- Also take a look at "Structural information"
- There is no crystal structures and the homology models from Modbase are not high quality. The homology model could be a guide, but I would not base an SBDD campaign against a *T.* brucei homology model.

## Structural information

## Modbase 3D models:

There are 3 models calculated for this protein. More info on these models, including the models themselves is available at: 
Modbase

Target Beg	Target End	Template	Template Beg	Template End	Identity	Evalue	Model Score	MPQS	zDope
32	156	1zoy (B)	16	151	46.00	0	1	0.950829	0.33
32	156	4ysx (B)	40	175	46.00	0	0.99	0.956829	0.26
168	229	5i9f (A)	348	409	23.00	0.53	0.98	0.708498	-1.37

+ Help me make sense of these data.

Target Beg: first modeled residue
Target End: last modeled residue

Template: template structure used for modelling (PDB accession and chain)

Template Beg: first template residue in target-template alignment Template End: last template residue in target-template alignment

Identity: sequence identity

Evalue: E value for target-template hit

Model Score: GA341 score (>0.7 for reliable model)

MPQS: ModPipe Quality Score (>1.1 for reliable model)

- The structure we have used as a basis for modeling studies has been 1NEK. I tried this in the PockeTome (<a href="http://ablab.ucsd.edu/POCKETOME/">http://ablab.ucsd.edu/POCKETOME/</a>). There was no record for it.
- Try a search for "succinate". Did you find SDH?

