



wwPDB NMR Structure Validation Summary Report ⓘ

Apr 26, 2016 – 03:43 PM BST

PDB ID : 1O2F
Title : COMPLEX OF ENZYME IIAGLC AND IIBGLC PHOSPHOCARRIER
PROTEIN HPR FROM ESCHERICHIA COLI NMR, RESTRAINED REG-
ULARIZED MEAN STRUCTURE
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Deposited on : 2003-03-11

This is a wwPDB NMR Structure Validation Summary Report for a publicly released PDB entry.
We welcome your comments at validation@mail.wwpdb.org
A user guide is available at
<http://wwpdb.org/validation/2016/NMRValidationReportHelp>
with specific help available everywhere you see the ⓘ symbol.

The following versions of software and data (see [references ⓘ](#)) were used in the production of this report:

Cyrange : **NOT EXECUTED**
NmrClust : **NOT EXECUTED**
MolProbity : **NOT EXECUTED**
Mogul : 1.7.1 (RC1), CSD as537be (2016)
Percentile statistics : **NOT EXECUTED**
RCI : **NOT EXECUTED**
PANAV : **NOT EXECUTED**
ShiftChecker : **NOT EXECUTED**
Ideal geometry (proteins) : Engh & Huber (2001)
Ideal geometry (DNA, RNA) : Parkinson et al. (1996)
Validation Pipeline (wwPDB-VP) : rb-20027457

1 Overall quality at a glance

The following experimental techniques were used to determine the structure:

SOLUTION NMR

The overall completeness of chemical shifts assignment was not calculated.

There are no percentiles available for this entry.

The sequence quality summary graphics cannot be shown.

2 Ensemble composition and analysis

This entry contains 3 models. The atoms present in the NMR models are not consistent. Some calculations may have failed as a result. All residues are included in the validation scores.

Cyrange was unable to find well-defined residues.

Error message: Cyrange did not run

NmrClust was unable to cluster the ensemble.

Error message: NmrClust did not run

ENTRY-COMPOSITION INFOmissingINFO

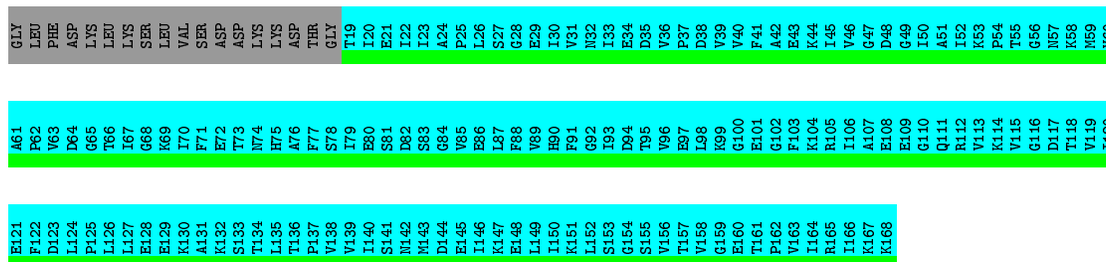
3 Residue-property plots

3.1 Average score per residue in the NMR ensemble

These plots are provided for all protein, RNA and DNA chains in the entry. The first graphic is the same as shown in the summary in section 1 of this report. The second graphic shows the sequence where residues are colour-coded according to the number of geometric quality criteria for which they contain at least one outlier: green = 0, yellow = 1, orange = 2 and red = 3 or more. Stretches of 2 or more consecutive residues without any outliers are shown as green connectors. Residues which are classified as ill-defined in the NMR ensemble, are shown in cyan with an underline colour-coded according to the previous scheme. Residues which were present in the experimental sample, but not modelled in the final structure are shown in grey.

