

## Report on BUSTER refinement run in directory 1pmq\_03\_short

## Contents

<b>1</b>	<b>Run overview</b>	<b>2</b>
1.1	Run conditions . . . . .	2
1.2	Refinement vital statistics . . . . .	2
<b>2</b>	<b>RSCC, R-factor, LLG and geometry evolution over the refine</b>	<b>3</b>
2.1	Reciprocal space correlation coefficient plots . . . . .	3
2.1.1	Initial RSCC plot . . . . .	3
2.1.2	Final RSCC plot . . . . .	3
2.2	R-factor behaviour during refinement . . . . .	4
2.3	LLG behaviour during refinement . . . . .	4
2.4	Geometry behaviour during optimisation . . . . .	5
2.4.1	Graph of RMS(bond) against cycle of refinement . . . . .	5
2.4.2	Graph of RMS(angle) against cycle of refinement . . . . .	5
<b>3</b>	<b>MolProbity analysis</b>	<b>6</b>
3.1	Summary statistics . . . . .	6
3.2	Ramachandran plot . . . . .	7
<b>4</b>	<b>Ligand analysis</b>	<b>8</b>
4.1	880 A 501 . . . . .	8
4.1.1	Statistics for ligand . . . . .	8
4.1.2	Picture of ligand in electron density . . . . .	8
4.1.3	Mogul analysis for 880 A 501 . . . . .	9
4.2	ANP A 502 . . . . .	15
4.2.1	Statistics for ligand . . . . .	15
4.2.2	Picture of ligand in electron density . . . . .	15
4.2.3	Mogul analysis for ANP A 502 . . . . .	16
<b>5</b>	<b>X-ray statistics</b>	<b>22</b>
5.1	Scaling parameters in last cycle . . . . .	22
5.2	Wilson plots . . . . .	22
5.2.1	Wilson plot at start of refinement . . . . .	22
5.2.2	Wilson plot at end of refinement . . . . .	22
<b>6</b>	<b>Real-space correlations</b>	<b>23</b>
6.1	Side chains of chain A . . . . .	23
6.2	Mainchain of chain A . . . . .	23

# 1 Run overview

## 1.1 Run conditions

refine command	/mnt/scratch_fs1/osmart/autobuster/Server/- autoBUSTER/bin/linux64/refine -p 1pmq_02_- refine.report-coot-0.pdb -m 1pmq/1pmq.mtz -d 1pmq_03_short -l 880.grade_PDB_ligand.cif -M ShortRunVoid -report
BUSTER version, run at, by user in directory	2.13.0, Mon Jun 16 15:52:09 BST 2014, osmart /home/osmart/2014/06/erice_workshop/1pmq_tuto- rial
nthreads, hostname, OS buster-report command	6, hypatia, Ubuntu precise (12.04.4 LTS) /home/osmart/autobuster/Server/scripts/- buster-report -d 1pmq_03_short -dr 1pmq_- 03_short.report
buster-report version, run at, by user	1.1.4 <July 25 2015>, Sat Jul 25 18:59:29 2015, osmart
buster-report run on refine directory	/home/osmart/2014/06/erice_workshop/1pmq_- tutorial/1pmq_03_short
buster-report output directory	/home/osmart/2014/06/erice_workshop/1pmq_- tutorial/1pmq_03_short.report
final pdb coordinates	1pmq_03_short.report.pdb
final mtzfile	1pmq_03_short.report.mtz

For help on “Run conditions table” see BUSTER wiki page  
<http://www.globalphasing.com/buster/wiki/index.cgi?BRrunConditions>

## 1.2 Refinement vital statistics

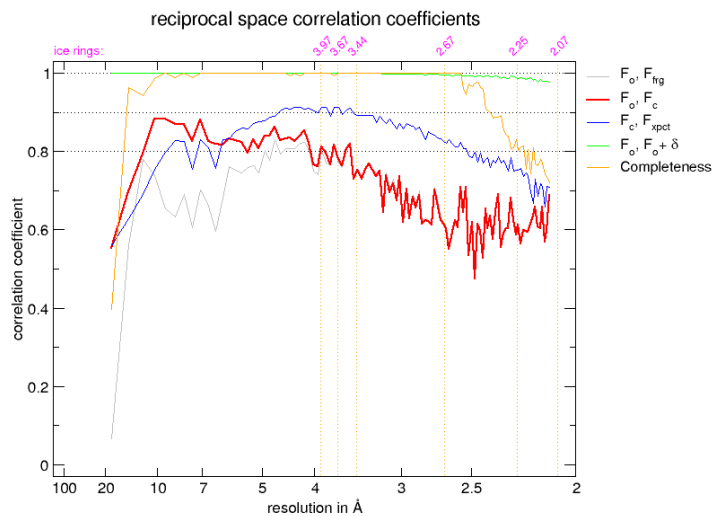
	start	final
<b>N<sub>cycles</sub> big</b>	0	2
<b>N<sub>cycles</sub> small</b>	0	21
<b>X-ray weight</b>	n/a	4.55
<b>R<sub>work</sub></b>	0.2213	0.2170
<b>R<sub>free</sub></b>	0.2438	0.2444
<b>100 (R<sub>free</sub> - R<sub>work</sub>)</b>	2.2%	2.7%
<b>LLG<sub>work</sub> (cumulative Log-Likelihood Gain, working set)</b>	0	0.0114
<b>LLG<sub>free</sub> (cumulative Log-Likelihood Gain, free set)</b>	0	-0.0156
<b>RMS bond in Å</b>	0.0095	0.0088
<b>RMS angle in degrees</b>	0.99	0.98
<b>High resolution limit in Å</b>	n/a	2.10
<b>Low resolution limit in Å</b>	n/a	16.94
<b>Number of waters</b>	54	54

For help on “Refinement vital statistics” see BUSTER wiki page  
<http://www.globalphasing.com/buster/wiki/index.cgi?BRTblVitalStats>

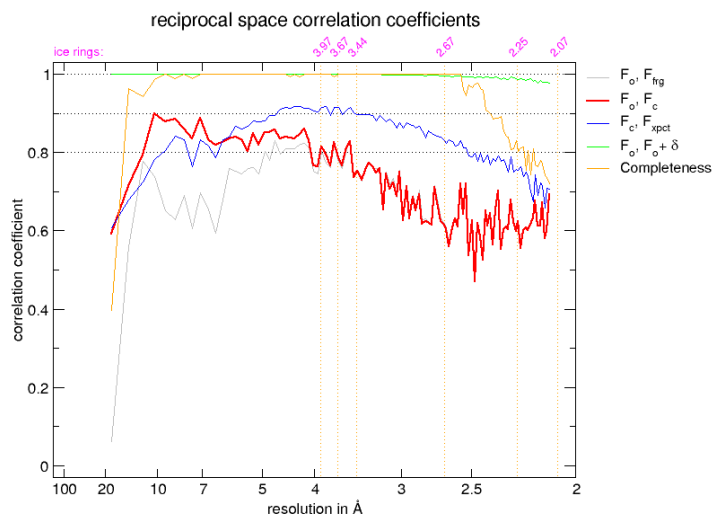
## 2 RSCC, R-factor, LLG and geometry evolution over the refine

