- Click on Tb927.8.3380 and scroll down to Essentiality Data
- Suppose that we discover a SDH inhibitor in our SBDD campaign against the ubiquinone binding site of *P. Aeruginosa*. Based on these data, which other species would be most worth testing its efficacy against? Which would be least worth testing?
- The inhibitor should be tested against *T. brucei* and *T. gondii*. It probably won't be effective against *M. tuberculosis* and *P. berghei*.

Essentiality Tb927.8.3380 has direct evidence of essentiality Gene/Ortholog Source Study Organism Phenotype mtu1581 Mycobacterium tuberculosis non-essential nmpdr mtu3379 Mycobacterium tuberculosis non-essential nmpdr Tb09.160.4380 Trypanosoma brucei no significant loss or gain of fitness in bloodstream forms (3 days) alsford significant gain of fitness in bloodstream forms (6 days) alsford Tb09.160.4380 Trypanosoma brucei no significant loss or gain of fitness in procyclic forms alsford Tb09.160.4380 Trypanosoma brucei significant gain of fitness in differentiation of procyclic to bloodstream forms alsford Tb09.160.4380 Trypanosoma brucei alsford Tb927.8.3380 this record significant loss of fitness in bloodstream forms (3 days) Trypanosoma brucei significant loss of fitness in bloodstream forms (6 days) alsford Tb927.8.3380 this record Trypanosoma brucei Tb927.8.3380 this record alsford significant loss of fitness in procyclic forms Trypanosoma brucei Tb927.8.3380 this record Trypanosoma brucei significant loss of fitness in differentiation of procyclic to bloodstream forms alsford ь0724 Escherichia coli non-essential goodall b4153 Escherichia coli goodall non-essential

- 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)