Model Building Report

This document lists the results for the homology modelling project "Hemoglobin b" submitted to SWISS-MODEL workspace on April 24, 2018, 9:53 a.m.. The submitted primary amino acid sequence is given in Table T1.

If you use any results in your research, please cite the relevant publications:

- Biasini, M., Bienert, S., Waterhouse, A., Arnold, K., Studer, G., Schmidt, T., Kiefer, F., Cassarino, T.G., Bertoni, M., Bordoli, L., Schwede, T. SWISS-MODEL: modelling protein tertiary and quaternary structure using evolutionary information. Nucleic
- Acids Res. 42, W252-W258 (2014). [M] [1015] • Guex, N., Peitsch, M.C., Schwede, T. Automated comparative protein structure modeling with SWISS-MODEL and Swiss-PdbViewer: A historical perspective. Electrophoresis 30, S162-S173 (2009). [Million 10]
- Bienert, S., Waterhouse, A., de Beer, T.A., Tauriello, G., Studer, G., Bordoli, L., Schwede, T. The SWISS-MODEL Repository -
- Bioinformatics 27, 343-350 (2011). [M] 11015 • Bertoni, M., Kiefer, F., Biasini, M., Bordoli, L., Schwede, T. Modeling protein quaternary structure of homo- and hetero-

• Benkert, P., Biasini, M., Schwede, T. Toward the estimation of the absolute quality of individual protein structure models.

- oligomers beyond binary interactions by homology. Scientific Reports 7 (2017). [[7] [7] [7]
- Results

The SWISS-MODEL template library (SMTL version 2018-04-18, PDB release 2018-04-13) was searched with BLAST (Camacho et al.) and HHBlits (Remmert et al.) for evolutionary related structures matching the target sequence in Table T1. For details on the template search, see Materials and Methods. Overall 1267 templates were found (Table T2).

Models

The following model was built (see Materials and Methods "Model Building"):

Model #01	File	Built with ProMod3 Version 1.1.0.	Oligo- State monomer	Ligands	GMQE	QMEAN -1.10
	PDB			1 x HEM: PROTOPORPHYRIN IX CONTAINING FE; 1 x OXY: OXYGEN MOLECULE;	0.98	
QMEAN Cβ All Atom Solvation Torsion	-1.10 -1.04 0.96 1.69 -1.33		Predicted Local Similarity to Target 7.0 8.0 9.0 7.0 7.0 7.0 7.0 7.0 7.0 7		Escorej = 2 = 2 - - 2 = 2 - - 2 2 - - - 2 = - - - - - - - - -	

Template Seq Identity Oligo-state Found by Method Resolution Seq Similarity Range Coverage Description 6bb5.1.A 100.00 monomer HHblits X-ray 2.28Å 0.60 1 - 137 1.00 Hemoglobin subunit alpha

Ligand	Added to Model	Description			
HEM	✓	PROTOPORPHYRIN IX CONTAINING FE			
OXY	✓	OXYGEN MOLECULE			
HEM	× - Binding site not conserved.	PROTOPORPHYRIN IX CONTAINING FE			
HEM	× - Binding site not conserved.	PROTOPORPHYRIN IX CONTAINING FE			
HEM	× - Binding site not conserved.	PROTOPORPHYRIN IX CONTAINING FE			
OXY	× - Binding site not conserved.	OXYGEN MOLECULE			
OXY	× - Binding site not conserved.	OXYGEN MOLECULE			
OXY	X - Binding site not conserved.	OXYGEN MOLECULE			

LSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHGKKVADALTNAVAHVDDMPNALS 6bb5.1.A LSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHGKKVADALTNAVAHVDDMPNALS

ALSDLHAHKLRVDPVNFKLLSHCLLVTLAAHLPAEFTPAVHASLDKFLASVSTVLTS 6bb5.1.A ALSDLHAHKLRVDPVNFKLLSHCLLVTLAAHLPAEFTPAVHASLDKFLASVSTVLTS

Materials and Methods

Template Search

Template search with BLAST and HHBlits has been performed against the SWISS-MODEL template library (SMTL, last update: 2018-04-18, last included PDB release: 2018-04-13).