### 2.1 Reciprocal space correlation coefficient plots

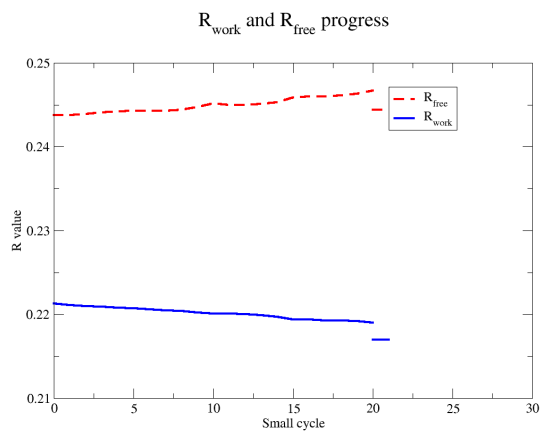
#### 2.1.1 Initial RSCC plot



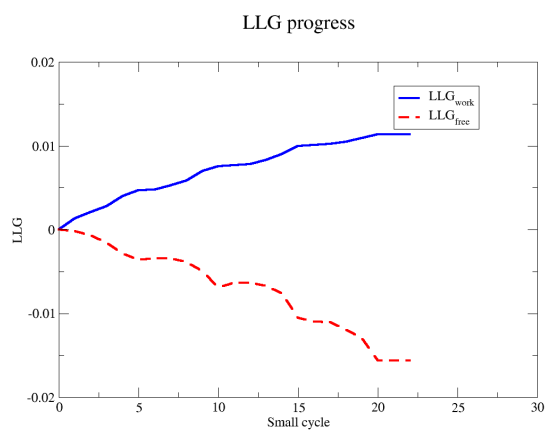
#### 2.1.2 Final RSCC plot



## 2.2 R-factor behaviour during refinement

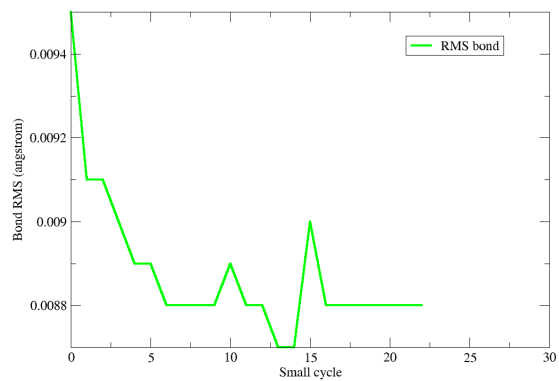


## 2.3 LLG behaviour during refinement

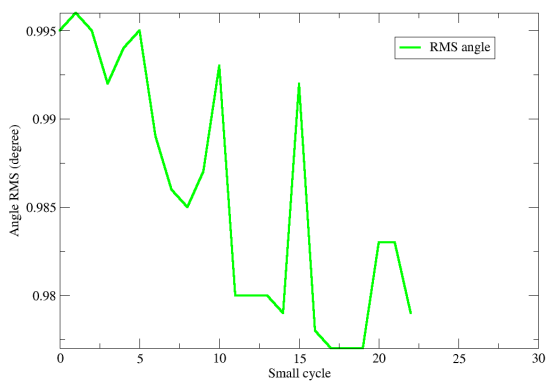


## 2.4 Geometry behaviour during optimisation

### 2.4.1 Graph of RMS(bond) against cycle of refinement



### 2.4.2 Graph of RMS(angle) against cycle of refinement



## 3 MolProbity analysis

### 3.1 Summary statistics

All-Atom Contacts	Clashscore, all atoms:	1.76		100 <sup>th</sup> percentile* N=456, 2.20Å ± 0.25Å
	Clashscore is the number of serious steric overlaps (> 0.4 Å) per 1000 atoms.			
Protein Geometry	Poor rotamers	8	2.59%	Goal: <1%
	Ramachandran outliers	2	0.58%	Goal: <0.05%
	Ramachandran favored	328	95.91%	Goal: >98%
	Cβ deviations >0.25Å	0	0.00%	Goal: 0
	MolProbity score <sup>†</sup>	1.53		98 <sup>th</sup> percentile* N=10167, 2.20Å ± 0.25Å
	Bad backbone bonds:	0 / 1389	0.00%	Goal: 0%
Bad backbone angles:	0 / 1731	0.00%	Goal: <0.1%	

In the two column results, the left column gives the raw count, right column gives the percentage.

