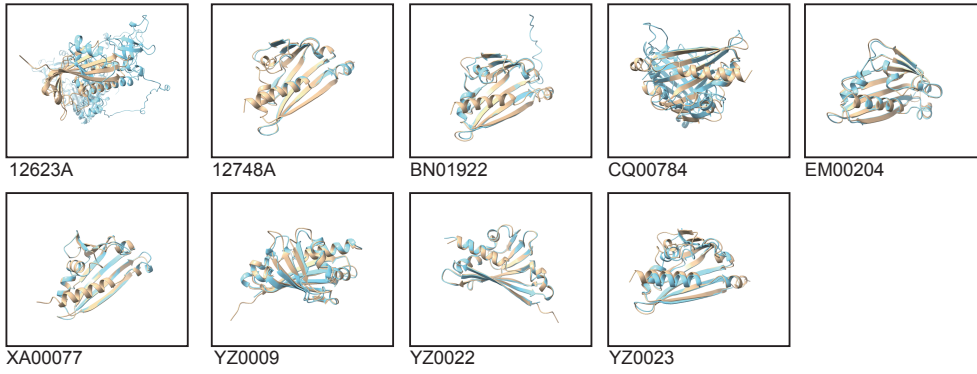
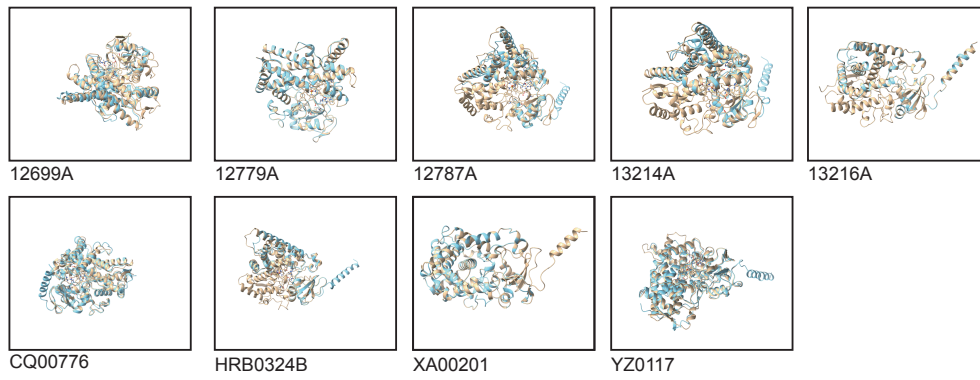


