# Using AlphaFold predictions for structure determination

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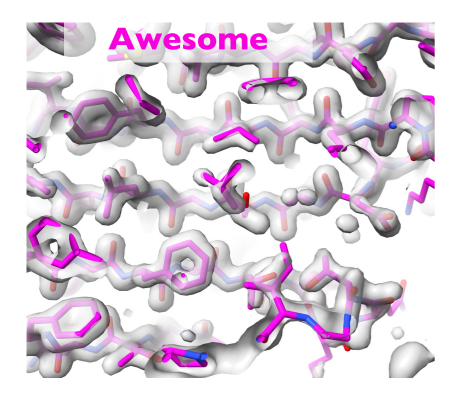


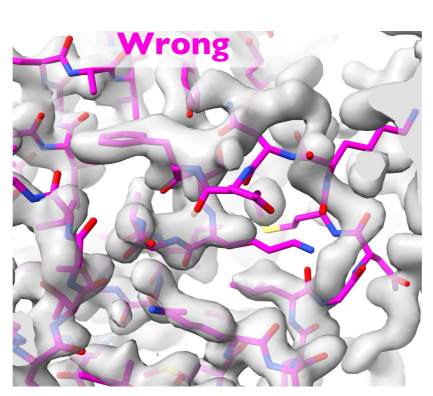


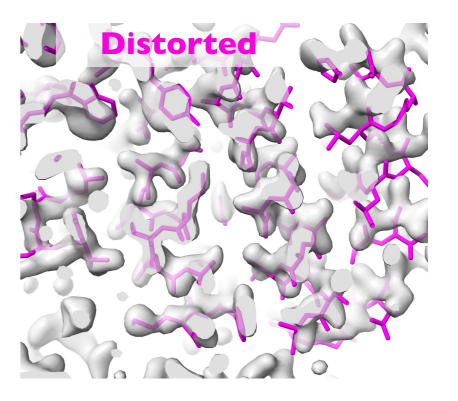


## AlphaFold predictions are great hypotheses

AlphaFold models can be....

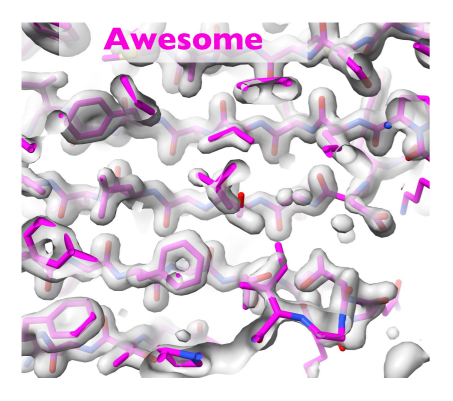




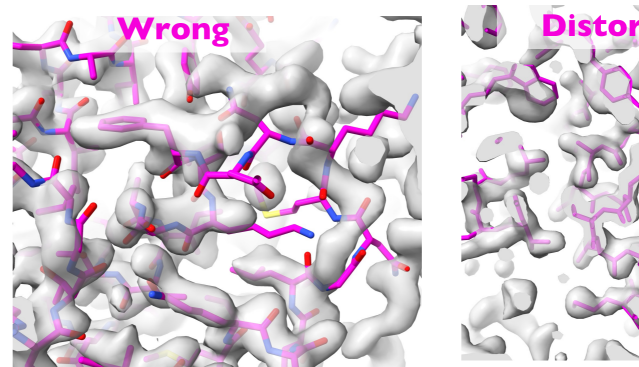


## AlphaFold predictions and confidence estimates

Residue-specific confidence (pLDDT) identifies where errors are more likely



AlphaFold confidence (pLDDT)	Median prediction error (Å)	Percentage with error over 2 Å
>90	0.6	10
80 - 90	1.1	22
70 - 80	1.5	33
<70	3.5	77



