

Analysis output: all-atom contacts and geometry for TJP2_511-887H.pdb

Summary statistics

All-Atom Contacts	Clashscore, all atoms:	19.4		34 th percentile* (N=1784, all resolutions)
	Clashscore is the number of serious steric overlaps (> 0.4 Å) per 1000 atoms.			
Protein Geometry	Poor rotamers	9	2.81%	Goal: <0.3%
	Favored rotamers	302	94.38%	Goal: >98%
	Ramachandran outliers	7	1.87%	Goal: <0.05%
	Ramachandran favored	353	94.13%	Goal: >98%
	MolProbity score [^]	2.52		46 th percentile* (N=27675, 0Å - 99Å)
	Cβ deviations >0.25Å	0	0.00%	Goal: 0
	Bad bonds:	11 / 3034	0.36%	Goal: 0%
	Bad angles:	15 / 4090	0.37%	Goal: <0.1%
Peptide Omegas	Cis Prolines:	0 / 12	0.00%	Expected: ≤1 per chain, or ≤5%
	Twisted Peptides:	1/376	0.27%	Goal: 0

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

By adding H to this model and allowing Asn/Gln/His flips, we could *automatically* fix 1 bad rotamers and improve your clashscore by 2.18 points.

Multi-criterion visualizations

^{* 100&}lt;sup>th</sup> percentile is the best among structures of comparable resolution; 0th percentile 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.



View (408 Kb)

Single-criterion visualizations

- Clash list (5 Kb): View
- Ramachandran plot kinemage (427 Kb): View in KiNG | Download
- Ramachandran plot PDF (1.7 Mb): View
- Cβ deviation scatter plot (30 Kb): View in KiNG | Download

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