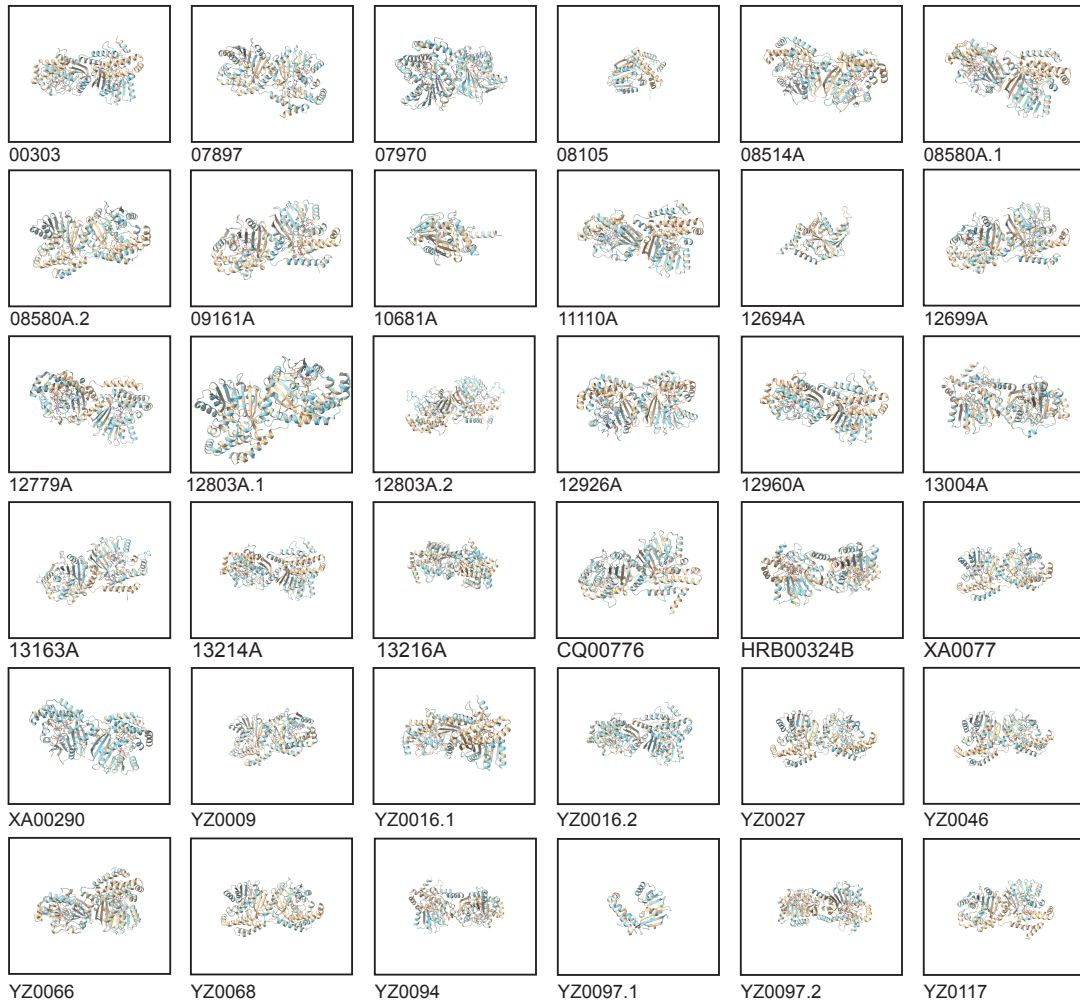
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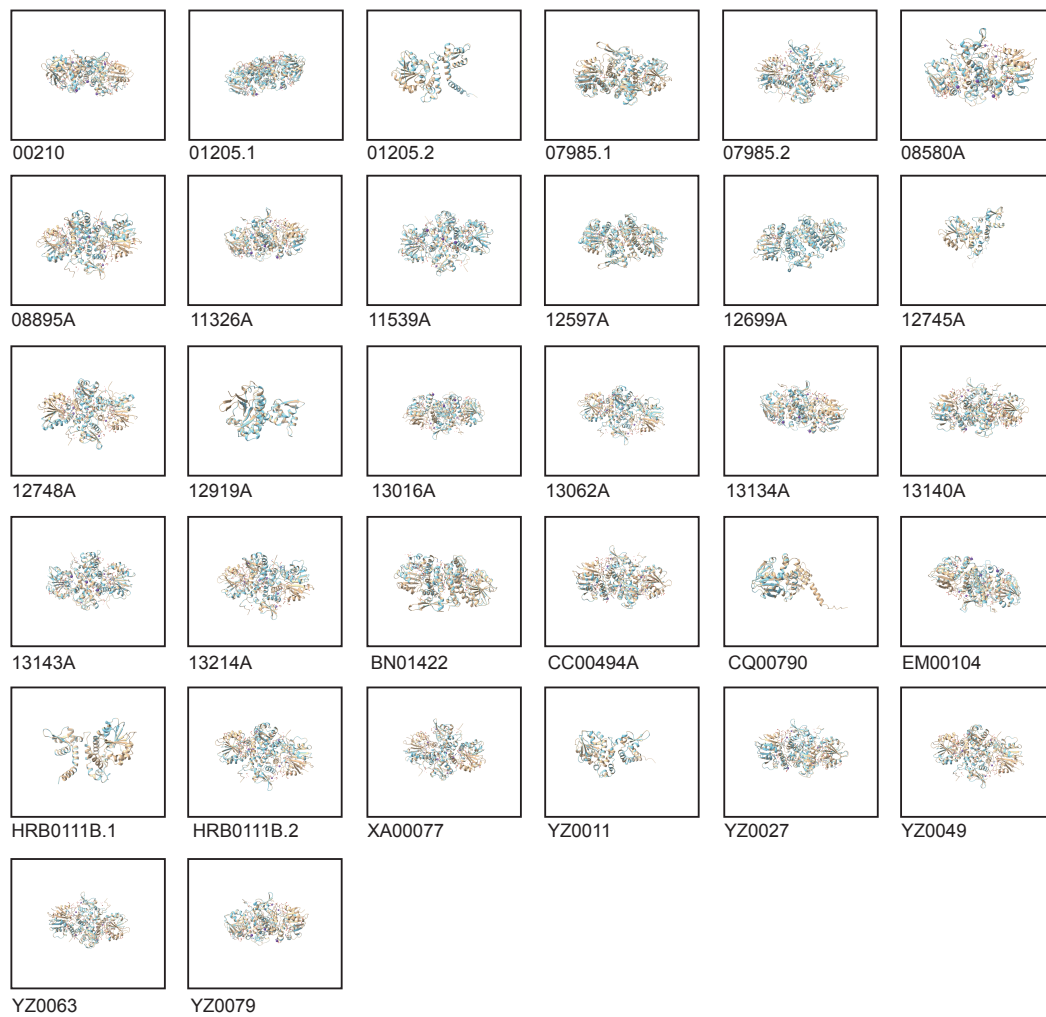
B



C



D



Supplementary Figure 9. Prediction of 3D structural models for positively selected sequences in key enzymes. The sequence code shown below each model corresponds to the entry in Supplementary Table S11. (A) Models for the nine positively selected NCS sequences (B) Models for the nine positively selected NMCH sequences. (C) Models for the Thirty-six positively selected CNMT sequences. (D) Models for the Thirty-two positively selected OMT sequences.