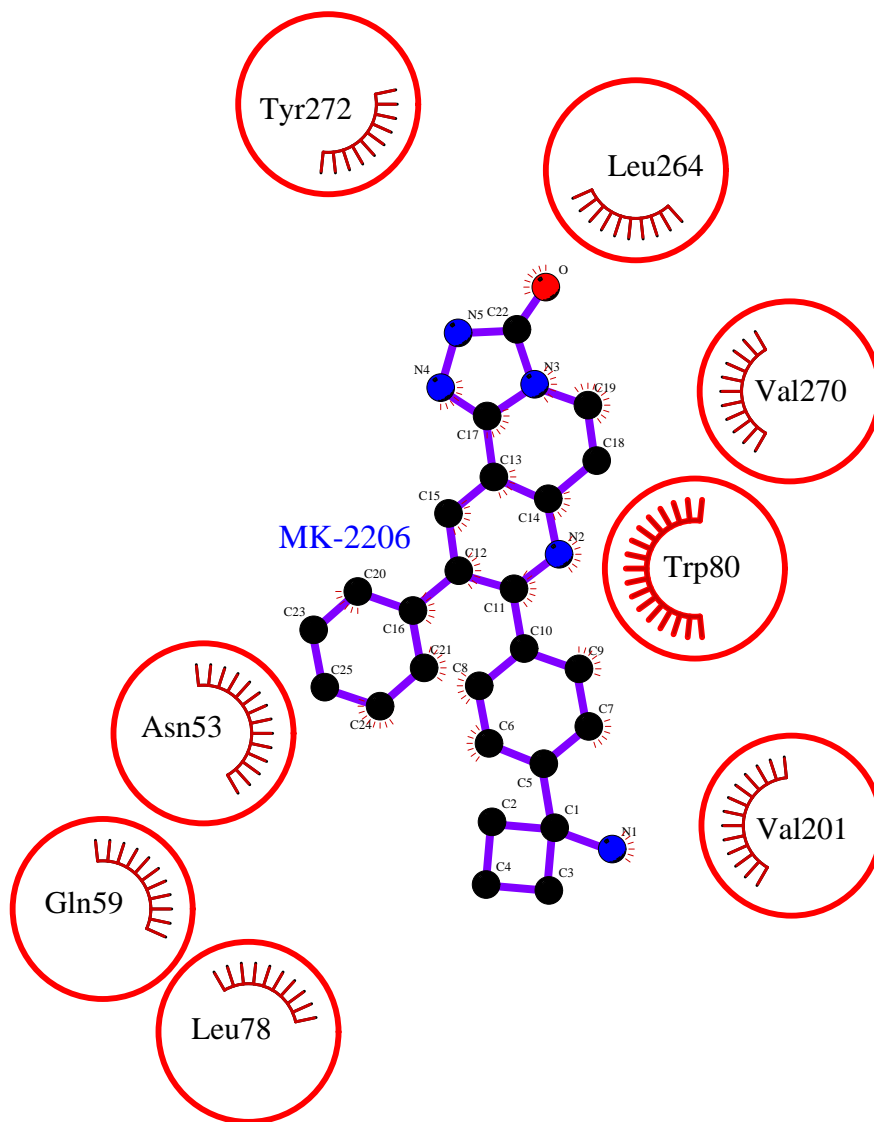
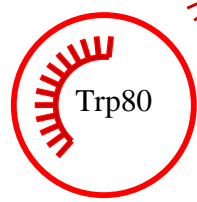
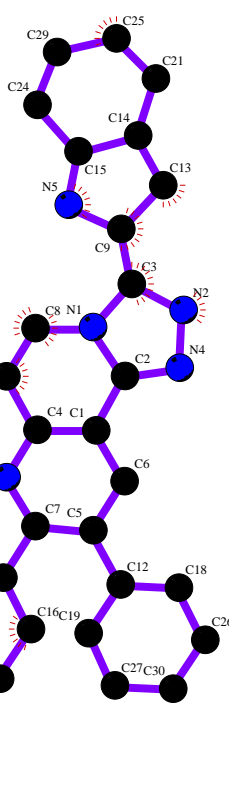
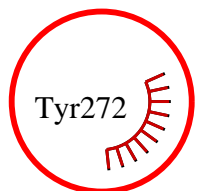


R₁R₂-analogs of MK-2206

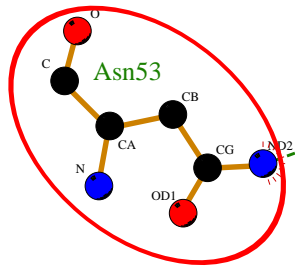
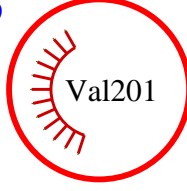
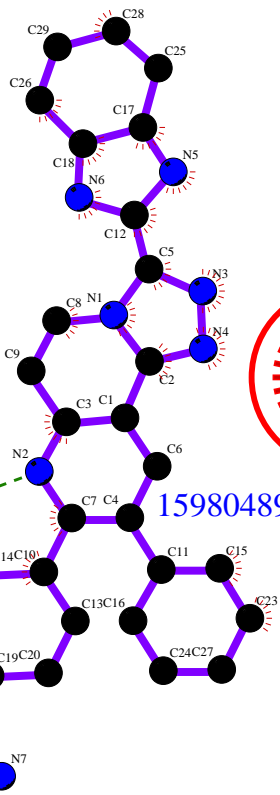
Ligand-interaction plots of R₁R₂-analogs of MK-2206 are shown. The MK-2206 analogs' names are indicated by PubChem ID. The hydrogen bonds are shown as green-dashed lines with indicated bond length and the residues involved in hydrophobic interactions are shown as red arcs. The residues which are common to the first compound, the drug MK-2206 are encircled.





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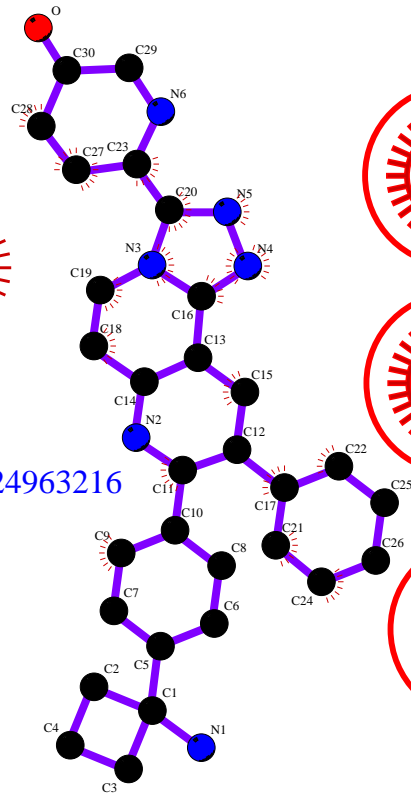


Gln79

Asn53

Lys268

24963216



Val270

Trp80

Val201