The target sequence was searched with BLAST against the primary amino acid sequence contained in the SMTL. A total of 504 templates were found.

An initial HHblits profile has been built using the procedure outlined in (Remmert et al.), followed by 1 iteration of HHblits against NR20. The obtained profile has then be searched against all profiles of the SMTL. A total of 941 templates were found.

Model Building

For each identified template, the template's quality has been predicted from features of the target-template alignment. The templates with the highest quality have then been selected for model building.

Models are built based on the target-template alignment using ProMod3. Coordinates which are conserved between the target and the template are copied from the template to the model. Insertions and deletions are remodelled using a fragment library. Side

Template Selection

chains are then rebuilt. Finally, the geometry of the resulting model is regularized by using a force field. In case loop modelling with ProMod3 fails, an alternative model is built with PROMOD-II (Guex et al.). **Model Quality Estimation**

The global and per-residue model quality has been assessed using the QMEAN scoring function (Benkert et al.). For improved performance, weights of the individual QMEAN terms have been trained specifically for SWISS-MODEL.

Ligand Modelling Ligands present in the template structure are transferred by homology to the model when the following criteria are met: (a) The

ligands are annotated as biologically relevant in the template library, (b) the ligand is in contact with the model, (c) the ligand is not clashing with the protein, (d) the residues in contact with the ligand are conserved between the target and the template. If any of these four criteria is not satisfied, a certain ligand will not be included in the model. The model summary includes information on why and which ligand has not been included. **Oligomeric State Conservation**

The quaternary structure annotation of the template is used to model the target sequence in its oligomeric form. The method (Bertoni et al.) is based on a supervised machine learning algorithm, Support Vector Machines (SVM), which combines interface conservation, structural clustering, and other template features to provide a quaternary structure quality estimate (QSQE). The QSQE score is a number between 0 and 1, reflecting the expected accuracy of the interchain contacts for a model built based a given alignment and template. Higher numbers indicate higher reliability. This complements the GMQE score which estimates the accuracy of the tertiary structure of the resulting model.

BLAST

References

Camacho, C., Coulouris, G., Avagyan, V., Ma, N., Papadopoulos, J., Bealer, K., Madden, T.L. BLAST+: architecture and

applications. BMC Bioinformatics 10, 421-430 (2009). HHblits

Remmert, M., Biegert, A., Hauser, A., Söding, J. HHblits: lightning-fast iterative protein sequence searching by HMM-HMM alignment. Nat Methods 9, 173-175 (2012). [[M] 1015

Table T1:

Primary amino acid sequence for which templates were searched and models were built. LSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHGKKVADALTNAVAHVDDMPNALSALSDLHAHKLRVDPVNFKLL

SHCLLVTLAAHLPAEFTPAVHASLDKFLASVSTVLTS

Table T2:

