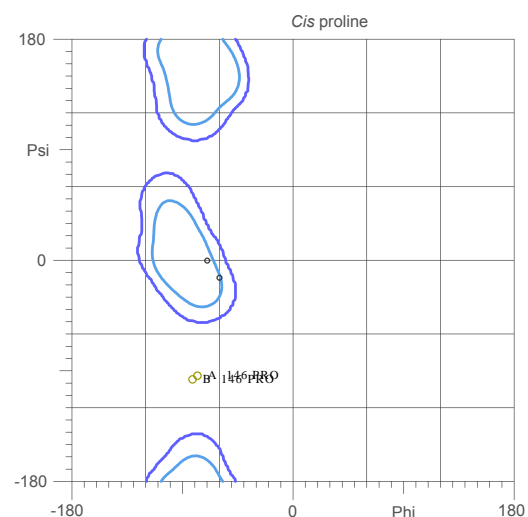
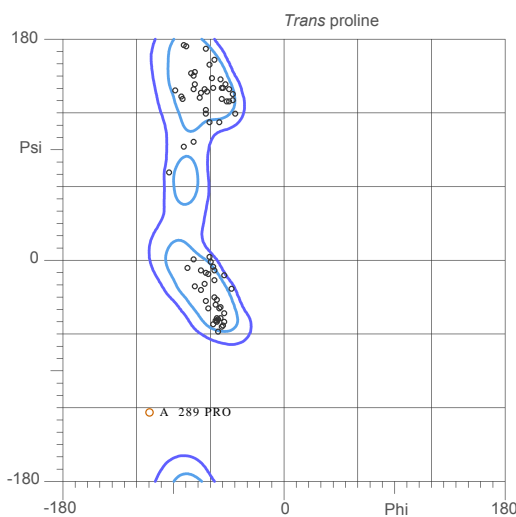
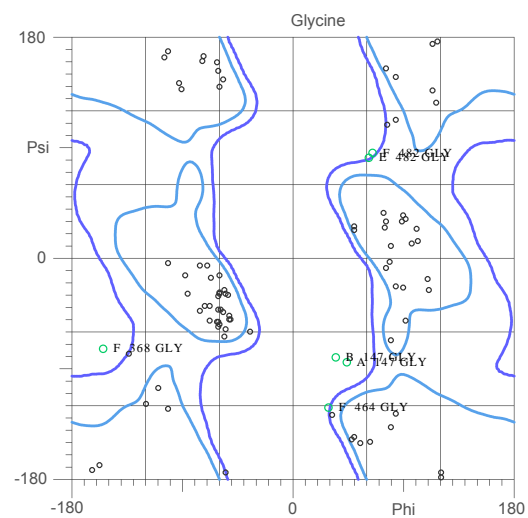
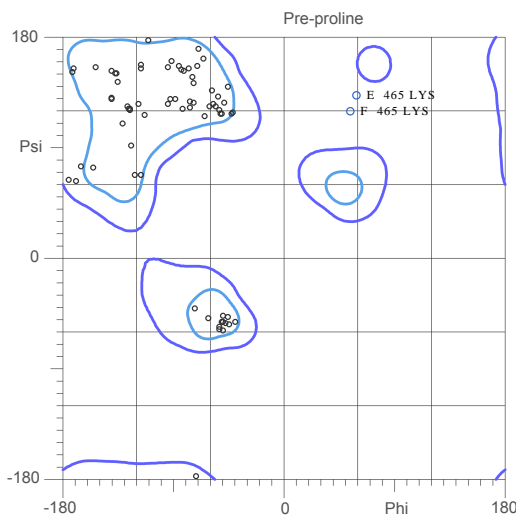
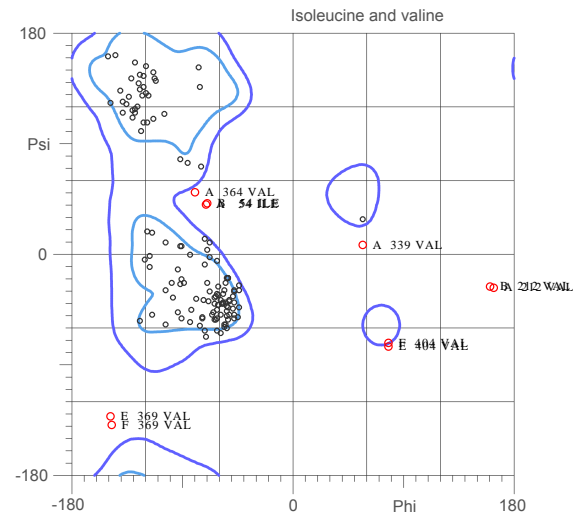
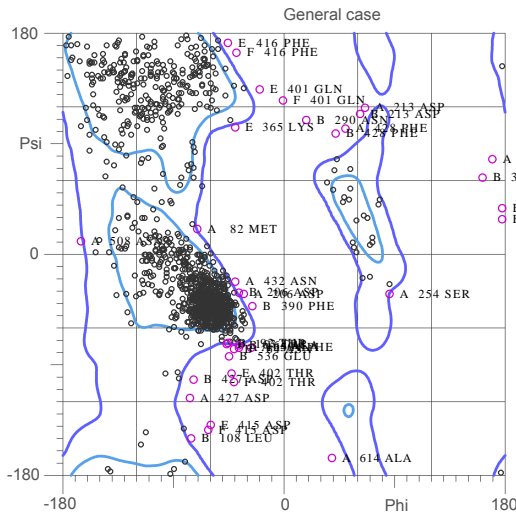