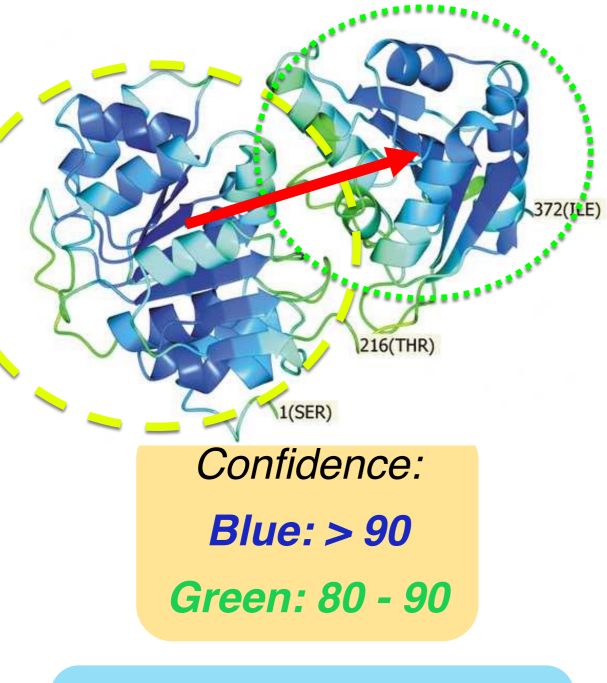
Distorted

Terwilliger et al. (2024), AlphaFold predictions are valuable hypotheses, and accelerate but do not replace experimental structure determination. Nature Methods 21, 110-116.

### AlphaFold confidence measure (pLDDT, Predicted difference distance test)

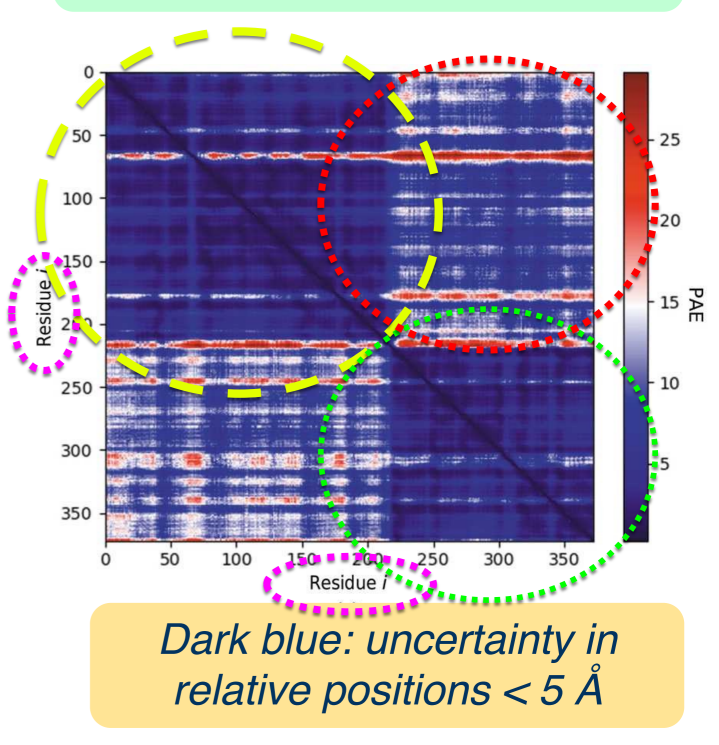
372(ILE)							
	1(SER)		AlphaFold	Median	Percentage with error		
	Confidence:		confidence (pLDDT)	prediction error (Å)	over 2 Å		
	<i>Blue: &gt; 90</i>		>90	0.6	10		
	Green: 80 - 90	-	80 - 90	(1.1	22		
AlphaFold prediction RNA helicase (PDB entry 6i5i)		for	70 - 80	1.5	33		
			<70	3.5	77		
				Oeffner et al. (2022). Acta	a Cryst. D78, 1303-1314		