Template	Seq Identity	Oligo- state	QSQE	Found by	Method	Resolution	Seq Similarity	Coverage	Description	
1bab.1.A	100.00	homo- dimer	0.67	HHblits	X-ray	1.50Å	0.60	1.00	HEMOGLOBIN THIONVILLE (DEOXY) (ALPHA CHAIN)	
6bb5.1.A	100.00	monomer		HHblits	X-ray	2.28Å	0.60	1.00	Hemoglobin subunit alpha	
5wog.1.A	100.00	monomer		HHblits	X-ray	1.54Å	0.60	1.00	Hemoglobin subunit alpha	
1y0a.1.A	100.00	homo- dimer	0.78	HHblits	X-ray	2.22Å	0.60	1.00	Hemoglobin alpha chain	
5sw7.1.A	100.00	homo- dimer	0.78	HHblits	X-ray	1.85Å	0.60	1.00	Hemoglobin subunit alpha	
1bzz.1.A	100.00	homo- dimer	0.73	HHblits	X-ray	1.59Å	0.60	1.00	PROTEIN (HEMOGLOBIN ALPH CHAIN)	
1y0d.1.A	100.00	monomer		HHblits	X-ray	2.10Å	0.60	1.00	Hemoglobin alpha chain	
3ia3.1.B	100.00	monomer		HHblits	X-ray	3.20Å	0.60	1.00	Hemoglobin subunit alpha	
4yu3.1.A	87.59	monomer		HHblits	X-ray	2.45Å	0.57	1.00	hemoglobin	
4yu4.1.C	87.59	homo- dimer	0.35	HHblits	X-ray	2.80Å	0.57	1.00	hemoglobin	
1fsx.1.C	87.59	homo- dimer	0.71	HHblits	X-ray	2.10Å	0.56	1.00	HEMOGLOBIN ALPHA CHAIN	
1fsx.1.A	87.59	homo- dimer	0.71	HHblits	X-ray	2.10Å	0.56	1.00	HEMOGLOBIN ALPHA CHAIN	
5c6e.1.C	87.59	homo- dimer	0.65	HHblits	neutron diff.	2.00Å	0.56	1.00	Hemoglobin subunit alpha	
1iwh.1.A	87.59	monomer		HHblits	X-ray	1.55Å	0.56	1.00	Hemoglobin alpha chain	
2dhb.1.A	86.86	homo- dimer	0.65	HHblits	X-ray	2.80Å	0.56	1.00	HEMOGLOBIN (DEOXY) (ALPH CHAIN)	
1g0a.1.A	87.59	homo- dimer	0.58	HHblits	X-ray	2.04Å	0.56	1.00	HEMOGLOBIN ALPHA CHAIN	
1g08.1.A	87.59	monomer		HHblits	X-ray	1.90Å	0.56	1.00	HEMOGLOBIN ALPHA CHAIN	
1g09.1.C	87.59	homo- dimer	0.49	HHblits	X-ray	2.04Å	0.56	1.00	HEMOGLOBIN ALPHA CHAIN	
1hda.1.A	87.59	homo- dimer	0.75	HHblits	X-ray	2.20Å	0.56	1.00	HEMOGLOBIN (DEOXY) (ALPH CHAIN)	
3fh9.1.A	86.13	homo- dimer	0.65	HHblits	X-ray	1.62Å	0.56	1.00	Hemoglobin alpha chain	
3hrw.1.C	85.40	monomer		HHblits	X-ray	2.80Å	0.56	1.00	Hemoglobin subunit alpha	
3hrw.1.A	85.40	monomer		HHblits	X-ray	2.80Å	0.56	1.00	Hemoglobin subunit alpha	
3gqp.1.A	84.67	homo- dimer	0.72	HHblits	X-ray	2.00Å	0.56	1.00	Hemoglobin subunit alpha	
3gqr.2.A	84.67	monomer		HHblits	X-ray	2.40Å	0.56	1.00	Hemoglobin subunit alpha	
