SET SIZE (%) : NULLREMARKB	REMARK	3	REFINEMENT.																																																																																																																				
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<tr><td>REMARK3BIN RESOLUTION RANGE LOW(A) : NULLREMARK3BIN COMPLETENESS (WORKING+TEST) (%) : NULLREMARK3REFLECTIONS IN BIN(WORKING SET) : NULLREMARK3BIN R VALUE(WORKING SET) : NULLREMARK3BIN FREE R VALUE: NULLREMARK3BIN FREE R VALUE TEST SET SIZE (%) : NULLREMARK3BIN FREE R VALUE TEST SET COUNT: NULLREMARK3BUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.REMARK3PROTEIN ATOMS: 0REMARK3NUCLEIC ACID ATOMS: 0REMARK3NUCLEIC ACID ATOMS: 132REMARK3SOLVENT ATOMS: 132REMARK3SOLVENT ATOMS: 132REMARK3B VALUES.(OVERALL, A**2) : 31.94REMARK3B VALUE(OVERALL, A**2) : 27.18REMARK3B11 (A**2) : NULLREMARK3B11 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL<td></td><td></td><td>TOTAL NUMBER OF BINS USED : NULL</td></td></tr> <tr><td>REMARK3BIN COMPLETENESS (WORKING+TEST) (%) : NULLREMARK3REFLECTIONS IN BIN (WORKING SET) : NULLREMARK3BIN R VALUE (WORKING SET) : NULLREMARK3BIN REE R VALUE (WORKING SET) : NULLREMARK3BIN FREE R VALUE TEST SET SIZE (%) : NULLREMARK3BIN FREE R VALUE TEST SET COUNT : NULLREMARK3ESTIMATED EROR OF BIN FREE R VALUE : NULLREMARK3NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.REMARK3NUCLEIC ACID ATOMS : 0REMARK3NUCLEIC ACID ATOMS : 0REMARK3SOLVENT ATOMS : 132REMARK3SOLVENT ATOMS : 132REMARK3B VALUES.REMARK3B VALUES.REMARK3B VALUES.REMARK3B11 (A**2) : NULLREMARK3B11 (A**2) : NULLREMARK3B22 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULL</td><td></td><td></td><td></td></tr> <tr><td>REMARK3BIN COMPLETENESS (WORKING+TEST) (%) : NULLREMARK3REFLECTIONS IN BIN (WORKING SET) : NULLREMARK3BIN R VALUE (WORKING SET) : NULLREMARK3BIN REE R VALUE (WORKING SET) : NULLREMARK3BIN FREE R VALUE TEST SET SIZE (%) : NULLREMARK3BIN FREE R VALUE TEST SET COUNT : NULLREMARK3ESTIMATED EROR OF BIN FREE R VALUE : NULLREMARK3NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.REMARK3NUCLEIC ACID ATOMS : 0REMARK3NUCLEIC ACID ATOMS : 0REMARK3SOLVENT ATOMS : 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NULLREMARK</td><td>REMARK</td><td>3</td><td>BIN COMPLETENESS (WORKING+TEST) (%) : NULL</td></tr> <tr><td>REMARK3BIN FREE R VALUE: NULLREMARK3BIN FREE R VALUE TEST SET SIZE (%) : NULLREMARK3BIN FREE R VALUE TEST SET COUNT: NULLREMARK3ESTIMATED ERROR OF BIN FREE R VALUE : NULLREMARK3REMARK3REMARK3NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.REMARK3PROTEIN ATOMS: 5711REMARK3NUCLEIC ACID ATOMS: 0REMARK3NUCLEIC ACID ATOMS: 132REMARK3SOLVENT ATOMS: 132REMARK3SOLVENT ATOMS: 27.18REMARK3FROM WILSON PLOT(A**2) : 31.94REMARK3MEAN B VALUE(OVERALL, A**2) : 27.18REMARK3B11 (A**2) : NULLREMARKREMARK3B22 (A**2) : NULLREMARKREMARK3B33 (A**2) : NULLREMARKREMARK3B12 (A**2) : NULLREMARKREMARK3B13 (A**2) : NULLREMARKREMARK3B13 (A**2) : NULLREMARKREMARK3B13 (A**2) : NULLREMARKREMARK3B13 (A**2) : NULLREMARK</td><td>REMARK</td><td>3</td><td>REFLECTIONS IN BIN (WORKING SET) : NULL</td></tr> <tr><td>REMARK3BIN FREE R VALUE: NULLREMARK3BIN FREE R VALUE TEST SET SIZE (%) : NULLREMARK3BIN FREE R VALUE TEST SET 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(A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL</td><td>REMARK</td><td>3</td><td></td></tr> <tr><td>REMARK3PROTEIN ATOMS: 5711REMARK3NUCLEIC ACID ATOMS: 0REMARK3HETEROGEN ATOMS: 0REMARK3SOLVENT ATOMS: 132REMARK3SOLVENT ATOMS: 132REMARK3B VALUES.REMARK3FROM WILSON PLOT(A**2) : 31.94REMARK3MEAN B VALUE(OVERALL, A**2) : 27.18REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A**2) : NULLREMARK3B22 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL</td><td>REMARK</td><td>3</td><td>NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.</td></tr> <tr><td>REMARK3NUCLEIC ACID ATOMS: 0REMARK3HETEROGEN ATOMS: 0REMARK3SOLVENT ATOMS: 132REMARK3BVALUES.REMARK3FROM WILSON PLOT(A**2) : 31.94REMARK3MEAN B VALUE(OVERALL, A**2) : 27.18REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A**2) : NULLREMARK3B22 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL</td><td>REMARK</td><td>3</td><td>PROTEIN ATOMS : 5711</td></tr> <tr><td>REMARK3HETEROGEN ATOMS: 0REMARK3SOLVENT ATOMS: 132REMARK3BVALUES.REMARK3FROM WILSON PLOT(A**2) : 31.94REMARK3MEAN B VALUE(OVERALL, A**2) : 27.18REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A**2) : NULLREMARK3B22 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL</td><td>REMARK</td><td>3</td><td>NUCLEIC ACID ATOMS : 0</td></tr> <tr><td>REMARK3SOLVENT ATOMS: 132REMARK3REMARK3B VALUES.REMARK3FROM WILSON PLOT(A**2) : 31.94REMARK3MEAN B VALUE(OVERALL, A**2) : 27.18REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A**2) : NULLREMARK3B22 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL</td><td>REMARK</td><td>3</td><td>HETEROGEN ATOMS : 0</td></tr> <tr><td>REMARK3REMARK3B VALUES.REMARK3FROM WILSON PLOT(A**2) : 31.94REMARK3MEAN B VALUE(OVERALL, A**2) : 27.18REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A**2) : NULLREMARK3B22 (A**2) : NULLREMARK3B33 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL</td><td>REMARK</td><td>3</td><td>SOLVENT ATOMS : 132</td></tr> <tr><td>REMARK3FROM WILSON PLOT<math>(A^{**2})</math>: 31.94REMARK3MEAN B VALUE<math>(OVERALL, A^{**2})</math>: 27.18REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A^{**2}) : NULLREMARK3B22 (A^{**2}) : NULLREMARK3B12 (A^{**2}) : NULLREMARK3B13 (A^{**2}) : NULLREMARK3B13 (A^{**2}) : NULLREMARK3B13 (A^{**2}) : NULLREMARK3B23 (A^{**2}) : NULL</td><td></td><td></td><td></td></tr> <tr><td>REMARK3FROM WILSON PLOT<math>(A^{**2})</math>: 31.94REMARK3MEAN B VALUE<math>(OVERALL, A^{**2})</math>: 27.18REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A^{**2}) : NULLREMARK3B22 (A^{**2}) : NULLREMARK3B12 (A^{**2}) : NULLREMARK3B13 (A^{**2}) : NULLREMARK3B13 (A^{**2}) : NULLREMARK3B13 (A^{**2}) : NULLREMARK3B23 (A^{**2}) : NULL</td><td>REMARK</td><td>3</td><td>B VALUES.</td></tr> <tr><td>REMARK3MEAN B VALUE(OVERALL, A**2) : 27.18REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A**2) : NULLREMARK3B22 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL</td><td></td><td>-</td><td></td></tr> <tr><td>REMARK3OVERALL ANISOTROPIC B VALUE.REMARK3B11 (A**2) : NULLREMARK3B22 (A**2) : NULLREMARK3B33 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL</td><td></td><td></td><td>MEAN B VALUE (OVERALL, A**2) : 27.18</td></tr> <tr><td>REMARK       3       B11 (A**2) : NULL         REMARK       3       B22 (A**2) : NULL         REMARK       3       B33 (A**2) : NULL         REMARK       3       B12 (A**2) : NULL         REMARK       3       B13 (A**2) : NULL         REMARK       3       B13 (A**2) : NULL         REMARK       3       B23 (A**2) : NULL</td><td></td><td></td><td></td></tr> <tr><td>REMARK       3       B22 (A**2) : NULL         REMARK       3       B33 (A**2) : NULL         REMARK       3       B12 (A**2) : NULL         REMARK       3       B13 (A**2) : NULL         REMARK       3       B23 (A**2) : NULL</td><td></td><td></td><td></td></tr> <tr><td>REMARK       3       B33 (A**2) : NULL         REMARK       3       B12 (A**2) : NULL         REMARK       3       B13 (A**2) : NULL         REMARK       3       B23 (A**2) : NULL</td><td></td><td></td><td></td></tr> <tr><td>REMARK       3       B12 (A**2) : NULL         REMARK       3       B13 (A**2) : NULL         REMARK       3       B23 (A**2) : NULL</td><td></td><td></td><td></td></tr> <tr><td>REMARK         3         B13 (A**2)         : NULL           REMARK         3         B23 (A**2)         : NULL</td><td></td><td></td><td></td></tr> <tr><td>REMARK 3 B23 (A**2) : NULL</td><td></td><td></td><td></td></tr> <tr><td></td><td></td><td>-</td><td></td></tr> <tr><td></td><td></td><td></td><td><u> </u></td></tr> <tr><td></td><td></td><td>-</td><td></td></tr>	REMARK	3	FIT IN THE HIGHEST RESOLUTION BIN.	REMARK3BIN RESOLUTION RANGE LOW(A) : NULLREMARK3BIN COMPLETENESS (WORKING+TEST) (%) : NULLREMARK3REFLECTIONS IN BIN(WORKING SET) : NULLREMARK3BIN R VALUE(WORKING SET) : NULLREMARK3BIN FREE R VALUE: NULLREMARK3BIN FREE R VALUE TEST SET SIZE (%) : NULLREMARK3BIN FREE R VALUE TEST SET COUNT: NULLREMARK3BUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.REMARK3PROTEIN ATOMS: 0REMARK3NUCLEIC ACID ATOMS: 0REMARK3NUCLEIC ACID ATOMS: 132REMARK3SOLVENT ATOMS: 132REMARK3SOLVENT ATOMS: 132REMARK3B VALUES.(OVERALL, A**2) : 31.94REMARK3B VALUE(OVERALL, A**2) : 27.18REMARK3B11 (A**2) : NULLREMARK3B11 (A**2) : NULLREMARK3B12 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B13 (A**2) : NULLREMARK3B23 (A**2) : NULL <td></td> <td></td> <td>TOTAL NUMBER OF BINS USED : NULL</td>			TOTAL NUMBER OF BINS USED : NULL	REMARK3BIN COMPLETENESS (WORKING+TEST) (%) : NULLREMARK3REFLECTIONS IN BIN (WORKING SET) : NULLREMARK3BIN R VALUE (WORKING SET) : NULLREMARK3BIN REE R 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		ESTIMATED COORDINATE ERROR.
REMARK	3	ESD FROM LUZZATI PLOT(A) : 0.24ESD FROM SIGMAA(A) : 0.25
REMARK	3	ESD FROM SIGMAA (A) : 0.25
REMARK	3	LOW RESOLUTION CUTOFF (A) : NULL
REMARK	3	
REMARK		CROSS-VALIDATED ESTIMATED COORDINATE ERROR.
		ESD FROM C-V LUZZATI PLOT (A) : NULL
REMARK		ESD FROM C-V SIGMAA (A) : NULL
REMARK	3	
REMARK	3	RMS DEVIATIONS FROM IDEAL VALUES.
REMARK	3	BOND LENGTHS(A) : 0.006BOND ANGLES(DEGREES) : 1.18
REMARK	3	BOND ANGLES (DEGREES) : 1.18
		DIHEDRAL ANGLES (DEGREES) : 27.95
REMARK	3	IMPROPER ANGLES (DEGREES) : NULL
REMARK		· · · · · · · · · · · · · · · · · · ·
REMARK	3	•
REMARK	3	VALUES/RESIDUE
REMARK	3	
REMARK		
REMARK	3	MAIN-CHAIN BOND (A**2) : NULL ; NULL
REMARK	3	MAIN-CHAIN ANGLE (A**2) : NULL ; NULL
REMARK	3	MAIN-CHAIN BOND(A**2) : NULL ; NULLMAIN-CHAIN ANGLE(A**2) : NULL ; NULLSIDE-CHAIN BOND(A**2) : NULL ; NULLSIDE-CHAIN ANGLE(A**2) : NULL ; NULL
REMARK	3	SIDE-CHAIN ANGLE (A**2) : NULL ; NULL
REMARK		NCS MODEL : NULL
REMARK REMARK		NCS MODEL : NULL
REMARK		NCS RESTRAINTS. RMS SIGMA/WEIGHT
REMARK	2	NCS RESTRAINTS. RMS SIGMA/WEIGHT GROUP 1 POSITIONAL (A) : NULL ; NULL
REMARK	3	GROUP 1 B-FACTOR (A**2) : NULL ; NULL
REMARK	3	GROUP I B-FACTOR (A**2) : NOLL ; NOLL
	-	PARAMETER FILE 1 : NULL
REMARK		
REMARK	3	
REMARK	-	OTHER REFINEMENT REMARKS: NULL
MERICIC	5	OTHER RELEASED TO THE THEFT TO THE THEFT TO THE THEFT THE THEFT THE THEFT THE THEFT THE THEFT THE THE THE THE THE THE THE THE THE TH

### **Refinement using CNS**

#### Template/example

3

REMARK

```
3 REFINEMENT.
REMARK
REMARK
        3
            PROGRAM
                        : CNS 1.2
REMARK
        3
            AUTHORS
                        : BRUNGER, ADAMS, CLORE, DELANO, GROS, GROSSE-
REMARK
        3
                        : KUNSTLEVE, JIANG, KUSZEWSKI, NILGES, PANNU,
       3
REMARK
                        : READ, RICE, SIMONSON, WARREN
       3
REMARK
REMARK
       3 REFINEMENT TARGET : ENGH & HUBER
REMARK
       3
REMARK
        3 DATA USED IN REFINEMENT.
REMARK
        3 RESOLUTION RANGE HIGH (ANGSTROMS) : 2.20
        3
            RESOLUTION RANGE LOW (ANGSTROMS) : 19.87
REMARK
            DATA CUTOFF
REMARK
        3
                                   (SIGMA(F)) : 2.0
REMARK
        3
            DATA CUTOFF HIGH
                                     (ABS(F)) : 89190.68
REMARK
        3
            DATA CUTOFF LOW
                                     (ABS(F)) : 0.0000
REMARK
        3
            COMPLETENESS (WORKING+TEST) (%) : 91.1
       3
REMARK
            NUMBER OF REFLECTIONS
                                            : 32745
REMARK
        3
        3 FIT TO DATA USED IN REFINEMENT.
REMARK
        3 CROSS-VALIDATION METHOD
                                            : THROUGHOUT
REMARK
REMARK
        3 FREE R VALUE TEST SET SELECTION : RANDOM
REMARK
       3 R VALUE
                               (WORKING SET) : 0.203
       3 FREE R VALUE
                                            : 0.237
REMARK
        3 FREE R VALUE TEST SET SIZE
                                         (%): 5.0
REMARK
REMARK
        3
            FREE R VALUE TEST SET COUNT
                                            : 1633
            ESTIMATED ERROR OF FREE R VALUE : 0.006
REMARK
        3
REMARK
        3
        3 FIT IN THE HIGHEST RESOLUTION BIN.
REMARK
        3 TOTAL NUMBER OF BINS USED
REMARK
                                               : 6
REMARK
        3 BIN RESOLUTION RANGE HIGH
                                           (A) : 2.00
        3 BIN RESOLUTION RANGE LOW
REMARK
                                           (A) : 2.13
REMARK
        3 BIN COMPLETENESS (WORKING+TEST) (%) : NULL
REMARK
       3 REFLECTIONS IN BIN
                                 (WORKING SET) : 0
REMARK
       3 BIN R VALUE
                                  (WORKING SET) : 0.237
        3 FREE R VALUE
REMARK
                                            : NULL
REMARK
        3
            FREE R VALUE TEST SET SIZE
                                         (%) : NULL
            FREE R VALUE TEST SET COUNT
REMARK
        3
                                            : NULL
REMARK
        3
            ESTIMATED ERROR OF FREE R VALUE : NULL
REMARK
        3
        3 NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
REMARK
REMARK
        3 PROTEIN ATOMS
                          : 2172
        3
            NUCLEIC ACID ATOMS
                                    : 0
REMARK
REMARK
        3 HETEROGEN ATOMS
                                    : 0
       3 SOLVENT ATOMS
REMARK
                                    : 127
REMARK
        3
        3 B VALUES.
REMARK
                                       (A**2) : 11.20
REMARK
        3
           FROM WILSON PLOT
                              (OVERALL, A**2) : 25.20
REMARK
        3
            MEAN B VALUE
REMARK
        3
           OVERALL ANISOTROPIC B VALUE.
      3 B11 (A**2) : 2.38000
REMARK
REMARK
        3
             B22 (A**2) : 2.38000
REMARK
       3
             B33 (A**2) : -4.76000
```

3 B12 (A\*\*2) : 0.00000 REMARK 3 B13 (A\*\*2) : 0.00000 REMARK B23 (A\*\*2) : 0.00000 REMARK 3 REMARK 3 REMARK 3 ESTIMATED COORDINATE ERROR. REMARK 3 ESD FROM LUZZATI PLOT (A) : 0.22 REMARK 3 ESD FROM SIGMAA (A) : 0.07 REMARK 3 LOW RESOLUTION CUTOFF (A) : 5.00 REMARK 3 REMARK 3 CROSS-VALIDATED ESTIMATED COORDINATE ERROR. REMARK 3 ESD FROM C-V LUZZATI PLOT (A) : 0.26 REMARK 3 ESD FROM C-V SIGMAA (A) : 0.18 REMARK 3 3 RMS DEVIATIONS FROM IDEAL VALUES. REMARK REMARK 3 REMARK 3 BOND LENGTHS (A) : 0.006 KAY:0.006REMARK3BOND ANGLES(DEGREES) : 1.30REMARK3DIHEDRAL ANGLES(DEGREES) : 24.30REMARK3IMPROPER ANGLES(DEGREES) : 0.82REMARK33IMPROPER ANGLES REMARK 3 REMARK 3 ISOTROPIC THERMAL MODEL : RESTRAINED REMARK 3 REMARK 3 ISOTROPIC THERMAL FACTOR RESTRAINTS. RMS SIGMA REMARK 3 MAIN-CHAIN BOND (A\*\*2) : NULL ; NULL REMARK 3 MAIN-CHAIN ANGLE (A\*\*2) : NULL ; NULL (A\*\*2) : NULL ; NULL (A\*\*2) : NULL ; NULL 3 SIDE-CHAIN BOND REMARK REMARK 3 SIDE-CHAIN ANGLE REMARK 3 REMARK 3 BULK SOLVENT MODELING. REMARK 3 METHOD USED : FLAT MODEL REMARK 3 KSOL : 0.45 REMARK 3 BSOL : 64.83 REMARK 3 REMARK 3 NCS MODEL : NULL REMARK 3 REMARK 3 NCS RESTRAINTS. RMS SIGMA/WEIGHT GROUP 1POSITIONAL(A) : NULL ; NULLGROUP 1B-FACTOR(A\*\*2) : NULL ; NULL REMARK 3 REMARK 3 REMARK 3 REMARK 3 PARAMETER FILE 1 : PROTEIN REP.PARAM REMARK 3 PARAMETER FILE 2 : WATER\_REP.PARAM REMARK 3 PARAMETER FILE 3 : ION.PARAM REMARK 3 PARAMETER FILE 4 : NULL REMARK 3 TOPOLOGY FILE 1 : PROTEIN.TOP REMARK 3 TOPOLOGY FILE 2 : WATER.TOP REMARK 3 TOPOLOGY FILE 3 : ION.TOP 3 TOPOLOGY FILE 4 : NULL REMARK 3 REMARK 3 OTHER REFINEMENT REMARKS: BULK SOLVENT MODEL USED REMARK

### **Refinement using CNX**

#### Template/example

3

REMARK

```
3 REFINEMENT.
REMARK
REMARK
         3
             PROGRAM
                         : CNX
REMARK
         3
             AUTHORS
                         : BRUNGER, ADAMS, CLORE, DELANO, GROS, GROSSE-
REMARK
         3
                         : KUNSTLEVE, JIANG, KUSZEWSKI, NILGES, PANNU,
REMARK
         3
                         : READ, RICE, SIMONSON, WARREN
         3
REMARK
REMARK
         3 DATA USED IN REFINEMENT.
         3 RESOLUTION RANGE HIGH (ANGSTROMS) : 3.00
REMARK
REMARK
         3
             RESOLUTION RANGE LOW (ANGSTROMS) : 50.00
                                     (SIGMA(F)) : 0.000
REMARK
         3
             DATA CUTOFF
             DATA CUTOFF HIGH
                                       (ABS(F)) : 1000.000
REMARK
         3
             DATA CUTOFF LOW
                                       (ABS(F)) : 0.0000
REMARK
         3
REMARK
         3
             COMPLETENESS (WORKING+TEST) (%) : 94.0
REMARK
         3
             NUMBER OF REFLECTIONS
                                                : 20693
REMARK
         3
REMARK
         3 FIT TO DATA USED IN REFINEMENT.
REMARK
         3
             CROSS-VALIDATION METHOD
                                               : NULL
             FREE R VALUE TEST SET SELECTION : NULL
REMARK
         3
                        (WORKING + TEST SET) : NULL
REMARK
         3
             R VALUE
REMARK
         3
             R VALUE
                                 (WORKING SET) : 0.219
REMARK
         3
             FREE R VALUE
                                               : 0.319
             FREE R VALUE TEST SET SIZE
                                           (%): 7.500
REMARK
         3
             FREE R VALUE TEST SET COUNT
REMARK
         3
                                               : 1643
REMARK
         3
             ESTIMATED ERROR OF FREE R VALUE
                                               : NULL
REMARK
         3
REMARK
         3 FIT/AGREEMENT OF MODEL WITH ALL DATA.
           R VALUE
                         (WORKING + TEST SET, NO CUTOFF) : NULL
REMARK
         3
             R VALUE
                                 (WORKING SET, NO CUTOFF) : NULL
REMARK
         3
REMARK
         3
           FREE R VALUE
                                              (NO CUTOFF) : NULL
         3
             FREE R VALUE TEST SET SIZE
                                           (%, NO CUTOFF) : NULL
REMARK
REMARK
         3
             FREE R VALUE TEST SET COUNT
                                              (NO CUTOFF) : NULL
REMARK
         3
             ESTIMATED ERROR OF FREE R VALUE (NO CUTOFF) : NULL
REMARK
         3
             TOTAL NUMBER OF REFLECTIONS
                                              (NO CUTOFF) : 20693
REMARK
         3
REMARK
         3 FIT IN THE HIGHEST RESOLUTION BIN.
             TOTAL NUMBER OF BINS USED
                                                  : NULL
REMARK
         3
REMARK
         3
             BIN RESOLUTION RANGE HIGH
                                              (A) : NULL
REMARK
         3
             BIN RESOLUTION RANGE LOW
                                              (A) : NULL
             BIN COMPLETENESS (WORKING+TEST) (%) : NULL
REMARK
         3
REMARK
         3
             REFLECTIONS IN BIN
                                    (WORKING SET) : NULL
         3
             BIN R VALUE
                                    (WORKING SET) : NULL
REMARK
REMARK
         3
             BIN FREE R VALUE
                                                  : NULL
                                            (%) : NULL
             BIN FREE R VALUE TEST SET SIZE
REMARK
         3
             BIN FREE R VALUE TEST SET COUNT
REMARK
         3
                                                  : NULL
             ESTIMATED ERROR OF BIN FREE R VALUE : NULL
REMARK
         3
REMARK
         3
            NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
REMARK
         3
REMARK
         3
             PROTEIN ATOMS
                                      : 7895
                                      : 0
REMARK
         3
             NUCLEIC ACID ATOMS
REMARK
         3
             HETEROGEN ATOMS
                                      : 276
REMARK
         3
             SOLVENT ATOMS
                                      : 0
```

REMARK 3 REMARK 3 B VALUES. REMARK 3 FROM WILSON PLOT (A\*\*2) : NULL REMARK 3 MEAN B VALUE (OVERALL, A\*\*2) : 43.13 REMARK 3 OVERALL ANISOTROPIC B VALUE. REMARK 3 B11 (A\*\*2) : -6.46200 REMARK 3 B22 (A\*\*2) : 0.93900 REMARK 3 B33 (A\*\*2) : 5.52300 REMARK 3 B12 (A\*\*2) : 0.00000 REMARK 3 B13 (A\*\*2) : 0.00000 REMARK 3 B23 (A\*\*2) : 0.00000 REMARK 3 REMARK 3 ESTIMATED COORDINATE ERROR. REMARK 3 ESD FROM LUZZATI PLOT (A) : NULL REMARK 3 REMARK 3 3 ESD FROM SIGMAA (A) : NULL LOW RESOLUTION CUTOFF (A) : NULL REMARK 3 REMARK 3 CROSS-VALIDATED ESTIMATED COORDINATE ERROR. REMARK 3 ESD FROM C-V LUZZATI PLOT (A) : NULL REMARK 3 ESD FROM C-V SIGMAA (A) : NULL REMARK 3 REMARK 3 RMS DEVIATIONS FROM IDEAL VALUES. REMARK 3 BOND LENGTHS (A) : 0.010 (DEGREES) : 1.85 (DEGREES) : NULL REMARK 3 BOND ANGLES 3 BOND ANGLES 3 DIHEDRAL ANGLES 3 IMPROPER ANGLES REMARK REMARK 3 (DEGREES) : NULL REMARK 3 REMARK 3 ISOTROPIC THERMAL MODEL : NULL REMARK 3 REMARK 3 ISOTROPIC THERMAL FACTOR RESTRAINTS. RMS SIGMA REMARK 3 MAIN-CHAIN BOND (A\*\*2) : 1.284 ; 1.500 REMARK 3 MAIN-CHAIN ANGLE (A\*\*2) : 2.239 ; 2.000 REMARK 3 SIDE-CHAIN BOND (A\*\*2) : 1.709 ; 2.000 REMARK 3 SIDE-CHAIN ANGLE (A\*\*2) : 2.698 ; 2.500 REMARK 3 3 BULK SOLVENT MODELING. REMARK REMARK 3 REMARK 3 3 METHOD USED : NULL : NULL KSOL REMARK 3 BSOL : NULL REMARK 3 REMARK 3 NCS MODEL : NULL REMARK 3 REMARK 3 PARAMETER FILE 1 : PROTEIN REP.PARAM REMARK 3 PARAMETER FILE 2 : DNA-RNA REP.PARAM REMARK 3 PARAMETER FILE 3 : WATER REP.PARAM REMARK 3 PARAMETER FILE 4 : N1234.XPRM REMARK 3 PARAMETER FILE 5 : LIG12AB.XPRM REMARK3TOPOLOGYFILE1: PROTEIN.TOPREMARK3TOPOLOGYFILE2: DNA-RNA.TOP REMARK 3 TOPOLOGY FILE 3 : WATER.TOP REMARK 3 TOPOLOGY FILE 4 : ION.TOP REMARK 3 TOPOLOGY FILE 5 : NULL REMARK 3 REMARK 3 OTHER REFINEMENT REMARKS: NULL

REMARK

# **Refinement using REFMAC**

## Template/example 1

3

REMARK	3	REFINEMENT.
REMARK	3	PROGRAM : REFMAC 5.3.0017 AUTHORS : MURSHUDOV,VAGIN,DODSON REFINEMENT TARGET : MAXIMUM LIKELIHOOD
REMARK	3	AUTHORS : MURSHUDOV, VAGIN, DODSON
REMARK	3	
REMARK	3	REFINEMENT TARGET : MAXIMUM LIKELIHOOD
REMARK	3	
		DATA USED IN REFINEMENT.
DEMADK	2	DEFAULTTION PANCE HIGH (ANCSTROMS) · 2 20
DEMARK	2	DESOLUTION DANCE LOW (ANCETROMS) . 2.20
DEMARK	2	RESOLUTION RANGE LOW (ANGSTROMS) : 50.00
REMARK	ວ ວ	RESOLUTION RANGE HIGH (ANGSTROMS) : 2.20RESOLUTION RANGE LOW (ANGSTROMS) : 50.00DATA CUTOFF (SIGMA(F)) : 0.000COMPLETENESS FOR RANGE (%) : 99.7NUMBER OF REFLECTIONS : 41377
DEMARK	ວ ວ	NUMPED OF DEFIECTIONS (3): 39.7
REMARK	ວ ວ	NOMBER OF REFLECTIONS : 415//
REMARK		
REMARK	3	FIT TO DATA USED IN REFINEMENT.
REMARK	3	CROSS-VALIDATION METHOD : THROUGHOUT
REMARK	3	FREE R VALUE TEST SET SELECTION : RANDOM
REMARK	3	R VALUE (WORKING + TEST SET) : 0.228
REMARK	3	CROSS-VALIDATION METHOD: THROUGHOUTFREE R VALUE TEST SET SELECTION: RANDOMR VALUE(WORKING + TEST SET): 0.228R VALUE(WORKING SET): 0.225FREE R VALUE: 0.283
REMARK	3	FREE R VALUE : 0.283 FREE R VALUE TEST SET SIZE (%) : 5.200 FREE R VALUE TEST SET COUNT : 2256
REMARK	3	FREE R VALUE TEST SET SIZE (%) : 5.200
REMARK	3	FREE R VALUE TEST SET COUNT : 2256
REMARK	3	
REMARK	3	FIT IN THE HIGHEST RESOLUTION BIN.
REMARK	3	TOTAL NUMBER OF BINS USED : 20
REMARK	3	FIT IN THE HIGHEST RESOLUTION BIN. TOTAL NUMBER OF BINS USED : 20 BIN RESOLUTION RANGE HIGH (A) : 2.20 BIN RESOLUTION RANGE LOW (A) : 2.26
REMARK	3	BIN RESOLUTION RANGE LOW (A) : 2.26
REMARK	3	BIN RESOLUTION RANGE HIGH(A) : 2.20BIN RESOLUTION RANGE LOW(A) : 2.26REFLECTION IN BIN(WORKING SET) : 2978BIN COMPLETENESS (WORKING+TEST) (%) : 98.65BIN R VALUE(WORKING SET) : 0.2840BIN FREE R VALUE SET COUNT: 161BIN FREE R VALUE: 0.3680
REMARK	3	BIN COMPLETENESS (WORKING+TEST) (%) : 98.65
REMARK	3	BIN R VALUE (WORKING SET) : 0.2840
REMARK	3	BIN FREE R VALUE SET COUNT : 161
REMARK	3	BIN FREE R VALUE : 0.3680
REMARK	- 3	
REMARK	3	NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
REMARK	3	PROTEIN ATOMS : 2043
REMARK	3	NUCLEIC ACID ATOMS : 0
REMARK	3	HETEROGEN ATOMS : 12
REMARK	3	PROTEIN ATOMS: 2043NUCLEIC ACID ATOMS: 0HETEROGEN ATOMS: 12SOLVENT ATOMS: 88
REMARK	3	
REMARK	3	B VALUES.
REMARK	3	B VALUE TYPE : LIKELY RESIDUAL
		FROM WILSON PLOT (A**2) : 41.60
REMARK		
REMARK	3	OVERALL ANISOTROPIC B VALUE.
REMARK		
REMARK		
REMARK	3	$B_{33}$ (A**2) : -0.44000
REMARK	3	B12 (A**2) : 0.00000
REMARK	3	B13 (A**2) : -0.05000
REMARK	3	B23 (A**2) : 0.00000
REMARK		
		ESTIMATED OVERALL COORDINATE ERROR.
REMARK		
	Ũ	

(A): 0.345

REMARK	3	ESU BASED ON FREE R VALUE ESU BASED ON MAXIMUM LIKELIHOOD	(.	A): 0.	
REMARK	3	ESU BASED ON MAXIMUM LIKELIHOOD	(	A): 0.	
REMARK	3	ESU FOR B VALUES BASED ON MAXIMUM LIKELI	100D (A**	2): 17	.897
REMARK	3				
REMARK	3	CORRELATION COEFFICIENTS.			
REMARK	3	CORRELATION COEFFICIENT FO-FC : 0.94	18		
REMARK	3	CORRELATION COEFFICIENT FO-FC FREE : 0.92	22		
REMARK	3				
REMARK	3	RMS DEVIATIONS FROM IDEAL VALUES CO	DUNT R	MS	WEIGHT
REMARK	3	BOND LENGTHS REFINED ATOMS (A): (	5752 ; 0.	012 ;	0.022
REMARK	3		487 ; O.		
REMARK	3	BOND ANGLES REFINED ATOMS (DEGREES):	9197 : 1.	419 :	1,960
REMARK	3	BOND ANGLES OTHERS (DEGREES): 10			
REMARK	3	TORSION ANGLES, PERIOD 1 (DEGREES):			
REMARK	3	TORSION ANGLES PERIOD 2 (DEGREES).	254 • 35	063 •2	2 724
REMARK	3	TORSION ANGLES, PERIOD 2(DEGREES):TORSION ANGLES, PERIOD 3(DEGREES):TORSION ANGLES, PERIOD 4(DEGREES):	1071 • 16	530 •1	5 000
REMARK	3	TORSION ANGLES, TERIOD 5 (DEGREES).	10,1,10.	210,1	5 000
REMARK	3	CHIRAL-CENTER RESTRAINTS (A**3):	LO31 ; 0.	210 ;1 002 ;	0.200
REMARK	3		7482 ; 0.		
REMARK	3		L286 ; 0.		
REMARK	3		L064 ; 0.		
REMARK	3	NON-BONDED CONTACTS OTHERS (A):	1210 ; 0.	•	
REMARK	3		3041 ; 0.		
REMARK	3	NON-BONDED TORSION OTHERS (A):	3676 ; 0.	087 ;	0.200
REMARK	3	H-BOND (XY) REFINED ATOMS (A):			
REMARK	3		NULL ; N		
REMARK	3	POTENTIAL METAL-ION REFINED ATOMS (A): 1	NULL ; N		
REMARK	3	POTENTIAL METAL-ION OTHERS (A): 1	NULL ; N		
REMARK	3	SYMMETRY VDW REFINED ATOMS (A):	7;0.	185 ;	0.200
REMARK	3	POTENTIAL METAL-ION OTHERS(A):SYMMETRY VDW REFINED ATOMS(A):SYMMETRY VDW OTHERS(A):	30;0.	167 ;	0.200
REMARK	3	SYMMETRY H-BOND REFINED ATOMS (A):	5;0.	189 ;	0.200
REMARK	3		NULL ; N		
REMARK	3	SYMMETRY METAL-ION REFINED ATOMS (A): 1	NULL ; N	ULL ;	NULL
REMARK	3		NULL ; N		
REMARK	3			•	
REMARK	3	ISOTROPIC THERMAL FACTOR RESTRAINTS.	COUNT R	MS	WEIGHT
REMARK	3		5474 ; 0.		
REMARK	3	MAIN-CHAIN BOND OTHER ATOMS (A**2):	1741 ; 0.		
REMARK	3		5943 ; 0.		
REMARK	3		2998; 1.		
REMARK	3				
REMARK	3			, 10	1.000
REMARK		ANISOTROPIC THERMAL FACTOR RESTRAINTS.	COUNT R	MS W	ETGHT
REMARK	3			ULL ;	
REMARK	3			,	
	3				
REMARK	3	SPHERICITI; BONDED ATOMS (A**2):	NULL; N	ULL ;	NULL
REMARK					
REMARK	3				
REMARK		NUMBER OF DIFFERENT NCS GROUPS : 5			
REMARK	3				
REMARK	3				
REMARK	3				
REMARK	3				
REMARK	3				
REMARK	3		6		
REMARK	3				
REMARK	3				
REMARK	3	LOOSE POSITIONAL 1 (A): 1461	; 0.25	; 5.0	0

REMARK	3	LOOSE THERMAL 1 (A**2): 1461 ; 0.73 ; 10.00
REMARK	3	
REMARK	3	NCS GROUP NUMBER : 2
REMARK	3	CHAIN NAMES : L A
REMARK	3	NUMBER OF COMPONENTS NCS GROUP : 1
REMARK	3	COMPONENT C SSSEQI TO C SSSEQI CODE
REMARK	3	1 L 108 L 211 6
REMARK	3	1 A 108 A 211 6
REMARK	3	GROUP CHAIN COUNT RMS WEIGHT
REMARK	3	LOOSE POSITIONAL 2 (A): 1359; 0.23; 5.00
REMARK	3	LOOSE THERMAL 2 (A**2): 1359 ; 1.29 ; 10.00
REMARK	3	
REMARK	3	NCS GROUP NUMBER : 3
REMARK	3	CHAIN NAMES : H B
REMARK	3	NUMBER OF COMPONENTS NCS GROUP : 1
REMARK	3	COMPONENT C SSSEQI TO C SSSEQI CODE
REMARK	3	1 H 1 H 113 6
REMARK	3	1 B 1 B 113 6
REMARK	3	GROUP CHAIN COUNT RMS WEIGHT
REMARK	3	GROUP CHAIN         COUNT         RMS         WEIGHT           LOOSE POSITIONAL         3         (A):         1547 ;         0.11 ;         5.00           LOOSE THERMAL         3         (A**2):         1547 ;         1.00 ;         10.00
REMARK	3	LOOSE THERMAL $3$ (A**2): 1547 : 1.00 : 10.00
REMARK	3	
REMARK	3	NCS GROUP NUMBER : 4
REMARK		NCS GROUP NUMBER : 4 CHAIN NAMES : H B
REMARK	3	NUMBER OF COMPONENTS NCS GROUP : 1
REMARK	3	COMPONENT C SSSEQI TO C SSSEQI CODE
REMARK	3	1 H 114 H 126 6
REMARK	3	1 B 114 B 126 6
REMARK	3	GROUP CHAIN COUNT RMS WEIGHT
REMARK	3	LOOSE POSITIONAL 4 (A): 155; 0.38; 5.00
REMARK	3	LOOSE THERMAL 4 $(A^{**2})$ : 155 ; 1.95 ; 10.00
REMARK	3	
REMARK	3	NCS GROUP NUMBER : 5
REMARK	3	NCS GROUP NUMBER : 5 CHAIN NAMES : H B
REMARK	3	NUMBER OF COMPONENTS NCS GROUP : 1
REMARK	3	COMPONENT C SSSEQI TO C SSSEQI CODE
REMARK	3	1 H 136 H 227 6
REMARK	3	1 B 136 B 227 6
REMARK	3	GROUP CHAIN COUNT RMS WEIGHT
REMARK	3	LOOSE POSITIONAL 5 (A): 973; 0.23; 5.00
REMARK	3	LOOSE THERMAL 5 (A**2): 973; 0.82; 10.00
REMARK	3	
REMARK	3	TLS DETAILS
REMARK	3	NUMBER OF TLS GROUPS : 4
REMARK	3	
REMARK	3	TLS GROUP : 1
REMARK	3	NUMBER OF COMPONENTS GROUP : 1
REMARK	3	
REMARK	3	T11: 0.2639 T22: 0.1856
REMARK	3	T33: 0.0412 T12: 0.0129
REMARK	3	T13: -0.0229 T23: 0.0075
REMARK	3	
REMARK	3	L11: 1.2476 L22: 18.8186
REMARK	3	
REMARK	3	
	U	

3 S TENSOR REMARK 

 S11:
 -0.1230
 S12:
 -0.1350
 S13:
 0.1070

 S21:
 0.1833
 S22:
 0.1989
 S23:
 -0.0673

 S31:
 0.2988
 S32:
 0.3017
 S33:
 -0.0759

 3 REMARK REMARK 3 3 REMARK REMARK 3 REMARK 3 TLS GROUP : 2 REMARK 3 NUMBER OF COMPONENTS GROUP : 1 REMARK 3 COMPONENTS C SSSEQI TO C SSSEQI REMARK 3 RESIDUE RANGE : B 1 B 227 
 REMARK
 3
 ORIGIN FOR THE GROUP (A):
 18.6717
 -2.2091
 -2.3508