## PAE matrix (Predicted aligned error)



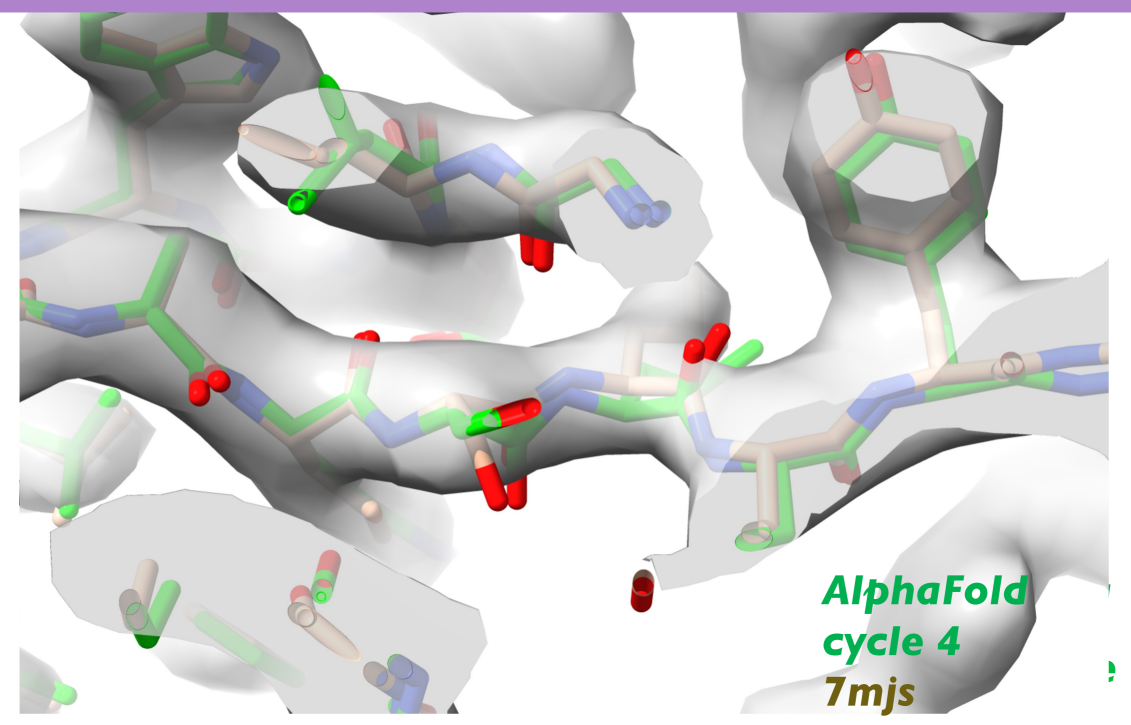
AlphaFold prediction for RNA helicase (PDB entry 6i5i)