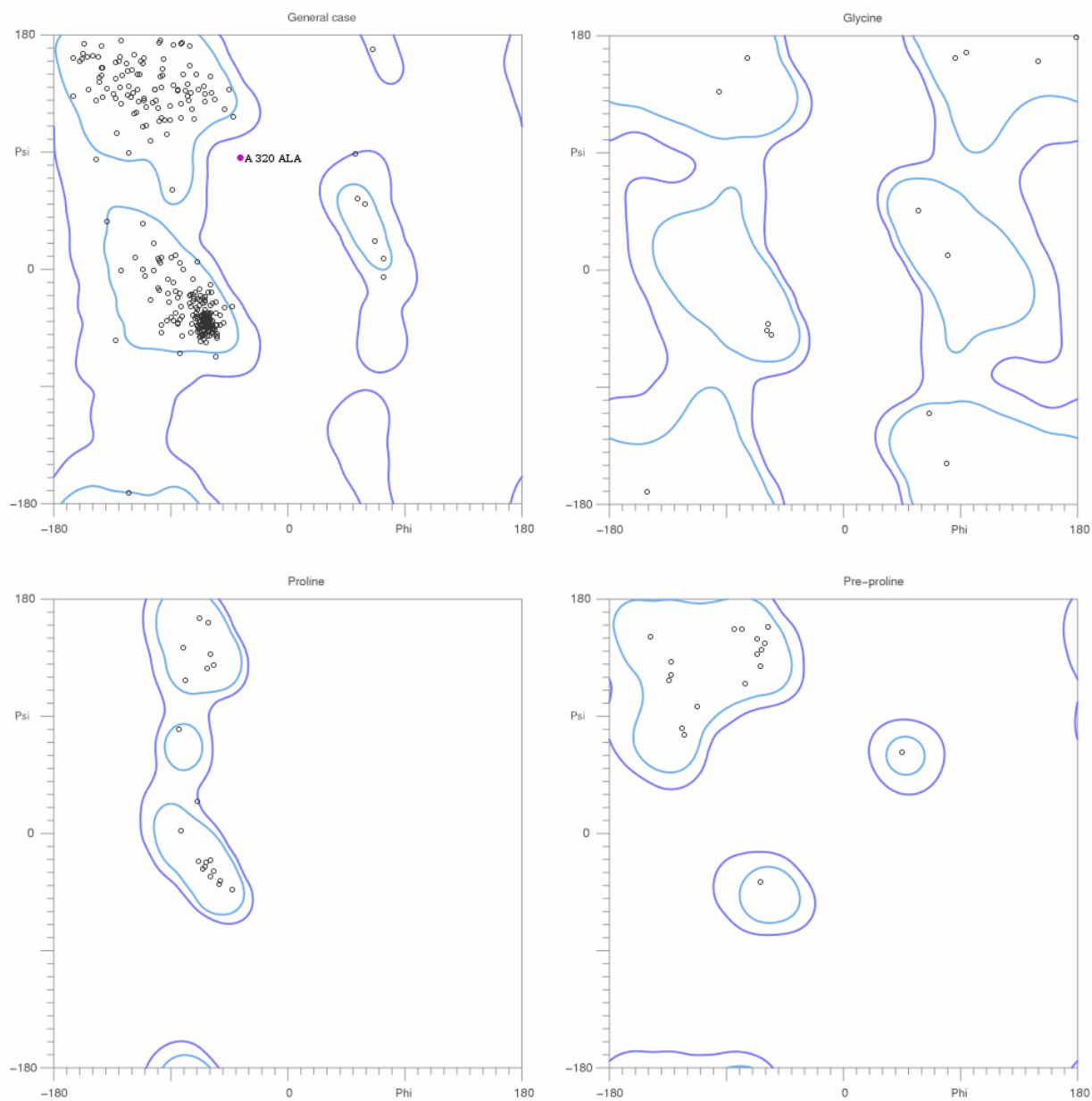
\* 100<sup>th</sup> percentile is the best among structures of comparable resolution; 0<sup>th</sup> is the worst " For clashscore the comparative set of structures was selected in 2004, for MolProbity score in 2006.

† MolProbity score combines the clashscore, rotamer, and Ramachandran evaluations into a single score, normalized to be on the same scale as X-ray resolution.

For more information see:

- MolProbity homepage: <http://molprobity.biochem.duke.edu/>
- MolProbity paper: Chen et al. (2010) "MolProbity: all-atom structure validation for macromolecular crystallography." *Acta Cryst.* **D66**: 12-21. <http://dx.doi.org/10.1107/S0907444909042073>
- MolProbity Ramachandran plot paper: Lovell et al. (2003) "Structure Validation by Cα Geometry: φ, ψ and Cβ Deviation." *Proteins: Struct Func Genet* **50**: 437-450. <http://dx.doi.org/10.1002/prot.10286>

## 3.2 Ramachandran plot



342 residues were evaluated in total for general, glycine, proline, and pre-pro.

96.49% of all residues were in favored (98%) regions. (330 residues)

99.71% of all residues were in allowed (>99.8%) regions. (341 residues)

There were 1 outliers:

A 320 ALA: (phi,psi) = (-35.24, 85.23)

## 4 Ligand analysis

### 4.1 880 A 501

#### 4.1.1 Statistics for ligand

Database ID	<a href="#">880 (PDB)</a>
3-letter code	880
CC(2mF <sub>o</sub> -DF <sub>c</sub> )	0.9615
min(B-factor)‡	30.0
avg(B-factor)‡	35.6
max(B-factor)‡	44.2
min(occupancy)‡	1.00
max(occupancy)‡	1.00
‡hydrogen atoms excluded	

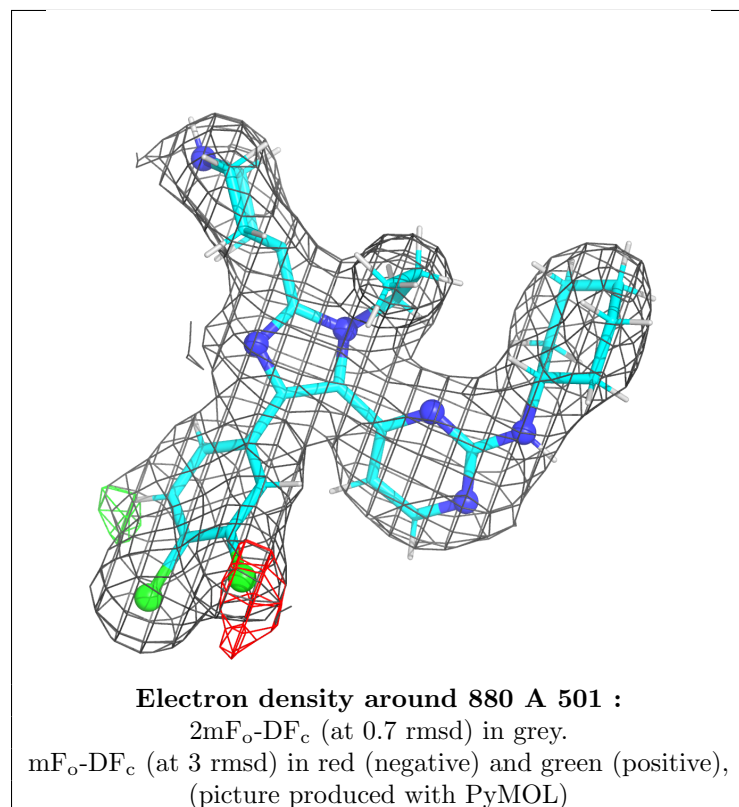
#### Restraints used

restraints for 880 (CYCLOHEXYL-4-5-3,4-DICHLOROPHENYL-2-PIPERIDIN-4-YL-3-PROPYL-3H-IMIDAZOL-4-YL-PYRIMIDIN-2-YLAMINE) from cif dictionary 880.grade\_PDB\_ligand.cif; generated by GRADE\_PDB\_LIGAND 1.2.9 (pre-release) (June 15 2014) using MOGUL 1.6.1(DEV7), CSD as535be, with quantum mechanics RM1