3d1a.1.C	86.13	monomer		HHblits	X-ray	2.61Å	0.55	1.00	Hemoglobin subunit alpha-1/2	
3d1a.1.A 3cy5.1.C	86.13 86.13	monomer		HHblits HHblits	X-ray X-ray	2.61Å 2.00Å	0.55 0.56	1.00	Hemoglobin subunit alpha-1/2 Hemoglobin subunit alpha-2	
3cy5.1.A	86.13	monomer		HHblits	X-ray	2.00Å	0.56	1.00	Hemoglobin subunit alpha-2	
3gys.1.C	84.67	monomer		HHblits	X-ray	2.90Å	0.56	1.00	Hemoglobin subunit alpha	
4h2l.1.A	84.67	monomer		HHblits	X-ray	1.78Å	0.56	1.00	Alpha-globin	
5eui.1.A	89.78	monomer		HHblits	X-ray	1.45Å	0.57	1.00	HBA protein	
5ker.2.A	84.67	homo- dimer	0.26	HHblits	X-ray	2.20Å	0.55	1.00	Alpha-globin	
5ker.1.C	84.67	homo- dimer	0.35	HHblits	X-ray	2.20Å	0.55	1.00	Alpha-globin	
5ker.1.A	84.67	homo- dimer	0.35	HHblits	X-ray	2.20Å	0.55	1.00	Alpha-globin	
2ri4.1.C	86.13	homo- dimer	0.33	HHblits	X-ray	2.70Å	0.55	1.00	Hemoglobin subunit alpha-1/2	
2ri4.3.A	86.13	monomer		HHblits	X-ray	2.70Å	0.55	1.00	Hemoglobin subunit alpha-1/2	
		homo-				_				
1y8i.1.C	86.86	dimer	0.52	HHblits	X-ray	2.60Å	0.56	1.00	Hemoglobin alpha chains	
1g0b.1.A	86.86	monomer		HHblits	X-ray	1.90Å	0.56	1.00	HEMOGLOBIN ALPHA CHAIN	
1y8i.1.A	86.86	homo- dimer	0.52	HHblits	X-ray	2.60Å	0.56	1.00	Hemoglobin alpha chains	
2zlx.2.A	86.86	monomer		HHblits	X-ray	2.80Å	0.56	1.00	Hemoglobin subunit alpha	
2zlw.1.A	86.86	homo- dimer	0.61	HHblits	X-ray	2.90Å	0.56	1.00	Hemoglobin subunit alpha	
3gdj.1.A	83.94	monomer		HHblits	X-ray	2.00Å	0.55	1.00	Hemoglobin subunit alpha	
2rao.1.A	81.75	homo- dimer	0.38	HHblits	X-ray	2.00Å	0.56	1.00	Hemoglobin subunit alpha-1/2	
2qu0.1.C	86.13	homo- dimer	0.55	HHblits	X-ray	2.70Å	0.55	1.00	Hemoglobin subunit alpha-1/2	
3eu1.1.A	86.13	monomer		HHblits	X-ray	3.00Å	0.55	1.00	Hemoglobin subunit alpha-1/2	
3eu1.1.C	86.13	monomer		HHblits	X-ray	3.00Å	0.55	1.00	Hemoglobin subunit alpha-1/2	
2qu0.1.A	86.13	homo- dimer	0.55	HHblits	X-ray	2.70Å	0.55	1.00	Hemoglobin subunit alpha-1/2	
3gou.1.A	83.21	homo- dimer	0.52	HHblits	X-ray	3.00Å	0.55	1.00	Hemoglobin subunit alpha	
3pel.1.A	83.21	homo- dimer	0.59	HHblits	X-ray	1.90Å	0.55	1.00	Hemoglobin subunit alpha	
3lqd.1.A	82.48	homo- dimer	0.58	HHblits	X-ray	2.80Å	0.56	1.00	Hemoglobin subunit alpha	