## PAE matrix identifies accurately-predicted domains



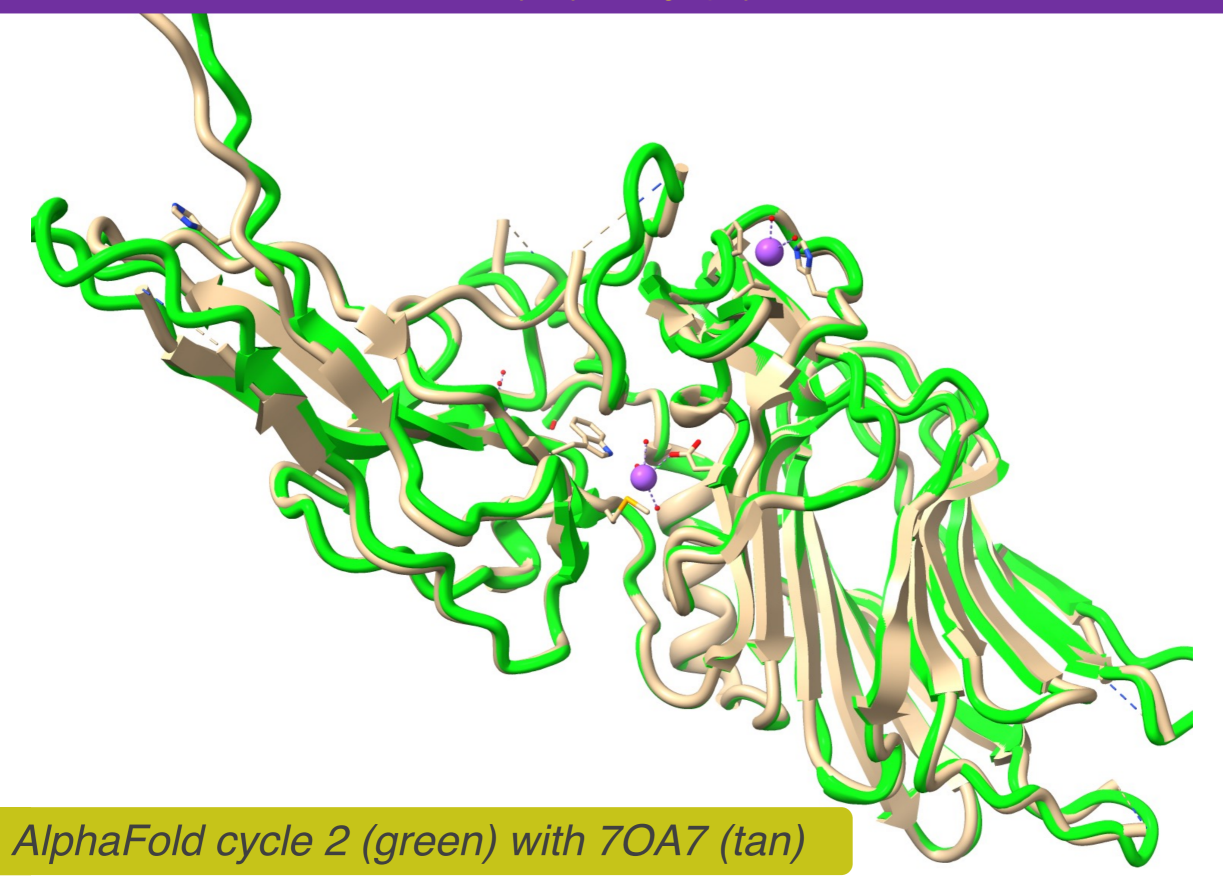
Why? Because your new prediction might be better than your model ... and better than your original AlphaFold prediction AlphaFold prediction New prediction Working model

