

CING; an integrated residue-based structure validation program suite

Supplementary Materials

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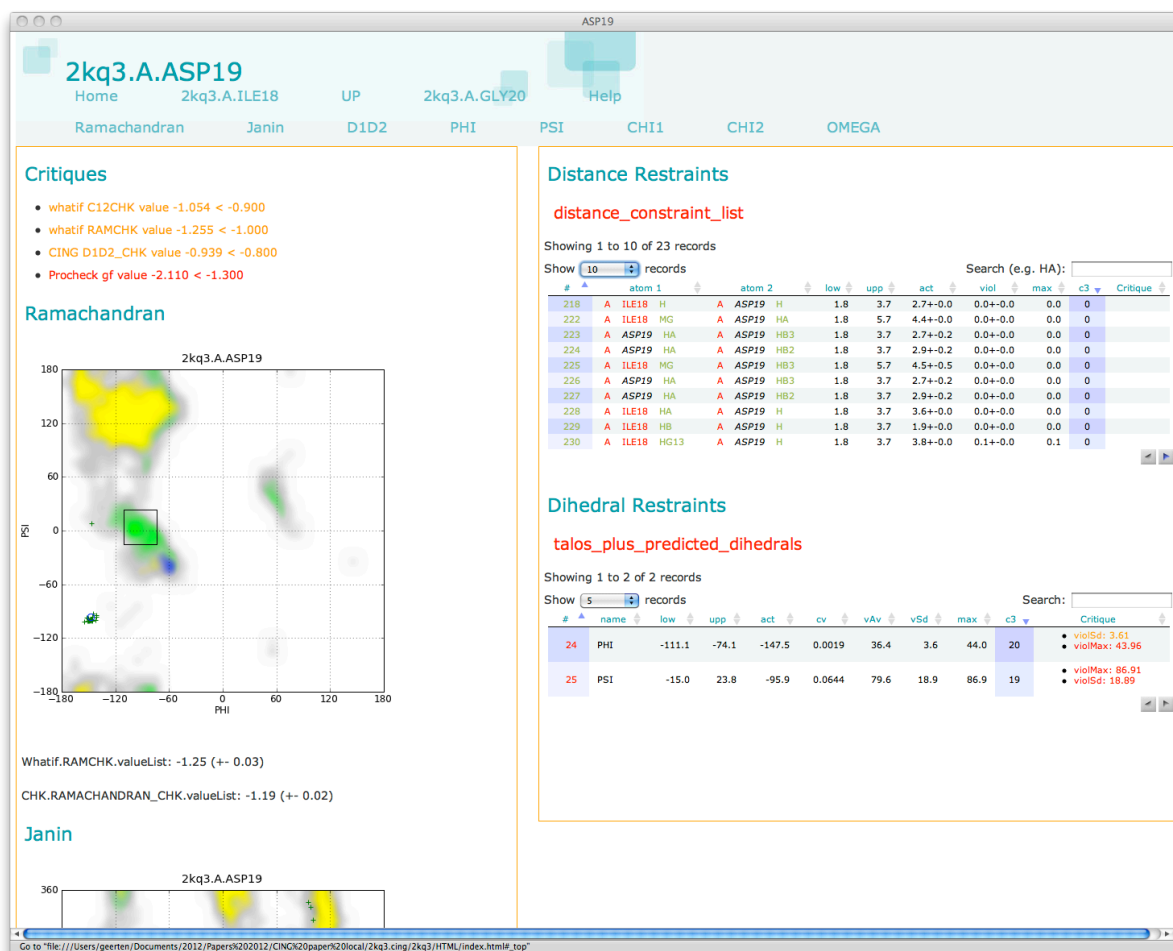
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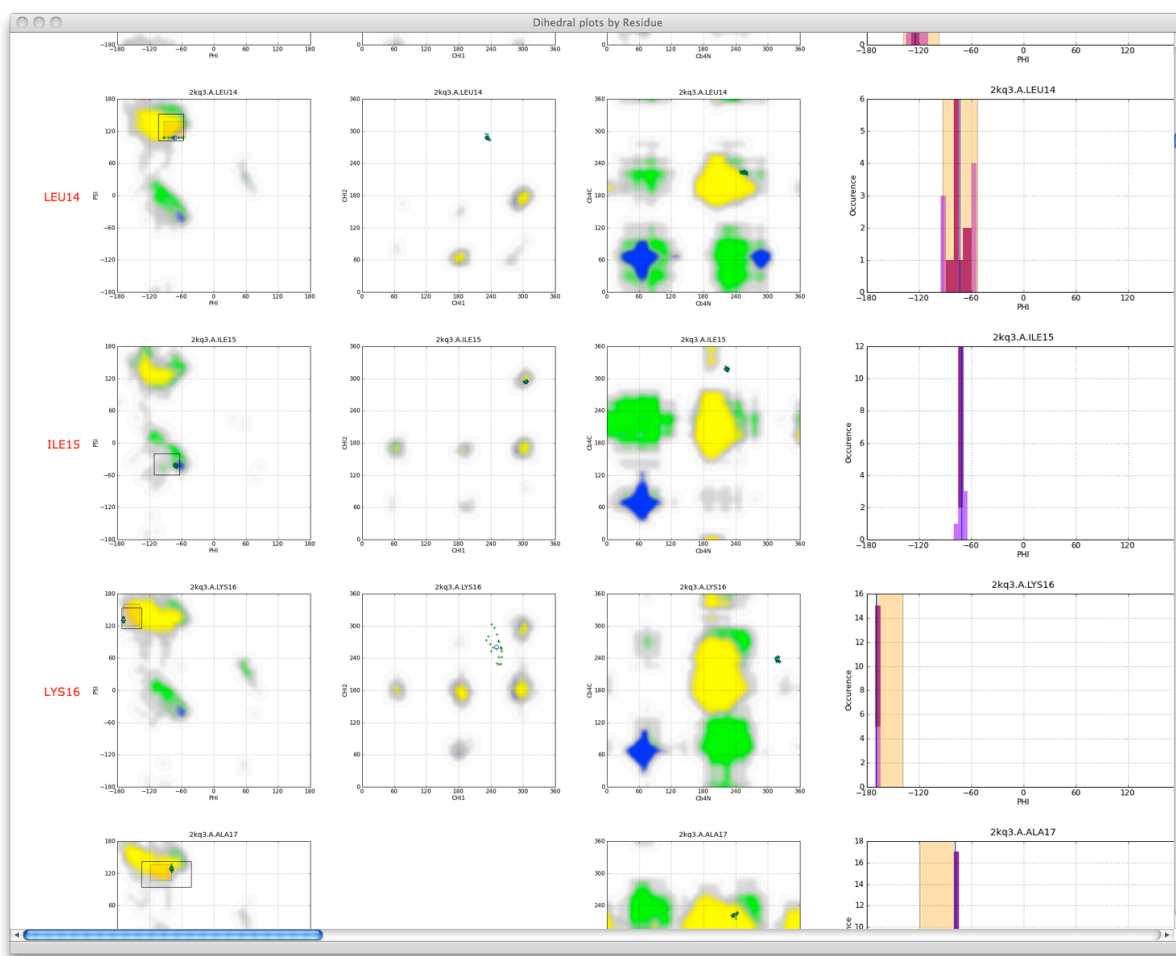
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Supplementary Figure S1.
CING Residue page for PDB entry 2kq3 residue Asp19.



Supplementary Figure S2.
CING 'Dihedral plots by residue' page for PDB entry 2kq3, residues Leu14-Lys16.