5m3l.1.A, 1h1x.1.A, 3o7n.1.A, 3s48.2.B, 5ojb.1.A, 1aby.1.A, 1aby.1.B, 3uhc.1.A, 1y0c.1.A, 1umo.1.A, 5ohe.4.A, 1hrm.1.A, 1mlj.1.A, 3mju.1.B, 1mni.1.A, 3pt8.1.B, 3mju.1.A, 4v93.124.A, 1x3k.1.A, 1abs.1.A, 1s0h.1.A, 1s56.1.A, 3nl7.1.B, 2qu0.1.D, 1lue.1.A, 5h3c.1.B, 1shr.1.D, 1shr.1.C, 1o1n.1.A, 1jra.1.A, 1v4w.1.D, 5hy8.2.B, 1dxv.1.B, 4xs0.1.A, 4xs0.1.B, 1or4.1.A, 4bja.1.A, 3ok5.1.B, 1yzi.1.B, 4v93.118.A, 3qzx.1.B, 3qzz.1.A, 1y4v.1.B, 1nej.1.B, 1m9p.1.A, 1m9p.1.B, 4mqk.1.A, 4mqk.1.B, 4mqk.1.C, 5v5r.1.A, 1jr8.1.A, 3bcq.1.B, 3bcq.1.C, 2gl3.1.A, 3bcq.1.A, 3qqr.1.A, 3bcq.1.D, 1bij.1.A, 1bij.1.B, 3zjh.1.A, 1ibe.1.A, 5jgg.1.B, 2nx0.1.A, 1oj6.2.A, 5e6e.1.B, 3zom.1.A, 4b4y.1.A, 1o1j.1.A, 1o1j.1.B, 5jgg.1.A, 3wtg.1.A, 1hdb.1.B, 3wtg.1.C, 2gnv.1.A, 2gnv.1.B, 4g1b.1.A, 4tyx.1.A, 1nz3.1.A, 4fwz.1.A, 2wy4.1.A, 1iop.1.A, 4f6d.1.A, 1cmy.1.B, 1cmy.1.A, 2v8a.1.A, 5yl3.1.A, 5gai.1.K, 3uh3.1.B, 1wmu.1.B, 3ugz.1.A, 3zhw.2.A, 3ubc.1.A, 1umo.1.B, 3eu1.1.D, 3eu1.1.B, 2vyy.1.A, 5o41.1.A, 4jac.1.A, 1mtk.1.A, 1or6.1.B, 5eet.1.A, 4mqj.2.D, 1y7g.1.B, 2nrl.1.A, 1ngk.1.B, 5eys.1.A, 2zlv.1.C, 2zlv.1.B, 1cg5.1.A, 5urc.1.D, 4mqc.1.A, 1hv4.1.D, 1v5h.1.A, 4f4o.1.A, 4f4o.1.B, 1yhu.1.B, 1myj.2.A, 1yhu.1.A, 2grz.1.B, 3wct.1.C, 1oj6.4.A, 1mnj.1.A, 3fs4.1.B, 1yih.1.B, 3vrg.1.A, 1it2.1.A, 1gbv.1.B, 1hbr.1.B, 1hbr.1.A, 1y7d.1.B, 1hbs.1.D, 1hbs.1.C, 1hbs.1.B, 1hbs.1.A, 1ch9.1.A, 3wfx.1.A, 1s56.2.A, 5ee4.1.C, 1ngk.1.H, 4u8u.1.D, 5x2s.2.C, 3vre.1.A, 3vre.1.B, 4mu5.1.A, 1y45.1.B, 1v4x.1.A, 1v4x.1.B, 5x2t.1.D, 5x2t.1.A, 5x2t.1.C, 1mlh.1.A, 4b3w.1.A, 5ile.1.A, 4zvb.1.A, 1y5j.1.B, 3ng6.1.B, 1y5k.1.B, 2ri4.1.D, 1cbl.1.D, 3gkv.1.A, 3gkv.1.B, 1v4u.1.B, 3s1i.1.A, 4o2g.1.A, 3s1i.1.B, 1c40.1.A, 5yzf.1.A, 1c40.1.B, 3w4u.1.B, 5tjf.1.A, 3w4u.1.A, 1y4g.1.B, 1moh.1.A, 5y6p.7.A, 3vhb.1.A, 3uh6.1.A, 2pgh.1.C, 1vxa.1.A, 2hbs.1.D, 2d6c.1.A, 3ciu.1.D, 1ewa.1.A, 1cpw.1.A, 3l39.1.A, 1y4q.1.B, 3wyo.1.B, 3wyo.1.A, 1yma.1.A, 3aq8.1.A, 2dc3.1.A, 2grf.1.A, 1lh1.1.A, 2dc3.1.B, 3hep.1.A, 2ohb.1.A, 4of9.1.A, 1urv.2.A, 2d6c.2.A, 1irc.1.A, 1y01.1.B, 1ch1.1.A, 