#### Improving AlphaFold prediction using partial models as templates (Cryo-EM)

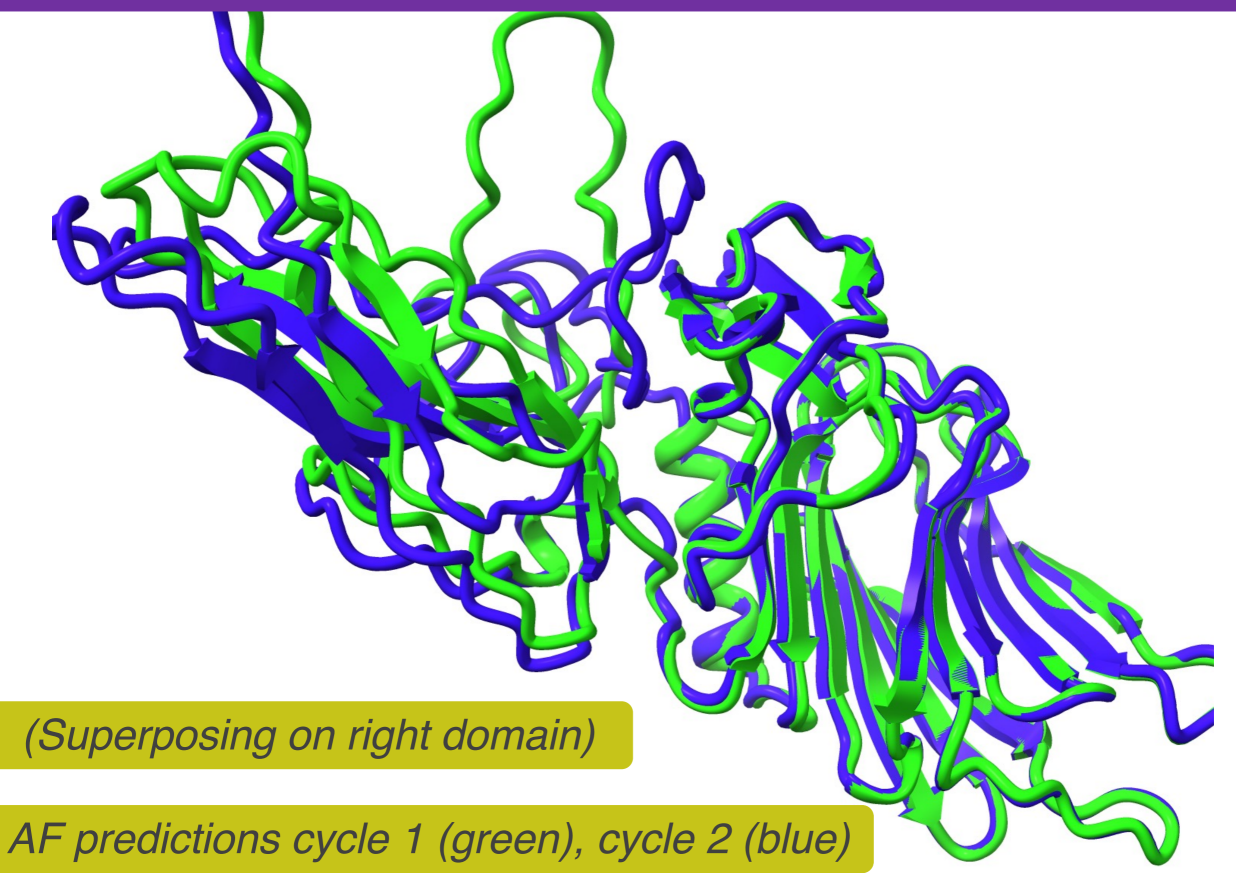


Data from 7mjs, Cater, R.J., et