 REMARK
 3
 T TENSOR

 REMARK
 3
 T TENSOR

 REMARK
 3
 T11:
 0.3169
 T22:
 0.0830

 REMARK
 3
 T33:
 0.0521
 T12:
 0.0175

 REMARK
 3
 T13:
 -0.0382
 T23:
 0.0060

 REMARK
 3
 L
 TENSOR

 REMARK
 3
 L11:
 2.8160
 L22:
 1.2951

 REMARK
 3
 L33:
 2.1804
 L12:
 0.8548

 REMARK
 3
 L13:
 -2.1037
 L23:
 -1.0227

 REMARK
 3
 S
 TENSOR
 S11:
 0.1656
 S12:
 0.1951

 REMARK
 3
 S11:
 0.1656
 S12:
 0.1951
 S13:
 0.1602

 REMARK
 3
 S21:
 -0.3132
 S22:
 0.0276
 S23:
 0.3597

 REMARK
 3
 S31:
 -0.0658
 S32:
 -0.1993
 S33:
 -0.1933

 REMARK 3 REMARK 3 TLS GROUP : 3 REMARK3ILS GROUP : 3REMARK3NUMBER OF COMPONENTS GROUP : 1REMARK3COMPONENTSC SSSEQIREMARK3RESIDUE RANGE : H1H227REMARK3ORIGIN FOR THE GROUP (A): 17.9538 -1.3383 43.2238REMARK3T TENSORDEMARK3T TINSOR 
 REMARK
 3
 T11:
 0.3108
 T22:
 0.1076

 REMARK
 3
 T33:
 0.0689
 T12:
 -0.0386

 REMARK
 3
 T13:
 0.0466
 T23:
 0.0367
 REMARK 3 L TENSOR 

 REMARK
 3
 L1 TENSOK

 REMARK
 3
 L11:
 0.7004 L22:
 1.2871

 REMARK
 3
 L33:
 2.0590 L12:
 -0.2673

 REMARK
 3
 L13:
 1.1558 L23:
 -0.0172

 REMARK
 3
 S TENSOR
 -0.1826

 REMARK
 3
 S11:
 0.0754 S12:
 -0.1826

 REMARK
 3
 S11:
 0.0754
 S12:
 -0.1826
 S13:
 -0.0304

 REMARK
 3
 S21:
 0.2142
 S22:
 0.1332
 S23:
 0.2512

 REMARK
 3
 S31:
 0.1977
 S32:
 -0.2560
 S33:
 -0.2086

 REMARK 3 REMARK 3 TLS GROUP : 4 REMARK 3 NUMBER OF COMPONENTS GROUP : 1 REMARK 3 COMPONENTS C SSSEQI TO C SSSEQI REMARK 3 RESIDUE RANGE : L 1 L 221 

 REMARK
 3
 RESIDCE RANGE:
 1
 1
 1
 221

 REMARK
 3
 ORIGIN FOR THE GROUP (A):
 36.2584
 -4.5702
 24.8879

 REMARK
 3
 T TENSOR
 7
 7
 7
 12:
 0.1946

 REMARK
 3
 T33:
 0.0079
 T12:
 0.0203
 7
 13:
 -0.0113
 T23:
 0.0538

 REMARK
 3
 L
 THEODE
 2
 L
 THEODE
 10:
 10:
 10:
 10:
 10:
 10:
 10:
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 <t REMARK 3 L TENSOR 

 REMARK
 3
 L11:
 2.9390
 L22:
 17.8781

 REMARK
 3
 L33:
 6.7012
 L12:
 4.8729

 REMARK
 3
 L13:
 1.9743
 L23:
 1.1500

 REMARK
 3
 S
 TENSOR
 1113.0
 111300

 REMARK
 3
 S
 TENSOR
 0.0598
 S13:
 0.3426

 REMARK
 3
 S21:
 0.2222
 S22:
 0.0581
 S23:
 0.7020

 REMARK
 3
 S31:
 0.0016
 S32:
 0.1934
 S33:
 0.0213

REMARK 3 REMARK 3 BULK SOLVENT MODELLING.	
REMARK 3 BOLK SOLVENT MODELLING.	
REMARK 3 METHOD USED : MASK	
REMARK 3 PARAMETERS FOR MASK CALCULATION	
REMARK 3 VDW PROBE RADIUS : 1.20	
REMARK 3 ION PROBE RADIUS : 0.80	
REMARK 3 SHRINKAGE RADIUS : 0.80	
REMARK 3	
REMARK 3 OTHER REFINEMENT REMARKS: HYDROGENS HAVE BEEN ADDED IN	$\Gamma HE$
REMARK 3 RIDING POSITIONS	

## Template/example 2

REMARK	3	
	-	REFINEMENT.
REMARK		
REMARK		AUTHORS : MURSHUDOV, VAGIN, DODSON
REMARK	3	
REMARK		REFINEMENT TARGET : MAXIMUM LIKELIHOOD
REMARK		
REMARK	3	DATA USED IN REFINEMENT.
REMARK		
REMARK	3	RESOLUTION RANGE HIGH (ANGSTROMS) : 1.40 RESOLUTION RANGE LOW (ANGSTROMS) : 38.27
REMARK	3	DATA CUTOFF (SIGMA(F)) : NULL
REMARK		COMPLETENESS FOR RANGE (%): 99.8
REMARK	3	NUMBER OF REFLECTIONS : 102082
REMARK	3	
REMARK	3	FIT TO DATA USED IN REFINEMENT.
REMARK		CROSS-VALIDATION METHOD : THROUGHOUT
REMARK	3	FREE R VALUE TEST SET SELECTION : RANDOM
REMARK	3	R VALUE (WORKING + TEST SET) : 0.179
REMARK	3	R VALUE(WORKING SET): 0.179FREE R VALUE: 0.193FREE R VALUE TEST SET SIZE(%): 5.000
REMARK	3	FREE R VALUE : 0.193
REMARK	-	
REMARK	3	
REMARK	3	
REMARK	3	FIT IN THE HIGHEST RESOLUTION BIN.
REMARK	3	
REMARK	3	BIN RESOLUTION RANGE HIGH (A) : 1.40
REMARK	3	BIN RESOLUTION RANGE LOW (A) : 1.44 REFLECTION IN BIN (WORKING SET) : 7804 BIN COMPLETENESS (WORKING+TEST) (%) : 103.34
REMARK	3	REFLECTION IN BIN (WORKING SET) : 7804
REMARK	3	BIN COMPLETENESS (WORKING+TEST) (%) : 103.34
REMARK		BIN R VALUE (WORKING SET) : 0.3830
REMARK	3	BIN FREE R VALUE SET COUNT : 404
REMARK	3	BIN FREE R VALUE : 0.3790
REMARK	3	
REMARK	3	
REMARK	3	PROTEIN ATOMS : 1043
REMARK	3	NUCLEIC ACID ATOMS: 0HETEROGEN ATOMS: 0SOLVENT ATOMS: 65
REMARK	3	HETEROGEN ATOMS : 0
REMARK	3	SOLVENT ATOMS : 65
REMARK	3	5
REMARK	3	
REMARK	3	FROM WILSON PLOT (A**2) : NULL

REMARK	3		4.92		
REMARK	3				
REMARK	3	B11 (A**2) : -7.82000			
REMARK		B22 (A**2) : -6.91000			
REMARK		B33 (A**2) : 14.73000			
REMARK		B12 (A**2) : 0.00000			
REMARK	3				
REMARK	3				
REMARK	3				
REMARK	3				
REMARK		ESU BASED ON R VALUE ESU BASED ON FREE R VALUE ESU BASED ON MAXIMUM LIKELIHOOD		(A):	
REMARK	3	ESU BASED ON FREE R VALUE		(A):	
REMARK	3	ESU BASED ON MAXIMUM LIKELIHOOD		(A):	0.037
REMARK	3		ELIHOOD	(A**2):	1.010
REMARK	3				
REMARK		CORRELATION COEFFICIENTS.			
REMARK	3	CORRELATION COEFFICIENT FO-FC :			
REMARK	3	CORRELATION COEFFICIENT FO-FC FREE :	0.965		
REMARK	3				
REMARK	3		COUNT	RMS	WEIGHT
REMARK	3	BOND LENGTHS REFINED ATOMS(A)BOND LENGTHS OTHERS(A)BOND ANGLES REFINED ATOMS(DEGREES)	: 4192	; 0.012	; 0.022
REMARK	3	BOND LENGTHS OTHERS (A)	: 2871	; 0.001	; 0.020
REMARK	3	BOND ANGLES REFINED ATOMS (DEGREES)	: 5694	; 1.405	; 1.992
REMARK	3	BOND ANGLES OTHERS (DEGREES)	: 7056	; 0.928	; 3.000
REMARK	3	TORSION ANGLES, PERIOD 1 (DEGREES)	: 558	; 5.807	; 5.000
REMARK	3		: 172	;35.581	;23.953
REMARK	3	TORSION ANGLES, PERIOD 3 (DEGREES)	: 773	;12.566	;15.000
REMARK	3	TORSION ANGLES, PERIOD 4 (DEGREES)	: 29	;12.738	;15.000
REMARK	3	CHIRAL-CENTER RESTRAINTS (A**3)	: 659	; 0.087	; 0.200
REMARK	3		: 4650	; 0.006	; 0.020
REMARK	3				
REMARK	3	. ,			
REMARK	3			; NULL	
REMARK	3			; NULL	
REMARK	3	NON-BONDED TORSTON OTHERS (A)	: NUT.T.		
REMARK	3	. ,	: NULL	-	
REMARK	3	H-BOND (XY) OTHERS (A)	: NULL	•	; NULL
REMARK	3	POTENTIAL METAL-ION REFINED ATOMS (A)	: NULL		-
REMARK					; NULL
REMARK	3	POTENTIAL METAL-ION OTHERS (A) SYMMETRY VDW REFINED ATOMS (A)			
REMARK	3			•	; NULL
REMARK	3				
REMARK	3			•	
REMARK	3				
REMARK	3	SYMMETRY METAL-ION OTHERS (A)			
REMARK	3	( )			
REMARK	3	ISOTROPIC THERMAL FACTOR RESTRAINTS.	COUN	r rms	WEIGHT
REMARK	3			; 0.801	
REMARK	3	· · · · · · · · · · · · · · · · · · ·		; 0.208	-
REMARK	3	· · · · · · · · · · · · · · · · · · ·		; 1.425	-
REMARK	3			; 2.325	
REMARK	3			; 3.654	
REMARK	3			,	,
REMARK		ANISOTROPIC THERMAL FACTOR RESTRAINTS.	COUN	r RMS	WEIGHT
REMARK	3				
REMARK	3	, , , , , , , , , , , , , , , , , , , ,		•	•
REMARK	3	,			
REMARK	3		. 110111	1 10111	1 110111
1/11/11/11/11/11/11/11/11/11/11/11/11/1	5				

3 NCS RESTRAINTS STATISTICS REMARK 3 NUMBER OF DIFFERENT NCS GROUPS : 1 REMARK 3 REMARK 3 NCS GROUP NUMBER REMARK : 1 : A B C D REMARK 3 CHAIN NAMES REMARK 3 NUMBER OF COMPONENTS NCS GROUP : 1 REMARK 3 COMPONENT C SSSEQI TO C SSSEQI CODE 1 A 3 A 300 6 REMARK 3 

 1
 B
 3
 B
 300
 0

 1
 C
 3
 C
 300
 6

 1
 D
 3
 D
 300
 6

 1
 D
 3
 D
 300
 6

 GROUP CHAIN
 COUNT
 RMS

 1
 C
 COUNT
 RMS

 REMARK 3 REMARK 3 REMARK 3 REMARK 3 REMARK 3 WEIGHT 

 REMARK
 3
 GROUP CHAIN
 COUNT
 RMS
 WEIGHT

 REMARK
 3
 LOOSE POSITIONAL
 1
 A
 (A):
 1265
 ;
 0.730
 ;
 5.000

 REMARK
 3
 LOOSE POSITIONAL
 1
 B
 (A):
 1265
 ;
 0.550
 ;
 5.000

 REMARK
 3
 LOOSE POSITIONAL
 1
 B
 (A):
 1265
 ;
 0.670
 ;
 5.000

 REMARK
 3
 LOOSE POSITIONAL
 1
 D
 (A):
 1265
 ;
 0.640
 ;
 5.000

 REMARK
 3
 LOOSE THERMAL
 1
 A
 (A\*\*2):
 1265
 ;
 5.080
 ;10.000

 REMARK
 3
 LOOSE THERMAL
 1
 B
 (A\*\*2):
 1265
 ;
 2.980
 ;10.000

 REMARK
 3
 LOOSE THERMAL
 1
 C
 (A\*\*2):
 1265
 ;
 3.660
 ;10.000

 REMARK
 3
 LOOSE THERMAL
 1
 D
 (A\*\*2):
 1265
 ;
 3.660
 ;10.000

 REMARK 3 REMARK 3 TWIN DETAILS 3 NUMBER OF TWIN DOMAINS : 2 REMARK 3 TWIN DOMAIN : 1 REMARK 3 TWIN OPENING REMARK TWIN OPERATOR : H,K,L REMARK3TWIN FRACTION : 0.867REMARK3TWIN DOMAIN : 2DEMAPK3TWIN OPERATOR : L, -K, J REMARK 3 TWIN OPERATOR : L,-K,H TWIN FRACTION : 0.133 REMARK 3 REMARK 3 REMARK 3 TLS DETAILS REMARK 3 NUMBER OF TLS GROUPS : 4 REMARK 3 REMARK 3 TLS GROUP : 1 REMARK3NUMBER OF COMPONENTS GROUP : 1REMARK3COMPONENTSC SSSEQIREMARK3COMPONENTSC SSSEQIREMARK3RESIDUE RANGE : A0A300REMARK3ORIGIN FOR THE GROUP (A):REMARK3T TENSORREMARK3T11: REMARK 3 T11: -0.1434 T22: -0.0225 

 REMARK
 3
 T33: -0.0349 T12: -0.0039

 REMARK
 3
 T13: -0.0011 T23: -0.0142

 REMARK
 3
 L TENSOR

 REMARK
 3
 L11:
 0.2499
 L22:
 2.6544

 REMARK
 3
 L33:
 3.1537
 L12:
 -0.0787

 REMARK
 3
 L13:
 -0.1230
 L23:
 -0.1410

 REMARK
 3
 S
 TENSOR

 3
 S11: -0.0939
 S12: -0.0135
 S13: 0.0635

 2
 S11: -0.0939
 S12: -0.0135
 S13: 0.0635
 REMARK 3 REMARK 3 REMARK3S21:0.4552S22:0.0720S23:0.3304REMARK3S31:-0.0916S32:-0.2462S33:0.0219 REMARK 3 REMARK 3 TLS GROUP : 2 REMARK 3 NUMBER OF COMPONENTS GROUP : 1 REMARK 3 COMPONENTS C SSSEQI TO C SSSEQI REMARK 3 RESIDUE RANGE : B 2 B 300 

 REMARK
 3
 ORIGIN FOR THE GROUP (A): 26.3124
 33.3641
 14.1724

 REMARK
 3
 T TENSOR

 REMARK
 3
 T11:
 0.0332
 T22:
 -0.0920

REMARK		T33: -0.0634 T12: -0.0201
REMARK	3	T13: 0.0037 T23: 0.0130
REMARK		L TENSOR
REMARK	3	L11: 0.8985 L22: 2.2480
REMARK	3	L33: 0.4623 L12: -1.0984
REMARK	3	L13: -0.0114 L23: 0.6608
REMARK	3	S TENSOR
REMARK	3	S11: -0.0856 S12: -0.0481 S13: -0.1305
REMARK	3	S21: 0.2230 S22: 0.0179 S23: 0.0879
REMARK	3	S31: 0.1878 S32: 0.0617 S33: 0.0677
REMARK	3	
REMARK	3	TLS GROUP : 3
REMARK	3	NUMBER OF COMPONENTS GROUP : 1
REMARK		COMPONENTS C SSSEQI TO C SSSEQI
REMARK	3	RESIDUE RANGE : C 3 C 300
REMARK		ORIGIN FOR THE GROUP (A): 31.0030 33.2958 50.0967
REMARK	3	T TENSOR
REMARK	3	T11: -0.1785 T22: -0.0337
REMARK	3	T33: -0.0199 T12: 0.0202
REMARK	3	T13: -0.0342 T23: -0.0065
REMARK	3 3	L TENSOR
REMARK		L11: 1.1097 L22: 4.1071
REMARK	3	L33: 2.4647 L12: 0.0878
REMARK	3	L13: 0.3839 L23: 0.9425
REMARK	3	S TENSOR
REMARK		S11: -0.0357 S12: 0.1026 S13: -0.1055
REMARK	3	S21: -0.2843 S22: 0.0934 S23: 0.3463
REMARK	3	S31: 0.4808 S32: -0.0374 S33: -0.0577
REMARK	3	
REMARK	3	TLS GROUP : 4
REMARK	3	NUMBER OF COMPONENTS GROUP : 1
REMARK	3	COMPONENTS C SSSEQI TO C SSSEQI
REMARK	3	RESIDUE RANGE : D 2 D 300
REMARK	3	ORIGIN FOR THE GROUP (A): 36.9044 51.7770 58.1371
REMARK	3	T TENSOR
REMARK	3	T11: 0.0198 T22: -0.0873
REMARK	3	T33: -0.0907 T12: -0.0053
REMARK	3	T13: 0.0045 T23: 0.0206
REMARK	3	L TENSOR
REMARK	3	L11: 0.6326 L22: 2.2127
REMARK	3	L33: 1.0826 L12: 0.2556
REMARK	3	L13: 0.1927 L23: 0.6195
REMARK		S TENSOR
REMARK	3	S11: -0.0674 S12: 0.0446 S13: 0.0174
REMARK	3	S21: -0.1848 S22: 0.0310 S23: 0.0198
REMARK	3	S31: -0.2108 S32: 0.0752 S33: 0.0364
REMARK	3	5510.2100 552. 0.0752 555. 0.0504
	3	BULK SOLVENT MODELLING.
REMARK REMARK	3	METHOD USED : BABINET MODEL WITH MASK
REMARK	3	PARAMETERS FOR MASK CALCULATION
REMARK	3	VDW PROBE RADIUS : 1.40
REMARK	3	ION PROBE RADIUS : 0.80
REMARK	3	SHRINKAGE RADIUS : 0.80
REMARK	3	
REMARK	3	
REMARK	3	RIDING POSITIONS

### **Refinement using NUCLSQ**

#### Template

3

REMARK

```
3 REFINEMENT.
REMARK
REMARK
        3
            PROGRAM
                      : NUCLSO
REMARK
        3
           AUTHORS
                      : WESTHOF, DUMAS, MORAS
REMARK
        3
      3 DATA USED IN REFINEMENT.
REMARK
      3 RESOLUTION RANGE HIGH (ANGSTROMS) :
REMARK
      3 RESOLUTION RANGE LOW (ANGSTROMS) :
REMARK
REMARK 3 DATA CUTOFF
                                  (SIGMA(F)) :
REMARK
      3 COMPLETENESS FOR RANGE
                                      (%) :
REMARK
      3 NUMBER OF REFLECTIONS
                                            :
       3
REMARK
        3 FIT TO DATA USED IN REFINEMENT.
REMARK
REMARK
        3
          CROSS-VALIDATION METHOD
                                           :
REMARK
        3
           FREE R VALUE TEST SET SELECTION
                                           :
          R VALUE
REMARK
        3
                     (WORKING + TEST SET)
                                           :
REMARK 3 R VALUE
                         (WORKING SET)
                                           :
REMARK 3 FREE R VALUE
                                           :
REMARK 3 FREE R VALUE TEST SET SIZE
                                        (%) :
REMARK 3 FREE R VALUE TEST SET COUNT
REMARK 3
REMARK
      3 FIT/AGREEMENT OF MODEL WITH ALL DATA.
      3 R VALUE (WORKING + TEST SET, NO CUTOFF) :
REMARK
      3 R VALUE
                            (WORKING SET, NO CUTOFF) :
REMARK
REMARK 3 FREE R VALUE
REMARK 3 FREE R VALUE
                                        (NO CUTOFF) :
        3 FREE R VALUE TEST SET SIZE (%, NO CUTOFF)
                                                   :
REMARK 3 FREE R VALUE TEST SET COUNT (NO CUTOFF) :
REMARK 3 TOTAL NUMBER OF REFLECTIONS
                                        (NO CUTOFF) :
REMARK 3
        3 NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
REMARK
        3 PROTEIN ATOMS
REMARK
                                   :
REMARK
        3 NUCLEIC ACID ATOMS
                                   :
REMARK
      3 HETEROGEN ATOMS
                                   :
      3 SOLVENT ATOMS
REMARK
                                   :
REMARK
        3
REMARK
        3 B VALUES.
          FROM WILSON PLOT
REMARK
        3
                                      (A**2) :
REMARK
                             (OVERALL, A**2) :
        3
          MEAN B VALUE
          OVERALL ANISOTROPIC B VALUE.
REMARK 3
REMARK 3 B11 (A**2) :
REMARK 3 B22 (A**2) :
REMARK 3 B33 (A**2) :
      3 B12 (A**2) :
REMARK
      3 B13 (A**2) :
REMARK
       3
           B23 (A**2) :
REMARK
REMARK
        3
        3 ESTIMATED COORDINATE ERROR.
REMARK
        3 ESD FROM LUZZATI PLOT
REMARK
                                       (A) :
REMARK
           ESD FROM SIGMAA
        3
                                       (A) :
REMARK 3 LOW RESOLUTION CUTOFF
                                       (A) :
REMARK
      3
REMARK
       3 RMS DEVIATIONS FROM IDEAL VALUES.
```

REMARK3SUGAR-BASE BOND DISTANCE(A) ::;REMARK3SUGAR-BASE BOND ANGLE DISTANCE(A) :;REMARK3PHOSPHATE BONDS DISTANCE(A) :;REMARK3PHOSPHATE BOND ANGLE, H-BOND(A) :;REMARK3PLANE RESTRAINT(A) :;REMARK3CHIRAL-CENTER RESTRAINT(A) :;REMARK3NON-BONDED CONTACT RESTRAINTS.;REMARK3SINGLE TORSION CONTACT(A) :;REMARK3MULTIPLE TORSION CONTACT(A) :;REMARK3ISOTROPIC THERMAL FACTOR RESTRAINTS.RMSSIGMA
REMARK3PHOSPHATE BONDS DISTANCE(A):;REMARK3PHOSPHATE BOND ANGLE, H-BOND(A):;REMARK3PLANE RESTRAINT(A):;REMARK3CHIRAL-CENTER RESTRAINT(A**3):;REMARK3NON-BONDED CONTACT RESTRAINTS.;REMARK3SINGLE TORSION CONTACT(A):;REMARK3MULTIPLE TORSION CONTACT(A):;REMARK3SINGLE TORSION CONTACT(A):;REMARK3MULTIPLE TORSION CONTACT(A):;REMARK3SINGLE TORSION CONTACT(A):;REMARK3SINGLE TORSION CONTACT(A):;REMARK3SINGLE TORSION CONTACT(A):;REMARK3SINGLE TORSION CONTACTSINGLE;REMARK3SINGLE TORSION CONTACTSINGLE;REMARK3SINGLE TORSION CONTACTSINGLE;REMARK3SINGLE TORSION CONTACTSINGLE;
REMARK3PHOSPHATE BOND ANGLE, H-BOND (A):;REMARK3;REMARK3PLANE RESTRAINT (A):;REMARK3CHIRAL-CENTER RESTRAINT (A**3):;REMARK3NON-BONDED CONTACT RESTRAINTS.REMARK3SINGLE TORSION CONTACT (A):;REMARK3MULTIPLE TORSION CONTACT (A):;REMARK3*********************************
REMARK3PLANE RESTRAINT(A):REMARK3PLANE RESTRAINT(A):REMARK3CHIRAL-CENTER RESTRAINT(A**3):REMARK3NON-BONDED CONTACT RESTRAINTS.REMARK3SINGLE TORSION CONTACT(A):REMARK3MULTIPLE TORSION CONTACT(A):REMARK3
REMARK3PLANE RESTRAINT(A):;REMARK3CHIRAL-CENTER RESTRAINT(A**3):;REMARK3NON-BONDED CONTACT RESTRAINTS.;REMARK3SINGLE TORSION CONTACT(A):;REMARK3MULTIPLE TORSION CONTACT(A):;REMARK3SINGLE TORSION CONTACT(A):;
REMARK3CHIRAL-CENTER RESTRAINT(A**3):;REMARK33;REMARK3NON-BONDED CONTACT RESTRAINTS.;REMARK3SINGLE TORSION CONTACT(A):;REMARK3MULTIPLE TORSION CONTACT(A):;REMARK3SINGLE TORSION CONTACT(A):;REMARK3SINGLE TORSION CONTACT(A):;
REMARK3REMARK3NON-BONDED CONTACT RESTRAINTS.REMARK3SINGLE TORSION CONTACT(A) : ;REMARK3MULTIPLE TORSION CONTACT(A) : ;REMARK3
REMARK3NON-BONDED CONTACT RESTRAINTS.REMARK3SINGLE TORSION CONTACT(A):REMARK3MULTIPLE TORSION CONTACT(A):REMARK3
REMARK3SINGLE TORSION CONTACT(A):;REMARK3MULTIPLE TORSION CONTACT(A):;REMARK3
REMARK 3 MULTIPLE TORSION CONTACT (A): ; REMARK 3
REMARK 3
ΡΕΜΑΡΚ 3 ΤΟΟ ΤΟ ΤΗ ΕΡΜΑΙ ΕΛΟΤΟΡ ΡΕΟΤΡΑΙΝΤΟ ΡΜΟ ΟΤΩΜΑ
REPARK 5 IDDITOTIC THERMAL FACTOR RESTRATIND. RMS SIGNA
REMARK 3 SUGAR-BASE BONDS (A**2) : ;
REMARK 3 SUGAR-BASE ANGLES (A**2) : ;
REMARK 3 PHOSPHATE BONDS (A**2) : ;
REMARK 3 PHOSPHATE BOND ANGLE, H-BOND (A**2) : ;
REMARK 3
REMARK 3 OTHER REFINEMENT REMARKS:

### Refinement using CCP4, PROFFT, GPRLSA, and related programs

#### Template/example

3

REMARK

REMARK 3 REFINEMENT. 3 PROGRAM : PROFFT 3 AUTHORS : KONNERT, HENDRICKSON, FINZEL REMARK 3 REMARK REMARK REMARK 3 DATA USED IN REFINEMENT. REMARK 3 RESOLUTION RANGE HIGH (ANGSTROMS) : 1.65 REMARK 3 RESOLUTION RANGE LOW (ANGSTROMS) : 5.00 REMARK 3 DATA CUTOFF (SIGMA(F)) : 2.000 (%) : NULL REMARK 3 COMPLETENESS FOR RANGE REMARK 3 NUMBER OF REFLECTIONS : 10699 REMARK 3 3 FIT TO DATA USED IN REFINEMENT. REMARK 3 CROSS-VALIDATION METHOD : NONE REMARK 3 FREE R VALUE TEST SET SELECTION : NULL REMARK REMARK 3 R VALUE (WORKING + TEST SET) : 0.180 REMARK 3 R VALUE (WORKING SET) : NULL REMARK 3 FREE R VALUE : NULL REMARK 3 FREE R VALUE TEST SET SIZE (%) : NULL REMARK 3 FREE R VALUE TEST SET COUNT : NULL REMARK 3 REMARK 3 FIT/AGREEMENT OF MODEL WITH ALL DATA. REMARK 3 R VALUE (WORKING + TEST SET, NO CUTOFF) : NULL REMARK 3 R VALUE (WORKING SET, NO CUTOFF) : NULL 3 FREE R VALUE REMARK (NO CUTOFF) : NULL 3 FREE R VALUE TEST SET SIZE (%, NO CUTOFF) : NULL REMARK 3 FREE R VALUE TEST SET COUNT (NO CUTOFF) : NULL REMARK TOTAL NUMBER OF REFLECTIONS REMARK 3 (NO CUTOFF) : NULL REMARK 3 REMARK 3 NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT. REMARK 3 PROTEIN ATOMS : 843 REMARK3NUCLEIC ACID ATOMS: 0REMARK3HETEROGEN ATOMS: 6REMARK3SOLVENT ATOMS: 8 REMARK 3 SOLVENT ATOMS : 85 REMARK 3 3 B VALUES. REMARK 3 FROM WILSON PLOT (A\*\*2) : NULL REMARK REMARK REMARK 3 MEAN B VALUE (OVERALL, A\*\*2) : 17.28 OVERALL ANISOTROPIC B VALUE. 3 REMARK 3 B11 (A\*\*2) : NULL REMARK 3 B22 (A\*\*2) : NULL REMARK 3 B33 (A\*\*2) : NULL REMARK 3 B12 (A\*\*2) : NULL REMARK 3 B13 (A\*\*2) : NULL REMARK 3 B23 (A\*\*2) : NULL REMARK 3 3 ESTIMATED COORDINATE ERROR. REMARK 3 ESD FROM LUZZATI PLOT (A) : 0.20 REMARK ESD FROM SIGMAA 3 (A) : NULL REMARK REMARK 3 LOW RESOLUTION CUTOFF (A) : NULL REMARK 3 REMARK 3 RMS DEVIATIONS FROM IDEAL VALUES. REMARK 3 DISTANCE RESTRAINTS. RMS SIGMA

REMARK	-	BOND LENGTH			0.012		
REMARK		ANGLE DISTANCE			NULL		
REMARK	-	INTRAPLANAR 1-4 DISTANCE	· · ·		NULL	•	
REMARK	3	H-BOND OR METAL COORDINAT	ION (A)	:	NULL	;	NULL
REMARK	3						
REMARK	3	PLANE RESTRAINT			NULL		
REMARK	3	CHIRAL-CENTER RESTRAINT	(A**3)	:	NULL	;	NULL
REMARK	3						
REMARK	3	NON-BONDED CONTACT RESTRAI	NTS.				
REMARK	3	SINGLE TORSION	(A)	:	NULL	;	NULL
REMARK	3	MULTIPLE TORSION	(A)	:	NULL	;	NULL
REMARK	3	H-BOND (XY)	(A)	:	NULL	;	NULL
REMARK	3	H-BOND (X- $HY$ )			NULL		
REMARK	3	· · · ·	. ,				
REMARK	3	CONFORMATIONAL TORSION ANG	LE RESTRAIN	IT:	s.		
REMARK	3	SPECIFIED	(DEGREES)	:	NULL	;	NULL
REMARK	3	PLANAR	(DEGREES)				
REMARK	3	STAGGERED	(DEGREES)				
REMARK	3	TRANSVERSE	(DEGREES)				
REMARK	3		,			•	
REMARK	3	ISOTROPIC THERMAL FACTOR RE	STRAINTS.		RMS	1	SIGMA
REMARK	3	MAIN-CHAIN BOND	(A**2)	:	NULL	:	NULL
REMARK	3	MAIN-CHAIN ANGLE	(A**2)				
REMARK	3	SIDE-CHAIN BOND	(A**2)				
REMARK	3	SIDE-CHAIN ANGLE	(A**2)				
REMARK	3		(11 2)	•	цопп	'	цопп
REMARK	3	OTHER REFINEMENT REMARKS: N	TIT T				
KEMARK	3	OTHER REFINEMENT REMARKS: N					

## **Refinement using SHELXL**

#### Template

3

REMARK

PROGRAM : SHELXL-97 AUTHORS : G.M.SHELDRICK REMARK 3 REMARK 3 AUTHORS REMARK 3 REMARK 3 DATA USED IN REFINEMENT. 3 RESOLUTION RANGE HIGH (ANGSTROMS) : 1.15 REMARK 3 RESOLUTION RANGE LOW (ANGSTROMS) : 30.00 REMARK REMARK 3 DATA CUTOFF (SIGMA(F)) : 0.000 REMARK 3 COMPLETENESS FOR RANGE (%) : 99.8 : FREE R REMARK 3 CROSS-VALIDATION METHOD REMARK 3 FREE R VALUE TEST SET SELECTION : RANDOM REMARK 3 REMARK 3 FIT TO DATA USED IN REFINEMENT (NO CUTOFF). REMARK 3 R VALUE (WORKING + TEST SET, NO CUTOFF) : 0.116 REMARK 3 R VALUE (WORKING SET, NO CUTOFF) : 0.116 REMARK 3 FREE R VALUE (NO CUTOFF) : 0.145 REMARK 3 FREE R VALUE TEST SET SIZE (%, NO CUTOFF) : 5.000 3 FREE R VALUE TEST SET COUNT (NO CUTOFF) : 4279 REMARK 3 TOTAL NUMBER OF REFLECTIONS REMARK (NO CUTOFF) : 85756 3 REMARK REMARK 3 FIT/AGREEMENT OF MODEL FOR DATA WITH F>4SIG(F). REMARK 3 R VALUE (WORKING + TEST SET, F>4SIG(F)) : 0.010 REMARK 3 R VALUE (WORKING SET, F>4SIG(F)) : 0.010 REMARK 3 FREE R VALUE (F>4SIG(F)) : 0.136 REMARK 3 FREE R VALUE TEST SET SIZE (%, F>4SIG(F)) : 5.000 REMARK 3 FREE R VALUE TEST SET COUNT (F>4SIG(F)) : 3859 REMARK 3 TOTAL NUMBER OF REFLECTIONS (F>4SIG(F)) : 77074 REMARK 3 REMARK 3 NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT. REMARK 3 PROTEIN ATOMS : 1439 NUCLEIC ACID ATOMS : 0 REMARK 3 HETEROGEN ATOMS : 0 REMARK 3 REMARK 3 SOLVENT ATOMS : 288 REMARK 3 REMARK 3 MODEL REFINEMENT. REMARK 3 OCCUPANCY SUM OF NON-HYDROGEN ATOMS : 1638.00 OCCUPANCY SUM OF HYDROGEN ATOMS REMARK 3 : 1406.00 REMARK 3 NUMBER OF DISCRETELY DISORDERED RESIDUES : 7 3 NUMBER OF LEAST-SQUARES PARAMETERS : 15553 REMARK 3 NUMBER OF RESTRAINTS : 19134 REMARK REMARK 3 REMARK 3 RMS DEVIATIONS FROM RESTRAINT TARGET VALUES. REMARK 3 BOND LENGTHS (A) : 0.015 (A) : 0.030 3 REMARK ANGLE DISTANCES REMARK 3 SIMILAR DISTANCES (NO TARGET VALUES) (A) : 0.000 REMARK 3 DISTANCES FROM RESTRAINT PLANES (A) : 0.030 REMARK 3 ZERO CHIRAL VOLUMES (A\*\*3) : 0.086 (A\*\*3) : 0.095 REMARK 3 NON-ZERO CHIRAL VOLUMES 3 ANTI-BUMPING DISTANCE RESTRAINTS (A) : 0.032 REMARK 3 RIGID-BOND ADP COMPONENTS (A\*\*2): 0.005 REMARK 3 SIMILAR ADP COMPONENTS SIMILAR ADP COMPONENTS(A\*\*2) : 0.048APPROXIMATELY ISOTROPIC ADPS(A\*\*2) : 0.107 REMARK 3 REMARK REMARK 3

REMARK	3	BULK SOLVENT MODELING.
REMARK	3	METHOD USED: MOEWS & KRETSINGER, J.MOL.BIOL.91(1973)201-228
REMARK	3	
REMARK	3	STEREOCHEMISTRY TARGET VALUES : ENGH & HUBER
REMARK	3	SPECIAL CASE: NULL
REMARK	3	
REMARK	3	OTHER REFINEMENT REMARKS: NULL

# **Refinement using TNT/BUSTER**

# Template/example 1

	n	
REMARK		
REMARK	<u>່</u>	REFINEMENT. PROGRAM : TNT
		AUTHORS : TRONRUD, TEN EYCK, MATTHEWS
REMARK		
REMARK		DATA USED IN REFINEMENT.
REMARK	3	RESOLUTION RANGE HIGH (ANGSTROMS) : 1.60 RESOLUTION RANGE LOW (ANGSTROMS) : 30.00
REMARK	3	RESOLUTION RANGE LOW (ANGSTROMS) : 30.00
REMARK	3	DATA CUTOFF (SIGMA(F)) : 0.000
REMARK	3	COMPLETENESS FOR RANGE (%): 93.6 NUMBER OF REFLECTIONS : 80952
REMARK	3	NUMBER OF REFLECTIONS : 80952
REMARK	3	
REMARK	3	USING DATA ABOVE SIGMA CUTOFF.
REMARK	3	CROSS-VALIDATION METHOD : THROUGHOUT
REMARK	3	USING DATA ABOVE SIGMA CUTOFF. CROSS-VALIDATION METHOD : THROUGHOUT FREE R VALUE TEST SET SELECTION : RANDOM
REMARK	3	R VALUE (WORKING + TEST SET) : 0.160
REMARK	3	R VALUE (WORKING SET) : 0.158
	2	
REMARK	3	FREE R VALUE TEST SET SIZE (%) : NULL
REMARK	3	FREE R VALUE       : 0.215         FREE R VALUE TEST SET SIZE       (%) : NULL         FREE R VALUE TEST SET COUNT       : 6164
REMARK	3	
REMARK	3	USING ALL DATA, NO SIGMA CUTOFF.
		R VALUE (WORKING + TEST SET, NO CUTOFF) : 0.1600
		R VALUE (WORKING SET, NO CUTOFF) : NULL
DEMYDR	3	EDEE D VALUE (WORKING BEL, NO COTOFF) . NULL
DEMADY	2	TREE & VALUE (NO COTOFF) . NULL
DEMADY	2 2	FREE R VALUE (NO CUTOFF) : NULL FREE R VALUE TEST SET SIZE (%, NO CUTOFF) : NULL FREE R VALUE TEST SET COUNT (NO CUTOFF) : NULL TOTAL NUMBER OF REFLECTIONS (NO CUTOFF) : 80952
REMARK	ა ა	TREE R VALUE TEST SET COUNT (NO CUTOFF) : NULL
REMARK	ა ა	TOTAL NUMBER OF REFLECTIONS (NO COTOFF) : 80952
REMARK	3	NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
REMARK	3	PROTEIN ATOMS : 5555
REMARK	3	PROTEIN ATOMS: 5555NUCLEIC ACID ATOMS: 0HETEROGEN ATOMS: 69SOLVENT ATOMS: 681
REMARK	3	HETEROGEN ATOMS : 69
		SOLVENT ATOMS : 681
REMARK		
REMARK		WILSON B VALUE (FROM FCALC, A**2) : NULL
REMARK	3	
REMARK	3	RMS DEVIATIONS FROM IDEAL VALUES.RMSWEIGHTCOUNTBOND LENGTHS(A) : NULL; NULL; NULLBOND ANGLES(DEGREES) : NULL; NULL; NULL
REMARK	3	BOND LENGTHS (A) : NULL ; NULL ; NULL
REMARK	3	
REMARK	3	TORSION ANGLES (DEGREES) : NULL ; NULL ; NULL
REMARK	3	PSEUDOROTATION ANGLES (DEGREES) : NULL ; NULL ; NULL
REMARK	3	TRIGONAL CARBON PLANES (A) : NULL ; NULL ; NULL
REMARK	3	GENERAL PLANES (A) : NULL ; NULL ; NULL
REMARK	3	ISOTROPIC THERMAL FACTORS (A**2) : NULL ; NULL ; NULL
REMARK	3	NON-BONDED CONTACTS (A) : NULL ; NULL ; NULL
REMARK	3	
REMARK	3	INCORRECT CHIRAL-CENTERS (COUNT) : NULL
REMARK	3	· · ·
REMARK	3	BULK SOLVENT MODELING.
REMARK	3	METHOD USED : NULL
REMARK	3	KSOL : NULL

REMARK3BSOL: NULLREMARK3RESTRAINT LIBRARIES.REMARK3STEREOCHEMISTRY : ENGH & HUBERREMARK3ISOTROPIC THERMAL FACTOR RESTRAINTS : NULLREMARK3REMARK3REMARK3OTHER REFINEMENT REMARKS: NULL

### **Template/example 2**

3

REMARK

REMARK		
REMARK	3	REFINEMENT. PROGRAM : BUSTER-TNT 2.1.1
REMARK	3	PROGRAM : BUSTER-TNT 2.1.1
REMARK	3	AUTHORS : BLANC, ROVERSI, VONRHEIN, BRICOGNE, TRONRUD,
REMARK		: TEN EYCK, MATTHEWS
REMARK		
REMARK	2	
REMARK	3	
REMARK	3	RESOLUTION RANGE HIGH (ANGSTROMS) : 1.75
REMARK	3	RESOLUTION RANGE LOW (ANGSTROMS) : 34.65
REMARK	3	DATA CUTOFF (SIGMA(F)) : 0.000
REMARK	3	COMPLETENESS FOR RANGE (%) : 97.4
REMARK	3	NUMBER OF REFLECTIONS : 53863
REMARK		
REMARK	3	USING DATA ABOVE SIGMA CUTOFF.
REMARK	3	USING DATA ABOVE SIGMA CUTOFF. CROSS-VALIDATION METHOD : THROUGHOUT
REMARK	3	FREE R VALUE TEST SET SELECTION : RANDOM
		R VALUE (WORKING + TEST SET) : 0.182
REMARK	2	R VALUE (WORKING SET) : 0.180
REMARK	3	FREE R VALUE : 0.218
REMARK	3	FREE R VALUE: 0.218FREE R VALUE TEST SET SIZE(%)FREE R VALUE TEST SET COUNT: 2691
REMARK	3	FREE R VALUE TEST SET COUNT : 2691
REMARK	- 3	
REMARK	3	FIT IN THE HIGHEST RESOLUTION BIN.
REMARK	3	TOTAL NUMBER OF BINS USED : 9
REMARK	3	BIN RESOLUTION RANGE HIGH (ANGSTROMS) : 1.75
REMARK	3	BIN RESOLUTION RANGE LOW (ANGSTROMS) : 1.86
REMARK	3	BIN RESOLUTION RANGE LOW (ANGSTROMS) : 1.86 BIN COMPLETENESS (WORKING+TEST) (%) : 97.43
REMARK	3	REFLECTIONS IN BIN (WORKING + TEST SET) : 7826
REMARK		
DEMADK	2	DEFIECTIONS IN DIN (MODELING SET) • 7447
DEMADK	2 2	REFLECTIONS IN BIN(WORKING SET) : 7447BIN R VALUE(WORKING SET) : 0.2376
REMARK	2	DIN R VALUE (WORKING SEI) : 0.2370
REMARK	3	BIN R VALUE(WORKTING SET)0.22370BIN FREE R VALUE TEST SET SIZE(%): 4.84BIN FREE R VALUE TEST SET COUNT: 379ESTIMATED ERROR OF FREE R VALUE: NULL
REMARK	3	BIN FREE R VALUE TEST SET SIZE (%) : 4.84
REMARK	3	BIN FREE R VALUE TEST SET COUNT : 379
REMARK	3	ESTIMATED ERROR OF FREE R VALUE : NULL
REMARK	3	
REMARK	3	NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
REMARK	3	PROTEIN ATOMS: 3944NUCLEIC ACID ATOMS: 0HETEROGEN ATOMS: 44SOLVENT ATOMS: 257
REMARK	3	NUCLEIC ACID ATOMS : 0
REMARK	3	HETEROGEN ATOMS : 44
REMARK	3	SOLVENT ATOMS : 257
REMARK	3	
REMARK		B VALUES.
REMARK		
	-	
REMARK		
REMARK		
REMARK		
REMARK	3	B22 (A**2) : 6.21317