MolProbity Ramachandran analysis

3d0i.H.pdb, model 1



87.1% (131/150) of all residues were in favored (98%) regions.
96.7% (143/147) of all residues were in allowed (100%) regions.

Shown here: 37 outliers (Phi, Psi):

A 340 GLN (17.4, 45.3)	B 340 GLN (17.4, 45.3)
A 82 MET (17.4, 45.3)	B 82 MET (17.4, 45.3)
A 401 GLN (17.4, 45.3)	B 401 GLN (17.4, 45.3)
A 402 THR (17.4, 45.3)	B 402 THR (17.4, 45.3)
A 427 ASP (17.4, 45.3)	B 427 ASP (17.4, 45.3)
A 432 ASN (17.4, 45.3)	B 432 ASN (17.4, 45.3)
A 465 LYS (17.4, 45.3)	B 465 LYS (17.4, 45.3)
A 536 GLU (17.4, 45.3)	B 536 GLU (17.4, 45.3)
A 614 ALA (17.4, 45.3)	B 614 ALA (17.4, 45.3)
A 129 PHE (17.4, 45.3)	B 129 PHE (17.4, 45.3)
A 206 ASP (17.4, 45.3)	B 206 ASP (17.4, 45.3)
A 289 PRO (17.4, 45.3)	B 289 PRO (17.4, 45.3)
A 364 VAL (17.4, 45.3)	B 364 VAL (17.4, 45.3)
A 369 VAL (17.4, 45.3)	B 369 VAL (17.4, 45.3)
A 370 SER (17.4, 45.3)	B 370 SER (17.4, 45.3)
A 415 ASP (17.4, 45.3)	B 415 ASP (17.4, 45.3)
A 416 PHE (17.4, 45.3)	B 416 PHE (17.4, 45.3)
A 429 THR (17.4, 45.3)	B 429 THR (17.4, 45.3)
A 431 THR (17.4, 45.3)	B 431 THR (17.4, 45.3)
A 433 THR (17.4, 45.3)	B 433 THR (17.4, 45.3)
A 434 THR (17.4, 45.3)	B 434 THR (17.4, 45.3)
A 435 THR (17.4, 45.3)	B 435 THR (17.4, 45.3)
A 436 THR (17.4, 45.3)	B 436 THR (17.4, 45.3)
A 437 THR (17.4, 45.3)	B 437 THR (17.4, 45.3)
A 438 THR (17.4, 45.3)	B 438 THR (17.4, 45.3)
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A 443 THR (17.4, 45.3)	B 443 THR (17.4, 45.3)
A 444 THR (17.4, 45.3)	B 444 THR (17.4, 45.3)
A 445 THR (17.4, 45.3)	B 445 THR (17.4, 45.3)
A 446 THR (17.4, 45.3)	B 446 THR (17.4, 45.3)
A 447 THR (17.4, 45.3)	B 447 THR (17.4, 45.3)
A 448 THR (17.4, 45.3)	B 448 THR (17.4, 45.3)
A 449 THR (17.4, 45.3)	B 449 THR (17.4, 45.3)
A 450 THR (17.4, 45.3)	B 450 THR (17.4, 45.3)
A 451 THR (17.4, 45.3)	B 451 THR (17.4, 45.3)
A 452 THR (17.4, 45.3)	B 452 THR (17.4, 45.3)
A 453 THR (17.4, 45.3)	B 453 THR (17.4, 45.3)
A 454 THR (17.4, 45.3)	B 454 THR (17.4, 45.3)
A 455 THR (17.4, 45.3)	B 455 THR (17.4, 45.3)
A 456 THR (17.4, 45.3)	B 456 THR (17.4, 45.3)
A 457 THR (17.4, 45.3)	B 457 THR (17.4, 45.3)
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A 473 THR (17.4, 45.3)	B 473 THR (17.4, 45.3)
A 474 THR (17.4, 45.3)	B 474 THR (17.4, 45.3)
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A 476 THR (17.4, 45.3)	B 476 THR (17.4, 45.3)
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A 496 THR (17.4, 45.3)	B 496 THR (17.4, 45.3)
A 497 THR (17.4, 45.3)	B 497 THR (17.4, 45.3)
A 498 THR (17.4, 45.3)	B 498 THR (17.4, 45.3)
A 499 THR (17.4, 45.3)	B 499 THR (17.4, 45.3)
A 500 THR (17.4, 45.3)	B 500 THR (17.4, 45.3)