al. (2021). Nature 595, 315–319

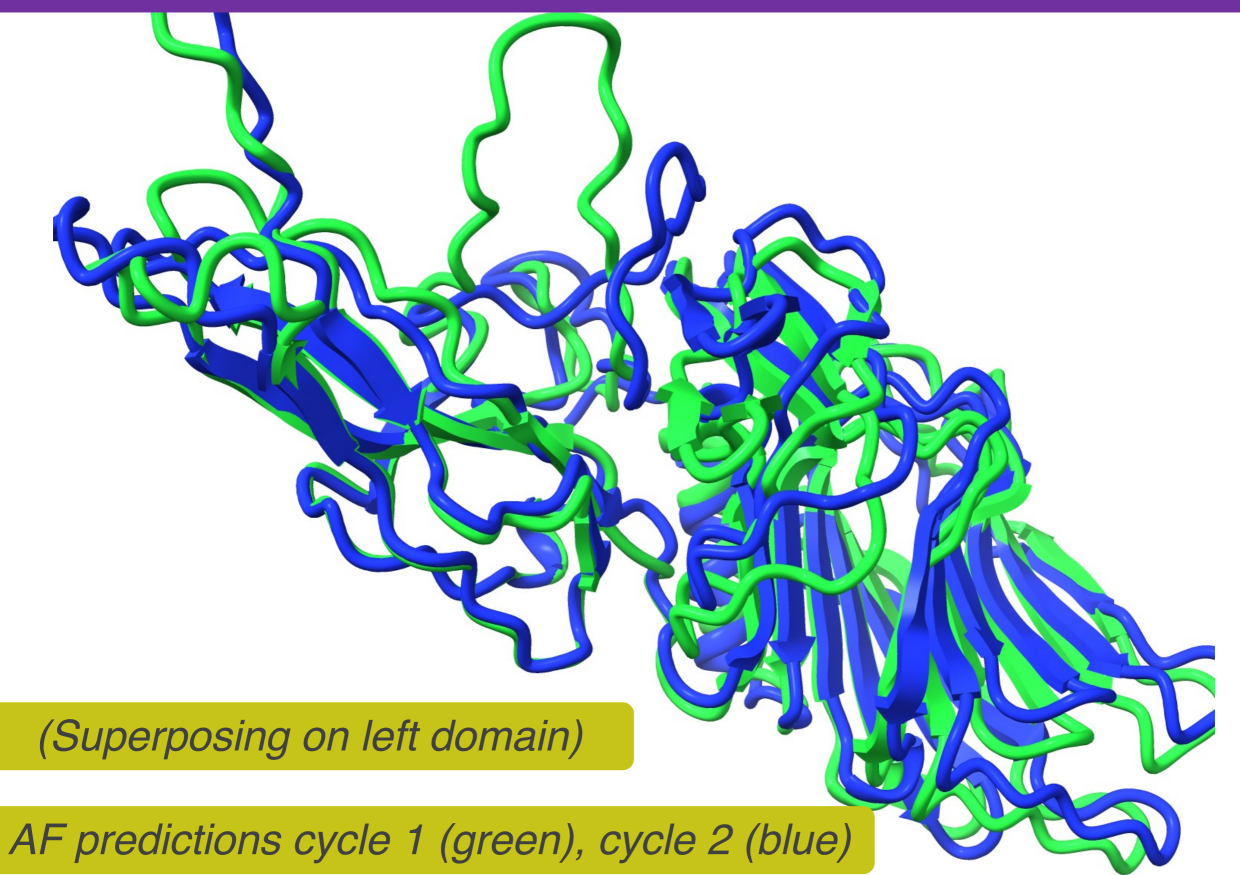
#### Improving AlphaFold prediction using partial models as templates (X-ray crystallography)