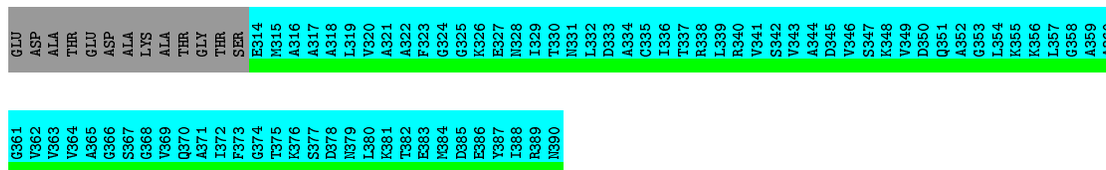
- Molecule 1: PTS system, glucose-specific IIA component

Chain A: 



- Molecule 2: PTS system, glucose-specific IIBC component

Chain B: 

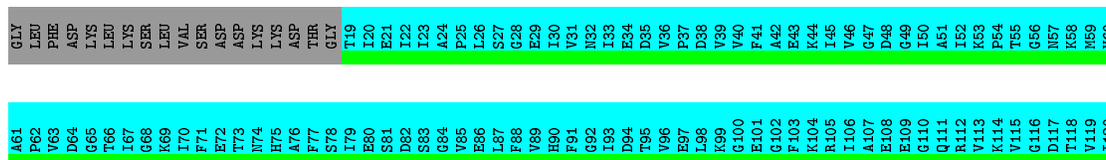


3.2 Residue scores for the first model from the NMR ensemble

No representative models were identified. Colouring as in section 3.1 above.

- Molecule 1: PTS system, glucose-specific IIA component

Chain A: 



E121
F122
D123
L124
P125
L126
L127
E128
E129
K130
A131
K132
S133
T134
L135
T136
P137
V138
V139
I140
S141
M142
M143
D144
E145
I146
K147
E148
L149
I150
K151
L152
S153
G154
S155
V156
T157
V158
G159
E160
T161
P162
V163
I164
M165
I166
K167
K168

- Molecule 2: PTS system, glucose-specific IIBC component

Chain B:

86%

14%

GLU
ASP
ALA
THR
GLU
ASP
ALA
LYS
ALA
THR
GLY
THR
SER
E314
M315
K316
A317
A318
I319
V320
A321
A322
F323
G324
G325
K326
E327
M328
I329
T330
N331
I332
D333
A334
C335
I336
T337
R338
L339
R340
V341
S342
V343
A344
D345
V346
S347
K348
V349
D350
Q351
A352
G353
L354
K355
K356
L357
G358
A359
A360

G361
V362
V363
V364
A365
G366
S367
G368
V369
Q370
A371
I372
F373
G374
T375
K376
S377
D378
N379
L380
K381
T382
E383
M384
D385
E386
T387
I388
R389
N390

4 Refinement protocol and experimental data overview

The models were refined using the following method: *CONJOINED RIGID BODY/TORSION ANGLE DYNAMICS*.

Of the 60 calculated structures, 3 were deposited, based on the following criterion: *REGULARIZED MEAN STRUCTURES*.

The following table shows the software used for structure solution, optimisation and refinement.

Software name	Classification	Version
X-PLOR NIH	refinement	(HTTP://NMR.CIT.NIH.GOV/XPLOR_NIH)

No chemical shift data was provided. No validations of the models with respect to experimental NMR restraints is performed at this time.

5 Model quality [i](#)

5.1 Standard geometry [i](#)

MolProbity was not executed - this section will have to be empty.

5.2 Too-close contacts [i](#)

MolProbity was not executed - this section will have to be empty.

5.3 Torsion angles [i](#)

5.3.1 Protein backbone [i](#)

MolProbity was not executed - this section will have to be empty.

5.3.2 Protein sidechains [i](#)

MolProbity was not executed - this section will have to be empty.

5.3.3 RNA [i](#)

MolProbity was not executed - this section will have to be empty.

5.4 Non-standard residues in protein, DNA, RNA chains [i](#)

MolProbity was not executed - this section will have to be empty.

5.5 Carbohydrates [i](#)

MolProbity was not executed - this section will have to be empty.

5.6 Ligand geometry [i](#)

MolProbity was not executed - this section will have to be empty.

5.7 Other polymers [i](#)

MolProbity was not executed - this section will have to be empty.

5.8 Polymer linkage issues

There are no chain breaks in this entry.

6 Chemical shift validation

No chemical shift data were provided