For help on “Ligand Statistics Table” see BUSTER wiki page

<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#statistics>

#### 4.1.2 Picture of ligand in electron density



For help on “Ligand Electron Density Picture” see BUSTER wiki page

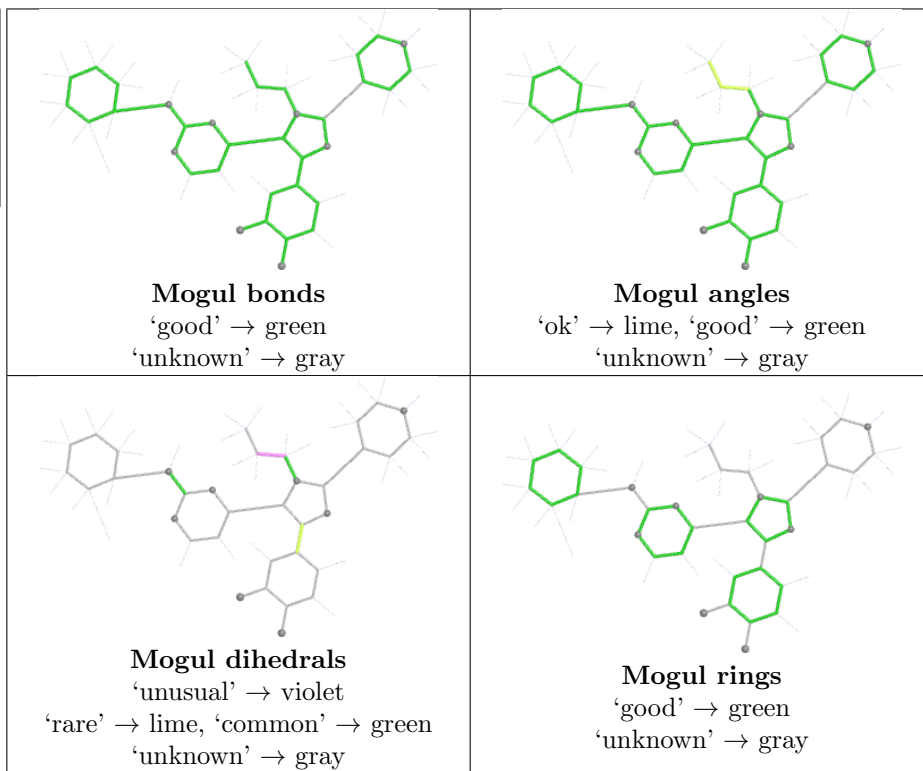
<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#density>



### 4.1.3 Mogul analysis for 880 A 501

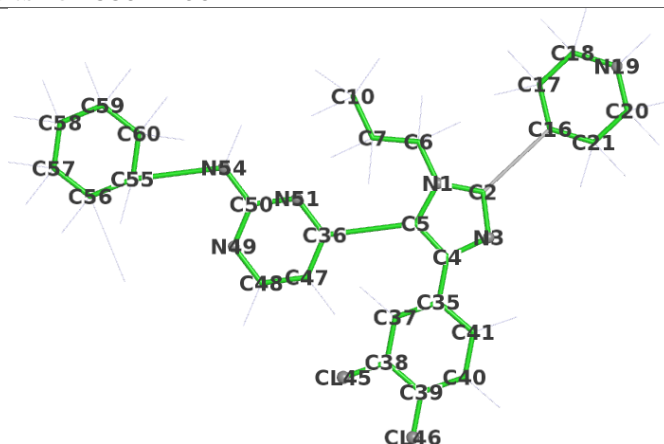
#### Summary

'bad' bonds	0/38
'bad' bond angles	0/43
'unusual' dihedrals	1/4
'bad' rings	0/4
bonds rms Z	0.585
angles rms Z	0.724



For help on "Ligand Mogul Analysis" see BUSTER wiki page  
<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#Mogul>

Mogul bond results for 880 A 501



Mogul bonds schematic

'good' → green (Z < 1.5)

'unknown' → gray (Mogul does not find sufficient CSD equivalents).

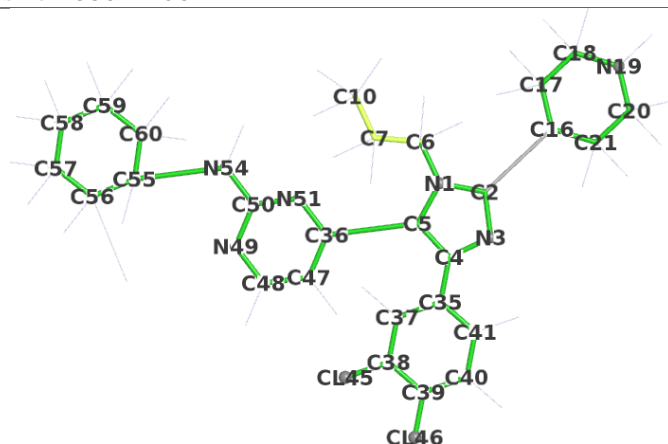
atoms	actual in Å	Mogul mean in Å	difference in Å	Mogul $\sigma$ in Å	Mogul # samples	Zscore
C38-CL45	1.719	1.734	-0.014	0.012	3693	1.23
C7-C6	1.542	1.514	0.028	0.023	243	1.20
C36-C5	1.486	1.478	0.008	0.007	10	1.13
C58-C57	1.535	1.516	0.019	0.018	1743	1.08
C50-N51	1.354	1.341	0.013	0.013	434	0.98
C2-N3	1.303	1.316	-0.013	0.015	17	0.88
C39-C38	1.378	1.388	-0.011	0.013	403	0.82
C17-C18	1.524	1.512	0.011	0.014	29	0.80
C59-C58	1.528	1.516	0.012	0.018	1743	0.68
C21-C20	1.521	1.512	0.009	0.014	29	0.65
C5-C4	1.386	1.377	0.009	0.013	125	0.64
C20-N19	1.476	1.467	0.010	0.018	139	0.56
C50-N49	1.346	1.340	0.006	0.011	146	0.53
C4-N3	1.378	1.385	-0.007	0.013	288	0.53
C41-C40	1.387	1.382	0.005	0.010	4486	0.51
C6-N1	1.473	1.467	0.006	0.012	575	0.50
C5-N1	1.396	1.389	0.007	0.014	79	0.49
C47-C48	1.386	1.381	0.006	0.011	2637	0.49
C40-C39	1.390	1.384	0.006	0.013	2673	0.46
C41-C35	1.396	1.391	0.006	0.013	4266	0.44

(table limited to 20 rows)

For help on "Ligand Mogul Analysis: Bonds" see BUSTER wiki page

<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#MogulBonds>

Mogul angle results for 880 A 501



Mogul angles schematic

'ok' → lime (1.5 < Z < 2.5)

'good' → green (Z < 1.5)

'unknown' → gray (Mogul does not find sufficient CSD equivalents).