	2	
REMARK		B33 (A**2) : -7.05095
		B12 (A**2) : 0.00000
		B13 (A**2) : -7.10165
REMARK	3	B23 (A**2) : 0.00000
REMARK	3	
REMARK	3	ESTIMATED COORDINATE ERROR.
REMARK	3	ESD FROM LUZZATI PLOT (A) : NULL
REMARK	3	
REMARK	3	CORRELATION COEFFICIENTS.
REMARK	3	CORRELATION COEFFICIENT FO-FC : NULL
REMARK	3	CORRELATION COEFFICIENT FO-FC FREE : NULL
REMARK	3	
REMARK		RMS DEVIATIONS FROM IDEAL VALUES RMS WEIGHT COUNT
REMARK	3	BOND LENGTHS (A) : 0.011 ; 2.000 ; 4060
REMARK	3	BOND LENGTHS(A) : 0.011 ; 2.000 ; 4060BOND ANGLES(DEGREES) : 1.186 ; 2.000 ; 5477
REMARK	3	TORSION ANGLES (DEGREES) : 16.973 ; 0.000 ; 827
REMARK		
REMARK		PSEUDOROTATION ANGLES (DEGREES) : NULL ; NULL ; NULL
REFIARE	3	PSEUDOROTATION ANGLES (DEGREES) : NULL ; NULL ; NULL TRIGONAL CARBON PLANES (A) : 0.012 ; 2.000 ; 101
REMARK	3 3	TRIGONAL CARBON PLANES       (A) : 0.012 ; 2.000 ; 101         GENERAL PLANES       (A) : 0.018 ; 5.000 ; 608
	3 3 3	PSEUDOROTATION ANGLES (DEGREES) : NULL ; NULL ; NULLTRIGONAL CARBON PLANES(A) : $0.012$ ; $2.000$ ; $101$ GENERAL PLANES(A) : $0.018$ ; $5.000$ ; $608$ ISOTROPIC THERMAL FACTORS (A**2) : $1.634$ ; $20.000$ ; $4060$
REMARK	3	TRIGONAL CARBON PLANES(A) : 0.012; 2.000; 101GENERAL PLANES(A) : 0.018; 5.000; 608ISOTROPIC THERMAL FACTORS(A**2) : 1.634; 20.000; 4060
REMARK REMARK	3	TRIGONAL CARBON PLANES       (A) : 0.012 ; 2.000 ; 101         GENERAL PLANES       (A) : 0.018 ; 5.000 ; 608
REMARK REMARK REMARK	3 3	TRIGONAL CARBON PLANES(A) : 0.012; 2.000; 101GENERAL PLANES(A) : 0.018; 5.000; 608ISOTROPIC THERMAL FACTORS(A**2) : 1.634; 20.000; 4060NON-BONDED CONTACTS(A) : 0.044; 5.000; 34
REMARK REMARK REMARK REMARK	3 3 3	TRIGONAL CARBON PLANES(A): 0.012; 2.000; 101GENERAL PLANES(A): 0.018; 5.000; 608ISOTROPIC THERMAL FACTORS(A**2): 1.634; 20.000; 4060NON-BONDED CONTACTS(A): 0.044; 5.000; 34
REMARK REMARK REMARK REMARK REMARK	3 3 3 3	TRIGONAL CARBON PLANES       (A) : 0.012 ; 2.000 ; 101         GENERAL PLANES       (A) : 0.018 ; 5.000 ; 608         ISOTROPIC THERMAL FACTORS (A**2) : 1.634 ; 20.000; 4060         NON-BONDED CONTACTS       (A) : 0.044 ; 5.000 ; 34         INCORRECT CHIRAL-CENTERS (COUNT) : NULL

REMARK

# **Refinement using PHENIX**

# Template/example 1

3

KEMAKK	5		
REMARK	3	REFINEMENT.	
REMARK	3	PROGRAM	: PHENIX (PHENIX.REFINE)
REMARK	3	AUTHORS	: PAUL ADAMS, PAVEL AFONINE, VICENT CHEN, IAN
REMARK	3		: DAVIS, KRESHNA GOPAL, RALF GROSSE-
REMARK	3		: KUNSTLEVE,LI-WEI HUNG,ROBERT IMMORMINO,
REMARK	3		: TOM IOERGER, AIRLIE MCCOY, ERIK MCKEE, NIGEL
REMARK	3		: MORIARTY, REETAL PAI, RANDY READ, JANE
REMARK	3		: RICHARDSON, DAVID RICHARDSON, TOD ROMO, JIM
REMARK	3		: SACCHETTINI, NICHOLAS SAUTER, JACOB SMITH,
REMARK	3		: LAURENT STORONI, TOM TERWILLIGER, PETER
REMARK	3		: ZWART
REMARK	3		
REMARK	3	REFINEMENT	TARGET : ML
REMARK	3		
REMARK	3	DATA USED IN	REFINEMENT.
REMARK	3		RANGE HIGH (ANGSTROMS) : 2.99
REMARK	3	RESOLUTION	· ,
REMARK	3	MIN(FOBS/SI	
REMARK	3	•	5 FOR RANGE (%) : 96.7
REMARK	3	NUMBER OF R	
REMARK	3		
REMARK	3	<b>ΕΤΤ ΤΟ DATA</b>	JSED IN REFINEMENT.
REMARK	3	R VALUE	(WORKING + TEST SET) : 0.293
REMARK	3	R VALUE	(WORKING SET) : 0.291
REMARK	3	FREE R VALU	
REMARK	3		E TEST SET SIZE (%) : 4.980
REMARK	3		E TEST SET COUNT : 12081
REMARK	3		
REMARK	3	בייעם איי ביי	JSED IN REFINEMENT (IN BINS).
REMARK	3		TION RANGE COMPL. NWORK NFREE RWORK RFREE
REMARK	3		- 9.2600 0.98 8197 419 0.1970 0.2050
REMARK	3	2 9.2600	
REMARK	3	3 7.3700	
REMARK	3		
REMARK	3	5 5.8500	
REMARK		6 5.4300	
REMARK	3	7 5.1200	
REMARK	3		- 4.6500         0.97         7613         423         0.2690         0.3260
REMARK	3	9 4.6500	- 4.4700 0.97 7737 394 0.2790 0.2920
REMARK	3	10 4.4700	
	3		
REMARK	3	11 4.3200 12 4.1800	
REMARK	3	13 4.0600	
REMARK	3	14 3.9500	
REMARK			
REMARK	3	15 3.8600	
REMARK	3	16 3.7700	
REMARK	3	17 3.6900	
REMARK	3	18 3.6200	
REMARK	3	19 3.5500	
REMARK	3	20 3.4800	
REMARK	3	21 3.4300	- 3.3700 0.98 7742 372 0.3200 0.3760

REMARK

REMARK	3	22  3.3700 = 3.3200  0.98  7667  411  0.3440  0.4360
REMARK	3	23       3.3200 - 3.2700       0.98       7700       414       0.3410       0.3840
REMARK	3	24 3.2700 - 3.2200 0.97 7667 411 0.3350 0.3870
REMARK	3	25 3.2200 - 3.1800 0.97 7541 419 0.3400 0.3790
REMARK	3	243.2700 -3.22000.9776674110.33500.3870253.2200 -3.18000.9775414190.34000.3790263.1800 -3.14000.9676374020.34600.4220
REMARK	3	27 3.1400 - 3.1000 0.96 7613 381 0.3580 0.3940
REMARK	3	28 3.1000 - 3.0600 0.96 7538 427 0.3790 0.4290
REMARK	3	29 3.0600 - 3.0300 0.95 7440 376 0.3760 0.4350
REMARK	3	283.1000 -3.06000.9675384270.37900.4290293.0600 -3.03000.9574403760.37600.4350303.0300 -2.99000.7760932900.39500.4490
REMARK	3	
REMARK		BULK SOLVENT MODELLING.
		METHOD USED : FLAT BULK SOLVENT MODEL
REMARK	ა ი	SOLVENT RADIUS : 1.11
REMARK	3	SHRINKAGE RADIUS : 0.90
REMARK	3	K_SOL : 0.30 B SOL : 56.99
REMARK		B_SOL : 56.99
REMARK	3	
REMARK		
REMARK	3	
REMARK	3	PHASE ERROR (DEGREES, MAXIMUM-LIKELIHOOD BASED) : 36.180
REMARK	3	
REMARK		B VALUES.
REMARK	3	FROM WILSON PLOT (A**2): 50.24
REMARK		MEAN B VALUE (OVERALL, A**2) : 62.67
REMARK		OVERALL ANISOTROPIC B VALUE.
REMARK		B11 (A**2) : -12.34000
REMARK	3	B22 (A**2) : -11.49000
REMARK	3	B22 (A**2) : -11.49000 B33 (A**2) : 23.84000 B12 (A**2) : 0.00000
DEMUDK	3	$B12 (A * * 2) \cdot 0.00000$
DEMADY	2	B12 $(A \times 2)$ : 0.00000 B13 $(A \times 2)$ : 0.00000
REMARK		
		$B25(A^{2}):0.00000$
REMARK	3	
REMARK	3	
REMARK	3	
REMARK	3	OPERATOR: -H,-K,L
REMARK	3	
REMARK	3	
REMARK	3	RMSD COUNT
REMARK	3	BOND : 0.059 59703
REMARK	3	
REMARK	3	CHIRALITY : 0.243 9800
REMARK	3	PLANARITY: 0.013 10535
REMARK	3	DIHEDRAL : 24.960 22449
REMARK	3	
REMARK	3	TLS DETAILS
REMARK	3	NUMBER OF TLS GROUPS : 4
REMARK	3	TLS GROUP : 1
REMARK	3	SELECTION: CHAIN A
REMARK	3	ORIGIN FOR THE GROUP (A): 34.3280 -44.3362 -33.2464
REMARK	3	T TENSOR
REMARK	3	T11: -0.3752 T22: -0.2836
REMARK	3	T33: -0.1972 T12: -0.0686
REMARK	3	T13: $0.0888$ T23: $-0.1454$
REMARK	3	L TENSOR
REMARK	3	L11: -0.0328 L22: -0.0264
REMARK	3	L33: -0.0458 L12: -0.0047
REMARK	3	L13: 0.0289 L23: -0.0435
REMARK	3	S TENSOR

	С	
REMARK	3	S11: -0.0220 S12: 0.7030 S13: 0.0888
REMARK	3	S21: -0.7542 S22: -0.0140 S23: -0.0265
REMARK	3	S31: -0.2584 S32: 0.0315 S33: 0.0000
REMARK	3	TLS GROUP : 2
REMARK	3	SELECTION: CHAIN B
REMARK	3	ORIGIN FOR THE GROUP (A): 45.1940 -82.3594 -33.3841
REMARK	3	T TENSOR
REMARK	3	T11: -0.0302 T22: -0.0378
REMARK	3	T33: -0.0668 T12: -0.0642
REMARK	3	T13: 0.1450 T23: -0.0811
REMARK	3	L TENSOR
REMARK	3	L11: 0.0154 L22: 0.0032
REMARK	3	L33: -0.0145 L12: 0.0014
REMARK	3	L13: 0.0441 L23: 0.0209
REMARK	3	S TENSOR
	3	
REMARK		S11: -0.0023 S12: 0.6259 S13: 0.1176 S21: -0.6677 S22: -0.0510 S23: 0.0868
REMARK	3	S21: -0.1498 S32: -0.2360 S33: 0.0000
REMARK	3	
REMARK	3	TLS GROUP : 3
REMARK	3	SELECTION: CHAIN C
REMARK	3	ORIGIN FOR THE GROUP (A): 81.9697 -97.2090 -33.4842
REMARK	3	T TENSOR
REMARK	3	T11: -0.2480 T22: -0.3700
REMARK	3	T33: -0.1970 T12: 0.0271
REMARK	3	T13: 0.0948 T23: 0.1261
REMARK	3	L TENSOR
REMARK	3	L11: -0.0431 L22: -0.0280
REMARK	3	L33: -0.0440 L12: 0.0175
REMARK	3	L13: 0.0465 L23: 0.0381
REMARK	3	S TENSOR
REMARK	3	S11: -0.0226 S12: 0.7182 S13: -0.0052
REMARK	3	S21: -0.7057 S22: -0.0069 S23: 0.1273
REMARK	3	S31: 0.0400 S32: -0.2722 S33: 0.0000
REMARK	3	TLS GROUP : 4
REMARK	3	SELECTION: CHAIN D
REMARK	3	ORIGIN FOR THE GROUP (A): 116.5141 -77.7951 -33.2613
REMARK	3	T TENSOR
REMARK	3	T11: -0.3864 T22: -0.1961
REMARK	3	T33: -0.1626 T12: 0.0163
REMARK	3	T13: 0.0020 T23: 0.1413
REMARK	3	L TENSOR
REMARK	3	L11: -0.0288 L22: -0.0282
REMARK	3	L33: -0.0395 L12: 0.0063
REMARK	3	L13: -0.0122 L23: 0.0401
REMARK	3	S TENSOR
REMARK	3	S11: -0.0178 S12: 0.6303 S13: -0.0642
REMARK	3	S21: -0.7512 S22: -0.0110 S23: 0.0796
REMARK	3	S31: 0.1866 S32: -0.1247 S33: 0.0000
REMARK	3	
REMARK	3	NCS DETAILS
REMARK		NUMBER OF NCS GROUPS : 3
		NUMBER OF NCS GROUPS : 3 NCS GROUP : 1
REMARK	3	
REMARK	3	NCS OPERATOR : 1
REMARK	3	REFERENCE SELECTION: CHAIN A AND (RESSEQ 2:525)
REMARK	3	SELECTION : CHAIN B AND (RESSEQ 2:525)
REMARK		ATOM PAIRS NUMBER : 3856
REMARK		RMSD : 0.214
REMARK	3	NCS OPERATOR : 2

REMARK	3	REFERENCE SELECTION:		(DESSED 2.525 )
REMARK		SELECTION :		
REMARK		ATOM PAIRS NUMBER :		(RESSEQ 2.525 )
REMARK		RMSD :		
REMARK	3	NCS OPERATOR : 3	0.214	
REMARK	3	REFERENCE SELECTION:		(DESSED 2.525 )
REMARK	-	SELECTION :		
REMARK		ATOM PAIRS NUMBER :		(RESSEQ 2.525 )
REMARK			0.186	
REMARK	3	NCS OPERATOR : 4	0.100	
REMARK	3	REFERENCE SELECTION:	СНАТИ А АИО	(RESSED 2.525 )
REMARK		SELECTION :	CHAIN F AND	(RESSEQ 2.525)
REMARK		ATOM PAIRS NUMBER :		(110010 2:323 )
REMARK	3		0.213	
REMARK		NCS OPERATOR : 5	0.215	
REMARK		REFERENCE SELECTION:	CHAIN A AND	(RESSEO 2:525 )
REMARK	3	SELECTION :	CHAIN F AND	(RESSEO 2.525)
REMARK		ATOM PAIRS NUMBER :	3856	(110010 2:323 )
REMARK			0.226	
REMARK	3	NCS OPERATOR : 6	0.220	
REMARK		REFERENCE SELECTION:	СНАТИ А АИО	(RESSED 2.525 )
REMARK		SELECTION :		
REMARK	3	ATOM PAIRS NUMBER :		(100010 2:020 )
REMARK	3		0.214	
REMARK		NCS GROUP : 2	0.214	
REMARK		NCS OPERATOR : 1		
REMARK		REFERENCE SELECTION:	CHAIN H AND	(RESSEO 2:525 )
REMARK		SELECTION :		
REMARK	-	ATOM PAIRS NUMBER :		(110010 2:525 )
REMARK	3		0.224	
REMARK	3	NCS OPERATOR : 2	0.224	
REMARK	-	REFERENCE SELECTION:	СНАТИ Н АМО	(RESSEO 2.525 )
REMARK		SELECTION :	CHAIN J AND	(RESSEO 2:525)
REMARK		ATOM PAIRS NUMBER :		(100010 2:020 )
REMARK	3		0.231	
REMARK	3	NCS OPERATOR : 3	0.201	
REMARK		REFERENCE SELECTION:	CHAIN H AND	(RESSEO 2:525 )
REMARK	3	SELECTION :		
REMARK	3	ATOM PAIRS NUMBER :		(1110019 11010 )
REMARK			0.203	
REMARK	3	NCS OPERATOR : 4		
REMARK		REFERENCE SELECTION:	CHATN H AND	(RESSEO 2:525)
REMARK	3	SELECTION :		
REMARK REMARK	3	ATOM PAIRS NUMBER :		(
REMARK	3		0.215	
REMARK		NCS OPERATOR : 5		
REMARK		REFERENCE SELECTION:	CHAIN H AND	(RESSEO 2:525 )
REMARK				
REMARK	3	ATOM PAIRS NUMBER :		( _ /
REMARK REMARK	3		0.239	
REMARK		NCS OPERATOR : 6		
REMARK		REFERENCE SELECTION:	CHAIN H AND	(RESSEQ 2:525 )
REMARK		SELECTION :		. ,
REMARK		ATOM PAIRS NUMBER :		/
REMARK	3		0.227	
REMARK	3	NCS GROUP : 3		
REMARK	3	NCS OPERATOR : 1		
REMARK	3	REFERENCE SELECTION:	CHAIN O AND	(RESSEQ 1:97 )
				/

REMARK REMARK	-	SELECTION : ATOM PAIRS NUMBER :		Ρ	AND	(RESSEQ	1 <b>:</b> 97	)
REMARK	-	RMSD :						
REMARK		NCS OPERATOR : 2						
REMARK	3	REFERENCE SELECTION:	CHAIN	0	AND	(RESSEQ	1:97	)
REMARK		SELECTION :						
REMARK	3	ATOM PAIRS NUMBER :	728					
REMARK	3	RMSD :	0.211					
REMARK	3	NCS OPERATOR : 3						
REMARK	3	REFERENCE SELECTION:	CHAIN	0	AND	(RESSEQ	1:97	)
REMARK	3	SELECTION :	CHAIN	R	AND	(RESSEQ	1:97	)
REMARK	3	ATOM PAIRS NUMBER :	728					
REMARK	3	RMSD :	0.196					
REMARK	3	NCS OPERATOR : 4						
REMARK	3	REFERENCE SELECTION:	CHAIN	0	AND	(RESSEQ	1:97	)
REMARK	3	SELECTION :	CHAIN	s	AND	(RESSEQ	1:97	)
REMARK	3	ATOM PAIRS NUMBER :	728					
REMARK	3	RMSD :	0.200					
REMARK	3	NCS OPERATOR : 5						
REMARK	3	REFERENCE SELECTION:	CHAIN	0	AND	(RESSEQ	1:97	)
REMARK	3	SELECTION :	CHAIN	т	AND	(RESSEQ	1:97	)
REMARK	3	ATOM PAIRS NUMBER :						
REMARK	3	RMSD :	0.214					
REMARK	3	NCS OPERATOR : 6						
REMARK	3	REFERENCE SELECTION:						
REMARK	3	SELECTION :	CHAIN	U	AND	(RESSEQ	1:97	)
REMARK	3	ATOM PAIRS NUMBER :	728					
REMARK		RMSD :	0.205					
REMARK	3							
REMARK	3	OTHER REFINEMENT REMARKS	S: NULI	5				

# Template/example 2 (Xray/Neutron hybrid)

REMARK	3	REFINEMENT.
REMARK	3	PROGRAM : PHENIX (PHENIX.REFINE)
REMARK	3	AUTHORS : PAUL ADAMS, PAVEL AFONINE, VICENT CHEN, IAN
REMARK	3	: DAVIS, KRESHNA GOPAL, RALF GROSSE-
REMARK	3	: KUNSTLEVE,LI-WEI HUNG,ROBERT IMMORMINO,
REMARK	3	: TOM IOERGER, AIRLIE MCCOY, ERIK MCKEE, NIGEL
REMARK	3	: MORIARTY, REETAL PAI, RANDY READ, JANE
REMARK	3	: RICHARDSON, DAVID RICHARDSON, TOD ROMO, JIM
REMARK	3	: SACCHETTINI, NICHOLAS SAUTER, JACOB SMITH,
REMARK	3	: LAURENT STORONI, TOM TERWILLIGER, PETER
REMARK	3	: ZWART
REMARK	3	
REMARK	3	X-RAY DATA.
REMARK	3	
REMARK	3	REFINEMENT TARGET : ML
REMARK	3	
REMARK	3	DATA USED IN REFINEMENT.
REMARK	3	RESOLUTION RANGE HIGH (ANGSTROMS) : 1.75
REMARK	3	RESOLUTION RANGE LOW (ANGSTROMS) : 33.56
REMARK	3	MIN(FOBS/SIGMA_FOBS) : 1.330
REMARK	3	COMPLETENESS FOR RANGE (%) : 98.8
REMARK	3	NUMBER OF REFLECTIONS : 31524
REMARK	3	
REMARK	3	FIT TO DATA USED IN REFINEMENT.

REMARK	3	R VALUE (WORKING + TEST SET) : 0.132
REMARK	3	R VALUE (WORKING SET) : 0.129
REMARK	3	FREE R VALUE : 0.166
REMARK	3	FREE R VALUE TEST SET SIZE (%) : 9.360
REMARK	3	FREE R VALUE TEST SET COUNT : 2952
REMARK	3	
REMARK	3	FIT TO DATA USED IN REFINEMENT (IN BINS).
REMARK	3	BIN RESOLUTION RANGE COMPL. NWORK NFREE RWORK RFREE
REMARK	3	1 33.5691 - 4.8295 0.99 1424 130 14.0900 17.9800
REMARK	3	2 4.8295 - 3.8351 1.00 1420 125 10.3500 11.3600
REMARK	3	3 3.8351 - 3.3508 1.00 1378 129 10.5000 12.4600
REMARK	3	4 3.3508 - 3.0447 1.00 1413 122 11.7500 13.7300
REMARK	3	5 3.0447 - 2.8265 1.00 1385 138 11.9800 16.9100
REMARK	3	6 2.8265 - 2.6600 1.00 1384 131 12.7800 16.5000
REMARK	3	7 2.6600 - 2.5268 1.00 1351 162 12.1800 17.9600
REMARK	3	8 2.5268 - 2.4168 1.00 1394 137 12.3800 15.7200
REMARK	3	9 2.4168 - 2.3238 1.00 1381 130 12.3100 15.4300
REMARK	3	10 2.3238 - 2.2437 1.00 1385 142 12.5900 17.5500
REMARK	3	11 2.2437 - 2.1735 1.00 1368 123 12.0600 18.7300
REMARK	3	12 2.1735 - 2.1114 1.00 1377 143 11.8600 15.5300
REMARK	3	13 2.1114 - 2.0558 1.00 1337 171 11.3000 16.9600
REMARK	3	14 2.0558 - 2.0057 1.00 1353 143 11.9200 15.9400
REMARK	3	15 2.0057 - 1.9601 0.99 1386 144 12.2000 16.0500
REMARK	3	16 1.9601 - 1.9184 1.00 1340 136 12.9800 19.2100
REMARK	3	17 1.9184 - 1.8800 1.00 1380 142 13.7700 21.1700
REMARK	3	18 1.8800 - 1.8445 1.00 1371 141 14.1400 18.2000
REMARK	3	19 1.8445 - 1.8116 1.00 1344 172 15.5900 22.0300
REMARK	3	20 1.8116 - 1.7809 0.99 1321 158 16.7800 22.1500
REMARK	3	21 1.7809 - 1.7522 0.79 1080 133 20.5300 25.6900
REMARK	3	
REMARK	3	BULK SOLVENT MODELLING.
REMARK	3	METHOD USED : FLAT BULK SOLVENT MODEL
REMARK	3	SOLVENT RADIUS : 1.11
REMARK	3	SHRINKAGE RADIUS : 0.90
REMARK	3	K_SOL : 0.35
REMARK	3	B_SOL : 29.39
REMARK	3	
REMARK	3	ERROR ESTIMATES.
REMARK	3	COORDINATE ERROR (MAXIMUM-LIKELIHOOD BASED) : 0.190
REMARK	3	PHASE ERROR (DEGREES, MAXIMUM-LIKELIHOOD BASED) : 15.390
REMARK	3	
REMARK	3	B VALUES.
REMARK	3	FROM WILSON PLOT (A**2) : 17.52
REMARK	3	MEAN B VALUE (OVERALL, A**2) : NULL
REMARK	3	OVERALL ANISOTROPIC B VALUE.
REMARK	3	B11 (A**2) : 5.33780
REMARK	3	B22 (A**2) : 5.45600
REMARK	3	B33 (A**2) : 4.24070
REMARK	3	B12 (A**2) : 0.00000
REMARK	3	B13 (A**2) : 0.43090
REMARK	3	B23 (A**2) : 0.00000
REMARK	3	
REMARK	3	TWINNING INFORMATION.
REMARK	3	FRACTION: NULL
REMARK	3	OPERATOR: NULL
REMARK	3	
REMARK		DEVIATIONS FROM IDEAL VALUES.
REMARK	3	RMSD COUNT

3

BOND : ANGLE : REMARK NULL 3 REMARK NULL NULL CHIRALITY : REMARK 3 NULL NULL REMARK 3 PLANARITY : NULL NULL REMARK 3 DIHEDRAL : NULL NULL REMARK 3 REMARK 3 TLS DETAILS 3 NUMBER OF TLS GROUPS : NULL REMARK REMARK 3 **3 NCS DETAILS** REMARK REMARK 3 NUMBER OF NCS GROUPS : NULL REMARK 3 3 OTHER REFINEMENT REMARKS: NULL REMARK REMARK 3 3 NEUTRON DATA. REMARK REMARK 3 3 REMARK REFINEMENT TARGET : ML 3 REMARK REMARK 3 DATA USED IN REFINEMENT. REMARK 3 RESOLUTION RANGE HIGH (ANGSTROMS) : 2.19 REMARK 3 RESOLUTION RANGE LOW (ANGSTROMS) : 40.11 : 1.530 REMARK 3 MIN(FOBS/SIGMA FOBS) 3 COMPLETENESS FOR RANGE (%) : 72.8 REMARK 3 NUMBER OF REFLECTIONS : 11884 REMARK REMARK 3 REMARK 3 FIT TO DATA USED IN REFINEMENT. (WORKING + TEST SET) : 0.260 3 R VALUE REMARK REMARK 3 R VALUE (WORKING SET) : 0.257 : 0.291 REMARK 3 FREE R VALUE REMARK 3 FREE R VALUE TEST SET SIZE (%): 8.350 REMARK 3 FREE R VALUE TEST SET COUNT : 992 REMARK 3 REMARK 3 FIT TO DATA USED IN REFINEMENT (IN BINS). REMARK 3 BIN RESOLUTION RANGE COMPL. NWORK NFREE RWORK RFREE 

 BIN
 RESOLUTION
 RANGE
 COMPL.
 NWORK
 NFREE
 RWORK
 RFREE

 1
 40.1164
 4.1959
 0.93
 2018
 176
 19.7800
 20.5900

 2
 4.1959
 3.3309
 0.89
 1915
 175
 21.1000
 25.3500

 3
 3.3309
 2.9100
 0.78
 1669
 152
 25.7100
 30.8200

 4
 2.9100
 2.6440
 0.70
 1475
 141
 26.7100
 30.7500

 5
 2.6440
 2.4545
 0.64
 1350
 133
 28.7700
 33.5100

 6
 2.4545
 2.3098
 0.60
 1269
 117
 30.9500
 33.6400

 7
 2.3098
 2.1942
 0.56
 1196
 98
 32.3200
 36.3200

 REMARK 3 1 40.1164 - 4.1959 REMARK 3 BULK SOLVENT MODELLING. REMARK 3 METHOD USED : FLAT BULK SOLVENT MODEL : 1.11 3 SOLVENT RADIUS REMARK 3 SHRINKAGE RADIUS : 0.90 REMARK 3 K\_SOL : 0.53 REMARK 3 B\_SOL : 82.90 REMARK REMARK 3 REMARK 3 ERROR ESTIMATES. 3 COORDINATE ERROR (MAXIMUM-LIKELIHOOD BASED) : 0.450 REMARK 3 PHASE ERROR (DEGREES, MAXIMUM-LIKELIHOOD BASED) : 26.920 REMARK REMARK 3 REMARK 3 B VALUES. (A\*\*2) : NULL REMARK 3 FROM WILSON PLOT REMARK 3 MEAN B VALUE (OVERALL, A\*\*2) : NULL 3 OVERALL ANISOTROPIC B VALUE. REMARK 3 B11 (A\*\*2) : -0.70340 REMARK

NUT.T.

REMARK	3	B22 (A**2) : -6.62540
REMARK	3	B33 (A**2) : -7.07190
REMARK	3	B12 (A**2) : -0.00000
REMARK	3	B13 (A**2) : -1.25130
REMARK	3	B23 (A**2) : -0.00000
REMARK	3	
REMARK	3	TWINNING INFORMATION.
REMARK	3	FRACTION: NULL
REMARK	3	OPERATOR: NULL
REMARK	3	
REMARK	3	DEVIATIONS FROM IDEAL VALUES.
REMARK	3	RMSD COUNT
REMARK	3	BOND : 0.018 5216
REMARK	3	ANGLE : 1.759 9458
REMARK	3	CHIRALITY : 0.160 392
REMARK	3	PLANARITY : 0.011 779
REMARK	3	DIHEDRAL : 25.100 1363
REMARK	3	
REMARK	3	TLS DETAILS
REMARK	3	NUMBER OF TLS GROUPS : NULL
REMARK	3	
REMARK	3	NCS DETAILS
REMARK	3	NUMBER OF NCS GROUPS : NULL
REMARK	3	
REMARK	3	OTHER REFINEMENT REMARKS: NULL

# Refinement using BUSTER-TNT

# Template/example

3

REMARK

KEPIAKK	5	
REMARK	3	REFINEMENT.
REMARK		PROGRAM : BUSTER-TNT 2.7.0
REMARK	3	AUTHORS : BRICOGNE, BLANC, BRANDL, FLENSBURG, KELLER,
REMARK	3	: PACIOREK, ROVERSI, SMART, VONRHEIN, WOMACK;
REMARK	3	: MATTHEWS, TEN EYCK, TRONRUD
REMARK	3	
REMARK	3	DATA USED IN REFINEMENT.
REMARK		RESOLUTION RANGE HIGH (ANGSTROMS) : 3.00
REMARK	3	
REMARK	3	
REMARK		COMPLETENESS FOR RANGE (%): 99.10
REMARK		COMPLETENESS FOR RANGE(%): 99.10NUMBER OF REFLECTIONS: 26306
REMARK		
REMARK		FIT TO DATA USED IN REFINEMENT.
REMARK		CROSS-VALIDATION METHOD : THROUGHOUT
REMARK		FREE R VALUE TEST SET SELECTION : RANDOM
REMARK		
REMARK		R VALUE(WORKING + TEST SET) : 0.1816R VALUE(WORKING SET) : 0.1750FREE R VALUE: 0.2442FREE R VALUE(%) : 0.64
REMARK		FREE R VALUE : 0.2442
REMARK	3	FREE R VALUE TEST SET SIZE (%) : 9.64
REMARK		
REMARK		ESTIMATED ERROR OF FREE R VALUE : NULL
REMARK		
REMARK		FIT IN THE HIGHEST RESOLUTION BIN.
REMARK		
REMARK		TOTAL NUMBER OF BINS USED : 13 BIN RESOLUTION RANGE HIGH (ANGSTROMS) : 3.00
REMARK	3	
REMARK		
REMARK		REFLECTIONS IN BIN (WORKING + TEST SET) : 2852
REMARK		BIN R VALUE (WORKING + TEST SET) $\cdot$ 0.2262
REMARK		REFLECTIONS IN BIN(WORKING SET) : 2544BIN R VALUE(WORKING SET) : 0.2148BIN FREE R VALUE: 0.3255
REMARK		BIN R VALUE (WORKING SET) : 0.2148
REMARK		BIN FREE R VALUE : 0.3255
REMARK	3	BIN FREE R VALUE TEST SET SIZE (%): 10.80
REMARK		BIN FREE R VALUE TEST SET COUNT : 308
REMARK		ESTIMATED ERROR OF BIN FREE R VALUE : NULL
REMARK	3	
REMARK		NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
REMARK	3	PROTEIN ATOMS : 7507
REMARK		NUCLEIC ACID ATOMS : 0
REMARK		
REMARK	-	
REMARK		
REMARK		B VALUES.
REMARK		FROM WILSON PLOT (A**2) : 71.99
REMARK		
REMARK		
REMARK		B11 (A**2) : 0.1789
REMARK		B22 (A**2) : 0.1789
REMARK		B22 (A**2) : 0.1789 B33 (A**2) : -0.3578
REMARK		B12 (A**2) : 0.0000
REMARK		B12 (A + 2) : 0.0000
	Ŭ	(·· <b>_</b> ) · · · · · · · · · · · · · · · · · · ·

3 B23 (A\*\*2) : 0.0000 REMARK REMARK 3 3 ESTIMATED COORDINATE ERROR. REMARK REMARK3DPI (BLOW EQ-10) BASED ON R VALUE(A) : 0.280REMARK3DPI (BLOW EQ-9) BASED ON FREE R VALUE(A) : 0.354REMARK3DPI (CRUICKSHANK) BASED ON R VALUE(A) : 0.216REMARK3DPI (CRUICKSHANK) BASED ON R VALUE(A) : 0.304REMARK3DPI (CRUICKSHANK) BASED ON R VALUE(A) : 0.304 (A) : 0.280 REMARK 3 ESD FROM LUZZATI PLOT DPI (CRUICKSHANK) BASED ON FREE R VALUE (A) : 0.209 REMARK 3 REMARK 3 3 REFERENCES: BLOW, D. (2002) ACTA CRYST D58, 792-797 REMARK CRUICKSHANK, D.W.J. (1999) ACTA CRYST D55, 583-601 REMARK 3 REMARK 3 3 CORRELATION COEFFICIENTS. REMARK CORRELATION COEFFICIENT FO-FC : NULL REMARK 3 3 CORRELATION COEFFICIENT FO-FC FREE : NULL REMARK REMARK 3 3 NUMBER OF GEOMETRIC FUNCTION TERMS DEFINED : 15 REMARK COUNT WEIGHT REMARK 3 TERM FUNCTION. : 7680 ; 2.00 ; HARMONIC REMARK 3 BOND LENGTHS REMARK 3 BOND ANGLES : 10408 ; 2.00 ; HARMONIC REMARK 3 TORSION ANGLES : 1732 ; 2.00 ; SINUSOIDAL REMARK3TRIGONAL CARBON PLANES:198;2.00;HARMONICREMARK3GENERAL PLANES:1079;5.00;HARMONIC 3 GENERAL PLANES REMARK3GENERAL PLANES:1079 ;5.00 ;HARMONICREMARK3ISOTROPIC THERMAL FACTORS :7680 ;20.00 ;HARMONICREMARK3BAD NON-BONDED CONTACTS :NULL ;NULL ;NULL ;REMARK3IMPROPER TORSIONS :NULL ;NULL ;NULL ;REMARK3PSEUDOROTATION ANGLES :NULL ;NULL ;NULL ;REMARK3CHIRAL IMPROPER TORSION :990 ;5.00 ;SEMIHARMONICREMARK3SUM OF OCCUPANCIES :NULL ;NULL ;NULL ;REMARK3UTILITY DISTANCES :NULL ;NULL ;NULL ; REMARK 3 UTILITY DISTANCES : NULL ; NUL ; NULL ; NULL ; NU : NULL ; NULL ; NULL REMARK 3 IDEAL-DIST CONTACT TERM : 9135 ; 4.00 ; SEMIHARMONIC REMARK 3 REMARK 3 RMS DEVIATIONS FROM IDEAL VALUES. REMARK 3 BOND LENGTHS (A) : 0.012 BOND ANGLES(DEGREES) :1.06PEPTIDE OMEGA TORSION ANGLES (DEGREES) :4.34 REMARK 3 3 REMARK (DEGREES) : 26.08 3 REMARK OTHER TORSION ANGLES REMARK 3 REMARK 3 TLS DETAILS. REMARK 3 NUMBER OF TLS GROUPS : 3 REMARK 3 TLS GROUP : 1 REMARK 3 SELECTION: (alphanumerical text here, { or | not allowed) 3 ORIGIN FOR THE GROUP (A): 77.0443 19.4729 -5.2350 REMARK 3 T TENSOR REMARK T11: 0.0000 T22: T33: 0.0000 T12: T13: 0.0000 T23. 3 0.0000 REMARK 0.0000 T12: 0.0000 3 REMARK 0.0000 T23: 0.0000 REMARK 3 REMARK 3 L TENSOR REMARK 3 L11: 0.1106 L22: 1.0233 
 REMARK
 3
 L33:
 0.6541
 L12:
 -0.4855

 REMARK
 3
 L13:
 0.0884
 L23:
 -0.7511
 REMARK 3 S TENSOR 
 REMARK
 3
 S11:
 -0.0061
 S12:
 -0.0010
 S13:
 0.0041

 REMARK
 3
 S21:
 0.0098
 S22:
 0.0044
 S23:
 -0.0029

 REMARK
 3
 S31:
 -0.0016
 S32:
 0.0018
 S33:
 0.0017

• • •

REMARK	3	OTHER REFINEMENT REMARKS: A MET-INHIBITION PROTOCOL WAS USED FOR
REMARK	3	SELENOMETHIONINE INCORPORATION DURING PROTEIN EXPRESSION. THE
REMARK	3	OCCUPANCY OF THE SE ATOMS IN THE MSE RESIDUES WAS REDUCED TO
REMARK	3	0.75 FOR THE REDUCED SCATTERING POWER DUE TO PARTIAL S-MET
REMARK	3	INCORPORATION.
REMARK	3	IDEAL-DIST CONTACT TERM CONTACT SETUP. RESIDUE TYPES WITHOUT
REMARK	3	CCP4 ATOM TYPE IN LIBRARY=CSS PQA. NUMBER OF ATOMS WITH PROPER
REMARK	3	CCP4 ATOM TYPE=9.
REMARK	3	NUMBER WITH APPROX DEFAULT CCP4 ATOM TYPE=34. NUMBER TREATED BY
REMARK	3	BAD NON-BONDED CONTACTS=1.