4ni0.1.A, 1ufj.1.A, 3zjl.1.A, 1ird.1.B, 3gou.1.B, 1vwt.1.A, 1y8i.1.D, 1y8i.1.B, 3d1k.1.A, 5f0b.1.A, 3uhh.1.A, 4qau.1.A, 3ugy.1.A, 4hrt.2.C, 1xz5.1.A, 1v07.1.A, 3aq5.1.A, 1cqx.1.A, 5k1l.1.B, 5eui.1.B, 1hds.1.A, 3tld.1.A, 1hds.1.C, 1hds.1.B, 1hds.1.D, 5hu6.1.B, 4iro.1.A, 1hbg.1.A, 4iro.1.C, 5hbi.1.B, 1myi.1.A, 1tu9.1.A, 2ig3.1.A, 3a0g.1.B, 1w92.1.A, 1dxd.1.A, 1buw.1.A, 1buw.1.B, 1buw.1.D, 2evp.1.A, 3qm7.1.A, 5ker.1.D, 1a00.1.B, 5ker.1.B, 4twv.1.A, 2ef2.1.A, 1hac.1.B, 1hac.1.C, 2r50.2.A, 4hsw.2.A, 2r50.2.B, 1vre.1.A, 3gqr.2.B, 2zs0.1.A, 2zs0.1.B, 2zs0.1.C, 3sdn.1.A, 1fsx.1.D, 1f6h.1.A, 2cmm.1.A, 3o89.1.A, 1mlm.1.A, 4xdi.1.A, 3zh0.1.A, 3wcu.1.C, 4esa.1.D, 3mjp.1.D, 3mjp.1.C, 5vmm.1.C, 3mjp.1.A, 2gkm.1.A, 7hbi.1.B, 2vjt.1.A, 5sw7.1.B, 3bj2.1.D, 5x2r.3.A, 5x2r.3.B, 5x2r.3.C, 5x2r.3.D, 3rgk.1.A, 3dht.1.A, 4o1t.1.A, 5x2r.1.C, 5x2r.1.A, 5x2r.1.D, 3gdj.1.B, 4rol.1.D, 1rse.1.A, 3g53.1.A, 5oj9.1.A, 1k1k.1.B, 1ufp.1.A, 1b33.1.A, 1d8u.1.B, 1ash.1.A, 3bj1.1.C, 3qjd.1.A, 1co9.1.A, 3ozu.1.A, 3pi4.1.B, 2eku.1.A, 1ycb.1.A, 1vxb.1.A, 1ch3.1.A, 5x2s.2.D, 3s65.1.B, 1fhj.1.A, 3s65.1.D, 5x2s.2.A, 1y09.1.A, 3szk.1.B, 3szk.1.A, 5ab8.1.A, 1mcy.1.A, 3v2v.1.A, 2mgf.1.A, 3gqg.1.B, 3gqg.1.A, 3myn.1.A, 2vyz.1.A, 2pgh.1.B, 3onz.1.B, 3fs4.1.A, 2pgh.1.A, 3uhd.1.A, 2pgh.1.D, 5m3l.1.L, 3heo.1.A, 5d1v.1.A, 1pmb.2.A, 4v93.42.A, 1abw.1.B, 1dm1.1.A, 1abw.1.A, 4g51.1.C, 1y7z.1.B, 1jl6.1.A, 5kdq.1.D, 2aup.1.A, 3mym.1.A, 3zhw.1.A, 1qpw.1.D, 1x46.1.A, 1myg.1.A, 3hyu.1.B, 3hyu.1.C, 4ns2.1.A, 3kuo.1.A, 1b33.2.E, 3h58.1.A, 1o16.1.A, 5ojc.1.A, 1jzk.1.A, 5utd.1.A, 4h07.1.A, 4yu3.1.B, 1rvw.1.A, 1rvw.1.B, 3uhr.1.A, 5ksj.1.B, 3whm.1.D, 4hrr.2.D, 1bvd.1.A, 1mob.1.A, 5v5q.1.A, 1hbs.2.B, 3qjc.1.B, 2r4w.1.A, 2oif.1.A, 1gli.1.A, 1ye1.1.B, 4f69.1.A, 1bbb.1.C, 5ohf.1.B, 1xch.1.A, 3gou.1.D, 2ig3.2.A, 4pqb.1.A, 3sdh.1.A, 5eu2.1.A, 3at6.1.A, 1ux9.2.A, 6hbw.1.D, 6hbw.1.B, 1fn3.1.D, 1dxu.1.B, 1dxu.1.A, 3kun.2.A, 3ozi.1.A, 4it8.1.A, 3k3u.2.B, 1xzv.1.A, 1z8u.2.B, 3oj1.1.A, 4uf6.1.C, 1y8w.1.A, 2aa1.1.A, 2aa1.1.B, 1ch7.1.A, 2w6w.1.A, 3at5.1.A, 