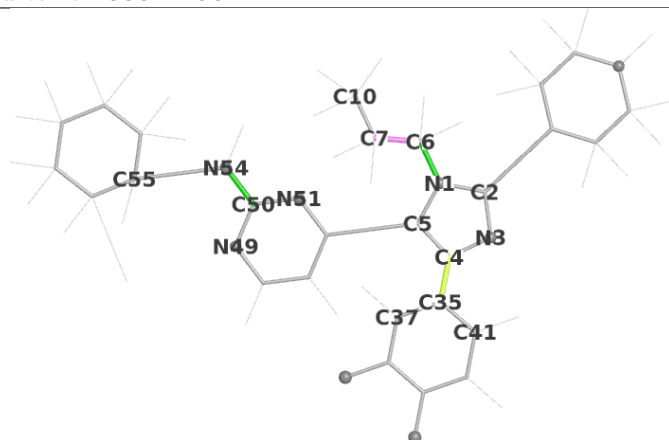
atoms	actual in °	Mogul mean in °	difference in °	Mogul $\sigma$ in °	Mogul # samples	Zscore
C10-C7-C6	117.3	111.8	5.4	2.5	217	2.16
C40-C39-CL46	120.3	118.4	2.0	1.3	2376	1.47
C35-C4-C5	131.8	130.0	1.8	1.3	92	1.35
C35-C4-N3	117.5	119.3	-1.8	1.4	227	1.27
C39-C38-CL45	119.9	120.9	-1.0	0.8	728	1.14
C6-N1-C5	128.2	126.6	1.5	1.4	14	1.09
C6-N1-C2	125.5	127.1	-1.7	1.6	26	1.06
C47-C36-N51	122.2	123.0	-0.8	0.8	691	1.01
C48-C47-C36	116.4	117.1	-0.7	0.7	16	1.01
C38-C39-CL46	120.1	120.9	-0.7	0.8	728	0.87
C37-C38-C39	121.0	120.2	0.8	1.0	476	0.85
C7-C6-N1	114.0	112.5	1.5	1.9	33	0.81
C47-C48-N49	123.4	124.1	-0.7	0.9	427	0.79
C4-C5-N1	104.8	105.4	-0.6	0.9	25	0.72
C60-C55-C56	110.1	110.7	-0.5	0.8	315	0.65
C60-C55-N54	109.5	110.6	-1.0	1.7	34	0.60
C41-C40-C39	119.7	120.2	-0.4	0.7	492	0.55
N54-C50-N51	117.2	116.5	0.7	1.4	22	0.50
N49-C50-N51	126.1	126.6	-0.5	1.0	20	0.48
C47-C36-C5	121.6	121.1	0.5	1.1	10	0.44

(table limited to 20 rows)

For help on "Ligand Mogul Analysis: Angles" see BUSTER wiki page

<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#MogulAngles>

Mogul dihedral results for 880 A 501



Mogul dihedrals schematic

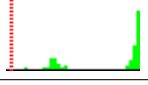
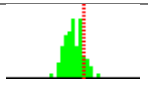
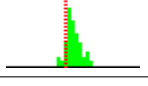
'unusual' → violet (some torsion angles have <0.5% of population within  $\pm 10^\circ$ )

'rare' → lime (all torsion angles have >0.5% of population within  $\pm 10^\circ$ )

'common' → green (all torsion angles have >10% of population within  $\pm 10^\circ$ )

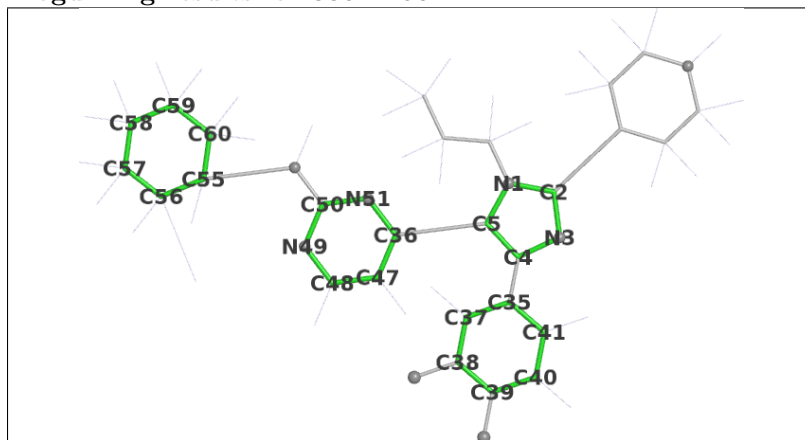
'unknown' → gray (Mogul does not find sufficient CSD equivalents).

atoms	actual torsion angle in $^\circ$	Mogul histogram	Mogul # samples	classification or % Mogul population within $\pm 10^\circ$
...-C35-C4-...				rare
C41-C35-C4-C5	-128.3		452	6%
C37-C35-C4-N3	-131.5		528	19%
C41-C35-C4-N3	47.8		528	16%
C37-C35-C4-C5	52.3		452	9%
...-C50-N54-...				common
N51-C50-N54-C55	1.7		22	45%
N49-C50-N54-C55	176.7		22	55%
...-C6-C7-...				unusual

atoms	actual torsion angle in °	Mogul histogram	Mogul # samples	classification or % Mogul population within $\pm 10^\circ$
N1-C6-C7-C10 ...-C6-N1-...	4.6		196	0% common
C7-C6-N1-C5	-103.0		97	41%
C7-C6-N1-C2	78.0		49	63%

For help on "Ligand Mogul Analysis: Dihedrals" see BUSTER wiki page  
<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#MogulDihedrals>

## Mogul ring results for 880 A 501



### Mogul rings schematic

'good' → green (ring strangeness score <math>< 3^\circ</math>)

'unknown' → gray (Mogul does not find sufficient CSD equivalents).

atoms	Mogul # samples	Ring strangeness score‡ in $^\circ$
C55-C56-C57-C58-C59-C60	148	0.3
N1-C2-N3-C4-C5	58	0.0
C36-C47-C48-N49-C50-N51	41	0.5
C35-C37-C38-C39-C40-C41	246	0.3

‡'ring strangeness score' is the RMS difference in torsion angles between the instance of the ring in the ligand in the model, and the nearest instance that mogul finds in the CSD.