Refinement using Electron Microscopy

#### Template/Example

3

REMARK

REMARK 3 REFINEMENT REMARK 3 SOFTWARE PACKAGES : SIMPLEX, PYPFT, EMFIT, O, XPLOR REMARK 3 RECONSTRUCTION SCHEMA : ICOSAHEDRAL REMARK 3 REMARK 3 EM MAP-MODEL FITTING AND REFINEMENT REMARK 3 PDB ENTRY : PDB ID 1HX6 REMARK 3 REFINEMENT SPACE : RECIPROCAL REMARK 3 REFINEMENT PROTOCOL : RIGID BODY REFINEMENT REMARK 3 REMARK 3 REFINEMENT TARGET : R-FACTOR OVERALL ANISOTROPIC B VALUE : NULL REMARK 3 REMARK 3 FITTING PROCEDURE : THE CRYSTAL STRUCTURE OF THE MAJOR COAT REMARK 3 PROTEIN P3 (PDB FILE 1HX6) WAS PLACED INTO THE CRYO-EM REMARK 3 DENSITY MAP. THE CAPSID PROTEIN WAS FIRST MANUALLY REMARK 3 POSITIONED INTO THE CRYO-EM DENSITY CORRESPONDING TO REMARK 3 POSITIONS OF THE FOUR INDEPENDENT TRIMERS IN THE REMARK 3 ICOSAHEDRAL ASYMMETRIC UNIT. THESE POSITIONS WERE THEN REMARK 3 REFINED BY RIGID BODY REFINEMENT IN RECIPROCAL SPACE WITH REMARK 3 THE PROGRAM XPLOR. REMARK 3 QUALITY OF THE FIT R-FACTOR= 0.339, CROSS-CORRELATION REMARK 3 COEFFICIENT 0.915, ATOMS OUTSIDE DENSITY PER ICOSAHEDRAL REMARK 3 ASYMMETRIC UNIT 527 (1.5%), ATOM CLASHES PER ICOSAHEDRAL REMARK 3 ASYMMETRIC UNIT 115 (0.3%) REMARK 3 REMARK 3 EM IMAGE RECONSTRUCTION STATISTICS REMARK 3 NOMINAL PIXEL SIZE (ANGSTROMS) : 3.68 REMARK 3 ACTUAL PIXEL SIZE (ANGSTROMS) : 3.44 REMARK 3 EFFECTIVE RESOLUTION (ANGSTROMS) : 14.0 REMARK 3 NUMBER OF PARTICLES : 1800 REMARK 3 CTF CORRECTION METHOD : NULL 3 REMARK REMARK 3 EM RECONSTRUCTION MAGNIFICATION CALIBRATION: THE PIXEL REMARK 3 SIZE OF THE CRYO-EM MAP WAS OBTAINED USING THE X-RAY REMARK 3 STRUCTURE OF THE P3 TRIMER AS A REFERENCE. AFTER AN INITIAL REMARK 3 FITTING USING THE NOMINAL PIXEL SIZE, THE P3 TRIMERS IN THE REMARK 3 ICOSAHEDRAL ASYMMETRIC UNIT WERE GRADUALLY TRANSLATED TOWARDS REMARK 3 THE CENTER OF THE PARTICLE UNTIL THE CRYSTALLOGRAPHIC R-FACTOR REMARK 3 WAS MINIMISED. 3 REMARK 3 OTHER DETAILS: THE ORIENTATIONS WERE REFINED BY THE CROSS REMARK 3 COMMON LINES LINES METHOD (SIMPLEX) AND THE POLAR FOURIER REMARK REMARK 3 TRANSFORM METHOD. MODEL-BASED, POLAR-FOURIER-IRANSFORM REMARK 3 (FULLER ET AL. 1996, J.STRUC.BIOL. 116, 48-55; BAKER AND REMARK 3 CHENG, 1996, J.STRUC.BIOL. 116, 120-130) MODEL-BASED CROSS REMARK 3 NATURE (LONDON) 226, 421-425; FULLER ET AL. 1996, REMARK 3 J.STRUC.BIOL. 116, 48-55; FERLENGHI ET AL. 1998, J.MOL.BIOL. REMARK 3 283, 71-81). THE EFFECTIVE RESOLUTION OF THE FINAL REMARK 3 RECONSTRUCTED DENSITY WAS DETERMINED TO BE AT LEAST 25 REMARK 3 ANGSTROMS, AS MEASURED BY RANDOMLY SPLITTING THE PARTICLES REMARK 3 INTO TWO SETS AND CALCULATING THE FOURIER SHELL CORRELATION 3 OBTAINED FROM SEPARATE RECONSTRUCTIONS (HARAUZ AND VAN HEEL REMARK

REMARK	3	1986, OPTIK 73, 146-156). THE EIGENVALUE SPECTRUM GAVE AN
REMARK	3	INDICATION OF THE RANDOMNESS OF THE DATA THAT WAS INCLUDED
REMARK	3	IN THE RECONSTRUCTION. THE COMPLETENESS OF THE DATA WAS
REMARK	3	VERIFIED IN THAT ALL EIGENVALUES EXCEEDED 100. THE COORDINATES
REMARK	3	ARE IN THE P, Q, R FRAME IN ANGSTROM UNITS AND CORRESPOND
REMARK	3	TO ICOSAHEDRAL SYMMETRY AXES. THE ORIGIN IS CHOSEN AT THE
REMARK	3	CENTER OF THE VIRUS WITH P, Q AND R ALONG MUTUALLY
REMARK	3	PERPENDICULAR TWO-FOLD AXES OF THE ICOSAHEDRON. THEY SHOULD
REMARK	3	REMAIN IN THAT FRAME FOR THE EASE OF THE USER IN CREATING
REMARK	3	THE BIOLOGICALLY SIGNIFICANT VIRAL COMPLEX PARTICLE USING
REMARK	3	THE 60 ICOSAHEDRAL SYMMETRY OPERATORS. RESIDUES NOT VISIBLE
REMARK	3	IN THE ORIGINAL CRYSTAL STRUCTURES ARE NOT INCLUDED IN THE
REMARK	3	CRYO-EM STRUCTURE MODEL.

## **Example for Solution Scattering**

```
REMARK3REMARK3REFINEMENT.REMARK3PROGRAM: INSIGHT II 98.0REMARK3AUTHORS: MSIREMARK3NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.REMARK3PROTEIN ATOMS: 1213REMARK3NUCLEIC ACID ATOMS: 0REMARK3HETEROGEN ATOMS: 0REMARK3SOLVENT ATOMS: 0REMARK3SOLVENT ATOMS: 0REMARK3OTHER REFINEMENT REMARKS: DISCOVER WAS USED FOR ENERGYREMARK3MINIMISATION
```

## **Non-diffraction studies**

Until standard refinement remarks are adopted for non-diffraction studies, refinement details will appear in REMARK 3 formatted in free text, beginning on the sixth line of the remark.

#### Template

```
2
                3
                     4
                           5
                                6
                                      7
     1
                                            8
REMARK 3
REMARK 3 REFINEMENT.
REMARK 3 PROGRAM
             :
REMARK 3
REMARK 3
      AUTHORS
             :
REMARK 3 FREE TEXT
```

#### Example

REMARK	3	
REMARK	3	REFINEMENT.
REMARK	3	PROGRAM : CNSSOLVE 1.1, X-PLOR 2.11.2, PROCHECK NMR 3.51,
REMARK	3	MOLPROBITY 3.01, QUEEN 1.1, PSVS 1.3
REMARK	3	AUTHORS : BRUNGER, ET. AL. (CNSSOLVE), CLORE ET. AL. (X-
REMARK	3	PLOR), LASKOWSKI, MACARTHUR (PROCHECK NMR),
REMARK	3	LOVELL, RICHARDSON ET. AL. (MOLPROBITY),
REMARK	3	NABUURS, VUISTER (QUEEN), BHATTACHARYA,
REMARK	3	MONTELIONE (PSVS)
REMARK	3	
REMARK	3	OTHER REFINEMENT REMARKS: NOESY ASSIGNMENT MADE WITH ITERATIVE
REMARK	3	METHOD USING CNS, HYPER (DIHEDRAL) AND DYANA FOLLOWED BY NIH-
REMARK	3	XPLOR FOR SIMMULATED ANNEALING MD. CONVERGED STRUCTURES WERE
REMARK	3	FURTHER MINIMIZED USING CNS IN EXPLICIT H2O SHELL (NILGES
REMARK	3	PROTOCOL). FULL LENGTH SEQUENCE WAS CARRIED THROUGH THE
REMARK	3	REFINEMENT PROTOCOL. COORDINATES FROM DISORDERED REGIONS,
REMARK	3	INCLUDING HEXHIS TAG, WERE NOT REPORTED. STRUCTURE IS BASED ON
REMARK	3	439 CONSTRAINTS (216 LONG RANGE), 43 DIHEDRAL AND 20 H-BOND.

## **REMARK 4 (updated), Format**

Remark 4 indicates the version of the PDB File Format used to generate the file.

#### Template

 1
 2
 3
 4
 5
 6
 7
 8

 12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890
 REMARK
 4
 XXXX COMPLIES WITH FORMAT V. N.MM, DD-MMM-YY
 8

XXXX refers to the ID code of the entry.

N.MM refers to the version number. The current version is 3.20. DD-MMM-YY refers to the release date of that version of the format. DD is the day of the month (a number 01 through 31), MMM is the English 3-letter abbreviation for the month, and YY is the year.

#### Example

REMARK 4 REMARK 4 1ABC COMPLIES WITH FORMAT V. 3.20, 01-DEC-08

## **REMARK 5 (updated), Obsolete Statement**

This REMARK describes the reason for structure obsolete in case that the structure is incorrect and the author obsoletes the entry without new coordinates.

## REMARKs 6 - 99

#### **Overview**

REMARKs following the format REMARK 4 consist of free text annotation, pre-defined templates, and token: value pair-styled templates. Presented here are examples of REMARK sections.

### **Record Format and Details**

- \* As with all other REMARKs, the first line of each REMARK is empty and is used as a spacer.
- \* REMARKs 6-99 are no longer for use of free text annotation.

## **REMARK 100 (updated), Deposition or Processing Site**

This REMARK indicates PDB process site: RCSB, PDBe, PDBj or BNL. This remark also contains process date and site id code with exception of BNL entries.

#### Template

1 2 3 4 5 6 7 8 1234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 100 THIS ENTRY HAS BEEN PROCESSED BY XXXX on DD-MMM-YY. REMARK 100 THE XXXX ID CODE IS VVVVVVVV.

XXXX is the process site. VVVVVVVVV is the site id code. The format of the date in this remark is DD-MMM-YY. DD is the day of the month (a number 01 through 31), MMM is the English 3-letter abbreviation for the month, and YY is the year.

#### **Examples**

REMARK 100 REMARK 100 THIS ENTRY HAS BEEN PROCESSED BY RCSB on 10-MAR-06. REMARK 100 THE RCSB ID CODE IS RCSB036809. REMARK 100 THIS ENTRY HAS BEEN PROCESSED BY PDBE on 13-FEB-07. REMARK 100 THE PDBE ID CODE IS EBI-28843. REMARK 100 THIS ENTRY HAS BEEN PROCESSED BY PDBJ on 21-MAR-05. REMARK 100 THE RCSB ID CODE IS RCSB026278. REMARK 100 THIS ENTRY HAS BEEN PROCESSED BY BNL.

## **REMARKs 200-265, Experimental Details**

REMARKs in this range present the data collection details for the data which resulted in the refinement statistics of REMARK 3. They provide information on the structure determination experiment, which may have been done by diffraction, NMR or some other technique.

The "NULL" value will be used if the data for a token is not supplied by the depositor.

## **REMARK 200 (updated), X-ray Diffraction Experimental Details**

REMARK 200 is mandatory if single crystal, fiber, or polycrystalline X-ray diffraction experiments were performed. The format of date in this remark is DD-MMM-YY. DD is the day of the month (a number 01 through 31), MMM is the English 3-letter abbreviation for the month, and YY is the year.

#### Template

100150	1	2 3 4					7		8
123456 REMARK		12345678901234567890123456789012	34:	5678901	234567	8901234	4567890	123456	57890
		EXPERIMENTAL DETAILS							
				X-RAY	DIFFRA	CTTON			
REMARK	200	DATE OF DATA COLLECTION	:	71 IUII	DIIIMI	CIION			
REMARK	200	TEMPERATURE (KELVIN)	:						
REMARK		· · · · · · · · · · · · · · · · · · ·	:						
REMARK	200	NUMBER OF CRYSTALS USED	:						
REMARK	200								
REMARK	200	SYNCHROTRON (Y/N)	:						
REMARK	200	RADIATION SOURCE	:						
REMARK	200	BEAMLINE	:						
REMARK	200		:						
REMARK									
REMARK			:						
REMARK			:						
REMARK		OPTICS	:						
REMARK									
REMARK		DETECTOR TYPE	:						
REMARK			:						
REMARK									
REMARK		DATA SCALING SOFTWARE	:						
REMARK REMARK		NUMBER OF UNIQUE REFLECTIONS							
REMARK									
REMARK									
REMARK									
REMARK			•						
		OVERALL.							
		COMPLETENESS FOR RANGE (%)	:						
		DATA REDUNDANCY	:						
REMARK	200	R MERGE (I)	:						
REMARK	200								
REMARK	200	<i sigma(i)=""> FOR THE DATA SET</i>	:						
REMARK	200								
REMARK	200	IN THE HIGHEST RESOLUTION SHELL	•						

REMARK 200 HIGHEST RESOLUTION SHELL, RANGE HIGH (A) : REMARK 200 HIGHEST RESOLUTION SHELL, RANGE LOW (A) : REMARK 200 COMPLETENESS FOR SHELL REMARK 200 DATA REDUNDANCY IN SHELL (%): : REMARK 200 R MERGE FOR SHELL (I) : (I) : REMARK 200 R SYM FOR SHELL REMARK 200 <I/SIGMA(I)> FOR SHELL : REMARK 200 REMARK 200 METHOD USED TO DETERMINE THE STRUCTURE: **REMARK 200 SOFTWARE USED:** REMARK 200 STARTING MODEL: REMARK 200 REMARK 200 REMARK:

#### Examples

The following **example** illustrates the how REMARK 200 will be used in cases in which multiple data collections are described. In this example, data items corresponding to different data collection sessions are separated by semi-colons. Multiple data values within a single session (e.g. wavelength) are separated by commas.

REMARK				
REMARK	200	EXPERIMENTAL DETAILS		
REMARK	200	EXPERIMENT TYPE DATE OF DATA COLLECTION	:	X-RAY DIFFRACTION
REMARK	200	DATE OF DATA COLLECTION	:	17-MAR-02; 17-MAR-02
REMARK	200	TEMPERATURE (KELVIN)	:	100; 100
REPARK	200	1 11	:	8.00
REMARK		NUMBER OF CRYSTALS USED	:	2
REMARK	200			
REMARK	200	SYNCHROTRON (Y/N) RADIATION SOURCE BEAMLINE X-RAY GENERATOR MODEL MONOCHROMATIC OR LAUE (M/L) WAVELENGTH OR RANGE (A)	:	Y; Y
REMARK	200	RADIATION SOURCE	:	APS ; APS
REMARK	200	BEAMLINE	:	17ID; 17ID
REMARK	200	X-RAY GENERATOR MODEL	:	NULL
REMARK	200	MONOCHROMATIC OR LAUE (M/L)	:	M; M
TODI II II (II (	200	WAVELENGTH OR RANGE (A) MONOCHROMATOR OPTICS	:	1.5545; 1.0720, 1.0723,
REMARK				1.0543
REMARK		MONOCHROMATOR	:	SI (111); SI (111)
REMARK	200	OPTICS	:	NULL
REMARK	200	DETECTOR TYPE DETECTOR MANUFACTURER		
REMARK	200	DETECTOR TYPE	:	CCD; CCD
REMARK	200	DETECTOR MANUFACTURER	:	ADSC QUANTUM 210; ADSC
REMARK	200			QUANTUM 210
REMARK	200	INTENSITI-INTEGRATION SOFTWARE	÷	DENZO
REMARK	200	DATA SCALING SOFTWARE	:	HKL
REMARK	200			
		NUMBER OF UNIQUE REFLECTIONS		
REMARK	200	RESOLUTION RANGE HIGH (A)	:	1.900
		RESOLUTION RANGE LOW (A)		
REMARK		REJECTION CRITERIA (SIGMA(I))	:	0.000
REMARK				
REMARK	200	OVERALL.		
REMARK	200	COMPLETENESS FOR RANGE(%)DATA REDUNDANCY(I)R MERGE(I)R SYM(I)	:	98.3
REMARK	200	DATA REDUNDANCY	:	19.800
REMARK	200	R MERGE (I)	:	NULL
REMARK	200	R SYM (I)	:	0.07500
REMARK	200	<i sigma(i)=""> FOR THE DATA SET</i>	:	17.0000

REMARK 200 REMARK 200 IN THE HIGHEST RESOLUTION SHELL. REMARK 200 IN THE HIGHEST RESOLUTION SHELL, RANGE HIGH (A) : 1.90 REMARK 200 HIGHEST RESOLUTION SHELL, RANGE LOW (A) : 1.97 REMARK 200 COMPLETENESS FOR SHELL (%) : 83.4 REMARK 200 DATA REDUNDANCY IN SHELL : 3.00 : 3.00 REMARK 200 R MERGE FOR SHELL (I) : NULL REMARK 200 R SYM FOR SHELL (I) : 0.65000 REMARK 200 <1/SIGMA(I)> FOR SHELL : 1.500 REMARK 200 REMARK 200 DIFFRACTION PROTOCOL: SINGLE WAVELENGTH; MAD REMARK 200 METHOD USED TO DETERMINE THE STRUCTURE: MAD REMARK 200 SOFTWARE USED: SOLVE 2.02 REMARK 200 STARTING MODEL: NULL REMARK 200 REMARK 200 REMARK: NULL

# **REMARK 205, Fiber Diffraction, Fiber Sample Experiment Details**

REMARK 205 is mandatory if data was obtained from a fiber diffraction - non-crystalline sample study

### Template

1 2 3 4 5 6 7 8 12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 205 REMARK 205 THESE COORDINATES WERE GENERATED FROM FIBER DIFFRACTION REMARK 205 DATA. PROTEIN DATA BANK CONVENTIONS REQUIRE THAT CRYST1 REMARK 205 AND SCALE RECORDS BE INCLUDED, BUT THE VALUES OF THESE REMARK 205 RECORDS ARE MEANINGLESS.

# **REMARKs 210 and 215/217, NMR Experiment Details**

Remark 210 is mandatory if data was obtained from an NMR experiment.

### Template

1	2 3 4		5	6	7 8
1234567890	1234567890123456789012345678901	2345678	9012345678	9012345678	901234567890
REMARK 210					
REMARK 210	EXPERIMENTAL DETAILS				
REMARK 210	EXPERIMENT TYPE	:			
REMARK 210	TEMPERATURE (KELVIN	) :			
REMARK 210	PH	:			
REMARK 210	IONIC STRENGTH	:			
REMARK 210	PRESSURE	:			
	SAMPLE CONTENTS	:			
REMARK 210					
REMARK 210		:			
	SPECTROMETER FIELD STRENGTH	:			
REMARK 210		:			
	SPECTROMETER MANUFACTURER	:			
REMARK 210					
	STRUCTURE DETERMINATION.				
	SOFTWARE USED	:			
	METHOD USED	:			
REMARK 210					
	CONFORMERS, NUMBER CALCULATED	:			
	CONFORMERS, NUMBER SUBMITTED	:			
	CONFORMERS, SELECTION CRITERIA	:			
REMARK 210					
REMARK 210					
	BEST REPRESENTATIVE CONFORMER	IN THIS	ENSEMBLE	:	
REMARK 210					
REMARK 210	REMARK:				

### Example

REMARK 210 REMARK 210 EXPERIMENTAL DETAILS REMARK 210 EXPERIMENT TYPE : NMR REMARK 210 TEMPERATURE (KELVIN) : 293 : 7.0 REMARK 210 PH REMARK 210 IONIC STRENGTH : NULL REMARK 210 PRESSURE : AMBIENT REMARK 210 SAMPLE CONTENTS : 4.0 MM PHYLLOSEPTIN-2, REMARK 210 TRIFLUOROETHANOL/WATER (60%/ REMARK 210 40%) REMARK 210 REMARK 210 NMR EXPERIMENTS CONDUCTED : 2D 1H-1H TOCSY, 2D 1H-1H REMARK 210 NOESY, 2D 1H-13C HSQC, 2D 1H-REMARK 210 15N HSQC REMARK 210SPECTROMETER FIELD STRENGTH: 600; 800REMARK 210SPECTROMETER MODEL: DMX600; DRX800REMARK 210SPECTROMETER MANUFACTURER: BRUKER REMARK 210 REMARK 210 STRUCTURE DETERMINATION. : XWINNMR, NMRPIPE, NMRVIEW REMARK 210 SOFTWARE USED REMARK 210 5.0.4, X-PLOR NIH 2.17.0, REMARK 210 MOLMOL 2K.2, PROCHECK 3.5.4 REMARK 210 METHOD USED : SIMULATED ANNEALING REMARK 210 REMARK 210 CONFORMERS, NUMBER CALCULATED : 200 REMARK 210 CONFORMERS, NUMBER SUBMITTED : 10 REMARK 210 CONFORMERS, SELECTION CRITERIA : STRUCTURES WITH THE LOWEST REMARK 210 ENERGY REMARK 210 REMARK 210 BEST REPRESENTATIVE CONFORMER IN THIS ENSEMBLE : 1 REMARK 210 REMARK 210 REMARK: NULL

#### **REMARK 215** is necessary if data was obtained from a solution NMR experiment.

### Template

123456781234567890REMARK 215NMR STUDYREMARK 215NMR DATA.PROTEIN DATA BANK CONVENTIONS REQUIRE THATREMARK 215CRYST1 AND SCALE RECORDS BE INCLUDED, BUT THE VALUES ONREMARK 215THESE RECORDS ARE MEANINGLESS.

```
Remark 217 is used in place of REMARK 215 if a Solid State NMR experiment was performed.
```

### Template

1	2	3	4	5	6	7	8

REMARK 217 REMARK 217 SOLID STATE NMR STUDY REMARK 217 THE COORDINATES IN THIS ENTRY WERE GENERATED FROM SOLID REMARK 217 STATE NMR DATA. PROTEIN DATA BANK CONVENTIONS REQUIRE THAT REMARK 217 CRYST1 AND SCALE RECORDS BE INCLUDED, BUT THE VALUES ON REMARK 217 THESE RECORDS ARE MEANINGLESS.

# **REMARK 230, Neutron Diffraction Experiment Details**

REMARK 230 is mandatory if data was obtained from a neutron diffraction study. The format of the date in this remark is DD-MMM-YY. DD is the day of the month (a number 01 through 31), MMM is the English 3-letter abbreviation for the month, and YY is the year.

### Template

	1	2 3 4		5	6	7	8
1234567	7890	12345678901234567890123456789012	345	567890123	34567890123456	6789012345678	890
REMARK	230						
REMARK	230	EXPERIMENTAL DETAILS					
REMARK	230			NEUTRON	DIFFRACTION		
		DATE OF DATA COLLECTION					
REMARK	230	TEMPERATURE (KELVIN)	:				
REMARK	230		:				
REMARK	230	NUMBER OF CRYSTALS USED	:				
REMARK	230						
		NEUTRON SOURCE	:				
		BEAMLINE	:				
		WAVELENGTH OR RANGE (A)	:				
		MONOCHROMATOR	:				
		OPTICS	:				
REMARK							
		DETECTOR TYPE	:				
		DETECTOR MANUFACTURER					
		INTENSITY-INTEGRATION SOFTWARE	:				
REMARK			:				
REMARK							
		NUMBER OF UNIQUE REFLECTIONS					
		RESOLUTION RANGE HIGH (A)					
		RESOLUTION RANGE LOW (A)					
REMARK		REJECTION CRITERIA (SIGMA(I))	:				
REMARK							
		OVERALL.					
REMARK	230	COMPLETENESS FOR RANGE (%)	:				
REMARK	230	DATA REDUNDANCY R MERGE (1)	:				
REMARK	230	R MERGE (I) R SYM (I)	:				
REMARK		<i sigma(i)=""> FOR THE DATA SET</i>	:				
REMARK		TN MUE UTCHECH DECOLUMION CHEIT					
		IN THE HIGHEST RESOLUTION SHELL			•		
		HIGHEST RESOLUTION SHELL, RANG					
KEPIARK DEMADV	230	HIGHEST RESOLUTION SHELL, RANG COMPLETENESS FOR SHELL (%)	с I •	LOW (A)	·		
KEMARK	230	COMPTETENESS LOK SUETT (2)	•				

```
REMARK 230 DATA REDUNDANCY IN SHELL :

REMARK 230 R MERGE FOR SHELL (I) :

REMARK 230 R SYM FOR SHELL (I) :

REMARK 230 

        REMARK 230

        REMARK 230

        REMARK 230

        REMARK 230

        REMARK 230

        REMARK 230

        REMARK 230

        REMARK 230

        REMARK 230

        REMARK 230

        SOFTWARE USED TO DETERMINE THE STRUCTURE:

        REMARK 230

        STARTING MODEL:

        REMARK 230

        REMARK 230

        REMARK 230

        REMARK 230
```

### Example

```
REMARK 230
REMARK 230 EXPERIMENTAL DETAILS
REMARK 230 EXPERIMENT TYPE
                                          : NEUTRON DIFFRACTION
REMARK 230 EXPERIMENT TIPE : NEUTRON D
REMARK 230 DATE OF DATA COLLECTION : 05-MAY-03
                         (KELVIN) : 293.0
REMARK 230 TEMPERATURE
REMARK 230 PH : 5.30
REMARK 230 NUMBER OF CRYSTALS USED : 1
REMARK 230
REMARK 230
                                         : NULL
REMARK 230 NEUTRON SOURCE
REMARK 230 BEAMLINE
                                          : NULL
REMARK 230 WAVELENGTH OR RANGE (A) : 2.88
                                      : ELLASTICALLY BENT SILICON
REMARK 230 MONOCHROMATOR
REMARK 230 OPTICS
                                          : MONOCHROMATOR
REMARK 230
REMARK 230 DETECTOR TYPE
                                          : NEUTRON IMAGING PLATE
REMARK 230 DETECTOR MANUFACTURER
                                           : BIX-3
REMARK 230 INTENSITY-INTEGRATION SOFTWARE : DENZO
REMARK 230 DATA SCALING SOFTWARE
                                           : SCALEPACK
REMARK 230
REMARK 230 NUMBER OF UNIQUE REFLECTIONS : 7001
REMARK 230 RESOLUTION RANGE HIGH (A) : 2.400
REMARK 230 RESOLUTION RANGE LOW
                                      (A) : 100.000
REMARK 230 REJECTION CRITERIA (SIGMA(I)) : 1.000
REMARK 230
REMARK 230 OVERALL.
REMARK 230 COMPLETENESS FOR RANGE (%) : 92.5
REMARK 230 DATA REDUNDANCY
                                           : NULL
REMARK 230 R MERGE
                                       (I) : 0.14300
REMARK 230 R SYM
                                       (I) : NULL
REMARK 230 <1/SIGMA(1)> FOR THE DATA SET : NULL
REMARK 230
REMARK 230 IN THE HIGHEST RESOLUTION SHELL.
REMARK 230 HIGHEST RESOLUTION SHELL, RANGE HIGH (A) : 2.40
REMARK 230 HIGHEST RESOLUTION SHELL, RANGE LOW (A) : 2.49
REMARK 230 COMPLETENESS FOR SHELL (%) : 82.1
REMARK 230 DATA REDUNDANCY IN SHELL
                                           : NULL
REMARK 230R MERGE FOR SHELL(I): 0.39500REMARK 230R SYM FOR SHELL(I): NULL
REMARK 230SIFT FOR SHELL(I) : NULLREMARK 230<I/SIGMA(I)> FOR SHELL: 2.30REMARK 230
                                           : 2.300
REMARK 230 METHOD USED TO DETERMINE THE STRUCTURE: MOLECULAR REPLACEMENT
REMARK 230 SOFTWARE USED : CNS
REMARK 230 STARTING MODEL: PDB ENTRY 1UCR
REMARK 230
REMARK 230 REMARK: NULL
```

PDB File Format v. 3.2

# **REMARK 240 (updated), Electron Crystallography Experiment Details**

REMARK 240 is mandatory if data was obtained from an electron crystallography study. The format of the date in this remark is DD-MMM-YY. DD is the day of the month (a number 01 through 31), MMM is the English 3-letter abbreviation for the month, and YY is the year.

### Template

1	2 3	=	-		7	8
	345678901234567890123	34567890123456	78901234	56789012345	67890123456	7890
REMARK 240						
	XPERIMENTAL DETAILS					
	RECONSTRUCTION METHO	DD :				
	SAMPLE TYPE	:				
	SPECIMEN TYPE	:				
	DATA ACQUISITION					
	DATE OF DATA COLLECT					
	TEMPERATURE	(KELVIN) :				
REMARK 240		:				
	NUMBER OF CRYSTALS U	JSED :				
	MICROSCOPE MODEL	:				
	DETECTOR TYPE	:				
	ACCELERATION VOLTAGE					
	NUMBER OF UNIQUE REP					
	RESOLUTION RANGE HIC					
REMARK 240	RESOLUTION RANGE LOW	V (A):				
	DATA SCALING SOFTWAR					
REMARK 240	COMPLETENESS FOR RAN	NGE (%):				
REMARK 240	DATA REDUNDANCY	:				
REMARK 240	IN THE HIGHEST RESO	LUTION SHELL				
	HIGHEST RESOLUTION S	-	• •			
REMARK 240	HIGHEST RESOLUTION S	SHELL, RANGE LO	OW (A)	:		
REMARK 240	COMPLETENESS FOR SHE	ELL (%):				
REMARK 240	DATA REDUNDANCY IN S	SHELL :				
REMARK 240	R MERGE FOR SHELL	(I) :				
REMARK 240	METHOD USED TO DETER	RMINE THE STRUC	CTURE:			
REMARK 240	SOFTWARE USED	:				
REMARK 240	STARTING MODEL	:				
Example						

REMARK	240				
REMARK	240	EXPERIMENTAL DETAILS			
REMARK	240	RECONSTRUCTION METHOD		:	CRYSTALLOGRAPHY
REMARK	240	SAMPLE TYPE		:	2D CRYSTAL
REMARK	240	SPECIMEN TYPE		:	VITREOUS ICE (CRYO EM)
REMARK	240	DATA ACQUISITION			
REMARK	240	DATE OF DATA COLLECTION		:	01-DEC-03
REMARK	240	TEMPERATURE (1	KELVIN)	:	300.0
REMARK	240	PH		:	6.00
REMARK	240	NUMBER OF CRYSTALS USED		:	286
REMARK	240	MICROSCOPE MODEL		:	JEM3000SFF
REMARK	240	DETECTOR TYPE		:	CCD
REMARK	240	ACCELERATION VOLTAGE (K	V)	:	300
REMARK	240	NUMBER OF UNIQUE REFLEC	TIONS	:	22293

REMARK 240	RESOLUTION RANGE HIGH	(A) : 1.9
REMARK 240	RESOLUTION RANGE LOW	(A) : 20.000
REMARK 240	DATA SCALING SOFTWARE	: SOFTWARE
REMARK 240	COMPLETENESS FOR RANGE	(%) : 80.0
REMARK 240	DATA REDUNDANCY	: 5.700
REMARK 240	IN THE HIGHEST RESOLUTION	SHELL.
REMARK 240	HIGHEST RESOLUTION SHELL,	RANGE HIGH (A) : 1.90
REMARK 240	HIGHEST RESOLUTION SHELL,	RANGE LOW (A) : 2.0
REMARK 240	COMPLETENESS FOR SHELL	(%) : 82.0
REMARK 240	DATA REDUNDANCY IN SHELL	: 5.70
REMARK 240	R MERGE FOR SHELL	(I) : 0.166
REMARK 240	METHOD USED TO DETERMINE	THE STRUCTURE: MOLECULAR
REMARK 240		REPLACEMENT
REMARK 240	SOFTWARE USED	: CNS
REMARK 240	STARTING MODEL	: PDB ENTRY 1SOR

### **REMARK 245 (updated), Electron Microscopy Experiment Details**

REMARK 245 is mandatory if data was obtained from a EM study. The format of the date in this remark is DD-MMM-YY. DD is the day of the month (a number 01 through 31), MMM is the English 3-letter abbreviation for the month, and YY is the year.

### Template

2 3 4 5 6 7 8 1 1234567890012345678900123456789001234567890000REMARK 245 **REMARK 245 EXPERIMENTAL DETAILS** REMARK 245 RECONSTRUCTION METHOD : REMARK 245 SPECIMEN TYPE : REMARK 245 REMARK 245 ELECTRON MICROSCOPE SAMPLE REMARK 245 SAMPLE TYPE : REMARK 245 PARTICLE TYPE : REMARK 245 NAME OF SAMPLE : REMARK 245 SAMPLE CONCENTRATION (MG ML-1) : REMARK 245 SAMPLE SUPPORT DETAILS : REMARK 245 SAMPLE VITRIFICATION DETAILS : REMARK 245 SAMPLE BUFFER : REMARK 245 PH : REMARK 245 SAMPLE DETAILS : REMARK 245 REMARK 245 DATA ACQUISITION REMARK 245 DATE OF EXPERIMENT : REMARK 245 NUMBER OF MICROGRAPHS-IMAGES : REMARK 245 TEMPERATURE (KELVIN) : REMARK 245 MICROSCOPE MODEL : REMARK 245 DETECTOR TYPE : REMARK 245 MINIMUM DEFOCUS (NM) : REMARK 245 MAXIMUM DEFOCUS (NM) : REMARK 245 MINIMUM TILT ANGLE (DEGREES) : REMARK 245 MAXIMUM TILT ANGLE (DEGREES) : REMARK 245 NOMINAL CS : REMARK 245 IMAGING MODE : REMARK 245 ELECTRON DOSE (ELECTRONS NM\*\*-2) : REMARK 245 ILLUMINATION MODE : NOMINAL MAGNIFICATION REMARK 245 : REMARK 245 CALIBRATED MAGNIFICATION : REMARK 245 SOURCE : REMARK 245 ACCELERATION VOLTAGE (KV) : REMARK 245 IMAGING DETAILS

REMARK	245			
REMARK	245	EXPERIMENTAL DETAILS		
REMARK	245	RECONSTRUCTION METHOD SPECIMEN TYPE	:	SINGLE PARTICLE
REMARK	245	SPECIMEN TYPE	:	VITREOUS ICE (CRYO EM)
REMARK	245			
REMARK	245	ELECTRON MICROSCOPE SAMPLE		
REMARK	245	SAMPLE TYPE PARTICLE TYPE NAME OF SAMPLE	:	PARTICLE
REMARK	245	PARTICLE TYPE	:	MIXED SYMMETRY
REMARK	245	NAME OF SAMPLE	:	BACTERIOPHAGE T4
REMARK	245	SAMPLE CONCENTRATION (MG ML-1)	:	20.00
REMARK	245	SAMPLE SUPPORT DETAILS	:	NULL
REMARK	245	SAMPLE VITRIFICATION DETAILS	:	NULL
REMARK	245	SAMPLE BUFFER	:	H2O
REMARK	245	SAMPLE VITRIFICATION DETAILS SAMPLE BUFFER PH	:	7.50
REMARK	245	PH SAMPLE DETAILS	:	PHAGE
REMARK	245			
REMARK	245	DATA ACQUISITION DATE OF EXPERIMENT NUMBER OF MICROGRAPHS-IMAGES TEMPERATURE (KELVIN) MICROSCOPE MODEL DETECTOR TYPE MINIMUM DEFOCUS (NM) MAXIMUM DEFOCUS (NM) MINIMUM TILT ANGLE (DEGREES) MAXIMUM TILT ANGLE (DEGREES) NOMINAL CS IMAGING MODE ELECTRON DOSE (ELECTRONS NM**-2		
REMARK	245	DATE OF EXPERIMENT		: 06-JAN-02
REMARK	245	NUMBER OF MICROGRAPHS-IMAGES		: NULL
REMARK	245	TEMPERATURE (KELVIN)		: 100.00
REMARK	245	MICROSCOPE MODEL		: FEI/PHILIPS CM300FEG/T
REMARK	245	DETECTOR TYPE		: NULL
REMARK	245	MINIMUM DEFOCUS (NM)		: 500.00
REMARK	245	MAXIMUM DEFOCUS (NM)		: 3400.00
REMARK	245	MINIMUM TILT ANGLE (DEGREES)		: 0.00
REMARK	245	MAXIMUM TILT ANGLE (DEGREES)		: 0.00
REMARK	245	NOMINAL CS		: 1.40
REMARK	245	IMAGING MODE		: BRIGHT FIELD
REMARK	245	ELECTRON DOSE (ELECTRONS NM**-2	2)	: 20.00
REMARK	245	ILLUMINATION MODE NOMINAL MAGNIFICATION		: SPOT SCAN
REMARK	245	NOMINAL MAGNIFICATION		: 45000
REMARK	245	CALIBRATED MAGNIFICATION		: 47000
REMARK	245	SOURCE		: FIELD EMISSION GUN
REMARK	245	SOURCE ACCELERATION VOLTAGE (KV)		: 300

# **REMARK 247, Electron Microscopy details**

REMARK 247 is mandatory if data was obtained from an EM study.

### Template

123456781234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890REMARK 247REMARK 247ELECTRON MICROSCOPYREMARK 247THE COORDINATES IN THIS ENTRY WERE GENERATED FROMREMARK 247ELECTRON MICROSCOPY DATA. PROTEIN DATA BANK CONVENTIONSREMARK 247REQUIRE THAT CRYST1 AND SCALE RECORDS BE INCLUDED,REMARK 247BUT THE VALUES ON THESE RECORDS ARE MEANINGLESSREMARK 247EXCEPT FOR THE CALCULATION OF THE STRUCTURE FACTORS

# **REMARK 250, Other Type of Experiment Details**

REMARKs specific to other kinds of studies, not listed above. REMARK 250 is mandatory if other than X-ray, NMR, neutron, or electron study. The format of the date in this remark is DD-MMM-YY. DD is the day of the month (a number 01 through 31), MMM is the English 3-letter abbreviation for the month, and YY is the year.