3at5.1.B, 3bj2.1.A, 5d5r.1.A, 1y8w.1.C, 1a3o.1.A, 1pbx.1.A, 1pbx.1.B, 4nzi.1.A, 2gln.1.B, 1c7d.1.A, 2gln.1.A, 1gcv.1.B, 1gcv.1.A, 1or6.1.A, 1ch5.1.A, 2w6y.1.A, 1gcw.1.D, 1gcw.1.C, 1gcw.1.B, 1gcw.1.A, 2h8d.1.D, 6bmg.1.A, 1mgn.1.A, 1o1m.1.A, 3wtg.1.B, 1dlw.1.A, 3gkt.1.A, 5ni1.1.B, 1it3.1.A, 3uhb.1.A, 2bkm.1.A, 5h3c.1.A, 1y83.1.B, 1fdh.1.B, 2wth.1.A, 1r1y.1.B, 4zva.1.A, 1r1y.1.A, 5x2s.3.C, 4n7o.1.A, 3dhr.4.B, 3dut.1.D, 2zfo.1.A, 2zfo.1.B, 2zfo.1.C, 1o1k.1.B, 1jzl.1.B, 2qsp.1.D, 3dhr.2.C, 5x2t.2.A, 5x2t.2.D, 2zlw.1.B, 1it2.2.A, 3mn0.1.A, 2vee.1.A, 2mgc.1.A, 1yg5.1.B, 1cik.1.A, 1ebt.1.A, 4mxl.1.A, 1gvh.1.A, 5oja.1.A, 2hhe.1.B, 1f5o.1.A, 1ofk.1.A, 2vml.1.A, 4v93.65.A, 1spg.1.B, 1spg.1.A, 1o1o.1.A, 1o1o.1.B, 4uiq.2.A, 2oh8.1.A, 1y5f.1.B, 1y4r.1.B, 3a2g.1.A, 1r1x.1.B, 5vlx.1.A, 2g3h.1.A, 1r1x.1.A, 5hlq.1.A, 1hbs.2.A, 1yeo.1.A, 1hbs.2.C, 1hbs.2.D, 5o18.1.A, 4wjg.2.G, 2e2y.1.A, 2d5x.1.B, 1mba.1.A, 5x2u.3.B, 2oha.1.A, 4u8u.1.C, 4u8u.1.B, 3zji.1.A, 3s48.1.B, 1urv.1.A, 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4hrt.1.B, 2gtl.1.A, 2gtl.1.B, 1do3.1.A, 3uhi.1.A, 1xq5.1.A, 3gys.2.B, 5m3l.1.F, 5m3l.1.G, 5c6y.1.A, 3pt7.1.B, 2vyw.1.A, 2qfn.1.A, 2in4.1.A, 4g1v.1.A, 3hrw.1.B, 3hrw.1.D, 1hlm.1.A, 2qrw.1.B, 1a6g.1.A, 2qrw.1.A, 2r80.5.B, 2jho.1.A, 1ye2.1.B, 3uhe.1.A, 3wyo.2.B, 2rao.1.B, 1ofj.1.A, 1ngk.1.G, 1ngk.1.D, 3w4u.2.A, 1ngk.1.A, 3wct.1.B, 3ogb.1.A, 3wct.1.A, 3wct.1.D, 5ker.2.C, 4v93.102.A, 5ker.2.D, 1v4u.1.A, 1hho.1.B, 1jy7.1.B, 1y2z.1.B, 5ufj.1.D, 5ufj.1.B, 2ekt.1.A, 2qrw.4.A, 5vmm.1.B, 1aj9.1.A, 4ni0.1.B, 5vmm.1.A, 1uc3.1.A, 2vhb.1.B, 5vmm.1.D, 5kd1.1.A, 3ozv.3.A, 1yen.1.A, 2bwh.1.A, 4kn3.1.A, 4h2l.1.B, 5f2a.1.A, 1moc.1.A, 1m6c.1.A, 3vrf.1.B, 2w31.1.B, 3vrf.1.A, 4v93.3.A, 5x2r.2.D, 5x2r.2.A, 5x2r.2.C, 3mjp.1.B, 1ux9.1.A, 1mlr.1.A, 5gai.1.V, 2r1h.1.B, 2r1h.1.C, 3zjr.1.B, 1dwr.1.A, 1fsl.2.A, 1la6.1.B, 1mbs.1.A, 3qjb.1.A, 4hrr.1.B, 1kfr.1.A, 2c0k.1.A, 1i3d.1.A, 1yvq.1.D, 1hjt.1.A, 4uii.1.B, 1vhb.1.A, 4v93.1.A, 4uii.1.A, 1ye0.1.B, 1mlu.1.A, 5x2s.3.A, 2oh9.1.A, 6bb5.1.D, 5x2s.3.B,