For help on "Ligand Mogul Analysis: Rings" see BUSTER wiki page

<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#MogulRings>

## 4.2 ANP A 502

### 4.2.1 Statistics for ligand

Database ID	ANP (PDB)
3-letter code	ANP
CC( $2mF_o-DF_c$ )	0.7893
min(B-factor) $\ddagger$	37.9
avg(B-factor) $\ddagger$	46.3
max(B-factor) $\ddagger$	53.4
min(occupancy) $\ddagger$	0.50
max(occupancy) $\ddagger$	0.50
$\ddagger$ hydrogen atoms excluded	

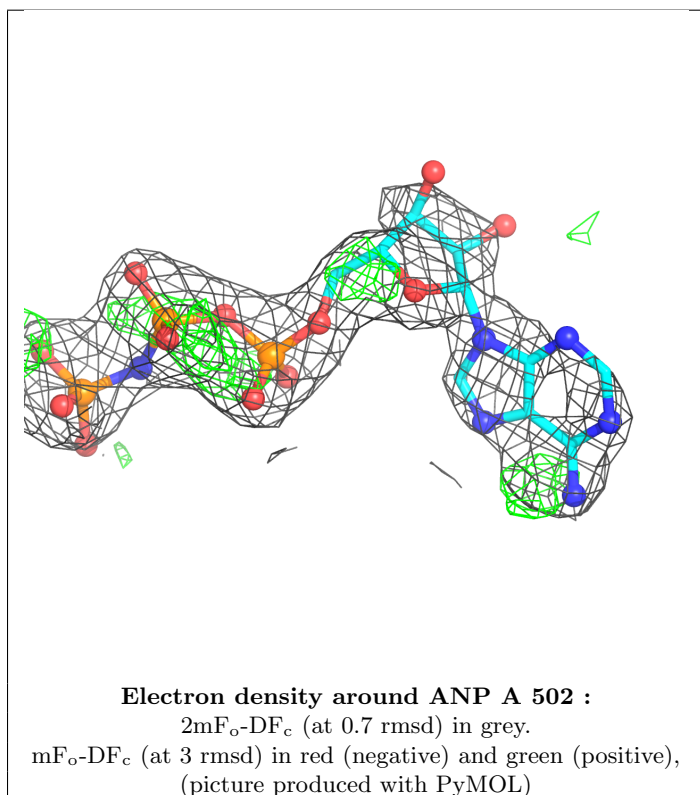
### Restraints used

restraints for ANP (PHOSPHOAMINOPHOSPHONIC ACID-ADENYLATE ESTER) from cif dictionary ANP.cif; bUSTER common-compounds v 3.0, Generated by GRADE\_PDB\_LIGAND 1.2.9pre (June 16 2014) using MOGUL 1.6.1(DEV7), CSD as535be, with quantum mechanics RM1

For help on “Ligand Statistics Table” see BUSTER wiki page

<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#statistics>

### 4.2.2 Picture of ligand in electron density



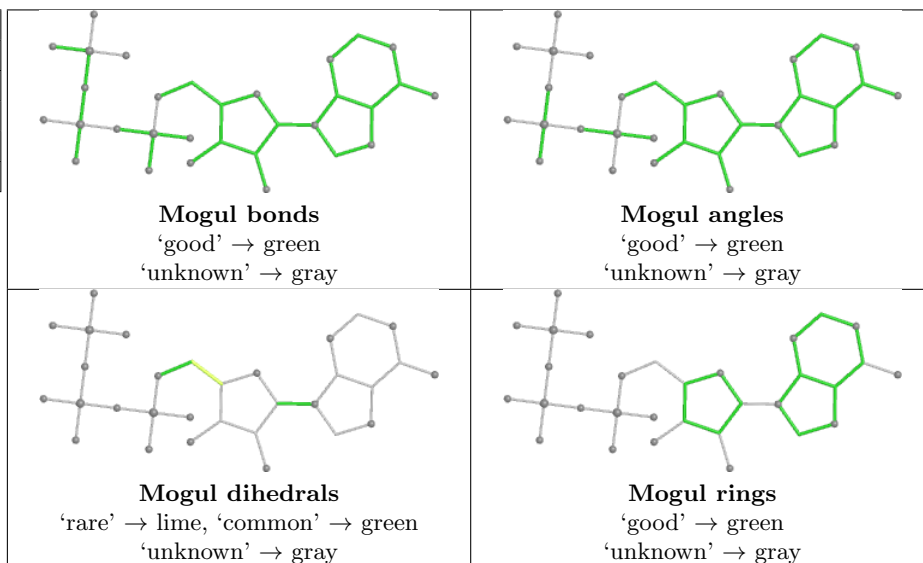
For help on “Ligand Electron Density Picture” see BUSTER wiki page

<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#density>

### 4.2.3 Mogul analysis for ANP A 502

#### Summary

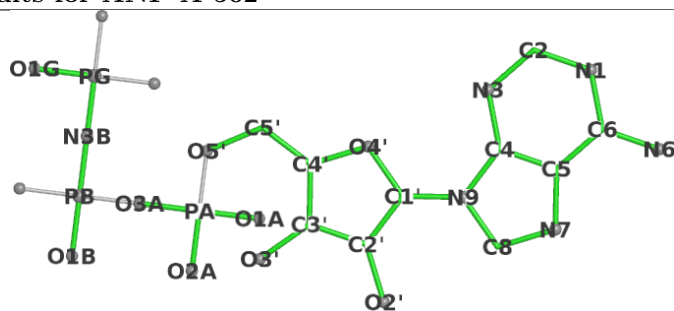
'bad' bonds	0/28
'bad' bond angles	0/33
'unusual' dihedrals	0/3
'bad' rings	0/3
bonds rms $Z$	0.252
angles rms $Z$	0.338



For help on "Ligand Mogul Analysis" see BUSTER wiki page  
<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#Mogul>



Mogul bond results for ANP A 502



Mogul bonds schematic

'good' → green ( $Z < 1.5$ )

'unknown' → gray (Mogul does not find sufficient CSD equivalents).