### Template

7 1 2 3 4 5 6 8 REMARK 250 REMARK 250 EXPERIMENTAL DETAILS REMARK 250 EXPERIMENT TYPE : REMARK 250 DATE OF DATA COLLECTION : REMARK 250 **REMARK 250 REMARK:** 

# **REMARK 265, Solution Scattering Experiment Details**

# Examples

REMARK				
		EXPERIMENTAL DETAILS		
REMARK				10
		EXPERIMENT TYPE : SMALL ANGLE X-RAY SCATTER		
REMARK	265	DATA ACQUISITION		
REMARK	265	RADIATION/NEUTRON SOURCE	:	SRS BEAMLINE 2.1
REMARK	265	SYNCHROTRON (Y/N)	:	Y
REMARK	265	BEAMLINE	:	2.1
REMARK	265	BEAMLINE INSTRUMENT	:	NULL
REMARK	265	DETECTOR TYPE	:	500-CHANNEL QUADRANT
REMARK	265	DETECTOR MANUFACTURER DETAILS	:	NULL
REMARK	265	DATA ACQUISITION RADIATION/NEUTRON SOURCE SYNCHROTRON (Y/N) BEAMLINE BEAMLINE INSTRUMENT DETECTOR TYPE DETECTOR MANUFACTURER DETAILS TEMPERATURE (KELVIN) PH NUMBER OF TIME FRAMES USED PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM)	:	288
REMARK	265	PH	:	NULL
REMARK	265	NUMBER OF TIME FRAMES USED	:	10
REMARK	265	PROTEIN CONCENTRATION RANGE (MG/ML)	:	0.7 - 14
REMARK	265	SAMPLE BUFFER	:	TRIS
REMARK	265	DATA REDUCTION SOFTWARE	:	ОТОКО
			•	
REMARK	265	SIGMA MEAN RADIUS OF GYRATION	:	0.4
REMARK	265	R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM)	:	4.4
REMARK	265	R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII	:	0.2
REMARK	265	R(XS-2) MEAN CROSS SECTIONAL RADII (NM)	:	1.7
REMARK	265	R(XS-2) SIGMA MEAN CROSS SECTIONAL RADII		
REMARK		P(R) PROTEIN LENGTH (NM)	:	40
REMARK	265			
REMARK	265	EXPERIMENT TYPE : SMALL ANGLE NEUTRON SCATT	ΓEI	RING
REMARK	265	DATA ACQUISITION		
REMARK	265	RADIATION/NEUTRON SOURCE	:	ILL
REMARK	265	SYNCHROTRON (Y/N)	:	N
REMARK	265	BEAMLINE TYPE	:	NULL
REMARK	265	BEAMLINE INSTRUMENT	:	D11, D22
REMARK	265	DETECTOR TYPE	:	AREA
REMARK	265	DETECTOR MANUFACTURER DETAILS	:	NULL
REMARK	265	TEMPERATURE (KELVIN)	:	NULL
REMARK	265	וות	•	
REMARK		PH	:	NULL
	265	DATA ACQUISITION RADIATION/NEUTRON SOURCE SYNCHROTRON (Y/N) BEAMLINE TYPE BEAMLINE INSTRUMENT DETECTOR TYPE DETECTOR MANUFACTURER DETAILS TEMPERATURE (KELVIN) PH NUMBER OF TIME FRAMES USED	:	NULL NULL
REMARK	265 265	NUMBER OF TIME FRAMES USED PROTEIN CONCENTRATION RANGE (MG/ML)	::	NULL NULL 0.4 - 9.6
REMARK REMARK	265 265 265	NUMBER OF TIME FRAMES USED PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER	::	NULL NULL 0.4 - 9.6 PBS IN 99.9% D20
REMARK REMARK REMARK	265 265 265 265	NUMBER OF TIME FRAMES USED PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE	::	NULL NULL 0.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY
REMARK REMARK REMARK REMARK	265 265 265 265 265	NUMBER OF TIME FRAMES USED PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE	::	0.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY SCTPL5, GNOM
REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265	PH NUMBER OF TIME FRAMES USED PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM)	::	NULL NULL 0.4 - 9.6 PBS IN 99.9% D2O DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3
REMARK REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM)	: : :	0.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY SCTPL5, GNOM
REMARK	265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION	• • • •	0.4 - 9.6 PBS IN 99.9% D2O DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4
REMARK REMARK	265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM)	•	0.4 - 9.6 PBS IN 99.9% D2O DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9
REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM)		0.4 - 9.6 PBS IN 99.9% D2O DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51
REMARK REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII		0.4 - 9.6 PBS IN 99.9% D2O DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51
REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM)	• • • • • • • •	0.4 - 9.6 PBS IN 99.9% D2O DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51
REMARK REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM) R(XS-2) SIGMA MEAN CROSS SECTIONAL RADII	• • • • • • • •	0.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51 0.06
REMARK REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM) R(XS-2) SIGMA MEAN CROSS SECTIONAL RADII	• • • • • • • •	0.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51 0.06
REMARK REMARK REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM) R(XS-2) SIGMA MEAN CROSS SECTIONAL RADII P(R) PROTEIN LENGTH (NM)	• • • • • • • • • •	0.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51 0.06
REMARK REMARK REMARK REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM) R(XS-2) SIGMA MEAN CROSS SECTIONAL RADII P(R) PROTEIN LENGTH (NM) DATA ACQUISITION RADIATION/NEUTRON SOURCE		0.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51 0.06 37 - 39
REMARK REMARK REMARK REMARK REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM) R(XS-2) SIGMA MEAN CROSS SECTIONAL RADII P(R) PROTEIN LENGTH (NM) DATA ACQUISITION RADIATION/NEUTRON SOURCE SYNCHROTRON (Y/N) BEAMLINE TYPE		No.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51 0.06 37 - 39 ISIS
REMARK REMARK REMARK REMARK REMARK REMARK REMARK REMARK REMARK REMARK	265 265 265 265 265 265 265 265 265 265	PROTEIN CONCENTRATION RANGE (MG/ML) SAMPLE BUFFER DATA REDUCTION SOFTWARE DATA ANALYSIS SOFTWARE GUINIER MEAN RADIUS OF GYRATION (NM) SIGMA MEAN RADIUS OF GYRATION R(XS-1) MEAN CROSS SECTIONAL RADII (NM) R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII R(XS-2) MEAN CROSS SECTIONAL RADII (NM) R(XS-2) SIGMA MEAN CROSS SECTIONAL RADII P(R) PROTEIN LENGTH (NM) DATA ACQUISITION RADIATION/NEUTRON SOURCE SYNCHROTRON (Y/N) BEAMLINE TYPE		No.4 - 9.6 PBS IN 99.9% D20 DETEC, RNILS, SPOLLY SCTPL5, GNOM 11.3 0.4 3.9 0.2 1.51 0.06 37 - 39 ISIS N

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REMARK 265 DETECTOR TYPE : AREA (TIME-OF-FLIGHT) REMARK 265 TEMPERATURE (KELVIN) : NULL REMARK 265 : NULL  $\mathbf{PH}$ REMARK 265 NUMBER OF TIME FRAMES USED : NULL PROTEIN CONCENTRATION RANGE (MG/ML) REMARK 265 : 3.7, 6.1 : PBS IN 99.9% D20 REMARK 265 SAMPLE BUFFER REMARK 265 DATA REDUCTION SOFTWARE : COLLETTE REMARK 265 DATA ANALYSIS SOFTWARE : SCTPL5, GNOM REMARK 265 GUINIER MEAN RADIUS OF GYRATION (NM) : 11.7 REMARK 265 SIGMA MEAN RADIUS OF GYRATION : 0.5 REMARK 265 R(XS-1) MEAN CROSS SECTIONAL RADII (NM) : NULL REMARK 265 R(XS-1) SIGMA MEAN CROSS SECTIONAL RADII : NULL REMARK 265 R(XS-2) MEAN CROSS SECTIONAL RADII (NM) : NULL REMARK 265 R(XS-2) SIGMA MEAN CROSS SECTIONAL RADII : NULL REMARK 265 P(R) PROTEIN LENGTH (NM) : 40 REMARK 265 REMARK 265 DATA ANALYSIS AND MODEL FITTING: REMARK 265 METHOD USED TO DETERMINE THE STRUCTURE: CONSTRAINED SCATTERING REMARK 265 FITTING OF HOMOLOGY REMARK 265 MODELS REMARK 265 SOFTWARE USED : INSIGHT II, HOMOLOGY, DISCOVERY, REMARK 265 BIOPOLYMER, DELPHI, SCTPL5, GNOM REMARK 265 SOFTWARE AUTHORS : MSI REMARK 265 STARTING MODEL : PDB CODE 1HFI, 1HCC, 1HFH, 1VCC REMARK 265 REMARK 265 CONFORMERS, NUMBER CALCULATED : 2010 REMARK 265 CONFORMERS, NUMBER SUBMITTED : 4 REMARK 265 CONFORMERS, SELECTION CRITERIA : THE MODELLED SCATTERING REMARK 265 CURVES WERE ASSESSED BY CALCULATION OF THE REMARK 265 RG, RSX-1 AND RXS-2 VALUES IN THE SAME Q RANGES REMARK 265 USED IN THE EXPERIMENTAL GUINIER FITS. MODELS WERE REMARK 265 THEN RANKED USING A GOODNESS-OF-FIT R-FACTOR REMARK 265 DEFINED BY ANALOGY WITH PROTEIN CRYSTALLOGRAPHY REMARK 265 AND BASED ON THE EXPERIMENTAL CURVES IN THE Q RANGE REMARK 265 EXTENDING TO 1.4 NM-1. REMARK 265 REMARK 265 REPRESENTATIVE CONFORMER IN THIS ENSEMBLE : 1 REMARK 265 REMARK 265 OTHER DETAILS: HOMOLOGY MODELS WERE BUILT FOR REMARK 265 THE 17 SCR DOMAINS AND ENERGY MINIMISATIONS WERE REMARK 265 PERFORMED TO IMPROVE THE CONNECTIVITY IN THE FH MODEL. REMARK 265 TRIANTENNARY COMPLEX-TYPE CARBOHYDRATE STRUCTURES REMARK 265 (MAN3GLCNAC6GAL3FUC3NEUNAC1) WERE ADDED TO EACH OF THE REMARK 265 N-LINKED GLYCOSYLATION SITES. A LIBRARY OF LINKER PEPTIDE REMARK 265 CONFORMATIONS WAS USED IN DOMAIN MODELLING CONSTRAINED REMARK 265 BY THE SOLUTION SCATTERING FITS. MODELLING WITH THE REMARK 265 SCATTERING DATA WAS ALSO CARRIED OUT BY ROTATIONAL REMARK 265 SEARCH METHODS. THE X-RAY AND NEUTRON SCATTERING CURVE I(O) WAS CALCULATED ASSUMING A UNIFORM SCATTERING DENSITY REMARK 265 REMARK 265 FOR THE SPHERES USING THE DEBYE EQUATION AS ADAPTED TO REMARK 265 SPHERES. X-RAY CURVES WERE CALCULATED FROM THE HYDRATED REMARK 265 SPHERE MODELS WITHOUT CORRECTIONS FOR WAVELENGTH SPREAD OR REMARK 265 BEAM DIVERGENCE, WHILE THESE CORRECTIONS WERE APPLIED FOR REMARK 265 THE NEUTRON CURVES BUT NOW USING UNHYDRATED MODELS.

# **REMARKs 280-290, Crystallographic Details**

# **REMARK 280, Crystal**

REMARK 280 presents information about the crystal. The solvent content and Matthews coefficient are provided for protein and polypeptide crystals. Crystallization conditions are in free text.

REMARK 280 is mandatory for single crystal studies.

### Template

```
5
                                                   7
      1
              2
                     3
                             4
                                           6
                                                          8
REMARK 280
REMARK 280 CRYSTAL
REMARK 280 SOLVENT CONTENT, VS
                        (%):
REMARK 280 MATTHEWS COEFFICIENT, VM (ANGSTROMS**3/DA):
REMARK 280
REMARK 280 CRYSTALLIZATION CONDITIONS: FREE TEXT GOES HERE.
```

### Example

```
REMARK 280 CRYSTAL
REMARK 280 SOLVENT CONTENT, VS (%): 36.85
REMARK 280 MATTHEWS COEFFICIENT, VM (ANGSTROMS**3/DA): 1.79
REMARK 280
REMARK 280 CRYSTALLIZATION CONDITIONS: 1.4M SODIUM ACETATE,
REMARK 280 0.1M MES PH 6.5
```

# **REMARK 285, CRYST1**

REMARK 285 presents information about the unit cell.

### Template

1 2 3 4 5 6 7 8 1234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 285 REMARK 285 CRYST1 REMARK 285 FREE TEXT GOES HERE.

### Example

REMARK 285 REMARK 285 CRYST1 REMARK 285 TEXT TO EXPLAIN UNUSUAL UNIT-CELL DATA: THE DATA WAS REMARK 285 COLLECTED ON TWO-DIMENSIONAL CRYSTALS AND HENCE THE REMARK 285 C-AXIS REPEAT DOES NOT CORRESPOND TO A REAL REPEAT, BUT REMARK 285 INSTEAD REFERS TO THE SAMPLING THAT IS USED TO DESCRIBE REMARK 285 THE CONTINUOUS TRANSFORM. THE C VALUE OF 100.9 IS REMARK 285 THEREFORE THE VALUE WHICH SHOULD BE USED IN REMARK 285 INTERPRETING THE MEANING OF THE L INDEX.

# **REMARK 290, Crystallographic Symmetry**

REMARK 290 is mandatory for crystalline studies. The REMARK is automatically generated.

### Example

2 5 7 3 4 6 8 1 REMARK 290 REMARK 290 CRYSTALLOGRAPHIC SYMMETRY REMARK 290 SYMMETRY OPERATORS FOR SPACE GROUP: P 21 21 21 REMARK 290 SYMOP SYMMETRY REMARK 290 REMARK 290 NNNMMM OPERATOR REMARK 290 1555 X,Y,Z REMARK 290 2555 1/2-X,-Y,1/2+Z REMARK 290 3555 -X, 1/2+Y, 1/2-Z4555 REMARK 290 1/2+X, 1/2-Y, -ZREMARK 290 WHERE NNN -> OPERATOR NUMBER REMARK 290 REMARK 290 MMM -> TRANSLATION VECTOR REMARK 290 REMARK 290 CRYSTALLOGRAPHIC SYMMETRY TRANSFORMATIONS REMARK 290 THE FOLLOWING TRANSFORMATIONS OPERATE ON THE ATOM/HETATM REMARK 290 RECORDS IN THIS ENTRY TO PRODUCE CRYSTALLOGRAPHICALLY REMARK 290 RELATED MOLECULES. REMARK 290 SMTRY1 1 1.000000 0.000000 0.000000 0.00000 REMARK 290SMTRY210.0000001.0000000.000000REMARK 290SMTRY310.0000000.0000001.000000 0.00000 0.00000 

 SMTRY3
 1
 0.000000
 0.000000
 1.000000

 SMTRY1
 2
 -1.000000
 0.000000
 0.000000

 SMTRY2
 2
 0.000000
 -1.000000
 0.000000

 SMTRY3
 2
 0.000000
 0.000000
 1.000000

 SMTRY1
 3
 -1.000000
 0.000000
 0.000000

 SMTRY2
 3
 0.000000
 1.000000
 0.000000

 36.30027 REMARK 290 REMARK 290 0.00000 REMARK 290 59.50256 REMARK 290 0.00000 REMARK 290 46.45545 REMARK 290 SMTRY3 3 0.000000 0.000000 -1.000000 59.50256 REMARK 290 SMTRY1 4 1.000000 0.000000 0.000000 36.30027 46.45545 REMARK 290 SMTRY2 4 0.000000 -1.000000 0.000000 REMARK 290 SMTRY3 4 0.000000 0.000000 -1.000000 0.00000 REMARK 290

### REMARK 300 (updated), Biomolecule

Description of the biologically functional molecule (biomolecule) in free text. Remark 300 is mandatory if REMARK 350 is provided.

### Template

2 3 4 5 6 7 8 REMARK 300 REMARK 300 BIOMOLECULE: 1 REMARK 300 SEE REMARK 350 FOR THE AUTHOR PROVIDED AND/OR PROGRAM REMARK 300 GENERATED ASSEMBLY INFORMATION FOR THE STRUCTURE IN REMARK 300 THIS ENTRY. THE REMARK MAY ALSO PROVIDE INFORMATION ON REMARK 300 BURIED SURFACE AREA. REMARK 300 FREE TEXT GOES HERE.

### Examples

REMARK 300 REMARK 300 BIOMOLECULE: 1, 2, 3, 4, 5, 6, 7, 8 REMARK 300 SEE REMARK 350 FOR THE AUTHOR PROVIDED AND/OR PROGRAM REMARK 300 GENERATED ASSEMBLY INFORMATION FOR THE STRUCTURE IN REMARK 300 THIS ENTRY. THE REMARK MAY ALSO PROVIDE INFORMATION ON REMARK 300 BURIED SURFACE AREA.

REMARK 300 REMARK 300 BIOMOLECULE: 1, 2, 3 REMARK 300 SEE REMARK 350 FOR THE AUTHOR PROVIDED AND/OR PROGRAM REMARK 300 GENERATED ASSEMBLY INFORMATION FOR THE STRUCTURE IN REMARK 300 THIS ENTRY. THE REMARK MAY ALSO PROVIDE INFORMATION ON REMARK 300 BURIED SURFACE AREA. REMARK 300 DETAILS: THE CATALYTIC SUBUNIT OF LIVER ALCOHOL DEHYDROGENASE FROM REMARK 300 EQUUS CABALLUS IS A HOMODIMER.

#### Example - Icosahedral virus

REMARK 300 REMARK 300 BIOMOLECULE: 1 REMARK 300 SEE REMARK 350 FOR THE AUTHOR PROVIDED AND/OR PROGRAM REMARK 300 GENERATED ASSEMBLY INFORMATION FOR THE STRUCTURE IN REMARK 300 THIS ENTRY. THE REMARK MAY ALSO PROVIDE INFORMATION ON REMARK 300 BURIED SURFACE AREA. REMARK 300 DETAILS: THE ASSEMBLY REPRESENTED IN THIS ENTRY HAS REGULAR REMARK 300 ICOSAHEDRAL POINT SYMMETRY (SCHOENFLIES SYMBOL = I).

#### Example - Helical viruses

REMARK 300 REMARK 300 BIOMOLECULE: 1 REMARK 300 SEE REMARK 350 FOR THE AUTHOR PROVIDED AND/OR PROGRAM REMARK 300 GENERATED ASSEMBLY INFORMATION FOR THE STRUCTURE IN REMARK 300 THIS ENTRY. THE REMARK MAY ALSO PROVIDE INFORMATION ON REMARK 300 BURIED SURFACE AREA. REMARK 300 DETAILS: THE ASSEMBLY REPRESENTED IN THIS ENTRY HAS REGULAR REMARK 300 HELICAL SYMMETRY WITH THE FOLLOWING PARAMETERS: REMARK 300 ROTATION PER SUBUNIT (TWIST) = -33.23 DEGREES REMARK 300 RISE PER SUBUNIT (HEIGHT) = 16.00 ANGSTROMS REMARK 300 IN ADDITION, THERE IS 5-FOLD CIRCULAR REMARK 300 SYMMETRY AROUND THE HELIX AXIS

### Example - point symmetry crystal structure

REMARK 300 REMARK 300 BIOMOLECULE: 1 REMARK 300 SEE REMARK 350 FOR THE AUTHOR PROVIDED AND/OR PROGRAM REMARK 300 GENERATED ASSEMBLY INFORMATION FOR THE STRUCTURE IN REMARK 300 THIS ENTRY. THE REMARK MAY ALSO PROVIDE INFORMATION ON REMARK 300 BURIED SURFACE AREA. REMARK 300 DETAILS: THE ASSEMBLY REPRESENTED IN THIS ENTRY HAS REMARK 300 REGULAR DIHEDRAL POINT SYMMETRY (SCHOENFLIES SYMBOL = D17).

### **REMARK 350 (updated), Generating the Biomolecule**

REMARK 350 presents all transformations, both crystallographic and non-crystallographic, needed to generate the biomolecule. These transformations operate on the coordinates in the entry. Both author and computational descriptions of assemblies are provided, if applicable.

### Template

1 2 3 4 5 6 7 8 REMARK 350 COORDINATES FOR A COMPLETE MULTIMER REPRESENTING THE KNOWN REMARK 350 BIOLOGICALLY SIGNIFICANT OLIGOMERIZATION STATE OF THE REMARK 350 MOLECULE CAN BE GENERATED BY APPLYING BIOMT TRANSFORMATIONS REMARK 350 GIVEN BELOW. BOTH NON-CRYSTALLOGRAPHIC AND REMARK 350 CRYSTALLOGRAPHIC OPERATIONS ARE GIVEN. REMARK 350 REMARK 350 BIOMOLECULE: 1 REMARK 350 AUTHOR DETERMINED BIOLOGICAL UNIT: DODECAMERIC REMARK 350 SOFTWARE DETERMINED QUATERNARY STRUCTURE: DODECAMERIC REMARK 350 SOFTWARE USED: PISA REMARK 350 TOTAL BURIED SURFACE AREA: 2990 ANGSTROM\*\*2 REMARK 350 SURFACE AREA OF THE COMPLEX: 9330 ANGSTROM\*\*2 REMARK 350 CHANGE IN SOLVENT FREE ENERGY: -40.0 KCAL/MOL REMARK 350 APPLY THE FOLLOWING TO CHAINS: A, B, C, D, E, F, G, H, I, REMARK 350 AND CHAINS: J, K, L REMARK 350 BIOMT1 1 1.000000 0.000000 0.000000 0.00000 REMARK 350 BIOMT2 1 0.000000 1.000000 0.000000 0.00000 BIOMT3 1 0.000000 0.000000 1.000000 REMARK 350 0.00000

Note: If entry is part of a SPLIT record (larger multi-protein complex), REMARK 350 represents only the quaternary structure of that split entry.

```
REMARK 350 COORDINATES FOR A COMPLETE MULTIMER REPRESENTING THE KNOWN
REMARK 350 BIOLOGICALLY SIGNIFICANT OLIGOMERIZATION STATE OF THE
REMARK 350 MOLECULE CAN BE GENERATED BY APPLYING BIOMT TRANSFORMATIONS
REMARK 350 GIVEN BELOW. BOTH NON-CRYSTALLOGRAPHIC AND
REMARK 350 CRYSTALLOGRAPHIC OPERATIONS ARE GIVEN.
REMARK 350
REMARK 350 BIOMOLECULE: 1
REMARK 350 QUATERNARY STRUCTURE FOR THIS ENTRY: 21MERIC
REMARK 350 APPLY THE FOLLOWING TO CHAINS: A, B, C, D, E, F, G, H, I,
REMARK 350
                            AND CHAINS: J, K, L, M, N, O, P, Q, T,
REMARK 350
                            AND CHAINS: S, T, U
REMARK 350 BIOMT1 1 1.000000 0.000000 0.000000
                                                          0.00000
REMARK 350 BIOMT2 1 0.000000 1.000000 0.000000
                                                          0.00000
REMARK 350 BIOMT3 1 0.000000 0.000000 1.000000
                                                          0.00000
```

### Example – Author and computed assembly predictions agree

REMARK 350 COORDINATES FOR A COMPLETE MULTIMER REPRESENTING THE KNOWN REMARK 350 BIOLOGICALLY SIGNIFICANT OLIGOMERIZATION STATE OF THE REMARK 350 MOLECULE CAN BE GENERATED BY APPLYING BIOMT TRANSFORMATIONS REMARK 350 GIVEN BELOW. BOTH NON-CRYSTALLOGRAPHIC AND REMARK 350 CRYSTALLOGRAPHIC OPERATIONS ARE GIVEN. REMARK 350 REMARK 350 BIOMOLECULE: 1 REMARK 350 AUTHOR DETERMINED BIOLOGICAL UNIT: DODECAMERIC REMARK 350 SOFTWARE DETERMINED QUATERNARY STRUCTURE: DODECAMERIC REMARK 350 SOFTWARE USED: PISA REMARK 350 TOTAL BURIED SURFACE AREA: 2990 ANGSTROM\*\*2 REMARK 350 SURFACE AREA OF THE COMPLEX: 9330 ANGSTROM\*\*2 REMARK 350 CHANGE IN SOLVENT FREE ENERGY: -40.0 KCAL/MOL REMARK 350 APPLY THE FOLLOWING TO CHAINS: A, B, C, D, E, F, G, H, I, REMARK 350 AND CHAINS: J, K, L REMARK 350 BIOMT1 1 1.000000 0.000000 0.000000 0.00000 REMARK 350 BIOMT2 1 0.000000 1.000000 0.000000 0.00000 REMARK 350 BIOMT3 1 0.000000 0.000000 1.000000 0.00000

Note: The value for the average buried surface area will be round to the nearest 10.

### Example – Author and computed assembly predictions differ

REMARK 350 COORDINATES FOR A COMPLETE MULTIMER REPRESENTING THE KNOWN REMARK 350 BIOLOGICALLY SIGNIFICANT OLIGOMERIZATION STATE OF THE REMARK 350 MOLECULE CAN BE GENERATED BY APPLYING BIOMT TRANSFORMATIONS REMARK 350 GIVEN BELOW. BOTH NON-CRYSTALLOGRAPHIC AND REMARK 350 CRYSTALLOGRAPHIC OPERATIONS ARE GIVEN. REMARK 350 REMARK 350 BIOMOLECULE: 1 REMARK 350 AUTHOR DETERMINED BIOLOGICAL UNIT: HEXAMERIC REMARK 350 APPLY THE FOLLOWING TO CHAINS: A, B, C, D, E, F 
 REMARK 350
 BIOMT1
 1
 1.000000
 0.000000
 0.000000

 REMARK 350
 BIOMT2
 1
 0.000000
 1.000000
 0.000000

 REMARK 350
 BIOMT2
 1
 0.000000
 1.000000
 0.000000

 REMARK 350
 BIOMT3
 1
 0.000000
 0.000000
 1.000000
 0.00000 0.00000 0.00000 REMARK 350 REMARK 350 BIOMOLECULE: 2 REMARK 350 AUTHOR DETERMINED BIOLOGICAL UNIT: HEXAMERIC REMARK 350 APPLY THE FOLLOWING TO CHAINS: G, H, I, J, K, L REMARK 350 BIOMT1 1 1.000000 0.000000 0.000000 0.00000 REMARK 350 BIOMT2 1 0.000000 1.000000 0.000000 0.00000 REMARK 350 BIOMT3 1 0.000000 0.000000 1.000000 0.00000 REMARK 350 REMARK 350 BIOMOLECULE: 3 REMARK 350 SOFTWARE DETERMINED QUATERNARY STRUCTURE: DODECAMERIC REMARK 350 SOFTWARE USED: PISA REMARK 350 TOTAL BURIED SURFACE AREA: 2990 ANGSTROM\*\*2 REMARK 350 SURFACE AREA OF THE COMPLEX: 9330 ANGSTROM\*\*2 REMARK 350 CHANGE IN SOLVENT FREE ENERGY: -40.0 KCAL/MOL REMARK 350 APPLY THE FOLLOWING TO CHAINS: A, B, C, D, E, F, G, H, I, REMARK 350 AND CHAINS: J, K, L REMARK 350 BIOMT1 1 1.000000 0.000000 0.000000 0.00000 REMARK 350 BIOMT2 1 0.000000 1.000000 0.000000 0.00000 0.00000 REMARK 350 BIOMT3 1 0.000000 0.000000 1.000000

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### Example – When there are no quaternary assemblies provided by either author or software

REMARK 350 COORDINATES FOR A COMPLETE MULTIMER REPRESENTING THE KNOWN REMARK 350 BIOLOGICALLY SIGNIFICANT OLIGOMERIZATION STATE OF THE REMARK 350 MOLECULE CAN BE GENERATED BY APPLYING BIOMT TRANSFORMATIONS REMARK 350 GIVEN BELOW. BOTH NON-CRYSTALLOGRAPHIC AND REMARK 350 CRYSTALLOGRAPHIC OPERATIONS ARE GIVEN. REMARK 350 REMARK 350 BIOMOLECULE: 1 REMARK 350 SOFTWARE DETERMINED QUATERNARY STRUCTURE: MONOMERIC REMARK 350 SOFTWARE USED: PISA REMARK 350 APPLY THE FOLLOWING TO CHAINS: A REMARK 350 BIOMT1 1 1.000000 0.000000 0.000000 0.00000 REMARK 350 BIOMT2 1 0.000000 1.000000 0.000000 0.00000 REMARK 350 BIOMT3 1 0.000000 0.000000 1.000000 0.00000

Note that the average buried surface area is not included in this example because the quaternary structure is a monomer.

# Example – When software predicts multiple quaternary assemblies

For example, the author states the biological unit to be a dimer, but software predicts the quaternary structure to be either a dimer or a tetramer:

```
REMARK 300
REMARK 300
REMARK 300 BIOMOLECULE: 1, 2
REMARK 300 SEE REMARK 350 FOR THE AUTHOR PROVIDED AND/OR PROGRAM
REMARK 300 GENERATED ASSEMBLY INFORMATION FOR THE STRUCTURE IN
                       THE REMARK MAY ALSO PROVIDE INFORMATION ON
REMARK 300 THIS ENTRY.
REMARK 300 BURIED SURFACE AREA.
REMARK 300
REMARK 350 COORDINATES FOR A COMPLETE MULTIMER REPRESENTING THE KNOWN
REMARK 350 BIOLOGICALLY SIGNIFICANT OLIGOMERIZATION STATE OF THE
REMARK 350 MOLECULE CAN BE GENERATED BY APPLYING BIOMT TRANSFORMATIONS
REMARK 350 GIVEN BELOW. BOTH NON-CRYSTALLOGRAPHIC AND
REMARK 350 CRYSTALLOGRAPHIC OPERATIONS ARE GIVEN.
REMARK 350
REMARK 350 BIOMOLECULE:
                        1
REMARK 350 AUTHOR DETERMINED BIOLOGICAL UNIT: DIMERIC
REMARK 350 SOFTWARE DETERMINED QUATERNARY STRUCTURE: DIMERIC
REMARK 350 SOFTWARE USED: PISA
REMARK 350 TOTAL BURIED SURFACE AREA: 1460 ANGSTROM**2
REMARK 350 SURFACE AREA OF THE COMPLEX: 9330 ANGSTROM**2
REMARK 350 CHANGE IN SOLVENT FREE ENERGY: -40.0 KCAL/MOL
REMARK 350 APPLY THE FOLLOWING TO CHAINS: A, B
REMARK 350
            BIOMT1 1 1.000000 0.000000 0.000000
                                                             0.00000
            BIOMT2 1 0.000000 1.000000 0.000000
REMARK 350
                                                             0.00000
            BIOMT3 1 0.000000 0.000000 1.000000
REMARK 350
                                                             0.00000
REMARK 350
REMARK 350 BIOMOLECULE:
                         2
REMARK 350 SOFTWARE DETERMINED QUATERNARY STRUCTURE: TETRAMERIC
REMARK 350 SOFTWARE USED: PISA
REMARK 350 TOTAL BURIED SURFACE AREA: 2860 ANGSTROM**2
```

REMARK 350	SURFACE ARE	ΞA	OF THE COM	PLEX: 1233	0 ANGSTROM**2	
REMARK 350	GAIN IN SOL	LVE	NT FREE EN	ERGY: -20.	5 KCAL/MOL	
REMARK 350	APPLY THE F	FOL	LOWING TO	CHAINS: A,	В	
REMARK 350	BIOMT1	1	1.000000	0.00000	0.000000	0.00000
REMARK 350	BIOMT2	1	0.000000	1.000000	0.000000	0.00000
REMARK 350	BIOMT3	1	0.000000	0.00000	1.000000	0.00000
REMARK 350	BIOMT1	2	-1.000000	0.000000	0.000000	0.00000
REMARK 350	BIOMT2	2	0.000000	1.000000	0.000000	0.00000
REMARK 350	BIOMT3	2	0.000000	0.000000	-1.000000	0.00000

# **REMARK 375 (updated), Special Position**

REMARK 375 specifies atoms which lie within 0.15A of a symmetry-related atom and therefore, are considered to be on a special position, with cumulative occupancies of such atoms not exceeding 1.0.

### Template

1 2 3 4 5 6 7 8 1234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 375 REMARK 375 SPECIAL POSITION REMARK 375 FREE TEXT GOES HERE.

### Example

REMARK 375 REMARK 375 SPECIAL POSITION REMARK 375 HOH A 301 LIES ON A SPECIAL POSITION. REMARK 375 REMARK 375 SPECIAL POSITION REMARK 375 HOH A 13 LIES ON A SPECIAL POSITION. REMARK 375 HOH A 28 LIES ON A SPECIAL POSITION. REMARK 375 HOH A 36 LIES ON A SPECIAL POSITION.

### **REMARK 400, Compound**

Further details about the macromolecular contents of the entry.

### Template

1 2 3 4 5 6 7 8 1234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 400 REMARK 400 COMPOUND REMARK 400 FREE TEXT GOES HERE.

### Examples

REMARK 400 COMPOUND REMARK 400 THE PRD1 SUS1 MUTANT LACKS THE PACKAGING PROTEIN P9 REMARK 400 AND PRODUCES ONLY EMPTY PARTICLES, WHICH REPRESENT REMARK 400 AN ASSEMBLY INTERMEDIATE REMARK 400 COMPOUND REMARK 400 COMPONENT OF NAPHTHALENE DIOXYGENASE (NDO) REMARK 400 MULTICOMPONENT ENZYME SYSTEM WHICH CATALYZES THE INCORPORATION REMARK 400 OF BOTH ATOMS OF MOLECULAR OXYGEN INTO NAPHTHALENE TO FORM REMARK 400 CIS-NAPHTHALENE DIHYDRODIOL.

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### **REMARK 450, Source**

Further details about the biological source of the macromolecular contents of the entry.

### Template

1 2 3 4 5 6 7 8 1234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 450 REMARK 450 SOURCE REMARK 450 FREE TEXT GOES HERE.

# **REMARK 465 (updated), Missing residues**

REMARK 465 lists the residues that are present in the SEQRES records but are completely absent from the coordinates section.

#### Template for non NMR entries

1 2 3 4 5 6 7 8 REMARK 465 REMARK 465 MISSING RESIDUES REMARK 465 THE FOLLOWING RESIDUES WERE NOT LOCATED IN THE REMARK 465 EXPERIMENT. (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN REMARK 465 IDENTIFIER; SSSEQ=SEQUENCE NUMBER; I=INSERTION CODE.) REMARK 465 REMARK 465 M RES C SSSEQI

### Example

REMARK 465 **REMARK 465 MISSING RESIDUES** REMARK 465 THE FOLLOWING RESIDUES WERE NOT LOCATED IN THE REMARK 465 EXPERIMENT. (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN REMARK 465 IDENTIFIER; SSSEQ=SEQUENCE NUMBER; I=INSERTION CODE.) REMARK 465 M RES C SSSEQI REMARK 465 REMARK 465 ARG A 46 GLY A 47 REMARK 465 48 REMARK 465 ALA A REMARK 465 ARG A 49 REMARK 465 MET A 50

### Template for NMR entries (added)

1 2 3 4 5 6 7 8 12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 465 REMARK 465 MISSING RESIDUES REMARK 465 THE FOLLOWING RESIDUES WERE NOT LOCATED IN THE REMARK 465 EXPERIMENT. (RES=RESIDUE NAME; C=CHAIN IDENTIFIER; REMARK 465 SSSEQ=SEQUENCE NUMBER; I=INSERTION CODE.) REMARK 465 MODELS X-YYY REMARK 465 RES C SSSEQI

The models is listed as a range, X-YYY.

### Example

REMARK 465 REMARK 465 MISSING RESIDUES REMARK 465 THE FOLLOWING RESIDUES WERE NOT LOCATED IN THE REMARK 465 EXPERIMENT. (RES=RESIDUE NAME; C=CHAIN IDENTIFIER; REMARK 465 SSEQ=SEQUENCE NUMBER; I=INSERTION CODE.) REMARK 465 MODELS 1-20 REMARK 465 RES C SSSEQI REMARK 465 MET A 1 REMARK 465 GLY A 2

# REMARK 470 (updated), Missing Atom(s)

Non-hydrogen atoms of standard residues which are missing from the coordinates are listed. Missing HETATMs (atoms within hetetrogen groups) are not listed here.

#### Template for non NMR entries

2 3 4 5 7 8 1 6 REMARK 470 REMARK 470 MISSING ATOM REMARK 470 THE FOLLOWING RESIDUES HAVE MISSING ATOMS (M=MODEL NUMBER; REMARK 470 RES=RESIDUE NAME; C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; REMARK 470 I=INSERTION CODE): REMARK 470 M RES CSSEQI ATOMS

### Example

REMARK	470									
REMARK	470	MISSING	A	гом						
REMARK	470	THE FOLI	JOI	VING F	RESIDUE	S HAV	VE MIS	SING	ATOMS	(M=MODEL NUMBER;
REMARK	470	RES=RESI	DU	JE NAM	IE; C=C	HAIN	IDENT	IFIER	; SSE	Q=SEQUENCE NUMBER;
REMARK	470	I=INSER	CI(	ON COL	DE):					
REMARK	470	M RES	CS	SSEQI	ATOMS					
REMARK	470	ARG	А	412	CG	CD	NE	CZ	NH1	NH2
REMARK	470	ARG	А	456	CG	CD	NE	CZ	NH1	NH2
REMARK	470	GLU	А	486	CG	CD	OE1	OE2		
REMARK	470	GLU	А	547	CG	CD	OE1	OE2		
REMARK	470	GLU	А	548	CG	CD	OE1	OE2		
REMARK	470	LYS	А	606	CG	CD	CE	NZ		
REMARK	470	ARG	в	456	CG	CD	NE	CZ	NH1	NH2
REMARK	470	ASP	в	484	CG	OD1	OD2			
REMARK	470	GLN	в	485	CG	CD	OE1	NE2		
REMARK	470	GLU	В	486	CG	CD	OE1	OE2		
REMARK	470	ARG	в	490	CG	CD	NE	CZ	NH1	NH2
REMARK	470	GLU	в	522	CG	CD	OE1	OE2		
REMARK	470	ARG	В	576	CG	CD	NE	CZ	NH1	NH2

REMARK 470 ASP B 599 CG OD1 OD2

### Template for NMR entries (added)

2 3 4 5 6 7 8 1 **REMARK 470 MISSING ATOM** REMARK 470 THE FOLLOWING RESIDUES HAVE MISSING ATOMS (RES=RESIDUE NAME; REMARK 470 C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE): MODELS X-YYY REMARK 470 REMARK 470 RES CSSEQI ATOMS

The models is listed as a range, X-YYY.

### Example

REMARK 470 MISSING ATOM REMARK 470 THE FOLLOWING RESIDUES HAVE MISSING ATOMS (RES=RESIDUE NAME; REMARK 470 C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE): REMARK 470 MODELS 1-25 REMARK 470 RES CSSEQI ATOMS REMARK 470 ILE A 20 CD1 REMARK 470 THR A 59 CG2

### REMARK 475 (added), Residues modeled with zero occupancy

REMARK 475 enumerates residues modeled with zero occupancy.

#### Template

1 2 3 4 5 6 7 8 REMARK 475 REMARK 475 ZERO OCCUPANCY RESIDUES REMARK 475 THE FOLLOWING RESIDUES WERE MODELED WITH ZERO OCCUPANCY. REMARK 475 THE LOCATION AND PROPERTIES OF THESE RESIDUES MAY NOT REMARK 475 BE RELIABLE. (M=MODEL NUMBER; RES=RESIDUE NAME; REMARK 475 C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE) REMARK 475 M RES C SSEQI

#### Examples

REMARK 475 REMARK 475 REMARK 475 ZERO OCCUPANCY RESIDUES REMARK 475 THE FOLLOWING RESIDUES WERE MODELED WITH ZERO OCCUPANCY. REMARK 475 THE LOCATION AND PROPERTIES OF THESE RESIDUES MAY NOT REMARK 475 BE RELIABLE. (M=MODEL NUMBER; RES=RESIDUE NAME; REMARK 475 C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE) REMARK 475 M RES C SSEQI REMARK 475 DG D 4 REMARK 475 ZERO OCCUPANCY RESIDUES REMARK 475 THE FOLLOWING RESIDUES WERE MODELED WITH ZERO OCCUPANCY.

REMARK 475 THE LOCATION AND PROPERTIES OF THESE RESIDUES MAY NOT

REMARK 475 BE RELIABLE. (M=MODEL NUMBER; RES=RESIDUE NAME; REMARK 475 C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE) REMARK 475 M RES C SSEQI REMARK 475 GLY A 24

### REMARK 480 (added), Polymer atoms modeled with zero occupancy

REMARK 480 enumerates non-hydrogen atoms in residues modeled with zero occupancy.

### Template

5 7 2 3 1 4 6 8 REMARK 480 REMARK 480 ZERO OCCUPANCY ATOM REMARK 480 THE FOLLOWING RESIDUES HAVE ATOMS MODELED WITH ZERO REMARK 480 OCCUPANCY. THE LOCATION AND PROPERTIES OF THESE ATOMS REMARK 480 MAY NOT BE RELIABLE. (M=MODEL NUMBER; RES=RESIDUE NAME; REMARK 480 C=CHAIN IDENTIFIER; SSEO=SEOUENCE NUMBER; I=INSERTION CODE): M RES C SSEQI ATOMS REMARK 480

### Examples

REMARK 480

HIS B

REMARK 480 GLU B 52 CD

26

CB

REMARK 480 REMARK 480 ZERO OCCUPANCY ATOM REMARK 480 THE FOLLOWING RESIDUES HAVE ATOMS MODELED WITH ZERO REMARK 480 OCCUPANCY. THE LOCATION AND PROPERTIES OF THESE ATOMS REMARK 480 MAY NOT BE RELIABLE. (M=MODEL NUMBER; RES=RESIDUE NAME; REMARK 480 C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE): REMARK 480 M RES C SSEQI ATOMS REMARK 480 DC D C4 ' 04' C1' C3' 3 03' REMARK 480 REMARK 480 ZERO OCCUPANCY ATOM REMARK 480 THE FOLLOWING RESIDUES HAVE ATOMS MODELED WITH ZERO REMARK 480 OCCUPANCY. THE LOCATION AND PROPERTIES OF THESE ATOMS REMARK 480 MAY NOT BE RELIABLE. (M=MODEL NUMBER; REMARK 480 RES=RESIDUE NAME; C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; REMARK 480 I=INSERTION CODE): REMARK 480 M RES C SSEQI ATOMS REMARK 480 HIS A 26 CG ND1 CD2 CE1 NE2

CG

OE1 OE2

ND1 CD2 CE1

NE2

# **REMARK 500 (updated), Geometry and Stereochemistry**

REMARK 500 provides further details about the stereochemistry of the structure. This REMARK is generated automatically and may incorporate comments provided by the author. It is currently divided into the subtopics:

- CLOSE CONTACTS IN SAME ASYMMETRIC UNIT,
- CLOSE CONTACTS,
- COVALENT BOND LENGTHS,
- COVALENT BOND ANGLES,
- TORSION ANGLES,
- NON-CIS & NON-TRANS,
- PLANAR GROUPS,
- MAIN CHAIN PLANARITY,
- CHIRAL CENTERS.

Additional subtopics may be added as needed. For close contacts, the cutoff limit is 2.2 Angstroms for non-hydrogen atoms and is 1.6 Angstroms for H and D atoms. These distances are listed in the REMARK 500 for close contacts symmetry.

All the calculations on RMSD deviations include all the atoms present in the coordinates including atoms with zero occupancy.

The calculation of bond and angle deviations for protein entries will be based on the updated Engh & Huber amino acid target values<sup>1</sup>. For nucleic acids, the Parkinson et al., statistics will be used for these calculations<sup>2</sup>. All bonds and angles that deviate more than 6 times from their standard target values will be flagged as a deviation. The PHI/PSI values are based on the Kleywegt-Jones calculations<sup>3</sup>.

The improper CA-C-CB-N angles for chiral centers are calculated and are defined below with 10 degree allowed deviations.

+35 for L amino acids -35 for D amino acids

- +25 to +45 degree range is defined as sp3, L.
  - If D is expected, it gives "WRONG HAND" in the details. If the calculated value is positive and outside this range, it gives "OUTSIDE RANGE" in the details.
- -10 to +10 degree range is defined as sp2, planar.

If it is expected to be sp2 and the value is outside this range, it gives "EXPECTING PLANAR" in the details. If it is expected to be sp3 and the value is within this range, it gives "EXPECTING SP3" in the details.

• -45 to -25 degree range is defined as sp3, D.

<sup>&</sup>lt;sup>1</sup> Structure quality and target parameters. R. A. Engh and R. Huber. International Tables for Crystallography (2006). Vol. F, ch. 18.3, pp. 382-392 <sup>2</sup> "New Parameters for the Refinement of Nucleic Acid Containing Structures." G. Parkinson, J. Vojtechovsky, L. Clowney, A. Brunger\*, and H. M. Berman. (1996) Acta Cryst. D 52, 57-64

<sup>&</sup>lt;sup>3</sup> "PHI/PSI- Chology: Ramachandran revisited. "G.J. Kleywegt and T.A. Jones (1996) Structure 4, 1395-1400.

If L is expected, it gives "WRONG HAND" in the details. If the calculated value is negative and outside this range, it gives "OUTSIDE RANGE" in the details.

The improper CA-C-CB-N angles for chiral centers are calculated for all the alternate conformations. However alt id will not be listed for outliers where residue involves alternate conformations.

### Template

1 2 3 4 5 6 7 8 12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY REMARK 500 SUBTOPIC: REMARK 500 REMARK 500 FREE TEXT GOES HERE.

