ConSurf Job Information

Go to Results of Job 1678114938 (until they are deleted from the server).

The information below is obtained from the header of PDB file AF_vglut1NAT_rank1_with_seqres_consurf_firstglance.pdb

This PDB file can be saved to your computer.

Job Parameters

Job 1678114938 by ConSurf version 3.0 on 2023-03-06.

- PDB ID: AF_VGLUT1NAT_RANK1_WITH_SEQRES
- Chain processed: A
- Chains in the deposited model identical in sequence to chain processed: **A**
- Multiple sequence alignment source: Generated by ConSurf
- Multiple sequence alignment method: MAFFT
- Phylogenetic tree source: Generated by ConSurf

- Homologous sequences obtained from: UNIREF90
- PSI Blast E value cutoff: **0.0001**
- PSI Blast Iterations: 1
- Maximum homologous sequences to use: **150**
- Conservation score method: **Bayes**
- Amino acid substitution model: LG

For an explanation of the Job Parameters and Results, please see the on-line documentation for the <u>ConSurf</u> Server.

Job Results

Number of amino acids at each conservation grade, 1-9:

Black: excluding amino acids with insufficient data.

Yellow: including amino acids with insufficient data, shown only if different.

- SEQRES Length: **560** amino acids in experimental protein.
- Amino acids with coordinates (ATOM records): **294**
- Unique homologous sequences found: 2067
- Sequences used in the MSA: 150

- 1. 13 Variable
- 2. **6**
- 3. **20**
- 4. 40
- 5. **35**
- *6*. 36
- *7*. 53
- *8*. 53
- 9. 38 Conserved

Sum 294 294

- + 0 (Amino acids with insufficient data)
- = 294 (0% with insufficient data)

Close