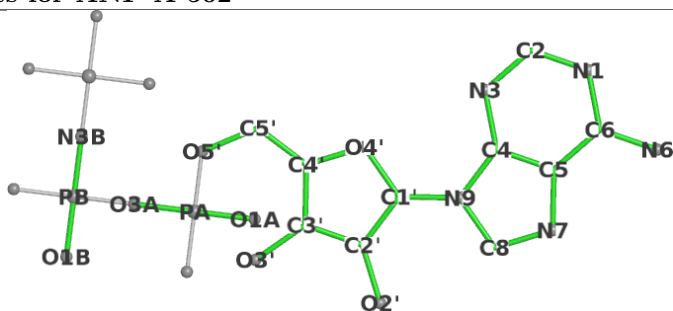
atoms	actual in Å	Mogul mean in Å	difference in Å	Mogul $\sigma$ in Å	Mogul # samples	Zscore
C8-N9	1.373	1.370	0.004	0.008	190	0.48
C5-N7	1.384	1.387	-0.003	0.007	359	0.44
C6-N1	1.354	1.350	0.004	0.010	239	0.43
PA-O2A	1.516	1.527	-0.012	0.030	30	0.39
O4'-C4'	1.448	1.444	0.004	0.011	1633	0.39
C2-N1	1.339	1.335	0.004	0.011	829	0.37
C2-N3	1.338	1.335	0.004	0.011	829	0.36
PA-O1A	1.475	1.482	-0.006	0.018	31	0.34
PG-N3B	1.637	1.631	0.006	0.022	23	0.28
PB-N3B	1.637	1.631	0.006	0.022	23	0.28
C4-N3	1.342	1.339	0.003	0.012	542	0.26
C6-N6	1.333	1.336	-0.003	0.014	531	0.23
C4-N9	1.376	1.374	0.002	0.008	181	0.22
C3'-C4'	1.528	1.526	0.003	0.014	577	0.20
O4'-C1'	1.417	1.415	0.002	0.011	592	0.19
C5'-C4'	1.510	1.508	0.002	0.012	1182	0.16
C8-N7	1.310	1.311	-0.001	0.008	447	0.12
O5'-C5'	1.441	1.443	-0.002	0.015	124	0.11
C1'-N9	1.460	1.459	0.001	0.012	78	0.09
PG-O1G	1.455	1.454	0.001	0.009	68	0.09

(table limited to 20 rows)

For help on "Ligand Mogul Analysis: Bonds" see BUSTER wiki page

[http://www.globalphasing.com/buster/wiki/index.cgi?BR\\_LigandReportAfter201507#MogulBonds](http://www.globalphasing.com/buster/wiki/index.cgi?BR_LigandReportAfter201507#MogulBonds)

Mogul angle results for ANP A 502



Mogul angles schematic

'good' → green (Z < 1.5)

'unknown' → gray (Mogul does not find sufficient CSD equivalents).

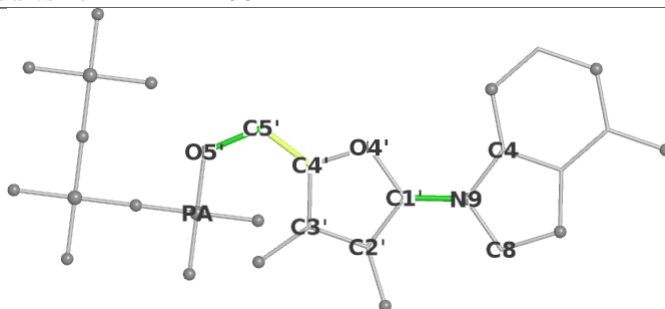
atoms	actual in °	Mogul mean in °	difference in °	Mogul $\sigma$ in °	Mogul # samples	Zscore
O5'-C5'-C4'	110.5	109.1	1.5	2.3	84	0.65
O3'-C3'-C2'	113.6	111.9	1.6	2.6	938	0.63
O2'-C2'-C3'	113.6	111.9	1.6	2.6	938	0.62
C4-N9-C8	105.4	105.7	-0.3	0.5	150	0.60
C2'-C1'-N9	115.0	114.1	0.9	1.5	59	0.58
O4'-C4'-C5'	110.3	109.6	0.7	1.5	204	0.48
C4'-O4'-C1'	110.2	109.5	0.7	1.4	434	0.47
C5'-C4'-C3'	114.5	115.3	-0.8	1.8	70	0.46
O4'-C4'-C3'	105.7	105.3	0.5	1.3	448	0.38
O1B-PB-N3B	111.5	112.1	-0.6	1.7	15	0.37
C1'-N9-C8	127.4	126.8	0.7	1.9	65	0.35
C6-C5-C4	116.8	117.0	-0.2	0.7	127	0.30
C2-N3-C4	110.9	111.5	-0.6	2.0	224	0.29
C3'-C2'-C1'	101.2	101.5	-0.3	1.2	273	0.29
O2'-C2'-C1'	111.3	110.6	0.8	2.8	323	0.27
C6-C5-N7	132.4	132.1	0.3	1.2	118	0.27
O4'-C1'-N9	108.7	108.4	0.3	1.3	66	0.25
N3-C4-N9	127.3	127.1	0.3	1.2	126	0.24
C5-C6-N1	117.8	117.6	0.2	0.9	133	0.22
C2-N1-C6	118.7	118.5	0.2	1.0	154	0.20

(table limited to 20 rows)

For help on "Ligand Mogul Analysis: Angles" see BUSTER wiki page

<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#MogulAngles>

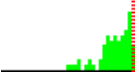
Mogul dihedral results for ANP A 502



Mogul dihedrals schematic

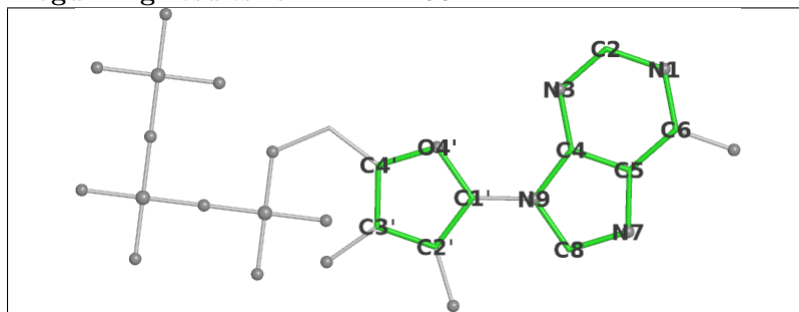
'rare' → lime (all torsion angles have >0.5% of population within  $\pm 10^\circ$ )  
 'common' → green (all torsion angles have >10% of population within  $\pm 10^\circ$ )  
 'unknown' → gray (Mogul does not find sufficient CSD equivalents).