### Example – Close Contacts in the same asymmetric unit

7 2 5 1 3 4 6 8 REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY REMARK 500 SUBTOPIC: CLOSE CONTACTS IN SAME ASYMMETRIC UNIT REMARK 500 REMARK 500 THE FOLLOWING ATOMS ARE IN CLOSE CONTACT. REMARK 500 REMARK 500 ATM1 RES C SSEOI ATM2 RES C SSEOI DISTANCE REMARK 500 PHE 1 OD2 ASP 1 2.17 N 8 31 REMARK 500 OD2 ASP 1 31 PHE 1 8 2.17 Ν REMARK 500 REMARK 500 THIS ENTRY HAS 104 CLOSE CONTACTS REMARK 500 REMARK 500 REMARK: NULL

### Example – Close Contacts

REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY REMARK 500 SUBTOPIC: CLOSE CONTACTS REMARK 500 REMARK 500 THE FOLLOWING ATOMS THAT ARE RELATED BY CRYSTALLOGRAPHIC REMARK 500 SYMMETRY ARE IN CLOSE CONTACT. AN ATOM LOCATED WITHIN 0.15 REMARK 500 ANGSTROMS OF A SYMMETRY RELATED ATOM IS ASSUMED TO BE ON A REMARK 500 SPECIAL POSITION AND IS, THEREFORE, LISTED IN REMARK 375 REMARK 500 INSTEAD OF REMARK 500. ATOMS WITH NON-BLANK ALTERNATE REMARK 500 LOCATION INDICATORS ARE NOT INCLUDED IN THE CALCULATIONS. REMARK 500 **REMARK 500 DISTANCE CUTOFF:** REMARK 500 2.2 ANGSTROMS FOR CONTACTS NOT INVOLVING HYDROGEN ATOMS REMARK 500 1.6 ANGSTROMS FOR CONTACTS INVOLVING HYDROGEN ATOMS REMARK 500 REMARK 500 ATM1 RES C SSEQI ATM2 RES C SSEQI SSYMOP DISTANCE

2565 REMARK 500 O ALA G 153 OD1 ASP H 46 1.84 REMARK 500 CB ALA G OD1 ASP H 153 46 2565 2.18 REMARK 500 REMARK 500 THIS ENTRY HAS 64 SYMMETRY CONTACTS REMARK 500 REMARK 500 REMARK: NULL

### Example – Covalent bond lengths

2 3 4 5 7 6 REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY REMARK 500 SUBTOPIC: COVALENT BOND LENGTHS REMARK 500 REMARK 500 THE STEREOCHEMICAL PARAMETERS OF THE FOLLOWING RESIDUES REMARK 500 HAVE VALUES WHICH DEVIATE FROM EXPECTED VALUES BY MORE REMARK 500 THAN 6\*RMSD (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN REMARK 500 IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE). REMARK 500 **REMARK 500 STANDARD TABLE:** REMARK 500 FORMAT: (10X, I3, 1X, 2(A3, 1X, A1, I4, A1, 1X, A4, 3X), 1X, F6.3) REMARK 500 REMARK 500 EXPECTED VALUES PROTEIN: ENGH AND HUBER, 1999 REMARK 500 EXPECTED VALUES NUCLEIC ACID: CLOWNEY ET AL 1996 REMARK 500 REMARK 500 M RES CSSEQI ATM1 RES CSSEQI ATM2 DEVIATION REMARK 500 ASP B 117 O 0.129 ASN B 117 C REMARK 500 CYS J 29 CB CYS J 29 SG -0.111 REMARK 500 REMARK 500 REMARK: NULL

### Example – Covalent bond angles

```
REMARK 500
REMARK 500 GEOMETRY AND STEREOCHEMISTRY
REMARK 500 SUBTOPIC: COVALENT BOND ANGLES
REMARK 500
REMARK 500 THE STEREOCHEMICAL PARAMETERS OF THE FOLLOWING RESIDUES
REMARK 500 HAVE VALUES WHICH DEVIATE FROM EXPECTED VALUES BY MORE
REMARK 500 THAN 6*RMSD (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN
REMARK 500 IDENTIFIER; SSEO=SEOUENCE NUMBER; I=INSERTION CODE).
REMARK 500
REMARK 500 STANDARD TABLE:
REMARK 500 FORMAT: (10X,I3,1X,A3,1X,A1,I4,A1,3(1X,A4,2X),12X,F5.1)
REMARK 500
REMARK 500 EXPECTED VALUES: ENGH AND HUBER, 1999
REMARK 500 EXPECTED VALUES NUCLEIC ACID: CLOWNEY ET AL 1996
REMARK 500
REMARK 500 M RES CSSEQI ATM1
                               ATM2
                                      ATM3
REMARK 500 VAL A 124 CB - CA - C
                                           ANGL. DEV. = -12.0 DEGREES
REMARK 500
             PRO B 109
                        CA - N - CD ANGL. DEV. = -3.7 DEGREES
REMARK 500
REMARK 500 REMARK: NULL
```

4 2 1 3 5 6 7 REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY **REMARK 500 SUBTOPIC: TORSION ANGLES** REMARK 500 REMARK 500 TORSION ANGLES OUTSIDE THE EXPECTED RAMACHANDRAN REGIONS: REMARK 500 (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN IDENTIFIER; REMARK 500 SSEQ=SEQUENCE NUMBER; I=INSERTION CODE). REMARK 500 **REMARK 500 STANDARD TABLE:** REMARK 500 FORMAT: (10X, I3, 1X, A3, 1X, A1, I4, A1, 4X, F7.2, 3X, F7.2) REMARK 500 REMARK 500 EXPECTED VALUES: GJ KLEYWEGT AND TA JONES (1996). PHI/PSI-REMARK 500 CHOLOGY: RAMACHANDRAN REVISITED. STRUCTURE 4, 1395 - 1400 REMARK 500 REMARK 500 M RES CSSEQI PSI PHI ASN A 100 -110.87 REMARK 500 -163.72 REMARK 500 ILE A 166 -28.81 -31.64 REMARK 500 REMARK 500 THIS ENTRY HAS 108 RAMACHANDRAN OUTLIERS. REMARK 500 REMARK 500 REMARK: NULL

### Example – Cis/Trans geometry

REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY REMARK 500 SUBTOPIC: NON-CIS, NON-TRANS REMARK 500 REMARK 500 THE FOLLOWING PEPTIDE BONDS DEVIATE SIGNIFICANTLY FROM BOTH REMARK 500 CIS AND TRANS CONFORMATION. CIS BONDS, IF ANY, ARE LISTED REMARK 500 ON CISPEP RECORDS. TRANS IS DEFINED AS 180 +/- 30 AND REMARK 500 CIS IS DEFINED AS 0 +/- 30 DEGREES. REMARK 500 MODEL OMEGA ASP A 414 ASN B 289 REMARK 500 ARG A 413 1 147.84 REMARK 500 ALA B 288 2 -39.12REMARK 500 REMARK 500 REMARK: NULL

### Example – Planar groups

REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY REMARK 500 SUBTOPIC: PLANAR GROUPS REMARK 500 REMARK 500 PLANAR GROUPS IN THE FOLLOWING RESIDUES HAVE A TOTAL REMARK 500 RMS DISTANCE OF ALL ATOMS FROM THE BEST-FIT PLANE REMARK 500 BY MORE THAN AN EXPECTED VALUE OF 6\*RMSD, WITH AN REMARK 500 RMSD 0.02 ANGSTROMS, OR AT LEAST ONE ATOM HAS REMARK 500 AN RMSD GREATER THAN THIS VALUE REMARK 500 (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN IDENTIFIER; REMARK 500 SSEQ=SEQUENCE NUMBER; I=INSERTION CODE). REMARK 500 RMS REMARK 500 M RES CSSEQI TYPE 0.08 REMARK 500 TYR A 36 SIDE CHAIN 0.08 SIDE CHAIN REMARK 500 TYR A 104 REMARK 500 REMARK 500 REMARK: NULL

#### Example – Main chain planarity

2 3 4 5 6 7 8 1 REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY REMARK 500 SUBTOPIC: MAIN CHAIN PLANARITY REMARK 500 REMARK 500 THE FOLLOWING RESIDUES HAVE A PSEUDO PLANARITY REMARK 500 TORSION ANGLE, C(I) - CA(I) - N(I+1) - O(I), GREATER REMARK 500 10.0 DEGREES. (M=MODEL NUMBER; RES=RESIDUE NAME; REMARK 500 C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; REMARK 500 I=INSERTION CODE). REMARK 500 REMARK 500 M RES CSSEQI ANGLE REMARK 500 1 GLY A 289 -10.28 REMARK 500 REMARK 500 REMARK: NULL

4 2 3 5 6 7 REMARK 500 REMARK 500 GEOMETRY AND STEREOCHEMISTRY REMARK 500 SUBTOPIC: CHIRAL CENTERS REMARK 500 REMARK 500 UNEXPECTED CONFIGURATION OF THE FOLLOWING CHIRAL REMARK 500 CENTER(S) USING IMPROPER C--N--CA--CB CHIRALITY REMARK 500 FOR AMINO ACIDS AND C1'--O4'--N1(N9)--C2' FOR REMARK 500 NUCLEIC ACIDS OR EQUIVALENT ANGLE REMARK 500 M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN REMARK 500 IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE REMARK 500 REMARK 500 STANDARD TABLE: REMARK 500 FORMAT: (10X, I3, 1X, A3, 1X, A1, I4, A1, 6X, F5.1, 6X, A1, 10X, A1, 3X, A16) REMARK 500 REMARK 500 M RES CSSEQI IMPROPER EXPECTED FOUND DETAILS L D REMARK 500 16 LEU A 20 0.1 EXPECTING SP3 D -96.0LDOUTSIDE RANGE-54.1LDOUTSIDE RANGE-42.0LDWRONG HAND-96.9LDOUTSIDE RANGE-133.0LDOUTSIDE RANGE53.6LLOUTSIDE RANGE-45.4LDOUTSIDE RANGE-41.3LDWRONG HAND-43.2LDWRONG HAND -96.0 REMARK 500 16 VAL A 21 L OUTSIDE RANGE REMARK 500 16 GLN A 22 REMARK 500 16 THR A 24 REMARK 500 16 LYS A 26 REMARK 500 16 ARG A 29 REMARK 500 16 LEU A 31 REMARK 500 16 LYS A 32 REMARK 500 16 GLU A 33 REMARK 500 16 ASP A 34 REMARK 500 REMARK 500 REMARK: NULL

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# **REMARK 525 (updated), Distant Solvent Atoms**

REMARK 525 lists solvent atoms more than 5 Angstroms from any polymer chain.

### Template

1 2 3 4 5 6 7 8 12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 525 REMARK 525 REMARK 525 formatted text.

### Example

REMARK 525 REMARK 525 SOLVENT REMARK 525 REMARK 525 THE SOLVENT MOLECULES HAVE CHAIN IDENTIFIERS THAT REMARK 525 INDICATE THE POLYMER CHAIN WITH WHICH THEY ARE MOST REMARK 525 CLOSELY ASSOCIATED. THE REMARK LISTS ALL THE SOLVENT REMARK 525 MOLECULES WHICH ARE MORE THAN 5A AWAY FROM THE REMARK 525 NEAREST POLYMER CHAIN (M=MODEL NUMBER; REMARK 525 RES=RESIDUE NAME; C=CHAIN IDENTIFIER; SSEQ=SEQUENCE REMARK 525 NUMBER; I=INSERTION CODE): REMARK 525 REMARK 525 M RES CSSEQI DISTANCE = 6.29 ANGSTROMS REMARK 525 HOH B 89 REMARK 525 HOH B 94 DISTANCE = 5.58 ANGSTROMS

### **REMARK 600, Heterogen**

Further details on the heterogens in the entry.

### Template

```
1 2 3 4 5 6 7 8
12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890
REMARK 600
REMARK 600
REMARK 600
REMARK 600 FREE TEXT GOES HERE.
```

### Example

REMARK 600 REMARK 600 HETEROGEN REMARK 600 REMARK 600 CHAIN A ENDOTHIAPEPSIN: REMARK 600 RESIDUES ASP 54 AND GLY 55 HAVE CYCLISED REMARK 600 TO FORM A SUCCINIMIDE (RESIDUE SUI 54) REMARK 600 REMARK 600 CHAIN B IN THIS PDB ENTRY IS THE REMARK 600 GEM-DIOL INHIBITOR PD-135.040

# REMARK 610 and REMARK 615 (added)

Ligands or hetgroups that are not part of any polymer (protein or nucleic acid) in the structure may also have missing atoms or atoms with zero occupancy. In such instances the name of the hetgroup or ligand, chain ID and model number (if applicable) will be listed in REMARK 610 (for missing atoms) or REMARK 615 (for atoms with 0.00 occupancy). As the list of specific atoms missing from a hetgroup may be really large, they will not listed in the remarks described above. The list of all missing atoms from the ligands may be easily derived by comparing the coordinates of the hetgroup to its definition in the ligand dictionary.

# **REMARK 610, Non-polymer residues with missing atoms**

REMARK 610 enumerates non-polymer residues with missing atoms.

### Example

5 7 2 3 4 1 6 8 REMARK 610 **REMARK 610 MISSING HETEROATOM** REMARK 610 THE FOLLOWING RESIDUES HAVE MISSING ATOMS (M=MODEL NUMBER; REMARK 610 RES=RESIDUE NAME; C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; REMARK 610 I=INSERTION CODE): REMARK 610 M RES C SSEQI REMARK 610 PPI 438

# **REMARK 615, Non-polymer residues containing atoms with zero occupancy**

REMARK 615 enumerates non-polymer residues containing atoms modeled with zero occupancy.

### Example

7 3 4 5 8 1 2 6 REMARK 615 REMARK 615 ZERO OCCUPANCY ATOM REMARK 615 THE FOLLOWING RESIDUES HAVE ATOMS MODELED WITH ZERO REMARK 615 OCCUPANCY. THE LOCATION AND PROPERTIES OF THESE ATOMS REMARK 615 MAY NOT BE RELIABLE. (M=MODEL NUMBER; RES=RESIDUE NAME; REMARK 615 C=CHAIN IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE): REMARK 615 M RES C SSEQI REMARK 615 PPI 438

### **REMARK 620 (added), Metal coordination**

Details of metal coordination are provided in REMARK 620. By default, coordination angles for any metal coordination and surrounding residues (if present) will be provided in this REMARK.

### Template:

2 3 4 5 6 7 8 1 REMARK 620 REMARK 620 METAL COORDINATION REMARK 620 (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN IDENTIFIER; REMARK 620 SSEQ=SEQUENCE NUMBER; I=INSERTION CODE): REMARK 620 REMARK 620 COORDINATION ANGLES FOR: M RES CSSEQI METAL REMARK 620 N RES C SSEQI ATOM

#### Example

REMARK 620 REMARK 620 METAL COORDINATION REMARK 620 (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN IDENTIFIER; REMARK 620 SSEQ=SEQUENCE NUMBER; I=INSERTION CODE): REMARK 620 REMARK 620 COORDINATION ANGLES FOR: M RES CSSEQI METAL REMARK 620 F3S A 107 FE1 REMARK 620 N RES CSSEQI ATOM REMARK 620 1 CYS A 39 SG REMARK 620 2 F3S A 107 FE3 142.2 REMARK 620 3 F3S A 107 FE4 154.3 59.7 REMARK 620 4 F3S A 107 S1 120.2 53.8 55.7 REMARK 620 5 F3S A 107 S2 113.0 103.5 54.3 106.6 REMARK 620 6 F3S A 107 S3 103.8 53.0 101.7 103.2 109.2 REMARK 620 N 1 2 3 4 5 REMARK 620 REMARK 620 COORDINATION ANGLES FOR: M RES CSSEQI METAL REMARK 620 F3S A 107 FE3 REMARK 620 N RES CSSEQI ATOM REMARK 620 1 F3S A 107 FE1 REMARK 620 2 F3S A 107 FE4 59.0 REMARK 620 3 F3S A 107 S1 52.7 55.1 S3 52.9 101.0 102.1 REMARK 620 4 F3S A 107 REMARK 620 5 CYS A 45 SG 146.5 146.2 115.6 112.8 REMARK 620 6 F3S A 107 S4 103.5 54.5 106.3 109.6 110.0 REMARK 620 N 2 1 3 4 5 REMARK 620 REMARK 620 COORDINATION ANGLES FOR: M RES CSSEQI METAL REMARK 620 F3S A 107 FE4 REMARK 620 N RES CSSEQI ATOM REMARK 620 1 F3S A 107 FE1 REMARK 620 2 F3S A 107 FE3 61.3 REMARK 620 3 F3S A 107 S1 53.4 53.9 REMARK 620 4 F3S A 107 S2 54.4 105.0 104.5 REMARK 620 5 CYS A 20 SG 142.7 140.2 109.0 114.5 REMARK 620 6 F3S A 107 S4 105.1 54.1 104.8 111.7 111.6 REMARK 620 N 1 2 3 4 5

REMARK 620 REMARK 620 COORDINATION ANGLES FOR: M RES CSSEQI METAL REMARK 620 F3S A 108 FE1 REMARK 620 N RES CSSEQI ATOM S3 REMARK 620 1 F3S A 108 REMARK 620 2 CYS A 16 SG 120.1 REMARK 620 3 F3S A 108 FE3 51.4 145.9 REMARK 620 4 F3S A 108 FE4 54.3 148.5 59.9 REMARK 620 5 F3S A 108 S1 98.3 110.0 50.6 101.5 REMARK 620 6 F3S A 108 S2 104.2 109.4 104.5 53.3 114.7 REMARK 620 N 1 2 3 4 5 REMARK 620

# **REMARK 630 (added), Inhibitor Description**

Details of inhibitor/peptide inhibitor which is presented as a chemical component (het group) are provided in REMARK 630. By default, molecule type and inhibitor's name will be provided in this REMARK.

#### Template:

3 5 7 1 2 4 6 8 REMARK 630 MOLECULE TYPE: REMARK 630 MOLECULE NAME: REMARK 630 (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN IDENTIFIER; SSSEQ=SEQUENCE NUMBER; I=INSERTION CODE.) REMARK 630 REMARK 630 REMARK 630 M RES C SSSEQI REMARK 630 SOURCE: REMARK 630 TAXONOMY: REMARK 630 SUBCOMP: **REMARK 630 DETAILS:** 

# **REMARK 650, Helix**

Further details on the helical portions of the entry.

#### Template

1 2 3 4 5 6 7 8 12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 650 REMARK 650 HELIX REMARK 650 FREE TEXT GOES HERE.

#### **Examples**

REMARK 650 REMARK 650 HELIX REMARK 650 DETERMINATION METHOD: KDSSP REMARK 650 DETERMINATION METHOD: KDSSP REMARK 650 BETA-BARREL DOMAINS ARE: "N" FOR N-TERMINAL DOMAIN, "B" FOR REMARK 650 BETA-BARREL DOMAIN, AND "C" FOR C-TERMINAL DOMAIN. "F" REMARK 650 REFERS TO THE ACTIVE SITE FLAP. ALPHA HELICES ARE NAMED REMARK 650 WITH TWO CHARACTERS, THE FIRST REFERRING TO THE DOMAIN REMARK 650 IN WHICH THEY OCCUR. REMARK 650 HELIX REMARK 650 DETERMINATION METHOD: AUTHOR PROVIDED.

## **REMARK 700, Sheet**

Further details on the sheet content of the structure. Several standard templates are shown.

#### Template

1 2 3 4 5 6 7 8 1234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 700 REMARK 700 SHEET REMARK 700 FREE TEXT GOES HERE.

#### Examples

REMARK 700 REMARK 700 SHEET **REMARK 700 DETERMINATION METHOD:** REMARK 700 THE SHEET STRUCTURE OF THIS MOLECULE IS BIFURCATED. ΤN REMARK 700 ORDER TO REPRESENT THIS FEATURE IN THE SHEET RECORDS BELOW, REMARK 700 TWO SHEETS ARE DEFINED. STRANDS N1, N2, N3 AND N4 OF SHEET REMARK 700 XXX AND XXX ARE IDENTICAL. REMARK 700 REMARK 700 SHEET **REMARK 700 DETERMINATION METHOD:** REMARK 700 THE SHEET PRESENTED AS XXX ON SHEET RECORDS BELOW IS REMARK 700 ACTUALLY AN N-STRANDED BETA-BARREL. THIS IS REMARK 700 REPRESENTED BY A N+1-STRANDED SHEET IN WHICH THE FIRST AND REMARK 700 LAST STRANDS ARE IDENTICAL. REMARK 700 REMARK 700 SHEET **REMARK 700 DETERMINATION METHOD:** REMARK 700 THERE ARE SEVERAL BIFURCATED SHEETS IN THIS STRUCTURE. REMARK 700 EACH IS REPRESENTED BY TWO SHEETS WHICH HAVE ONE OR MORE REMARK 700 IDENTICAL STRANDS. REMARK 700 SHEETS XXX AND XXX REPRESENT ONE BIFURCATED SHEET. REMARK 700 SHEETS XXX AND XXX REPRESENT ONE BIFURCATED SHEET.

N1, N2, N3 and N4 represent strand numbers, and XXX represents sheet identifiers.

When the remark for several bifurcated sheets is used, its last line is repeated for the appropriate number of bifurcated sheets, as shown in the last template above.

#### Examples

REMARK 700 REMARK 700 SHEET REMARK 700 THE SHEET STRUCTURE OF THIS MOLECULE IS BIFURCATED. IN REMARK 700 ORDER TO REPRESENT THIS FEATURE IN THE SHEET RECORDS BELOW, REMARK 700 TWO SHEETS are defined. STRANDS 3, 4, AND 5 REMARK 700 OF SHEET \*B2A\* AND \*B2B\* ARE IDENTICAL. STRANDS 3, 4, AND REMARK 700 5 OF SHEET \*B2C\* AND \*B2D\* ARE IDENTICAL. REMARK 700

REMARK 700 SHEET REMARK 700 STRANDS 1 TO 4 OF THE BETA-SHEET HAVE GREEK-KEY TOPOLOGY. REMARK 700 THE SHEET FORMS A FIVE-STRANDED BETA-BARREL WITH BULGES IN REMARK 700 STRANDS 3 AND 5. IN ORDER TO REPRESENT THIS FEATURE IN THE REMARK 700 SHEET RECORDS BELOW, TWO SHEETS ARE DEFINED.

REMARK 700 SHEET REMARK 700 THE SHEET PRESENTED AS S5 ON SHEET RECORDS BELOW IS REMARK 700 ACTUALLY A 6-STRANDED BETA-BARREL. THIS IS REMARK 700 REPRESENTED BY A 7-STRANDED SHEET IN WHICH THE FIRST AND REMARK 700 LAST STRANDS ARE IDENTICAL.

REMARK 700 REMARK 700 SHEET REMARK 700 DETERMINATION METHOD: AUTHOR PROVIDED.

# **REMARK 800 (updated), Important Sites**

Further details on important sites of the entry. REMARK 800 is mandatory if SITE records exist.

#### Template

1 2 3 4 5 6 7 8 12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 800 REMARK 800 SITE REMARK 800 SITE\_IDENTIFIER: FREE TEXT GOES HERE. REMARK 800 EVIDENCE\_CODE: (AUTHOR OR SOFTWARE OR UNKNOWN) REMARK 800 SITE DESCRIPTION: FREE TEXT GOES HERE.

\* Site identifiers are 3-letter codes in a character range of AC1-ZZ9 if it is software determined.

#### Examples

REMARK 800 REMARK 800 SITE REMARK 800 SITE IDENTIFIER: RCA REMARK 800 EVIDENCE CODE: AUTHOR REMARK 800 SITE DESCRIPTION: DESIGNATED RECOGNITION REGION IN PRIMARY REMARK 800 REFERENCE. PROPOSED TO AFFECT SUBSTRATE SPECIFICITY. REMARK 800 REMARK 800 SITE\_IDENTIFIER: RCB REMARK 800 EVIDENCE CODE: AUTHOR REMARK 800 SITE DESCRIPTION: DESIGNATED RECOGNITION REGION IN PRIMARY REMARK 800 REFERENCE. PROPOSED TO AFFECT SUBSTRATE SPECIFICITY. REMARK 800 REMARK 800 SITE REMARK 800 SITE IDENTIFIER: AC1 REMARK 800 EVIDENCE CODE: SOFTWARE REMARK 800 SITE DESCRIPTION: BINDING SITE FOR RESIDUE BAT A 19 REMARK 800 REMARK 800 SITE\_IDENTIFIER: AC2 REMARK 800 EVIDENCE CODE: SOFTWARE REMARK 800 SITE DESCRIPTION: BINDING SITE FOR RESIDUE CA A 1 REMARK 800 REMARK 800 SITE\_IDENTIFIER: AC3 REMARK 800 EVIDENCE CODE: SOFTWARE REMARK 800 SITE DESCRIPTION: BINDING SITE FOR RESIDUE BIL A 20

#### Relationship to other records:

Remark 800 is mandatory if site records exist.

#### **REMARK 900, Related Entries**

This REMARK provides information about other PDB entries related to the entry. These may include coordinate entries deposited as a related set, an EMDB identifier for the related EM map, a BMRB identifier for the related NMR chemical shifts, or a structural genomics target identifier.

#### Template

1 2 3 4 5 6 7 8 1234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 900 REMARK 900 RELATED ENTRIES REMARK 900 FREE TEXT GOES HERE.

#### Examples

REMARK 900
REMARK 900 RELATED ENTRIES
REMARK 900 RELATED ID: 2GB8 RELATED DB: PDB
REMARK 900 SOLUTION STRUCTURE OF WT CC-CCP COMPLEX
REMARK 900 RELATED ID: 2PCC RELATED DB: PDB
REMARK 900 CRYSTAL STRUCTURE OF WT CC-CCP COMPLEX
REMARK 900 RELATED ID: 1YCC RELATED DB: PDB
REMARK 900 CRYSTAL STRUCTURE OF YEAST ISO-1-CYTOCHROME C
REMARK 900 RELATED ID: 1ZBY RELATED DB: PDB
REMARK 900 HIGH-RESOLUTION CRYSTAL STRUCTURE OF YEAST CYTOCHROME C
REMARK 900 PEROXIDASE

REMARK 900 REMARK 900 RELATED ENTRIES REMARK 900 RELATED ID: STR82 RELATED DB: TARGETDB REMARK 900 RELATED ID: 15386 RELATED DB: BMRB

# **REMARK 999, Sequence**

This remark is a free text remark which describes anything unusual about a particular polymer sequence in SEQRES records.

For examples,

- 1. If the exact sequence of the sample is not known, due to, for example, proteolysis, the sequence should match the coordinates and a REMARK 999 can be added.
- 2. The information about a sequence region of a chimeric protein which does not match the UNP entry, such as a linker region, can be added to REMARK 999.
- 3. Sequence conflicts which are listed in the UNP reference can also be described in REMARK 999. A full explanation of the microheterogeneity for all residues at a particular residue number can be elaborated in REMARK 999.
- If the coordinates alignment with the sequence is unknown and the residue numbering is arbitrary. The sequence would be poly UNK. The sequence, if it is known, would be listed in the REMARK 999

### Template

1 2 3 4 5 6 7 8 1234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890 REMARK 999 REMARK 999 SEQUENCE REMARK 999 FREE TEXT GOES HERE.

## Example

REMARK 999 **REMARK 999 SEQUENCE** REMARK 999 THE N-TERMINAL 19 RESIDUES 'GSHMVPGQKQHYVQPTAAN' REMARK 999 CORRESPOND TO A PHAGE-DISPLAY DERIVED PEPTIDE, REMARK 999 WHICH IS FUSED TO THE SECRETION CHAPERONE PROTEIN REMARK 999 **REMARK 999 SEOUENCE** REMARK 999 THE SEQUENCE USED IS THAT PROVIDED BY THE CDNA, WHICH REMARK 999 CORRECTS SEVERAL ASP/ASN AND GLU/GLN MISASSIGNMENTS. REMARK 999 **REMARK 999 SEQUENCE** REMARK 999 THR AT POSITION 74 WAS FOUND BY WOLMAN ET AL., JOURNAL OF REMARK 999 BIOCHEMISTRY 263, 15506 (1988). **REMARK 999 SEQUENCE** REMARK 999 THE INSERTED RESIDUES AT THE N-TERMINUS OF THE PROTEIN REMARK 999 CORRESPOND TO A 32-RESIDUE DSE3 LANTHIDE-BINDING TAG REMARK 999 THE RESIDUES NUMBERED 66 TO 100 IN THIS ENTRY CORRESPOND

REMARK 999 TO RESIDUES -4 TO 13 AND -1' TO 15' IN THE PRIMARY CITATION.

# 3. Primary Structure Section

The primary structure section of a PDB formatted file contains the sequence of residues in each chain of the macromolecule(s). Embedded in these records are chain identifiers and sequence numbers that allow other records to link into the sequence.

# **DBREF** (standard format)

The DBREF record provides cross-reference links between PDB sequences (what appears in SEQRES record) and a corresponding database sequence.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"DBREF "	
8 - 11	IDcode	idCode	ID code of this entry.
13	Character	chainID	Chain identifier.
15 - 18	Integer	seqBegin	Initial sequence number of the PDB sequence segment.
19	AChar	insertBegin	Initial insertion code of the PDB sequence segment.
21 - 24	Integer	seqEnd	Ending sequence number of the PDB sequence segment.
25	AChar	insertEnd	Ending insertion code of the PDB sequence segment.
27 - 32	LString	database	Sequence database name.
34 - 41	LString	dbAccession	Sequence database accession code.
43 - 54	LString	dbIdCode	Sequence database identification code.
56 - 60	Integer	dbseqBegin	Initial sequence number of the database seqment.
61	AChar	idbnsBeg	Insertion code of initial residue of the segment, if PDB is the reference.
63 - 67	Integer	dbseqEnd	Ending sequence number of the database segment.
68	AChar	dbinsEnd	Insertion code of the ending residue of the segment, if PDB is the reference.

Note: By default this format is used as long as the information entered into these fields fits. For

sequence databases that use longer accession code or long sequence numbering, the new DBREF1/DBREF2 format can be used.

#### Details

\* PDB entries contain multi-chain molecules with sequences that may be wild type, variant, or synthetic. Sequences may also have been modified through site-directed mutagenesis experiments (engineered). A number of PDB entries report structures of individual domains cleaved from larger molecules.

The DBREF records present sequence correlations between PDB SEQRES records and corresponding GenBank (for nucleic acids) or UNIPROT/Norine (for proteins) entries. PDB entries containing heteropolymers are linked to different sequence database entries.

\* Database names and their abbreviations as used on DBREF records.

Database name	Database abbreviations (columns 27 - 32)
GenBank	GB
Protein Data Bank	PDB
UNIPROT	UNP
Norine	NORINE

\* wwPDB does not guarantee that all possible references to the listed databases will be provided. In most cases, only one reference to a sequence database will be provided.

\* If no reference is found in the sequence databases, then the PDB entry itself is given as the reference.

\* Selection of the appropriate sequence database entry or entries to be linked to a PDB entry is done on the basis of the sequence and its biological source. Questions on entry assignment that may arise are resolved by consultation with the database.

### Verification/Validation/Value Authority Control

The sequence database entry found during PDB's search is compared to that provided by the depositor and any differences are resolved or annotated.

All polymers in the entry will be assigned a DBREF record.

### **Relationships to Other Record Types**

DBREF represents the sequence as found in SEQRES records.

DBREF1/DBREF2 replaces DBREF when the accession codes or sequence numbering does not fit the DBREF format.

# Examples

•	1		2	3	4	5	6	7	8
123456	78901234	56789	012345	6789012	345678901	2345678901234	456789012	34567890	1234567890
DBREF	2JHQ A	1	226	UNP	Q9KPK8	UNG_VIBCH	1	226	
DBREF	ЗАКҮ А	1	219	UNP	P07170	KAD1_YEAST	3	221	
DBREF	1HAN A	2	298	UNP	P47228	BPHC_BURCE	1	297	
DBREF	3D3I A	0	760	UNP	P42592	YGJK_ECOLI	23	783	
DBREF	3D3I B	0	760	UNP	P42592	YGJK_ECOLI	23	783	
DBREF	3C2J A	1	8	PDB	3C2J	3C2J	1	8	
DBREF	3C2J B	101	108	PDB	3C2J	3C2J	101	108	
DBREF	1FFK 0	2	2923	GB	3377779	AF034620	2597	5518	
DBREF	1FFK 9	1	122	GB	3377779	AF034620	5658	5779	
DBREF	1UNJ X	6	11	NOR	NOR00228	NOR00228	6	11	

# DBREF1 / DBREF2 (added)

### Details

This updated two-line format is used when the accession code or sequence numbering does not fit the space allotted in the standard DBREF format. This includes some GenBank sequence numbering (greater than 5 characters) and UNIMES accession numbers (greater than 12 characters).

## **Record Format**

## DBREF1

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"DBREF1"	
8 - 11	IDcode	idCode	ID code of this entry.
13	Character	chainID	Chain identifier.
15 - 18	Integer	seqBegin	Initial sequence number of the PDB sequence segment, right justified.
19	AChar	insertBegin	Initial insertion code of the PDB sequence segment.
21 - 24	Integer	seqEnd	Ending sequence number of the PDB sequence segment, right justified.
25	AChar	insertEnd	Ending insertion code of the PDB sequence segment.
27 - 32	LString	database	Sequence database name.
48 - 67	LString	dbIdCode	Sequence database identification code, left justified.

## DBREF2

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"DBREF2"	
8 - 11	IDcode	idCode	ID code of this entry.
13	Character	chainID	Chain identifier.
19 - 40	LString	dbAccession	Sequence database accession code, left justified.
46 - 55	Integer	seqBegin	Initial sequence number of the Database segment, right justified.
58 - 67	Integer	seqEnd	Ending sequence number of the

Database segment, right justified.

### Details

\* The DBREF1/DBREF2 record presents sequence correlations between PDB SEQRES records and corresponding GenBank (for nucleic acids) or UNIMES (for proteins) entries. Several cases are easily represented by means of pointers between the databases using DBREF.

\* Database names and their abbreviations as used as in DBREF records.

Database name	Database abbreviations (columns 27 - 32)
GenBank	GB
UNIMES	UNIMES

\* wwPDB does not guarantee that all possible references to the listed databases will be provided. In most cases, only one reference to a sequence database will be provided.

### Verification/Validation/Value Authority Control

The sequence database entry found by wwPDB staff is compared to answers provided by the depositor; any differences are resolved or annotated appropriately.

### **Relationships to Other Record Types**

DBREF1/DBREF2 represents the sequence as found in SEQRES records.

### Template

1	2	3	4	5	6	7	8
123456789012	345678901234	5678901234	5678901234	567890123456	578901234	5678901234	567890
DBREF1 2J83	A 61 322	XXXXXX		YYYYYYYY	YYYYYYYY	YYY	
DBREF2 2J83	A ZZZZZZ	Z Z Z Z Z Z Z Z Z Z Z Z	ZZZZZZ	nnnnnnnn	mmmmmmm	mmm	

### Examples

1	2	3	4	5	6	7	8
123456789012345	67890123456	7890123	456789012345	6789012345	678901234	5678901234	567890
DBREF1 2J83 A	61 322	UNIMES		UPI00014	8A153		
DBREF2 2J83 A	MES00005	880000		61		322	
1	2	3	4	5	6	7	8
123456789012345	67890123456	7890123	456789012345	6789012345	678901234	5678901234	567890
DBREF1 2J83 A	61 322	GB		AE017221			
DBREF2 2J83 A	46197919	)		1534489	1537	377	

# SEQADV

**Overview** 

The SEQADV record identifies differences between sequence information in the SEQRES records of the PDB entry and the sequence database entry given in DBREF. Please note that these records were designed to identify differences and not errors. No assumption is made as to which database contains the correct data. A comment explaining any engineered differences in the sequence between the PDB and the sequence database may also be included here.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"SEQADV"	
8 - 11	IDcode	idCode	ID code of this entry.
13 - 15	Residue name	resName	Name of the PDB residue in conflict.
17	Character	chainID	PDB chain identifier.
19 - 22	Integer	seqNum	PDB sequence number.
23	AChar	iCode	PDB insertion code.
25 - 28	LString	database	
30 - 38	LString	dbIdCode	Sequence database accession number.
40 - 42	Residue name	dbRes	Sequence database residue name.
44 - 48	Integer	dbSeq	Sequence database sequence number.
50 - 70	LString	conflict	Conflict comment.

## Details

\* In a number of cases, conflicts between the sequences found in PDB entries and in sequence database reference entries have been noted. There are several possible reasons for these conflicts, including natural variants or engineered sequences (mutants), polymorphic sequences, or ambiguous or conflicting experimental results. These discrepancies are reported in SEQADV. Additional details may be included in remark 999.

\* When conflicts arise which are not classifiable by these terms, a reference to either a published paper, a PDB entry, or a REMARK within the entry is given.

\* The comment "SEE REMARK 999" is included when the explanation for the conflict is too long to fit the SEQADV record.

\* Some of the possible conflict comments:

- Cloning artifact
- Expression tag
- Conflict
- Engineered

- Variant
- Insertion
- Deletion
- Microheterogeneity
- Chromophore

\* Microheterogeneity is to be represented as a variant with one of the possible residues in the site being selected (arbitrarily) as the primary residue. The residues which do not match to the UNP reference will be listed in SEQADV records with the explanation of "microheterogeneity".

#### Verification/Validation/Value Authority Control

SEQADV records are automatically generated.

#### **Relationships to Other Record Types**

SEQADV refers to the sequence as found in the SEQRES records, and to the sequence database reference found on DBREF.

REMARK 999 contains text that explains discrepancies when the explanation is too lengthy to fit in SEQADV.

#### **Examples**

—	2 012345678	3 89012345678	4 9012345	5 6 7 8 56789012345678901234567890
SEQADV 3ABC MET A	-1 UNP	P10725		EXPRESSION TAG
SEQADV 3ABC GLY A	50 UNP	P10725	VAL	50 ENGINEERED
SEQADV 2QLE CRO A	66 UNP	P42212	SER	65 CHROMOPHORE
SEQADV 20KW LEU A	64 UNP	P42212	PHE	64 SEE REMARK 999
SEQADV 20KW LEU A	64 NOR	NOR00669	PHE	14 SEE REMARK 999

# **SEQRES** (updated)

### Overview

SEQRES records contain a listing of the consecutive chemical components covalently linked in a linear fashion to form a polymer. The chemical components included in this listing may be standard or modified amino acid and nucleic acid residues. It may also include other residues that are linked to the standard backbone in the polymer. Chemical components or groups covalently linked to side-chains (in peptides) or sugars and/or bases (in nucleic acid polymers) will not be listed here.

# **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"SEQRES"	
8 - 10	Integer	serNum	Serial number of the SEQRES record for the current chain. Starts at 1 and increments by one each line. Reset to 1 for each chain.
12	Character	chainID	Chain identifier. This may be any single legal character, including a blank which is is used if there is only one chain.
14 - 17	Integer	numRes	Number of residues in the chain. This value is repeated on every record.
20 - 22	Residue name	resName	Residue name.
24 - 26	Residue name	resName	Residue name.
28 - 30	Residue name	resName	Residue name.
32 - 34	Residue name	resName	Residue name.
36 - 38	Residue name	resName	Residue name.
40 - 42	Residue name	resName	Residue name.
44 - 46	Residue name	resName	Residue name.
48 - 50	Residue name	resName	Residue name.
52 <b>-</b> 54	Residue name	resName	Residue name.
56 <b>-</b> 58	Residue name	resName	Residue name.
60 - 62	Residue name	resName	Residue name.
64 - 66	Residue name	resName	Residue name.
68 - 70	Residue name	resName	Residue name.

### Verification/Validation/Value Authority Control

The residues presented in the ATOM records must agree with those on the SEQRES records.

The SEQRES records are checked using sequence databases and information provided by the depositor.

SEQRES is compared to the ATOM records during processing, and both are checked against the sequence databases. All discrepancies are either resolved or annotated appropriately in the entry.

The ribo- and deoxyribonucleotides in the SEQRES records are distinguished. The ribo- forms of these residues are identified with the residue names A, C, G, U and I. The deoxy- forms of these residues are identified with the residue names DA, DC, DG, DT and DI. Modified nucleotides in the sequence are identified by separate 3-letter residue codes. The *plus* character prefix to label modified nucleotides (e.g. +A, +C, +T) is no longer used.

### Example

	1		2		3			4		5			6		7		8
12345678	901234	15678	90123	34567	78901	L234	56789	90123	3456	78901	12345	56789	90123	34567	78901	234567	890
SEQRES	1 A	21	GLY	ILE	VAL	$\operatorname{GLU}$	$\operatorname{GLN}$	CYS	CYS	THR	SER	ILE	CYS	SER	LEU		
SEQRES	2 A	21	TYR	$\operatorname{GLN}$	LEU	GLU	ASN	TYR	CYS	ASN							
SEQRES	1 B	30	PHE	VAL	ASN	$\operatorname{GLN}$	HIS	LEU	CYS	GLY	SER	HIS	LEU	VAL	GLU		
SEQRES	2 B	30	ALA	LEU	TYR	LEU	VAL	CYS	GLY	GLU	ARG	GLY	PHE	PHE	TYR		
SEQRES	3 B	30	THR	PRO	LYS	ALA											
SEQRES	1 C	21	GLY	ILE	VAL	GLU	$\operatorname{GLN}$	CYS	CYS	THR	SER	ILE	CYS	SER	LEU		
SEQRES	2 C	21	TYR	GLN	LEU	GLU	ASN	TYR	CYS	ASN							
SEQRES	1 D	30	$\mathbf{PHE}$	VAL	ASN	$\operatorname{GLN}$	HIS	LEU	CYS	GLY	SER	HIS	LEU	VAL	GLU		
SEQRES	2 D	30	ALA	LEU	TYR	LEU	VAL	CYS	GLY	GLU	ARG	GLY	PHE	PHE	TYR		
SEQRES	3 D	30	THR	PRO	LYS	ALA											
SEQRES	1 A	8	DA	DA	DC	DC	DG	DG	DT	DT							
SEQRES	1 B	8	DA	DA	DC	DC	DG	DG	DT	DT							
SEQRES	1 X	39	U	С	С	С	С	С	G	U	G	С	С	С	А		
SEQRES	2 X	39	U	А	G	С	G	G	С	G	U	G	G	А	А		
SEQRES	3 X	39	C	С	A	С	С	С	G	U	U	С	С	С	A		

## **Known Problems**

Polysaccharides do not lend themselves to being represented in SEQRES.