3qje.1.B, 5x2s.3.D, 2hbs.2.B, 1gdl.1.A, 3ovu.1.C, 4o35.1.A, 5d1v.1.B, 5y6p.22.A, 2auq.1.A, 2r4x.1.A, 4u8u.1.A, 1jzm.1.A, 3k9z.1.A, 3at6.1.B, 1y8k.1.B, 1cbm.1.A, 1myf.1.A, 2zs0.1.D, 1a9w.1.B, 5b84.1.A, 1s5x.1.B, 2w6x.1.A, 3zh0.2.A, 2gtl.1.C, 1dxt.1.B, 1ch4.1.A, 1ch4.1.C, 3tm9.1.A, 3lqd.1.B, 4fh7.1.B, 2lhb.1.A, 3aq9.1.A, 3cy5.1.B, 3dut.2.B, 1bz1.1.B, 1bz1.1.A, 4pqc.1.A, 1sdl.1.D, 1moa.1.A,

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1g08.1.B, 1sdl.1.C, 2m6z.1.B, 3pel.1.B, 5x2t.3.D, 5x2t.3.C, 5x2t.3.A, 2yrs.1.B, 3lhb.1.A, 5llz.1.A, 1myt.1.A, 1xq5.1.B, 1o1l.1.A,

1y8h.1.B, 4f6i.1.A, 3ozv.3.B, 1ns6.1.A, 3bj3.1.A, 1si4.1.B, 3dhr.1.A, 3eok.1.B, 3eok.1.A, 2dhb.1.B

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4hsw.1.A, 1bje.1.A, 5m3l.1.E, 2qmb.1.D, 2eb9.1.A, 2qmb.1.A, 3zjj.1.A, 2bk9.1.A, 3gqp.1.D, 1qpw.1.B, 1qpw.1.C, 1a4f.1.A, 1a4f.1.B,

for modelling than the filtered list.