atoms	actual torsion angle in $^\circ$	Mogul histogram	Mogul # samples	classification or % Mogul population within $\pm 10^\circ$
...-C1'-N9-...				common
C2'-C1'-N9-C4	119.5		66	18%
O4'-C1'-N9-C8	54.2		64	16%
O4'-C1'-N9-C4	-121.2		68	18%
C2'-C1'-N9-C8	-65.2		63	17%
...-C4'-C5'-...				rare
O4'-C4'-C5'-O5'	36.4		63	3%
C3'-C4'-C5'-O5'	155.4		63	2%
...-C5'-O5'-...				common

atoms	actual torsion angle in °	Mogul histogram	Mogul # samples	classification or % Mogul population within $\pm 10^\circ$
C4'-C5'-O5'-PA	179.8		74	34%

For help on "Ligand Mogul Analysis: Dihedrals" see BUSTER wiki page  
<http://www.globalphasing.com/buster/wiki/index.cgi?BRLigandReportAfter201507#MogulDihedrals>

## Mogul ring results for ANP A 502



### Mogul rings schematic

'good' → green (ring strangeness score < 3 °)

'unknown' → gray (Mogul does not find sufficient CSD equivalents).

atoms	Mogul # samples	Ring strangeness score‡ in °
N9-C8-N7-C5-C4	186	0.0
C5-C6-N1-C2-N3-C4	184	0.2
C4'-O4'-C1'-C2'-C3'	191	0.4

‡'ring strangeness score' is the RMS difference in torsion angles between the instance of the ring in the ligand in the model, and the nearest instance that mogul finds in the CSD.

For help on "Ligand Mogul Analysis: Rings" see BUSTER wiki page

[http://www.globalphasing.com/buster/wiki/index.cgi?BR\\_LigandReportAfter201507#MogulRings](http://www.globalphasing.com/buster/wiki/index.cgi?BR_LigandReportAfter201507#MogulRings)

## 5 X-ray statistics

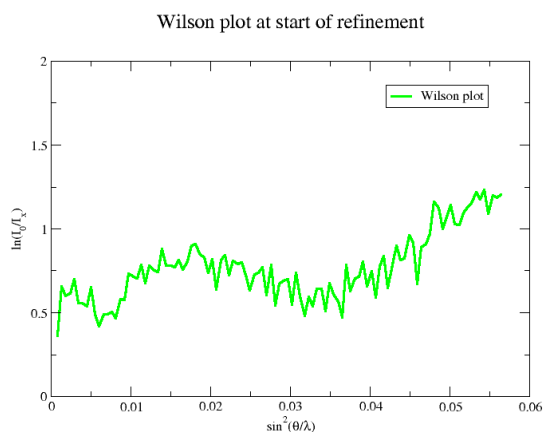
### 5.1 Scaling parameters in last cycle

Refined parameters		Unrefined parameters	
K_OVER	0.77251649	K_MISS	1.00000000
B_IMPF_FRAG	1.80352164	B_MISS	0.00000000
K_SOLV	0.73037453	K_IMPF_MISS	1.00000000
B_SOLV	51.90252837	B_IMPF_MISS	0.00000000
B_IMPF_SOLV	29.65066870	K_IMPF_SOLV	1.00000000
B_11	-34.24287568	B_OVER	0.00000000
B_22	13.96111574	B_12	0.00000000
B_33	20.28175994	B_13	0.00000000
		B_23	0.00000000
Anisotropic ratio		1.07	

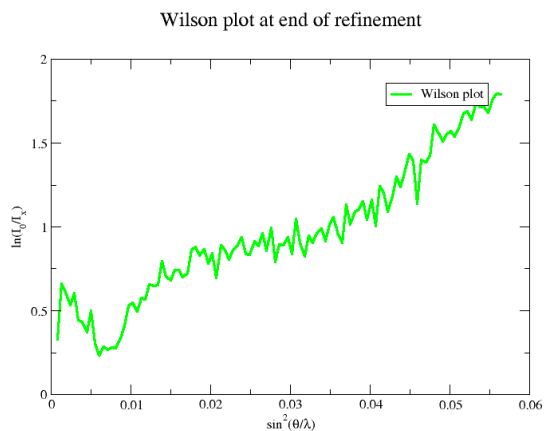
For help on “X-ray scaling parameters” see BUSTER wiki page  
<http://www.globalphasing.com/buster/wiki/index.cgi?BRScalingInfo>

### 5.2 Wilson plots

#### 5.2.1 Wilson plot at start of refinement

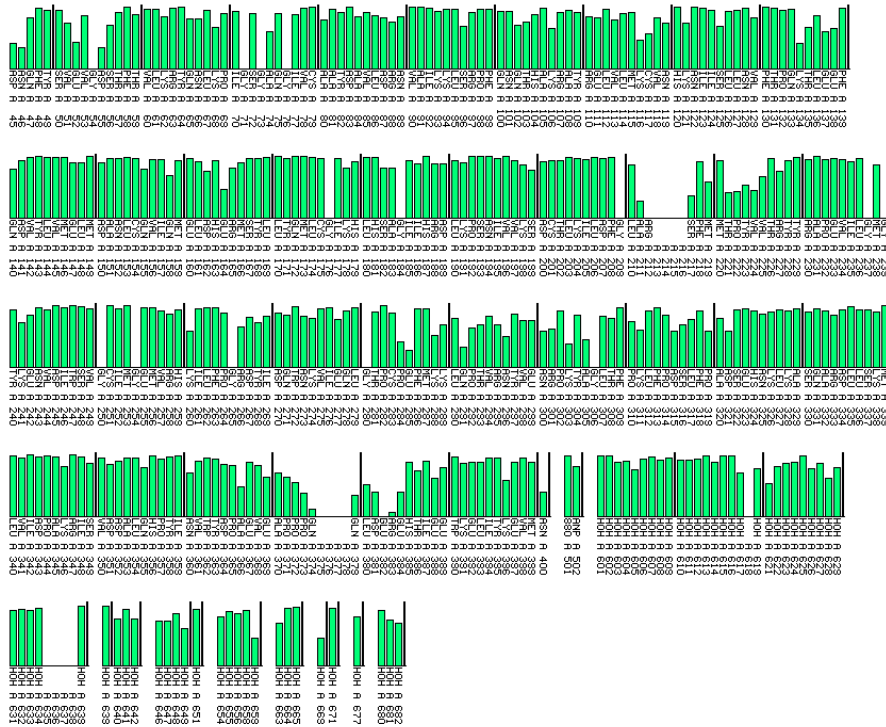


#### 5.2.2 Wilson plot at end of refinement



# 6 Real-space correlations

## 6.1 Side chains of chain A



## 6.2 Mainchain of chain A