There is no mechanism provided to describe the sequence order if their starting position is unknown.

For cyclic peptides, a residue is arbitrarily assigned as the N-terminus.

# **MODRES** (updated)

#### **Overview**

The MODRES record provides descriptions of modifications (e.g., chemical or post-translational) to protein and nucleic acid residues. Included are correlations between residue names given in a PDB entry and standard residues.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"MODRES"	
8 - 11	IDcode	idCode	ID code of this entry.
13 - 15	Residue name	resName	Residue name used in this entry.
17	Character	chainID	Chain identifier.
19 - 22	Integer	seqNum	Sequence number.
23	AChar	iCode	Insertion code.
25 - 27	Residue name	stdRes	Standard residue name.
30 - 70	String	comment	Description of the residue modification.

### Details

\* Residues modified post-translationally, enzymatically, or by design are described in MODRES records. In those cases where the wwPDB has opted to use a non-standard residue name for the residue, MODRES also correlates the new name to the precursor standard residue name.

\* Modified nucleotides in the sequence are now identified by separate 3-letter residue codes. The *plus* character prefix to label modified nucleotides (e.g. +A, +C, +T) is no longer used.

\* MODRES is mandatory when modified standard residues exist in the entry. Examples of some modification descriptions:

- Glycosylation site
- Post-translational modification
- Designed chemical modification
- Phosphorylation site
- D-configuration

\* A MODRES record is not required if coordinate records are not provided for the modified residue.

\* D-amino acids are given their own residue name (resName), i.e., DAL for D-alanine. This resName appears in the SEQRES records, and has the associated MODRES, HET, and FORMUL records. The coordinates are given as HETATMs within the ATOM records and occur in the correct order within the chain. This ordering is an exception to the stated Order of Records.

\* When a standard residue name is used to describe a modified site, resName (columns 13-15) and stdRES (columns 25-27) contain the same value.

#### Verification/Validation/Value Authority Control

MODRES is generated by the wwPDB.

### **Relationships to Other Record Types**

MODRES maps ATOM and HETATM records to the standard residue names. HET, and FORMUL may also appear.

#### Example

3 7 1 2 4 5 6 8 MODRES 2R0L ASN A 74 ASN GLYCOSYLATION SITE 1N-METHYLGUANOSINE-5'-MONOPHOSPHATE MODRES 11L2 1MG D 1937 G MODRES 4ABC MSE B 32 MET SELENOMETHIONINE

# 4. Heterogen Section (updated)

The heterogen section of a PDB formatted file contains the complete description of non-standard residues in the entry. Detailed chemical definitions of non-polymer chemical components are described in the Chemical Component Dictionary (<u>ftp://ftp.wwpdb.org/pub/pdb/data/monomers</u>)

# HET

HET records are used to describe non-standard residues, such as prosthetic groups, inhibitors, solvent molecules, and ions for which coordinates are supplied. Groups are considered HET if they are not part of a biological polymer described in SEQRES and considered to be a molecule bound to the polymer, or they are a chemical species that constitute part of a biological polymer and is not one of the following:

- standard amino acids, or
- standard nucleic acids (C, G, A, U, I, DC, DG, DA, DU, DT and DI), or
- unknown amino acid (UNK) or nucleic acid (N) where UNK and N are used to indicate the unknown residue name.

HET records also describe chemical components for which the chemical identity is unknown, in which case the group is assigned the hetID UNL (Unknown Ligand).

The heterogen section of a PDB formatted file contains the complete description of non-standard residues in the entry.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"HET "	
8 - 10	LString(3)	hetID	Het identifier, right-justified.
13	Character	ChainID	Chain identifier.
14 - 17	Integer	seqNum	Sequence number.
18	AChar	iCode	Insertion code.
21 - 25	Integer	numHetAtoms	Number of HETATM records for the group present in the entry.
31 - 70	String	text	Text describing Het group.

### Details

\* Each HET group is assigned a hetID of not more than three (3) alphanumeric characters. The sequence number, chain identifier, insertion code, and number of coordinate records are given for each occurrence of the HET group in the entry. The chemical name of the HET group is given in the HETNAM record and synonyms for the chemical name are given in the HETSYN records, see <a href="http://ftp.wwpdb.org/pub/pdb/data/monomers">http://ftp.wwpdb.org/pub/pdb/data/monomers</a>.

\* There is a separate HET record for each occurrence of the HET group in an entry.

\* A particular HET group is represented in the PDB archive with a unique hetID.

\* PDB entries do not have HET records for water molecules, deuterated water, or methanol (when used as solvent).

\* Unknown atoms or ions will be represented as UNX with the chemical formula X1. Unknown ligands are UNL; unknown amino acids are UNK.

#### Verification/Validation/Value Authority Control

For each het group that appears in the entry, the wwPDB checks that the corresponding HET, HETNAM, HETSYN, FORMUL, HETATM, and CONECT records appear, if applicable. The HET record is generated automatically using the Chemical Component Dictionary and information from the HETATM records.

Each unique hetID represents a unique molecule.

#### **Relationships to Other Record Types**

For each het group that appears in the entry, there must be corresponding HET, HETNAM, HETSYN, FORMUL, HETATM, and CONECT records. LINK records may also be created.

#### Example

3 2 4 5 6 7 1 HET TRS 975 8 HET UDP A1457 25 HET B3P A1458 19 15 HET NAG Y 3 HET FUC Y 4 10 NON Y HET5 12 HET UNK A 161 1

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# HETNAM

#### **Overview**

This record gives the chemical name of the compound with the given hetID.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"HETNAM"	
9 - 10	Continuation	continuation	Allows concatenation of multiple records.
12 - 14	LString(3)	hetID	Het identifier, right-justified.
16 - 70	String	text	Chemical name.

## Details

\* Each hetID is assigned a unique chemical name for the HETNAM record, see <a href="http://ftp.wwpdb.org/pub/pdb/data/monomers">http://ftp.wwpdb.org/pub/pdb/data/monomers</a> .

\* Other names for the group are given on HETSYN records.

\* PDB entries follow IUPAC/IUB naming conventions to describe groups systematically.

\* The special character "~" is used to indicate superscript in a heterogen name. For example:  $N^6$  will be listed in the HETNAM section as N~6~, with the ~ character indicating both the start and end of the superscript in the name, e.g.,

N- (BENZYLSULFONYL) SERYL-N~1~- { 4- [AMINO (IMINO) METHYL] BENZYL } GLYCINAMIDE

\* Continuation of chemical names onto subsequent records is allowed.

\* Only one HETNAM record is included for a given hetID, even if the same hetID appears on more than one HET record.

### Verification/Validation/Value Authority Control

For each het group that appears in the entry, the corresponding HET, HETNAM, FORMUL, HETATM, and CONECT records must appear. The HETNAM record is generated automatically using the Chemical Component Dictionary and information from HETATM records.

## **Relationships to Other Record Types**

For each het group that appears in the entry, there must be corresponding HET, HETNAM, FORMUL, HETATM, and CONECT records. HETSYN and LINK records may also be created.

# Example

	1		2	3	4	5	6	7	8
1234567	890	1234	567890123456789	012345678	901234567	8901234	56789012345	6789012345	67890
HETNAM		NAG	N-ACETYL-D-GLU	COSAMINE					
HETNAM		SAD	BETA-METHYLENE	SELENAZO	LE-4-CARE	OXAMIDE	ADENINE		
HETNAM	2	SAD	DINUCLEOTIDE						
HETNAM		UDP	URIDINE-5'-DIP	HOSPHATE					
HETNAM		UNX	UNKNOWN ATOM O	R ION					
HETNAM		UNL	UNKNOWN LIGAND	)					
HETNAM		B3P	2-[3-(2-HYDROX	Y-1,1-DIH	YDROXYMET	HYL-ETH	YLAMINO)-		
HETNAM	2	B3P	PROPYLAMINO]-	2-HYDROXY	METHYL-PR	OPANE-1	,3-DIOL		

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# HETSYN

#### **Overview**

This record provides synonyms, if any, for the compound in the corresponding (i.e., same hetID) HETNAM record. This is to allow greater flexibility in searching for HET groups.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"HETSYN"	
9 - 10	Continuation	continuation	Allows concatenation of multiple records.
12 - 14	LString(3)	hetID	Het identifier, right-justified.
16 - 70	SList	hetSynonyms	List of synonyms.

## Details

\* The wwPDB does not guarantee a complete list of possible synonyms. New synonyms may be added. The list can be continued onto additional HETSYN records. Even if the same hetID appears on more than one HET record, only one set of HETSYN records is included for the hetID.

### Verification/Validation/Value Authority Control

For each HETSYN record in the entry, the corresponding HET, HETNAM, FORMUL, HETATM, and CONECT records must appear.

### **Relationships to Other Record Types**

If there is a HETSYN record there must be corresponding HET, HETNAM, FORMUL, HETATM, and CONECT records. LINK records may also be created.

### Example

# FORMUL

#### **Overview**

The FORMUL record presents the chemical formula and charge of a non-standard group.

### **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"FORMUL"	
9 - 10	Integer	compNum	Component number.
13 - 15	LString(3)	hetID	Het identifier.
17 - 18	Integer	continuation	Continuation number.
19	Character	asterisk	"*" for water.
20 - 70	String	text	Chemical formula.

## Details

\* The elements of the chemical formula are given in the order following Hill ordering. The order of elements depends on whether carbon is present or not. If carbon is present, the order should be: C, then H, then the other elements in alphabetical order of their symbol. If carbon is not present, the elements are listed purely in alphabetic order of their symbol. This is the 'Hill' system used by Chemical Abstracts.

\* The number of each atom type present immediately follows its chemical symbol without an intervening blank space. There will be no number indicated if there is only one atom for a particular atom type.

\* Each set of SEQRES records and each HET group is assigned a component number in an entry. These numbers are assigned serially, beginning with 1 for the first set of SEQRES records. In addition:

- If a HET group is presented on a SEQRES record its FORMUL is assigned the component number of the chain in which it appears.
- If the HET group occurs more than once and is not presented on SEQRES records, the component number of its first occurrence is used.

\* All occurrences of the HET group within a chain are grouped together with a multiplier. The remaining occurrences are also grouped with a multiplier. The sum of the multipliers is the number equaling the number of times that HET group appears in the entry.

\* A continuation field is provided in the event that more space is needed for the formula. Columns 17 - 18 are used in order to maintain continuity with the existing format.

### Verification/Validation/Value Authority Control

For each het group that appears in the entry, the corresponding HET, HETNAM, FORMUL, HETATM, and CONECT records must appear. The FORMUL record is generated automatically by PDB processing programs using the het group template file and information from HETATM records. UNL, UNK and UNX will not be listed in FORMUL even though these het groups present in the coordinate section.

#### **Relationships to Other Record Types**

For each het group that appears in the entry, the corresponding HET, HETNAM, FORMUL, HETATM, and CONECT records must appear.

#### Example

3 5 7 1 2 4 6 8 FORMUL 3 MG 2(MG 2+) 5 SO4 6(04 S 2-) FORMUL FORMUL 13 HOH \*360(H2 O) 2(C21 H28 N7 O17 P3) FORMUL 3 NAP 4 FOL 2(C19 H19 N7 O6) FORMUL FORMUL 5 1PE C10 H22 O6 C14 H10 O2 CL2 S FORMUL 2 NX5

#### **Known Problems**

Partially deuterated centers are not well represented in this record.

# 5. Secondary Structure Section

The secondary structure section of a PDB formatted file describes helices, sheets, and turns found in protein and polypeptide structures.

# HELIX

#### **Overview**

HELIX records are used to identify the position of helices in the molecule. Helices are named, numbered, and classified by type. The residues where the helix begins and ends are noted, as well as the total length.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"HELIX "	
8 - 10	Integer	serNum	Serial number of the helix. This starts at 1 and increases incrementally.
12 - 14	LString(3)	helixID	Helix identifier. In addition to a serial number, each helix is given an alphanumeric character helix identifier.
16 - 18	Residue name	initResName	Name of the initial residue.
20	Character	initChainID	Chain identifier for the chain containing this helix.
22 - 25	Integer	initSeqNum	Sequence number of the initial residue.
26	AChar	initICode	Insertion code of the initial residue.
28 - 30	Residue name	endResName	Name of the terminal residue of the helix.
32	Character	endChainID	Chain identifier for the chain containing this helix.
34 - 37	Integer	endSeqNum	Sequence number of the terminal residue.
38	AChar	endICode	Insertion code of the terminal residue.
39 - 40	Integer	helixClass	Helix class (see below).
41 - 70	String	comment	Comment about this helix.
72 – 76 <b>Details</b>	Integer	length	Length of this helix.

\* Additional HELIX records with different serial numbers and identifiers occur if more than one helix is present.

\* The initial residue of the helix is the N-terminal residue.

\* Helices are classified as follows:

TYPE OF HELIX	CLASS NUMBER (COLUMNS 39 - 40)
Right-handed alpha (default) Right-handed omega Right-handed pi Right-handed gamma Right-handed 310 Left-handed alpha Left-handed omega Left-handed gamma 27 ribbon/helix Polyproline	1 2 3 4 5 6 7 8 9 10

# **Relationships to Other Record Types**

There may be related information in the REMARKs.

## Example

	1			2		3			4	5	5	6	7	8
1234567	8901	2345	56789	901	23456	78901	.23	45678	9012	234567890	012345	6789012345	67890123	4567890
HELIX	1	HA	$\operatorname{GLY}$	Α	86	GLY	А	94	1					9
HELIX	2	HB	GLY	В	86	GLY	В	94	1					9
HELIX	21	21	PRO	J	385	LEU	J	388	5					4
HELIX	22	22	PHE	J	397	PHE	J	402	5					6

# SHEET

### **Overview**

SHEET records are used to identify the position of sheets in the molecule. Sheets are both named and numbered. The residues where the sheet begins and ends are noted.

# **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"SHEET "	
8 - 10	Integer	strand	Strand number which starts at 1 for each strand within a sheet and increases by one.
12 - 14	LString(3)	sheetID	Sheet identifier.
15 - 16	Integer	numStrands	Number of strands in sheet.
18 - 20	Residue name	initResName	Residue name of initial residue.
22	Character	initChainID	Chain identifier of initial residue in strand.
23 - 26	Integer	initSeqNum	Sequence number of initial residue in strand.
27	AChar	initICode	Insertion code of initial residue in strand.
29 - 31	Residue name	endResName	Residue name of terminal residue.
33	Character	endChainID	Chain identifier of terminal residue.
34 - 37	Integer	endSeqNum	Sequence number of terminal residue.
38	AChar	endICode	Insertion code of terminal residue.
39 - 40	Integer	sense	Sense of strand with respect to previous strand in the sheet. 0 if first strand, 1 if parallel,and -1 if anti-parallel.
42 - 45	Atom	curAtom	Registration. Atom name in current strand.
46 - 48	Residue name	curResName	Registration. Residue name in current strand
50	Character	curChainId	Registration. Chain identifier in current strand.
51 - 54	Integer	curResSeq	Registration. Residue sequence number in current strand.
55	AChar	curICode	Registration. Insertion code in current strand.

57 - 60	Atom	prevAtom	Registration. Atom name in previous strand.
61 - 63	Residue name	prevResName	Registration. Residue name in previous strand.
65	Character	prevChainId	Registration. Chain identifier in previous strand.
66 - 69	Integer	prevResSeq	Registration. Residue sequence number in previous strand.
70	AChar	prevICode	Registration. Insertion code in previous strand.

#### Details

\* The initial residue for a strand is its N-terminus. Strand registration information is provided in columns 39 - 70. Strands are listed starting with one edge of the sheet and continuing to the spatially adjacent strand.

\* The sense in columns 39 - 40 indicates whether strand n is parallel (sense = 1) or anti-parallel (sense = -1) to strand n-1. Sense is equal to zero (0) for the first strand of a sheet.

\* The registration (columns 42 - 70) of strand n to strand n-1 may be specified by one hydrogen bond between each such pair of strands. This is done by providing the hydrogen bonding between the current and previous strands. No register information should be provided for the first strand.

\* Split strands, or strands with two or more runs of residues from discontinuous parts of the amino acid sequence, are explicitly listed. Detail description can be included in the REMARK 700.

#### **Relationships to Other Record Types**

If the entry contains bifurcated sheets or beta-barrels, the relevant REMARK 700 records must be provided. See the REMARK section for details.

#### Examples

	1			2			3			4			5			6		7		8
1234567	789012	345	567	78903	123	34567	89012	234	15678	3901	L234	56789	901	23456	5789	01234	156	57890	123456	57890
SHEET	1	А	5	THR	А	107	ARG	А	110	0										
SHEET	2	А	5	ILE	А	96	THR	А	99	-1	N	LYS	А	98	0	THR	А	107		
SHEET	3	А	5	ARG	А	87	SER	А	91	-1	Ν	LEU	А	89	0	TYR	А	97		
SHEET	4	А	5	TRP	А	71	ASP	А	75	-1	N	ALA	А	74	0	ILE	А	88		
SHEET	5	А	5	$\operatorname{GLY}$	А	52	PHE	А	56	-1	Ν	PHE	А	56	0	TRP	А	71		
SHEET	1	В	5	THR	В	107	ARG	В	110	0										
SHEET	2	В	5	ILE	В	96	THR	В	99	-1	Ν	LYS	В	98	0	THR	В	107		
SHEET	3	В	5	ARG	В	87	SER	В	91	-1	N	LEU	В	89	0	TYR	В	97		
SHEET	4	В	5	TRP	В	71	ASP	В	75	-1	Ν	ALA	В	74	0	ILE	В	88		
SHEET	5	В	5	$\operatorname{GLY}$	В	52	ILE	В	55	-1	Ν	ASP	В	54	0	GLU	В	73		

The sheet presented as BS1 below is an eight-stranded beta-barrel. This is represented by a ninestranded sheet in which the first and last strands are identical.

SHEET	1 BS1	9 VAL	13	ILE	17	0						
SHEET	2 BS1	9 ALA	70	ILE	73	1	0	TRP	72	Ν	ILE	17
SHEET	3 BS1	9 LYS	127	PHE	132	1	0	ILE	129	Ν	ILE	73
SHEET	4 BS1	9 GLY	221	ASP	225	1	0	GLY	221	Ν	ILE	130
SHEET	5 BS1	9 VAL	248	GLU	253	1	0	PHE	249	Ν	ILE	222
SHEET	6 BS1	9 LEU	276	ASP	278	1	Ν	LEU	277	0	GLY	252
SHEET	7 BS1	9 TYR	310	THR	318	1	0	VAL	317	Ν	ASP	278
SHEET	8 BS1	9 VAL	351	TYR	356	1	0	VAL	351	Ν	THR	318
SHEET	9 BS1	9 VAL	13	ILE	17	1	N	VAL	14	0	PRO	352

The sheet structure of this example is bifurcated. In order to represent this feature, two sheets are defined. Strands 2 and 3 of BS7 and BS8 are identical.

SHEET	1 BS7 3 HIS	662 THR	665 0				
SHEET	2 BS7 3 LYS	639 LYS	648 <b>-</b> 1 N	PHE	643	O HIS	662
SHEET	3 BS7 3 ASN	596 VAL	600 -1 N	TYR	598	O ILE	646
SHEET	1 BS8 3 ASN	653 TRP	656 0				
SHEET	2 BS8 3 LYS	639 LYS	648 <b>-</b> 1 N	LYS	647	O THR	655
SHEET	3 BS8 3 ASN	596 VAL	600 -1 N	TYR	598	O ILE	646

# 6. Connectivity Annotation Section

The connectivity annotation section allows the depositors to specify the existence and location of disulfide bonds and other linkages.

# **SSBOND** (updated)

The SSBOND record identifies each disulfide bond in protein and polypeptide structures by identifying the two residues involved in the bond.

The disulfide bond distance is included after the symmetry operations at the end of the SSBOND record.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"SSBOND"	
8 - 10	Integer	serNum	Serial number.
12 - 14	LString(3)	"CYS"	Residue name.
16	Character	chainID1	Chain identifier.
18 - 21	Integer	seqNuml	Residue sequence number.
22	AChar	icode1	Insertion code.
26 - 28	LString(3)	"CYS"	Residue name.
30	Character	chainID2	Chain identifier.
32 - 35	Integer	seqNum2	Residue sequence number.
36	AChar	icode2	Insertion code.
60 - 65	SymOP	syml	Symmetry operator for residue 1.
67 - 72	SymOP	sym2	Symmetry operator for residue 2.
74 - 78	Real(5.2)	Length	Disulfide bond distance

## Details

\* Bond distances between the sulfur atoms must be close to expected value.

\* sym1 and sym2 are right justified and are always given even when identity operator (no cell translation) is to be applied to the residue.

# Verification/Validation/Value Authority Control

wwPDB processing programs generate these records automatically

## **Relationships to Other Record Types**

CONECT records are generated for the disulfide bonds when SG atoms of both cysteines are present in the coordinate records.

### Example

	1		2	3		4	5	6	7	8
12345678	89012	3456'	789012	34567890	123456	57890123	45678901234	56789012345	67890123	4567890
SSBOND	1 C	YS A	6	CYS A	. 127			1555	1555	2.03
SSBOND	2 C	YS A	30	CYS A	. 115			1555	1555	2.07
SSBOND	3 C	YS A	64	CYS A	. 80			1555	1555	2.06
SSBOND	4 C	YS A	76	CYS A	94			1555	1555	2.04

## **Known Problems**

If SG of cysteine is disordered then there are possible alternate linkages. wwPDB practice is to put together all possible SSBOND records. This is problematic because the alternate location identifier is not specified in the SSBOND record.

# LINK (updated)

### Overview

The LINK records specify connectivity between residues that is not implied by the primary structure. Connectivity is expressed in terms of the atom names. They also include the distance associated with the each linkage following the symmetry operations at the end of each record.

This record supplements information given in CONECT records and is provided here for convenience in searching.

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"LINK "	
13 - 16	Atom	name1	Atom name.
17	Character	altLoc1	Alternate location indicator.
18 - 20	Residue name	resName1	Residue name.
22	Character	chainID1	Chain identifier.
23 - 26	Integer	resSeq1	Residue sequence number.
27	AChar	iCode1	Insertion code.
43 - 46	Atom	name2	Atom name.
47	Character	altLoc2	Alternate location indicator.
48 - 50	Residue name	resName2	Residue name.
52	Character	chainID2	Chain identifier.
53 - 56	Integer	resSeq2	Residue sequence number.
57	AChar	iCode2	Insertion code.
60 - 65	SymOP	sym1	Symmetry operator atom 1.
67 – 72	SymOP	sym2	Symmetry operator atom 2.
74 - 78	Real(5.2)	Length	Link distance

# **Record Format**

## Details

\* The atoms involved in bonds between HET groups or between a HET group and standard residue are listed.

- \* Inter-residue linkages not implied by the primary structure are listed (e.g., reduced peptide bond).
- \* Non-standard linkages between residues, e.g., side-chain to side-chain, are listed.

\* Each LINK record specifies one linkage.

\* These records do not specify connectivity within a HET group (see CONECT) or disulfide bridges (see SSBOND).

\* sym1 and sym2 are right justified and are given as blank when the identity operator (and no cell translation) is to be applied to the atom.

- For NMR entries, only one set (or model) of LINK records will be supplied.
- Coordinate bonds are also listed as LINKs.

### Verification/Validation/Value Authority Control

The distance between the pair of atoms listed must be consistent with the bonding.

### **Relationships to Other Record Types**

CONECT records are generated from LINKs when both atoms are present in the entry. If symmetry operators are given to generate one of the residues involved in the bond, REMARK 290 defines the symmetry transformation.

### Example

1	2	3	4	5	6	7	8
123456789012	3456789012	345678901234	567890123456	78901234567	8901234567	890123	4567890
LINK	O GLY A	49	NA	NA A6001	1555	1555	2.98
LINK	OG1 THR A	51	NA	NA A6001	1555	1555	2.72
LINK	OD2 ASP A	66	NA	NA A6001	1555	1555	2.72
LINK	NE ARG A	68	NA	NA A6001	1555	1555	2.93
LINK	C21 2EG A	7	C22	2EG B 19	1555	1555	1.56

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# CISPEP

#### **Overview**

CISPEP records specify the prolines and other peptides found to be in the *cis* conformation. Each cis peptide is listed on a separate line, with a consecutive numbering sequence.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"CISPEP"	
8 - 10	Integer	serNum	Record serial number.
12 - 14	LString(3)	pep1	Residue name.
16	Character	chainID1	Chain identifier.
18 - 21	Integer	seqNum1	Residue sequence number.
22	AChar	icode1	Insertion code.
26 - 28	LString(3)	pep2	Residue name.
30	Character	chainID2	Chain identifier.
32 - 35	Integer	seqNum2	Residue sequence number.
36	AChar	icode2	Insertion code.
44 - 46	Integer	modNum	Identifies the specific model.
54 - 59	Real(6.2)	measure	Angle measurement in degrees.

## Details

\* Cis peptides are those with omega angles of 0°±30°. Deviations larger than 30° are listed in REMARK 500.

# Verification/Validation/Value Authority Control

These records are generated automatically.

## **Relationships to Other Record Types**

Peptide bonds which deviate significantly from either the *cis* or *trans* conformation are annotated in REMARK 500.

Example							
1	2	3	4	5	6	7	8

12345678	3901234567	89012	345678901	23456	789012345678901234	456789012345678901234567890
CISPEP	1 SER A	58	GLY A	59	0	20.91
CISPEP	1 GLY A	116	GLY A	117	0	18.50
CISPEP	1 MET A	1	SER A	2	0	-3.69

# 7. Miscellaneous Features Section

The miscellaneous features section may describe properties in the molecule such as environments surrounding a non-standard residue or the assembly of an active site. Other features may be described in the remarks section but are not given a specific record type so far.

## SITE

## Overview

\* Site records specify residues comprising catalytic, co-factor, anti-codon, regulatory or other essential sites or environments surrounding ligands present in the structure.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"SITE "	
8 - 10	Integer	seqNum	Sequence number.
12 - 14	LString(3)	siteID	Site name.
16 - 17	Integer	numRes	Number of residues that compose the site.
19 - 21	Residue name	resName1	Residue name for first residue that creates the site.
23	Character	chainID1	Chain identifier for first residue of site.
24 - 27	Integer	seq1	Residue sequence number for first residue of the site.
28	AChar	iCode1	Insertion code for first residue of the site.
30 - 32	Residue name	resName2	Residue name for second residue that creates the site.
34	Character	chainID2	Chain identifier for second residue of the site.
35 - 38	Integer	seq2	Residue sequence number for second residue of the site.
39	AChar	iCode2	Insertion code for second residue of the site.
41 - 43	Residue name	resName3	Residue name for third residue that creates the site.
45	Character	chainID3	Chain identifier for third residue

#### of the site.

46 - 49	Integer	seq3	Residue sequence number for third residue of the site.
50	AChar	iCode3	Insertion code for third residue of the site.
52 - 54	Residue name	resName4	Residue name for fourth residue that creates the site.
56	Character	chainID4	Chain identifier for fourth residue of the site.
57 - 60	Integer	seq4	Residue sequence number for fourth residue of the site.
61	AChar	iCode4	Insertion code for fourth residue of the site.

#### Details

\* The sequence number (columns 8 - 10) is reset to 1 for each new site.

\* SITE identifiers (columns 12 - 14) should be fully explained in remark 800.

\* If a site is composed of more than four residues, these may be specified on additional records bearing the same site identifier.

\* SITE records can include HET groups.

## Verification/Validation/Value Authority Control

Every SITE must have a corresponding description in remark 800. The numbering of sequential SITE records and format of each one is verified, as well as the existence of each residue in the ATOM records.

#### **Relationships to Other Record Types**

Each listed SITE needs a corresponding REMARK 800 that details its significance.

	1	2	3	4	5	6	7	8
123456	78901234	5678901234	15678901234	1567890123	45678901234	56789012345	6789012345	567890
SITE	1 AC1	3 HIS A	94 HIS A	A 96 HIS	A 119			
SITE	1 AC2	5 ASN A	62 GLY A	A 63 HIS	А 64 НОН	A 328		
SITE	2 AC2	5 HOH A	634					
SITE	1 AC3	5 GLN A	136 GLN A	A 137 PRO	A 138 GLU	A 205		
SITE	2 AC3	5 CYS A	206					
SITE	1 AC4	11 HIS A	64 HIS A	A 94 HIS	A 96 HIS	A 119		
SITE	2 AC4	11 LEU A	198 THR A	A 199 THR	A 200 TRP	A 209		
SITE	3 AC4	11 HOH A	572 HOH A	А 582 НОН	A 635			

# 8. Crystallographic and Coordinate Transformation Section

This section describes the geometry of the crystallographic experiment and the coordinate system transformations.

## CRYST1

## **Overview**

The CRYST1 record presents the unit cell parameters, space group, and Z value. If the structure was not determined by crystallographic means, CRYST1 simply provides the unitary values, with an appropriate REMARK.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"CRYST1"	
7 – 15	Real(9.3)	a	a (Angstroms).
16 - 24	Real(9.3)	b	b (Angstroms).
25 - 33	Real(9.3)	С	c (Angstroms).
34 - 40	Real(7.2)	alpha	alpha (degrees).
41 - 47	Real(7.2)	beta	beta (degrees).
48 <b>-</b> 54	Real(7.2)	gamma	gamma (degrees).
56 - 66	LString	sGroup	Space group.
67 - 70	Integer	Z	Z value.

## Details

\* If the entry describes a structure determined by a technique other than X-ray crystallography, CRYST1 contains a = b = c = 1.0, alpha = beta = gamma = 90 degrees, space group = P 1, and Z = 1.

\* The Hermann-Mauguin space group symbol is given without parenthesis, e.g., P 43 21 2. Please note that the screw axis is described as a two digit number.

\* The full International Table's Hermann-Mauguin symbol is used, e.g., P 1 21 1 instead of P 21.

\* For a rhombohedral space group in the hexagonal setting, the lattice type symbol used is H.

\* The Z value is the number of polymeric chains in a unit cell. In the case of heteropolymers, Z is the number of occurrences of the most populous chain.

As an example, given two chains A and B, each with a different sequence, and the space group P 2 that has two equipoints in the standard unit cell, the following table gives the correct Z value.

Asymmetric Unit Content	Z value		
А	2		
AA	4		
AB	2		
AAB	4		
AABB	4		

\* In the case of a polycrystalline fiber diffraction study, CRYST1 and SCALE contain the normal unit cell data.

## Verification/Validation/Value Authority Control

The given space group and Z values are checked during processing for correctness and internal consistency. The calculated SCALE factor is compared to that supplied by the depositor. Packing is also computed, and close contacts of symmetry-related molecules are diagnosed.

## **Relationships to Other Record Types**

The unit cell parameters are used to calculate SCALE. If the EXPDTA record is NMR, Electron microscopy, or Fiber Diffraction, the CRYST1 record is predefined as in the last example (see below). In these cases, an explanatory REMARK must also appear in the entry. Some fiber diffraction structures will be done this way, while others will have a CRYST1 record containing measured values.

## Examples

3 4 5 6 7 2 1 CRYST1 52.000 58.600 61.900 90.00 90.00 90.00 P 21 21 21 8 CRYST1 42.544 69.085 50.950 90.00 95.55 90.00 P 1 21 1 2

## Example of experimental method other than X-ray crystallography or fiber diffraction

```
CRYST1 1.000 1.000 1.000 90.00 90.00 P 1 1
```

## **Known Problems**

No standard deviations are given.

# ORIGXn

#### **Overview**

The ORIGXn (n = 1, 2, or 3) records present the transformation from the orthogonal coordinates contained in the entry to the submitted coordinates.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"ORIGXn"	n=1, 2, or 3
11 - 20	Real(10.6)	o[n][1]	Onl
21 - 30	Real(10.6)	o[n][2]	On2
31 - 40	Real(10.6)	o[n][3]	On3
46 - 55	Real(10.5)	t[n]	Tn

## Details

\* This information is included in the file even if the transformation is an identity transformation (unitary matrix, null translation vector). See the SCALE section of this document for a definition of the default orthogonal Angstroms system.

\* If the original submitted coordinates are Xsub, Ysub, Zsub and the orthogonal Angstroms coordinates contained in the data entry are X, Y, Z, then:

Xsub = O11X + O12Y + O13Z + T1 Ysub = O21X + O22Y + O23Z + T2 Zsub = O31X + O32Y + O33Z + T3

## Verification/Validation/Value Authority Control

If the coordinates are submitted in the same orthogonal Angstrom coordinate frame as they appear in the entry (the usual case), then ORIGX is an identity matrix with a null translation vector. If the transformation is not an identity matrix with a null translation vector, then applying this transformation to the coordinates in the entry yields the coordinates of the original deposited file.

## **Relationships to Other Record Types**

ORIGX relates the coordinates in the ATOM and HETATM records to the coordinates in the file.

Example							
1	2	3	4	5	6	7	8

1234567890	012345678901	2345678901	23456789012345	67890123456789012345678901234567890
ORIGX1	0.963457	0.136613	0.230424	16.61000
ORIGX2	-0.158977	0.983924	0.081383	13.72000
ORIGX3	-0.215598	-0.115048	0.969683	37.65000

## SCALEn

## Overview

The SCALEn (n = 1, 2, or 3) records present the transformation from the orthogonal coordinates as contained in the entry to fractional crystallographic coordinates. Non-standard coordinate systems should be explained in the remarks.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"SCALEn"	n=1, 2, or 3
11 - 20	Real(10.6)	s[n][1]	Snl
21 - 30	Real(10.6)	s[n][2]	Sn2
31 - 40	Real(10.6)	s[n][3]	Sn3
46 - 55	Real(10.5)	u[n]	Un

## Details

\* The standard orthogonal Angstroms coordinate system used is related to the axial system of the unit cell supplied (CRYST1 record) by the following definition:

If vector a, vector b, vector c describe the crystallographic cell edges, and vector A, vector B, vector C are unit cell vectors in the default orthogonal Angstroms system, then vector A, vector B, vector C and vector a, vector b, vector c have the same origin; vector A is parallel to vector a, vector B is parallel to vector C times vector A, and vector C is parallel to vector a times vector b (i.e., vector c\*). \* If the orthogonal Angstroms coordinates are X, Y, Z, and the fractional cell coordinates are xfrac, yfrac, zfrac, then:

```
xfrac = S11X + S12Y + S13Z + U1
yfrac = S21X + S22Y + S23Z + U2
zfrac = S31X + S32Y + S33Z + U3
```

\* For NMR, fiber diffraction, and EM entries, SCALE is given as an identity matrix with no translation.

## Verification/Validation/Value Authority Control

The inverse of the determinant of the SCALE matrix equals the volume of the cell. This volume is calculated and compared to the SCALE matrix supplied by the depositor.

## **Relationships to Other Record Types**

The SCALE transformation is related to the CRYST1 record, as the inverse of the determinant of the SCALE matrix equals the cell volume.

## Example

1 2 3 4 5 6 7 8 SCALE1 0.019231 0.000000 0.000000 0.00000 SCALE2 0.000000 0.017065 0.000000 0.00000 0.000000 0.000000 0.016155 SCALE3 0.00000

# MTRIXn

## **Overview**

The MTRIXn (n = 1, 2, or 3) records present transformations expressing non-crystallographic symmetry. MTRIXn will appear only when such transformations are required to generate an entire asymmetric unit, such as a large viral structure.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"MTRIXn"	n=1, 2, or 3
8 - 10	Integer	serial	Serial number.
11 - 20	Real(10.6)	m[n][1]	Mn1
21 - 30	Real(10.6)	m[n][2]	Mn2
31 - 40	Real(10.6)	m[n][3]	Mn 3
46 <b>-</b> 55	Real(10.5)	v[n]	Vn
60	Integer	iGiven	1 if coordinates for the representations which are approximately related by the transformations of the molecule are contained in the entry. Otherwise, blank.

## Details

\* The MTRIX transformations operate on the coordinates in the entry to yield equivalent representations of the molecule in the same coordinate frame. One trio of MTRIX records with a constant serial number is given for each non-crystallographic symmetry operation defined. If coordinates for the representations which are approximately related by the given transformation are present in the file, the last "iGiven" field is set to 1. Otherwise, this field is blank.

## Verification/Validation/Value Authority Control

All MTRIX records are verified using records from the author and review.

## **Relationships to Other Record Types**

None. Example

2 3 5 6 7 1 4 MTRIX1 1 -1.000000 0.000000 0.000000 0.00000 1 1 0.000000 1.000000 0.000000 1 MTRIX2 0.00000 1 0.000000 0.000000 -1.000000 MTRIX3 0.00000 1

# 9. Coordinate Section

The Coordinate Section contains the collection of atomic coordinates as well as the MODEL and ENDMDL records.

## MODEL

#### **Overview**

The MODEL record specifies the model serial number when multiple models of the same structure are presented in a single coordinate entry, as is often the case with structures determined by NMR.

## Record Format

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"MODEL "	
11 - 14	Integer	serial	Model serial number.

## Details

\*This record is used only when more than one model appears in an entry. Generally, it is employed mainly for NMR structures. The chemical connectivity should be the same for each model. ATOM, HETATM, ANISOU, and TER records for each model structure and are interspersed as needed between MODEL and ENDMDL records.

\*The numbering of models is sequential, beginning with 1.

\* All models in a deposition should be superimposed in an appropriate author determined manner and only one superposition method should be used. Structures from different experiments, or different domains of a structure should not be superimposed and deposited as models of a deposition.

\* All models in an NMR ensemble should be homogeneous – each model should have the exact same atoms (hydrogen and heavy atoms), sequence and chemistry.

\*Deposition of minimized average structure (if available) must be accompanied with ensemble and must be homogeneous with ensemble.

\*A model cannot have more than 99,999 atoms. Where the entry does not contain an ensemble of models, then the entry cannot have more than 99,999 atoms. Entries that go beyond this atom limit must be split into multiple entries, each containing no more than the limits specified above.

## Verification/Validation/Value Authority Control

Entries with multiple models in the NUMMDL record are checked for corresponding pairs of MODEL/ ENDMDL records, and for consecutively numbered models.

## **Relationships to Other Record Types**

Each MODEL must have a corresponding ENDMDL record.

## Examples

	1 89012		2 57890:	123	34567	3 89012345678	4 901234567	5 89012345	6 67890	1234567	7 890123456	8 7890
MODEL ATOM	1	1 N	אד א	7	1	11 104	6.134	-6.504	1.00	0.00		NT
ATOM	2		ALA ALA		1	11.104 11.639	6.071	-0.504 -5.147	1.00	0.00		N C
	2	011	11111		-	11.009	0.071	5.117	1.00	0.00		C
ATOM		1HG	GLU		18		-4.847	0.361	1.00	0.00		Н
ATOM		2HG	GLU		18	-13.518	-3.769	0.084	1.00	0.00		Н
TER	295		GLU	A	18							
ENDMDL MODEL		2										
ATOM	296	N	ALA	А	1	10.883	6.779	-6.464	1.00	0.00		N
ATOM	297	CA	ALA		1	11.451	6.531	-5.142	1.00	0.00		C
ATOM		1HG	GLU		18	-13.363		-2.372	1.00	0.00		Η
ATOM		2HG	GLU		18	-12.634	-3.023	-3.475	1.00	0.00		Н
TER ENDMDL	590		GLU	А	18							
ENDRUL												
	1		•									
	L 1		2			3	4	5	6		7	8
1234567		23450	2 57890:	123	34567	3 89012345678				1234567		-
1234567 MODEL		23450 1		123	34567					1234567		-
MODEL ATOM	89012 1	1 N	57890: AALA	A	1	890123456789 72.883	901234567 57.697	89012345 56.410	67890: 0.50	83.80		7890 N
MODEL ATOM ATOM	89012 1 2	1 N CA	57890: AALA AALA	A A	1 1	890123456789 72.883 73.796	901234567 57.697 56.531	56.410 56.644	678903 0.50 0.50	83.80 84.78		7890 N C
MODEL ATOM ATOM ATOM	89012 1 2 3	1 N CA C	57890 AALA AALA AALA AALA	A A A	1 1 1	89012345678 72.883 73.796 74.549	901234567 57.697 56.531 56.551	89012345 56.410 56.644 57.997	678903 0.50 0.50 0.50	83.80 84.78 85.05		N C C
MODEL ATOM ATOM ATOM ATOM	89012 1 2	1 N CA C	57890: AALA AALA	A A A	1 1	890123456789 72.883 73.796	901234567 57.697 56.531	56.410 56.644	678903 0.50 0.50 0.50	83.80 84.78		7890 N C
MODEL ATOM ATOM ATOM	89012 1 2 3	1 N CA C	57890 AALA AALA AALA AALA	A A A	1 1 1	89012345678 72.883 73.796 74.549	901234567 57.697 56.531 56.551	89012345 56.410 56.644 57.997	678903 0.50 0.50 0.50	83.80 84.78 85.05		N C C
MODEL ATOM ATOM ATOM ATOM	89012 1 2 3 4	1 N CA C	57890 AALA AALA AALA AALA	A A A	1 1 1	89012345678 72.883 73.796 74.549 73.951	901234567 57.697 56.531 56.551	89012345 56.410 56.644 57.997	678903 0.50 0.50 0.50 0.50	83.80 84.78 85.05		N C C
MODEL ATOM ATOM ATOM ATOM 	89012 1 2 3 4 7900	1 N CA C O	57890: AALA AALA AALA AALA AALA	A A A	1 1 1	89012345678 72.883 73.796 74.549 73.951 -24.915	901234567 57.697 56.531 56.551 56.413	89012345 56.410 56.644 57.997 59.075	678903 0.50 0.50 0.50 0.50 0.50	83.80 84.78 85.05 84.77		N C C O
MODEL ATOM ATOM ATOM  HETATM3	89012 1 2 3 4 7900 7901	1 N CA C O	AALA AALA AALA AALA AALA AALA	A A A	1 1 1 1	89012345678 72.883 73.796 74.549 73.951 -24.915 -28.699	901234567 57.697 56.531 56.551 56.413 147.513	89012345 56.410 56.644 57.997 59.075 36.413	678903 0.50 0.50 0.50 0.50 0.50	83.80 84.78 85.05 84.77 41.86		7890 N C C O
MODEL ATOM ATOM ATOM  HETATM3 HETATM3 HETATM3 ENDMDL	89012 1 2 3 4 7900 7901	1 N CA C O O O	AALA AALA AALA AALA AALA AALA AALA AHOH	A A A	1 1 1 1 490 491	89012345678 72.883 73.796 74.549 73.951 -24.915 -28.699	901234567 57.697 56.531 56.551 56.413 147.513 130.471	89012345 56.410 56.644 57.997 59.075 36.413 22.248	678903 0.50 0.50 0.50 0.50 0.50	83.80 84.78 85.05 84.77 41.86 36.06		7890 N C C O O
MODEL ATOM ATOM ATOM  HETATM3 HETATM3 HETATM3 ENDMDL MODEL	8901: 1 2 3 4 7900 7901 7902	1 N CA C O O O O 2	AALA AALA AALA AALA AALA AALA AALA AHOH AHOH	A A A	1 1 1 1 490 491 492	89012345678 72.883 73.796 74.549 73.951 -24.915 -28.699 -33.309	901234567 57.697 56.531 56.551 56.413 147.513 130.471 184.488	89012345 56.410 56.644 57.997 59.075 36.413 22.248 26.176	678902 0.50 0.50 0.50 0.50 0.50 0.50 0.50	83.80 84.78 85.05 84.77 41.86 36.06 15.00		N C C O O O O
MODEL ATOM ATOM ATOM  HETATM3 HETATM3 HETATM3 ENDMDL MODEL ATOM	89012 1 2 3 4 7900 7901 7902 1	1 N CA C O O O O 2 N	AALA AALA AALA AALA AALA AALA AALA AHOH AHOH	A A A A	1 1 1 490 491 492	89012345678 72.883 73.796 74.549 73.951 -24.915 -28.699 -33.309 72.883	901234567 57.697 56.531 56.551 56.413 147.513 130.471 184.488 57.697	89012345 56.410 56.644 57.997 59.075 36.413 22.248 26.176 56.410	678902 0.50 0.50 0.50 0.50 0.50 0.50 0.50	83.80 84.78 85.05 84.77 41.86 36.06 15.00 83.80		N C C O O O O N
MODEL ATOM ATOM ATOM  HETATM3 HETATM3 HETATM3 ENDMDL MODEL ATOM ATOM	8901: 1 2 3 4 7900 7901 7902 1 2	1 N CA C 0 0 0 0 2 N CA	AALA AALA AALA AALA AALA AALA AALA AHOH AHOH	A A A A	1 1 1 1 490 491 492 1 1	89012345678 72.883 73.796 74.549 73.951 -24.915 -28.699 -33.309 72.883 73.796	901234567 57.697 56.531 56.551 56.413 147.513 130.471 184.488 57.697 56.531	89012345 56.410 56.644 57.997 59.075 36.413 22.248 26.176 56.410 56.644	678903 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	83.80 84.78 85.05 84.77 41.86 36.06 15.00 83.80 84.78		N C C O O O O O O O O O O
MODEL ATOM ATOM ATOM ATOM  HETATM3 HETATM3 HETATM3 ENDMDL MODEL ATOM ATOM ATOM	8901: 1 2 3 4 7900 7901 7902 1 2 3	1 N CA C O O O O 2 N	AALA AALA AALA AALA AALA AALA AALA AHOH AHOH	A A A A A A A	1 1 1 490 491 492	89012345678 72.883 73.796 74.549 73.951 -24.915 -28.699 -33.309 72.883 73.796 74.549	901234567 57.697 56.531 56.551 56.413 147.513 130.471 184.488 57.697 56.531 56.551	89012345 56.410 56.644 57.997 59.075 36.413 22.248 26.176 56.410 56.644 57.997	678903 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	83.80 84.78 85.05 84.77 41.86 36.06 15.00 83.80 84.78 85.05		N C C O O O O N
MODEL ATOM ATOM ATOM  HETATM3 HETATM3 HETATM3 ENDMDL MODEL ATOM ATOM	8901: 1 2 3 4 7900 7901 7902 1 2	1 N CA C O O O O O O O O O O O O O O O O O	AALA AALA AALA AALA AALA AALA AALA AHOH AHOH	A A A A A A A A	1 1 1 490 491 492 1 1 1	89012345678 72.883 73.796 74.549 73.951 -24.915 -28.699 -33.309 72.883 73.796	901234567 57.697 56.531 56.551 56.413 147.513 130.471 184.488 57.697 56.531	89012345 56.410 56.644 57.997 59.075 36.413 22.248 26.176 56.410 56.644	678903 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	83.80 84.78 85.05 84.77 41.86 36.06 15.00 83.80 84.78		7890 N C C O O O O O O O O O O C
MODEL ATOM ATOM ATOM ATOM  HETATM3 HETATM3 HETATM3 ENDMDL MODEL ATOM ATOM ATOM	8901: 1 2 3 4 7900 7901 7902 1 2 3 4	1 N CA C O O O O O O O O O O O O O O O O O	AALA AALA AALA AALA AALA AALA AALA AHOH AHOH	A A A A A A A A A A	1 1 1 490 491 492 1 1 1 1	89012345678 72.883 73.796 74.549 73.951 -24.915 -28.699 -33.309 72.883 73.796 74.549 73.951	901234567 57.697 56.531 56.551 56.413 147.513 130.471 184.488 57.697 56.531 56.551 56.413	89012345 56.410 56.644 57.997 59.075 36.413 22.248 26.176 56.410 56.644 57.997 59.075	678903 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	83.80 84.78 85.05 84.77 41.86 36.06 15.00 83.80 84.78 85.05 84.77		7890 N C C O O O O O O O O O O O O O O O O O

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## ATOM

## Overview

The ATOM records present the atomic coordinates for standard amino acids and nucleotides. They also present the occupancy and temperature factor for each atom. Non-polymer chemical coordinates use the HETATM record type. The element symbol is always present on each ATOM record; charge is optional.

Changes in ATOM/HETATM records result from the standardization atom and residue nomenclature. This nomenclature is described in the Chemical Component Dictionary (<u>ftp://ftp.wwpdb.org/pub/pdb/data/monomers</u>).

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"ATOM "	
7 - 11	Integer	serial	Atom serial number.
13 - 16	Atom	name	Atom name.
17	Character	altLoc	Alternate location indicator.
18 - 20	Residue name	resName	Residue name.
22	Character	chainID	Chain identifier.
23 - 26	Integer	resSeq	Residue sequence number.
27	AChar	iCode	Code for insertion of residues.
31 - 38	Real(8.3)	x	Orthogonal coordinates for X in Angstroms.
39 - 46	Real(8.3)	У	Orthogonal coordinates for Y in Angstroms.
47 - 54	Real(8.3)	Z	Orthogonal coordinates for Z in Angstroms.
55 - 60	Real(6.2)	occupancy	Occupancy.
61 - 66	Real(6.2)	tempFactor	Temperature factor.
77 – 78	LString(2)	element	Element symbol, right-justified.
79 - 80	LString(2)	charge	Charge on the atom.

## Details

\* ATOM records for proteins are listed from amino to carboxyl terminus.

\* Nucleic acid residues are listed from the 5'  $\rightarrow$  3' terminus.

\* No ordering is specified for polysaccharides.

\* Non-blank alphanumerical character is used for chain identifier.

\* The list of ATOM records in a chain is terminated by a TER record.

\* If more than one model is present in the entry, each model is delimited by MODEL and ENDMDL records.

\* AltLoc is the place holder to indicate alternate conformation. The alternate conformation can be in the entire polymer chain, or several residues or partial residue (several atoms within one residue). If an atom is provided in more than one position, then a non-blank alternate location indicator must be used for each of the atomic positions. Within a residue, all atoms that are associated with each other in a given conformation are assigned the same alternate position indicator. There are two ways of representing alternate conformation- either at atom level or at residue level (see examples).

\* For atoms that are in alternate sites indicated by the alternate site indicator, sorting of atoms in the ATOM/HETATM list uses the following general rules:

- In the simple case that involves a few atoms or a few residues with alternate sites, the coordinates occur one after the other in the entry.
- In the case of a large heterogen groups which are disordered, the atoms for each conformer are listed together.

\* Alphabet letters are commonly used for insertion code. The insertion code is used when two residues have the same numbering. The combination of residue numbering and insertion code defines the unique residue.

\* If the depositor provides the data, then the isotropic B value is given for the temperature factor.

\* If there are neither isotropic B values from the depositor, nor anisotropic temperature factors in ANISOU, then the default value of 0.0 is used for the temperature factor.

\* Columns 79 - 80 indicate any charge on the atom, e.g., 2+, 1-. In most cases, these are blank.

#### Verification/Validation/Value Authority Control

The ATOM/HETATM records are checked for PDB file format, sequence information, and packing.

## **Relationships to Other Record Types**

The ATOM records are compared to the corresponding sequence database. Sequence discrepancies appear in the SEQADV record. Missing atoms are annotated in the remarks. HETATM records are formatted in the same way as ATOM records. The sequence implied by ATOM records must be identical to that given in SEQRES, with the exception that residues that have no coordinates, e.g., due to disorder, must appear in SEQRES.

# Examples

	1		2		3	4	5	6		7 8
1234567	/89012	34567	3901	23456	5789012345678	3901234567	/89012345	67890	1234567	8901234567890
ATOM	32	N A	ARG	A -3	3 11.281	86.699	94.383	0.50	35.88	N
ATOM	33	N B	ARG	A -3	11.296		94.521	0.50	35.60	N
АТОМ	34	CA A	ARG	A -3	12.353	85.696	94.456	0.50	36.67	C
ATOM	35	CA B	ARG	A -3	12.333	85.862	95.041	0.50	36.42	С
ATOM	36	C A	ARG	A -3	13.559	86.257	95.222	0.50	37.37	С
ATOM	37	C B	ARG	A -3	12.759	86.530	96.365	0.50	36.39	C
ATOM	38	0 A.	ARG	A -3	3 13.753	8 87.471	95.270	0.50	37.74	0
ATOM	39	0 B.	ARG	A -3	12.924	87.757	96.420	0.50	37.26	0
ATOM	40	CB A			3 12.774	85.306	93.039	0.50	37.25	C
ATOM	41	CB B	ARG	A -3	13.428	8 85.746	93.980		36.60	C
ATOM	42	CG A					92.321		38.44	C
ATOM	43	CG B					92.651		37.31	C
ATOM	44	CD A					90.815		38.51	C
ATOM	45	CD B					91.406		37.66	C
ATOM	46	NE A					90.163		39.94	N
ATOM	47	NE B					90.195		38.24	N
ATOM	48	CZ A					88.850		40.03	С
ATOM	49	CZ B					88.947		39.55	C
ATOM	50	NH1A					88.007		40.76	N
ATOM	51	NH1B					88.706		40.23	N
ATOM	52	NH2A	ARG	A -3	3 14.421	84.308	88.373	0.50	40.45	N
	1		2		3	4	5	6		7 8
1234567		34567		23456					1234567	7 8 8901234567890
1234567 ATOM					5789012345678	8901234567		67890	L234567 35.88	
	789012		3901 ARG	A -3	5789012345678 3 11.281	8901234567 86.699	789012345	67890: 0.50		8901234567890
ATOM	789012 32	N A. CA A.	3901 ARG	A -3 A -3	5789012345678 3 11.281 3 12.353	8901234567 86.699 8 85.696	789012345 94.383	678903 0.50 0.50	35.88	8901234567890 N
АТОМ АТОМ	789012 32 33	N A. CA A. C A.	3901 ARG ARG	A -3 A -3 A -3	5789012345678 11.281 12.353 13.559	8901234567 86.699 85.696 86.257	789012345 94.383 94.456	678902 0.50 0.50 0.50	35.88 36.67	8901234567890 N C
АТОМ АТОМ АТОМ	789012 32 33 34	N A. CA A. C A.	3901 ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753	8901234567 86.699 85.696 86.257 887.471	789012345 94.383 94.456 95.222	678902 0.50 0.50 0.50 0.50	35.88 36.67 37.37	8901234567890 N C C
АТОМ АТОМ АТОМ АТОМ	789012 32 33 34 35	N AL CA AL C AL O AL	3901 ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774	8901234567 86.699 85.696 86.257 87.471 85.306	789012345 94.383 94.456 95.222 95.270	678903 0.50 0.50 0.50 0.50 0.50	35.88 36.67 37.37 37.74	8901234567890 N C C O
ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36	N A. CA A. C A. O A. CB A. CG A. CD A.	3901 ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678	789012345 94.383 94.456 95.222 95.270 93.039	678903 0.50 0.50 0.50 0.50 0.50 0.50 0.50	35.88 36.67 37.37 37.74 37.25 38.44 38.51	8901234567890 N C C O C
ATOM ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36 37 38 39	N A. CA A. C A. O A. CB A. CG A. CD A. NE A.	3901 ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698         12.984	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.447	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163	678902 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	35.88 36.67 37.37 37.74 37.25 38.44 38.51 39.94	8901234567890 N C C O C C C C N
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36 37 38 39 40	NA.CAA.CA.CBA.CGA.CDA.NEA.CZA.	3901 ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698         12.984         13.202	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.534	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850	678902 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	35.88 36.67 37.37 37.74 37.25 38.44 38.51 39.94 40.03	8901234567890 N C C O C C C C N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36 37 38 39 40 41	N A. CA A. C A. CB A. CB A. CD A. NE A. CZ A. NH1A.	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698         12.984         13.202         12.218	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.534         84.534         84.840	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007	678903 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	35.88 36.67 37.37 37.74 37.25 38.44 38.51 39.94 40.03 40.76	8901234567890 N C C O C C C C N
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36 37 38 39 40 41 42	N A. CA A. C A. CB A. CG A. CD A. NE A. CZ A. NH1A. NH2A.	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698         12.984         13.202         13.202         14.421	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.534         84.534         84.308	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373	678903 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	$\begin{array}{c} 35.88\\ 36.67\\ 37.37\\ 37.74\\ 37.25\\ 38.44\\ 38.51\\ 39.94\\ 40.03\\ 40.76\\ 40.45\\ \end{array}$	8901234567890 N C C O C C C N C N N N
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43	N A. CA A. C A. CB A. CB A. CD A. NE A. CZ A. NH1A. NH2A. N B.	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         11.754         11.698         12.984         13.202         12.18         12.218         14.421         11.296	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.534         84.534         84.308         84.308         86.721	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521	67890 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	$\begin{array}{c} 35.88\\ 36.67\\ 37.37\\ 37.74\\ 37.25\\ 38.44\\ 38.51\\ 39.94\\ 40.03\\ 40.76\\ 40.45\\ 35.60\\ \end{array}$	8901234567890 N C C O C C C N C N N N N N
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43 44	N A. CA A. C A. CB A. CG A. CD A. NE A. CZ A. NH1A. NH2A. N B. CA B.	3901 ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698         12.984         11.698         12.218         11.294         11.296         12.333	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.534         84.534         84.308         86.721         85.862	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041	678903 0.50 0.50 0.50 0.50 0.50 0.50 0.50 0.	$\begin{array}{c} 35.88\\ 36.67\\ 37.37\\ 37.74\\ 37.25\\ 38.44\\ 38.51\\ 39.94\\ 40.03\\ 40.76\\ 40.45\\ 35.60\\ 36.42 \end{array}$	8901234567890 N C C O C C C N C N N N N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43 44 45	N A. CA A. C A. CB A. CB A. CD A. NE A. CZ A. NH1A. NH2A. N B. CA B. C B.	3901 ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698         12.984         11.698         12.984         11.698         12.984         11.298         12.984         11.298         12.218         11.296         12.333         12.333         12.759	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.534         84.534         84.308         86.721         85.862         86.530	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041 96.365	67890 0.5	$\begin{array}{c} 35.88\\ 36.67\\ 37.37\\ 37.74\\ 37.25\\ 38.44\\ 38.51\\ 39.94\\ 40.03\\ 40.76\\ 40.45\\ 35.60\\ 36.42\\ 36.39 \end{array}$	8901234567890 N C C O C C C N N C N N C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46	N A. CA A. C A. CB A. CB A. CD A. NE A. CZ A. NH1A. NH2A. NH2A. N B. CA B. C B. O B.	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698         12.984         11.698         12.984         11.698         12.984         11.298         12.984         12.984         12.984         12.218         12.218         12.218         12.218         12.218         12.218         12.228         12.233         12.233         12.333         12.924	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.534         84.534         84.308         86.721         85.862         86.530         87.757	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041 96.365 96.420	67890 0.5	35.88 36.67 37.37 37.74 37.25 38.44 38.51 39.94 40.03 40.76 40.45 35.60 36.42 36.39 37.26	8901234567890 N C C O C C C N N C N N C N N C O O
ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47	N       A.         CA       A.         C       A.         CB       A.         CB       A.         CD       A.         CD       A.         CD       A.         CZ       A.         NE       A.         NH1A.       NH2A.         N       B.         CA       B.         C       B.         C       B.         C       B.         C       B.         C       B.         CB       B.	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         12.774         11.754         11.698         12.984         11.698         12.984         11.698         12.984         11.296         12.218         11.296         12.333         12.333         12.924         13.428	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.534         84.534         84.308         86.721         85.862         86.530         87.757         85.746	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041 96.365 96.420 93.980	67890 0.5	35.88 36.67 37.37 37.74 37.25 38.44 38.51 39.94 40.03 40.76 40.45 35.60 36.42 36.39 37.26 36.60	8901234567890 N C C O C C C N N C N N C N N C C O C C O C
ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48	N       A.         CA       A.         C       A.         CB       A.         CB       A.         CD       A.         CD       A.         CD       A.         CD       A.         CZ       A.         NE       A.         NH1A.       N.         NH2A.       N.         CA       B.         C       B.         C       B.         CB       B. <td>3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG</td> <td><math display="block"> \begin{array}{cccc} A &amp; -3 \\ A</math></td> <td>5789012345678         11.281         12.353         13.559         13.753         11.754         11.754         11.698         12.984         11.698         12.984         11.698         12.984         11.2984         12.984         12.984         12.984         12.984         12.984         12.984         12.924         12.924         13.428         12.866</td> <td>8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.447         84.534         84.534         84.308         86.721         85.862         86.530         87.757         85.746         85.172</td> <td>789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041 95.041 96.365 96.420 93.980 92.651</td> <td>67890 0.5</td> <td><math display="block">\begin{array}{c} 35.88\\ 36.67\\ 37.37\\ 37.74\\ 37.25\\ 38.44\\ 38.51\\ 39.94\\ 40.03\\ 40.76\\ 40.45\\ 35.60\\ 36.42\\ 36.39\\ 37.26\\ 36.60\\ 37.31 \end{array}</math></td> <td>8901234567890 N C C O C C C N N C N N C N N C C C C C</td>	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	$ \begin{array}{cccc} A & -3 \\ A$	5789012345678         11.281         12.353         13.559         13.753         11.754         11.754         11.698         12.984         11.698         12.984         11.698         12.984         11.2984         12.984         12.984         12.984         12.984         12.984         12.984         12.924         12.924         13.428         12.866	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.447         84.534         84.534         84.308         86.721         85.862         86.530         87.757         85.746         85.172	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041 95.041 96.365 96.420 93.980 92.651	67890 0.5	$\begin{array}{c} 35.88\\ 36.67\\ 37.37\\ 37.74\\ 37.25\\ 38.44\\ 38.51\\ 39.94\\ 40.03\\ 40.76\\ 40.45\\ 35.60\\ 36.42\\ 36.39\\ 37.26\\ 36.60\\ 37.31 \end{array}$	8901234567890 N C C O C C C N N C N N C N N C C C C C
ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48 49	N       A.         CA       A.         C       A.         CB       A.         CB       A.         CD       A.         CD       A.         CD       A.         CD       A.         NE       A.         NH1A.       NH2A.         N       B.         CA       B.         CB       B.         CB       B.         CB       B.         CB       B.         CCB	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         11.754         11.754         11.698         12.984         11.698         12.984         11.2984         12.984         11.298         12.984         12.984         12.984         12.984         12.984         12.984         12.924         12.924         13.428         12.866         13.374	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.432         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         85.862         86.530         87.757         85.746         85.172         85.886	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041 95.041 96.365 96.420 93.980 92.651 91.406	67890 0.5	35.88 36.67 37.37 37.74 37.25 38.44 38.51 39.94 40.03 40.76 40.45 35.60 36.42 36.39 37.26 36.60 37.31 37.66	8901234567890 N C C O C C C N N C N N C N N C C C C C
ATOM         ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50	N       A.         CA       A.         C       A.         CB       A.         CB       A.         CD       A.         CD       A.         CD       A.         CD       A.         NE       A.         NH1A       NH2A.         NH2A.       B.         CA       B.         CB       B.         CD       B.	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         13.753         11.754         11.754         11.698         12.984         11.698         12.984         11.2984         11.2984         11.2984         11.2984         11.2984         11.2984         11.2984         11.2984         11.2984         11.2984         11.2984         11.2984         12.333         12.924         13.428         13.374         12.866         13.374         12.644	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.432         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         84.534         85.862         86.530         87.757         85.746         85.886         85.487	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041 96.365 96.420 93.980 92.651 91.406 90.195	67890 0.5	35.88 36.67 37.37 37.74 37.25 38.44 38.51 39.94 40.03 40.76 40.45 35.60 36.42 36.39 37.26 36.60 37.31 37.66 38.24	8901234567890 N C C O C C C N N C N N N C C C C C C N N N N
ATOM	789012 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48 49	N       A.         CA       A.         C       A.         CB       A.         CB       A.         CD       A.         CD       A.         CD       A.         CD       A.         NE       A.         NH1A.       NH2A.         N       B.         CA       B.         CB       B.         CB       B.         CB       B.         CB       B.         CCB	3901 ARG ARG ARG ARG ARG ARG ARG ARG ARG ARG	A -3 A -3 A -3 A -3 A -3 A -3 A -3 A -3	5789012345678         11.281         12.353         13.559         13.753         13.753         11.754         11.698         12.984         11.698         12.984         13.202         11.298         11.2984         12.984         13.202         11.298         12.984         12.984         12.218         12.218         12.333         12.924         13.428         13.374         12.644         13.114	8901234567         86.699         85.696         86.257         87.471         85.306         84.432         84.678         84.432         84.534         84.534         84.534         84.534         84.534         84.534         85.862         86.530         87.757         85.746         85.172         85.886         85.487         85.582	789012345 94.383 94.456 95.222 95.270 93.039 92.321 90.815 90.163 88.850 88.007 88.373 94.521 95.041 95.041 96.365 96.420 93.980 92.651 91.406	67890 0.5	35.88 36.67 37.37 37.74 37.25 38.44 38.51 39.94 40.03 40.76 40.45 35.60 36.42 36.39 37.26 36.60 37.31 37.66	8901234567890 N C C O C C C N N C N N C N N C C C C C

## ANISOU

## Overview

The ANISOU records present the anisotropic temperature factors.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"ANISOU"	
7 - 11	Integer	serial	Atom serial number.
13 - 16	Atom	name	Atom name.
17	Character	altLoc	Alternate location indicator
18 - 20	Residue name	resName	Residue name.
22	Character	chainID	Chain identifier.
23 - 26	Integer	resSeq	Residue sequence number.
27	AChar	iCode	Insertion code.
29 - 35	Integer	u[0][0]	U(1,1)
36 - 42	Integer	u[1][1]	U(2,2)
43 - 49	Integer	u[2][2]	U(3,3)
50 - 56	Integer	u[0][1]	U(1,2)
57 - 63	Integer	u[0][2]	U(1,3)
64 - 70	Integer	u[1][2]	U(2,3)
77 – 78	LString(2)	element	Element symbol, right-justified.
79 - 80	LString(2)	charge	Charge on the atom.

## Details

\* Columns 7 - 27 and 73 - 80 are identical to the corresponding ATOM/HETATM record.

\* The anisotropic temperature factors (columns 29 - 70) are scaled by a factor of 10\*\*4 (Angstroms\*\*2) and are presented as integers.

\* The anisotropic temperature factors are stored in the same coordinate frame as the atomic coordinate records.

\* ANISOU values are listed only if they have been provided by the depositor.

## Verification/Validation/Value Authority Control

The depositor provides ANISOU records, and the wwPDB verifies their format.

## **Relationships to Other Record Types**

The anisotropic temperature factors are related to the corresponding ATOM/HETATM isotropic temperature factors as ,B(eq), as described in the ATOM and HETATM sections.

## Example

	1		2		3	4	5		6	7	8
1234567	89012	3456	7890123	4567	8901234567	890123	456789012	2345678	9012345	6789012	34567890
ATOM	107	Ν	GLY A	13	12.68	1 37.	302 -25.2	211 1.0	00 15.5	6	N
ANISOU	107	Ν	GLY A	13	2406	1892	1614	198	519	-328	N
ATOM	108	CA	GLY A	13	11.98	2 37.	996 -26.2	241 1.0	00 16.9	2	C
ANISOU	108	CA	GLY A	13	2748	2004	1679	-21	155	-419	C
ATOM	109	С	GLY A	13	11.67	8 39.	447 -26.0	008 1.0	00 15.7	3	С
ANISOU	109	С	GLY A	13	2555	1955	1468	87	357	-109	C
ATOM	110	0	GLY A	13	11.44	4 40.	201 -26.9	971 1.0	00 20.9	3	0
ANISOU	110	0	GLY A	13	3837	2505	1611	164	-121	189	0
ATOM	111	Ν	ASN A	14	11.60	8 39.	863 -24.7	755 1.0	00 13.6	8	Ν
ANISOU	111	N	ASN A	14	2059	1674	1462	27	244	-96	N

## **Relationships to Other Record Types**

The standard deviations for the anisotropic temperature factors are related to the corresponding ATOM/ HETATM ANISOU temperature factors.

	1		2		3	4	5		6	7	8
1234567	89012	3456	7890123	34567	89012345	67890123	34567890	1234567	8901234	567890123	34567890
ATOM	107	N	GLY A	13	12.	681 37	.302 -25	.211 1.	000 15.	56	N
ANISOU	107	Ν	GLY A	13	2406	1892	1614	198	519	-328	N
SIGUIJ	107	N	GLY A	13	10	10	10	10	10	10	N
ATOM	108	CA	GLY A	13	11.	982 37	.996 -26	.241 1.	000 16.	92	С
ANISOU	108	CA	GLY A	13	2748	2004	1679	-21	155	-419	С
SIGUIJ	108	CA	GLY A	13	10	10	10	10	10	10	С
ATOM	109	С	GLY A	13	11.	678 39	.447 -26	.008 1.	000 15.	73	С
ANISOU	109	С	GLY A	13	2555	1955	1468	87	357	-109	С
SIGUIJ	109	С	GLY A	13	10	10	10	10	10	10	С
ATOM	110	0	GLY A	13	11.	444 40	.201 -26	.971 1.	000 20.	93	0
ANISOU	110	0	GLY A	13	3837	2505	1611	164	-121	189	0
SIGUIJ	110	0	GLY A	13	10	10	10	10	10	10	0
ATOM	111	Ν	ASN A	14	11.	608 39	.863 -24	.755 1.	000 13.	68	N
ANISOU	111	N	ASN A	14	2059	1674	1462	27	244	-96	N
SIGUIJ	111	Ν	ASN A	14	10	10	10	10	10	10	N

#### TER

#### **Overview**

The TER record indicates the end of a list of ATOM/HETATM records for a chain.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"TER "	
7 - 11	Integer	serial	Serial number.
18 - 20	Residue name	resName	Residue name.
22	Character	chainID	Chain identifier.
23 - 26	Integer	resSeq	Residue sequence number.
27	AChar	iCode	Insertion code.

## Details

\* Every chain of ATOM/HETATM records presented on SEQRES records is terminated with a TER record.

\* The TER records occur in the coordinate section of the entry, and indicate the last residue presented for each polypeptide and/or nucleic acid chain for which there are determined coordinates. For proteins, the residue defined on the TER record is the carboxy-terminal residue; for nucleic acids it is the 3'-terminal residue.

\* For a cyclic molecule, the choice of termini is arbitrary.

\* Terminal oxygen atoms are presented as OXT for proteins, and as O5' or OP3 for nucleic acids. These atoms are present only if the last residue in the polymer is truly the last residue in the SEQRES.

\* The TER record has the same residue name, chain identifier, sequence number and insertion code as the terminal residue. The serial number of the TER record is one number greater than the serial number of the ATOM/HETATM preceding the TER.

## Verification/Validation/Value Authority Control

TER must appear at the terminal carboxyl end or 3' end of a chain. For proteins, there is usually a terminal oxygen, labeled OXT. The validation program checks for the occurrence of TER and OXT records.

## **Relationships to Other Record Types**

The residue name appearing on the TER record must be the same as the residue name of the immediately preceding ATOM or non-water HETATM record. **Example** 

1		2		3	4	5	6		7 8
123456789012	34567	78901	234567	89012345678	901234567	89012345	678901	23456789	01234567890
ATOM 601	N	LEU	A 75	-17.070	-16.002	2.409	1.00	55.63	N
ATOM 602	CA	LEU	A 75	-16.343	-16.746	3.444	1.00	55.50	C
ATOM 603	С	LEU	A 75	-16.499	-18.263	3.300	1.00	55.55	С
ATOM 604	0	LEU	A 75	-16.645	-18.789	2.195	1.00	55.50	0
ATOM 605	CB	LEU	A 75	-16.776	-16.283	4.844	1.00	55.51	C
TER 606		LEU	A 75						
ATOM 1185	0	LEU	B 75	26.292	-4.310	16.940	1.00	55.45	0
ATOM 1186	СВ	LEU	B 75	23.881	-1.551	16.797	1.00	55.32	C
TER 1187		LEU	B 75						
HETATM 1188	Н2	SRT	A1076	-17.263	11.260	28.634	1.00	59.62	Н
HETATM 1189	HA	SRT	A1076	-19.347	11.519	28.341	1.00	59.42	Н
HETATM 1190	HЗ	SRT	A1076	-17.157	14.303	28.677	1.00	58.00	Н
HETATM 1191	HB	SRT	A1076	-15.110	13.610	28.816	1.00	57.77	Н
HETATM 1192	01	SRT	A1076	-17.028	11.281	31.131	1.00	62.63	0
ATOM 295	HB2	ALA	A 18	4.601	-9.393	7.275	1.00	0.00	H
ATOM 296	HB3	ALA	A 18	3.340	-9.147	6.043	1.00	0.00	H
TER 297		ALA	A 18						
ENDMDL									

# HETATM

## Overview

Non-polymer or other "non-standard" chemical coordinates, such as water molecules or atoms presented in HET groups use the HETATM record type. They also present the occupancy and temperature factor for each atom. The ATOM records present the atomic coordinates for standard residues. The element symbol is always present on each HETATM record; charge is optional.

Changes in ATOM/HETATM records will require standardization in atom and residue nomenclature. This nomenclature is described in the Chemical Component Dictionary, <u>ftp://ftp.wwpdb.org/pub/pdb/data/monomers</u>.

Record F	ormat
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COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"HETATM"	
7 - 11	Integer	serial	Atom serial number.
13 - 16	Atom	name	Atom name.
17	Character	altLoc	Alternate location indicator.
18 - 20	Residue name	resName	Residue name.
22	Character	chainID	Chain identifier.
23 - 26	Integer	resSeq	Residue sequence number.
27	AChar	iCode	Code for insertion of residues.
31 - 38	Real(8.3)	x	Orthogonal coordinates for X.
39 <b>-</b> 46	Real(8.3)	У	Orthogonal coordinates for Y.
47 <b>-</b> 54	Real(8.3)	Z	Orthogonal coordinates for Z.
55 <b>-</b> 60	Real(6.2)	occupancy	Occupancy.
61 - 66	Real(6.2)	tempFactor	Temperature factor.
77 - 78	LString(2)	element	Element symbol; right-justified.
79 - 80	LString(2)	charge	Charge on the atom.

## Details

\* The x, y, z coordinates are in Angstrom units.

\* No ordering is specified for polysaccharides.

\* See the HET section of this document regarding naming of heterogens. See the Chemical Component Dictionary for residue names, formulas, and topology of the HET groups that have appeared so far in the PDB (see <a href="http://ftp.wwpdb.org/pub/pdb/data/monomers">ftp://ftp.wwpdb.org/pub/pdb/data/monomers</a> ).

\* If the depositor provides the data, then the isotropic B value is given for the temperature factor.

\* If there are neither isotropic B values provided by the depositor, nor anisotropic temperature factors in ANISOU, then the default value of 0.0 is used for the temperature factor.

\* Insertion codes and element naming are fully described in the ATOM section of this document.

#### Verification/Validation/Value Authority Control

Processing programs check ATOM/HETATM records for PDB file format, sequence information, and packing.

#### **Relationships to Other Record Types**

HETATM records must have corresponding HET, HETNAM, FORMUL and CONECT records, except for waters.

1	2	3 4	5 6	5 7	8
1234567890123456	7890123456789	01234567890123456	78901234567890	1234567890123456789	0
HETATM 8237 MG	MG A1001	13.872 -2.555	-29.045 1.00	0 27.36 MG	
HETATM 3835 FE	HEM A 1	17.140 3.115	15.066 1.00	) 14.14 FE	
HETATM 8238 S	SO4 A2001	10.885 -15.746	-14.404 1.00	) 47.84 S	
HETATM 8239 01	SO4 A2001	11.191 -14.833	-15.531 1.00	0 50 <b>.</b> 12 0	
HETATM 8240 O2	SO4 A2001	9.576 -16.338	-14.706 1.00	0 48.55 O	
HETATM 8241 O3	SO4 A2001	11.995 -16.703	-14.431 1.00	0 49.88 O	
HETATM 8242 O4	SO4 A2001	10.932 -15.073	-13.100 1.00	0 49.91 O	

## ENDMDL

#### **Overview**

The ENDMDL records are paired with MODEL records to group individual structures found in a coordinate entry.

#### **Record Format**

 COLUMNS
 DATA TYPE
 FIELD
 DEFINITION

 1 - 6
 Record name
 "ENDMDL"

#### Details

\* MODEL/ENDMDL records are used only when more than one structure is presented in the entry, as is often the case with NMR entries.

\* All the models in a multi-model entry must represent the same structure.

\* Every MODEL record has an associated ENDMDL record.

#### Verification/Validation/Value Authority Control

Entries with multiple structures in the NUMMDL record are checked for corresponding pairs of MODEL/ ENDMDL records, and for consecutively numbered models.

#### **Relationships to Other Record Types**

There must be a corresponding MODEL record.

In the case of an NMR entry, the NUMMDL record states the number of model structures that are present in the individual entry.

	1		2		3	4	5	6	7	8
12345	678901	23456	78901	2345678	90123456789	01234567	789012345	678901	234567890	1234567890
• • •										
• • •										
ATOM	14550	1  HG	GLU	122	-14.364	14.787	-14.258	1.00	0.00	Н
ATOM	14551	2HG	GLU	122	-13.794	13.738	-12.961	1.00	0.00	Н
TER	14552		GLU	122						
ENDMD	L									
MODEL	I	9								
ATOM	14553	N	SER	1	-28.280	1.567	12.004	1.00	0.00	N
ATOM	14554	CA	SER	1	-27.749	0.392	11.256	1.00	0.00	С
• • •										
• • •										

ATOM	16369 1HG	GLU	122	-3.757	18.546	-8.439	1.00	0.00	Η
ATOM	16370 2HG	GLU	122	-3.066	17.166	-7.584	1.00	0.00	Η
TER	16371	GLU	122						
ENDMD	L								

# **10. Connectivity Section**

This section provides information on atomic connectivity. LINK, SSBOND, and CISPEP are found in the Connectivity Annotation section.

## CONECT

## Overview

The CONECT records specify connectivity between atoms for which coordinates are supplied. The connectivity is described using the atom serial number as shown in the entry. CONECT records are mandatory for HET groups (excluding water) and for other bonds not specified in the standard residue connectivity table. These records are generated automatically.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 – 6	Record name	"CONECT"	
7 – 11	Integer	serial	Atom serial number
12 - 16	Integer	serial	Serial number of bonded atom
17 - 21	Integer	serial	Serial number of bonded atom
22 - 26	Integer	serial	Serial number of bonded atom
27 - 31	Integer	serial	Serial number of bonded atom

## Details

\* CONECT records are present for:

- Intra-residue connectivity within non-standard (HET) residues (excluding water).
- Inter-residue connectivity of HET groups to standard groups (including water) or to other HET groups.
- Disulfide bridges specified in the SSBOND records have corresponding records.

\* No differentiation is made between atoms with delocalized charges (excess negative or positive charge).

\* Atoms specified in the CONECT records have the same numbers as given in the coordinate section.

\* All atoms connected to the atom with serial number in columns 7 - 11 are listed in the remaining fields of the record.

\* If more than four fields are required for non-hydrogen and non-salt bridges, a second CONECT record with the same atom serial number in columns 7 - 11 will be used.

\* These CONECT records occur in increasing order of the atom serial numbers they carry in columns 7 - 11. The target-atom serial numbers carried on these records also occur in increasing order.

\* The connectivity list given here is redundant in that each bond indicated is given twice, once with each of the two atoms involved specified in columns 7 - 11.

\* For hydrogen bonds, when the hydrogen atom is present in the coordinates, a CONECT record between the hydrogen atom and its acceptor atom is generated.

\* For NMR entries, CONECT records for one model are generated describing heterogen connectivity and others for LINK records assuming that all models are homogeneous models.

## Verification/Validation/Value Authority Control

Connectivity is checked for unusual bond lengths.

## **Relationships to Other Record Types**

CONECT records must be present in an entry that contains either non-standard groups or disulfide bonds.

## Example

```
1 2 3 4 5 6 7 8
12345678901234567890123456789012345678901234567890123456789012345678901234567890123456789012345678901234567890
CONECT 1179 746 1184 1195 1203
CONECT 1179 1211 1222
CONECT 1021 544 1017 1020 1022
```

## **Known Problems**

CONECT records involving atoms for which the coordinates are not present in the entry (e.g., symmetry-generated) are not given.

CONECT records involving atoms for which the coordinates are missing due to disorder, are also not provided.

# 11. Bookkeeping Section

The Bookkeeping Section provides some final information about the file itself.

## MASTER

#### Overview

The MASTER record is a control record for bookkeeping. It lists the number of lines in the coordinate entry or file for selected record types. MASTER records only the first model when there are multiple models in the coordinates.

## **Record Format**

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"MASTER"	
11 - 15	Integer	numRemark	Number of REMARK records
16 - 20	Integer	" 0 "	
21 - 25	Integer	numHet	Number of HET records
26 - 30	Integer	numHelix	Number of HELIX records
31 - 35	Integer	numSheet	Number of SHEET records
36 - 40	Integer	numTurn	deprecated
41 - 45	Integer	numSite	Number of SITE records
46 - 50	Integer	numXform	Number of coordinate transformation records (ORIGX+SCALE+MTRIX)
51 - 55	Integer	numCoord	Number of atomic coordinate records records (ATOM+HETATM)
56 - 60	Integer	numTer	Number of TER records
61 - 65	Integer	numConect	Number of CONECT records
66 - 70	Integer	numSeq	Number of SEQRES records

## Details

\* MASTER gives checksums of the number of records in the entry, for selected record types.

\* MASTER records only the first model when there are multiple models in the coordinates.

## Verification/Validation/Value Authority Control

The MASTER line is automatically generated.

## **Relationships to Other Record Types**

MASTER presents a checksum of the lines present for each of the record types listed above.

1		2		3		4		5	6		7	8
1234567890	1234567	89012	34567	89012	34567	89012	34567	89012345	67890123	4567	7890123	4567890
MASTER	40	0	0	0	0	0	0	6 2930	2	0	29	

## END

#### **Overview**

The END record marks the end of the PDB file.

## **Record Format**

 COLUMNS
 DATA TYPE
 FIELD
 DEFINITION

 1 - 6
 Record name
 "END "

## Details

\* END is the final record of a coordinate entry.

## Verification/Validation/Value Authority Control

END must appear in every coordinate entry.

## **Relationships to Other Record Types**

This is the final record in the entry.