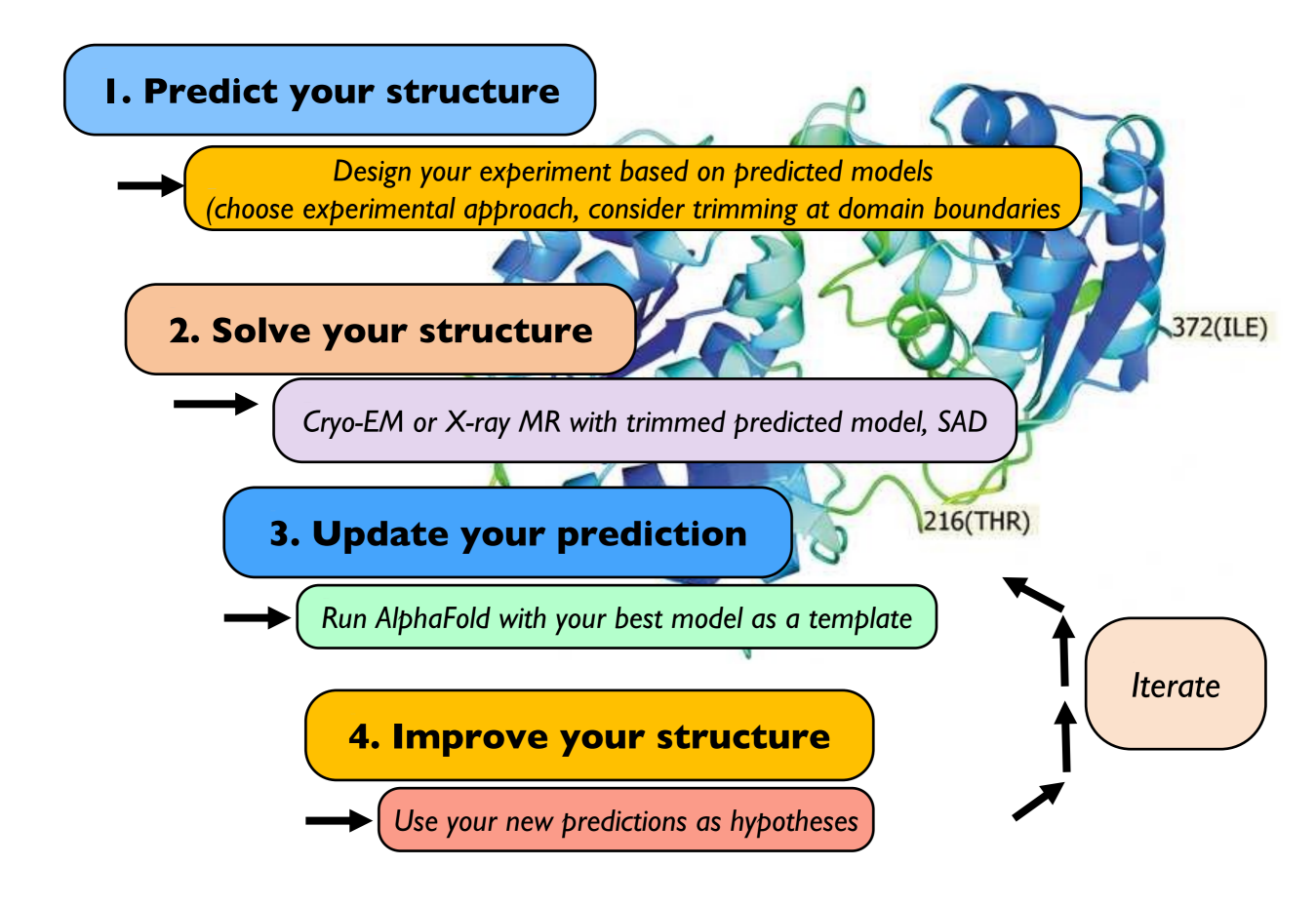
Improving AlphaFold prediction using partial models as templates (X-ray crystallography)



Improving AlphaFold prediction using partial models as templates (X-ray crystallography)



#### Strategy for structure determination in the AlphaFold era



#### Phenix AlphaFold prediction server

Available from the Phenix GUI

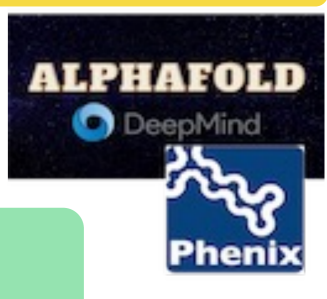
Predicts structures of protein chains (one at a time)

Can use a template to guide the prediction

You do not need an MSA (multiple sequence alignment) if you supply a template

The template should not be an AlphaFold model

Many thanks for AlphaFold, ColabFold scripts, and the MMseqs2 server for MSAs



Process predicted model

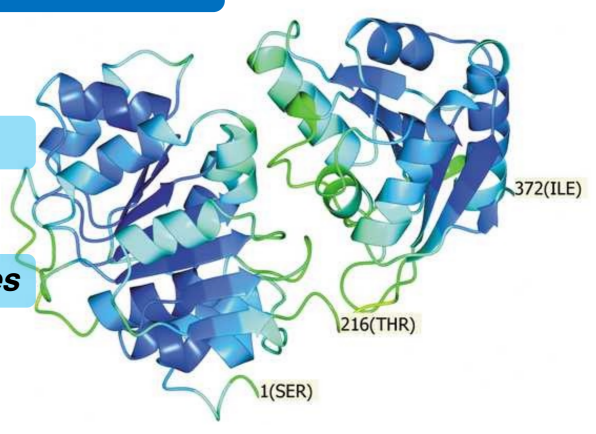
Convert pLDDT to B-value

Trim low-confidence parts of model

Identify high-confidence domains

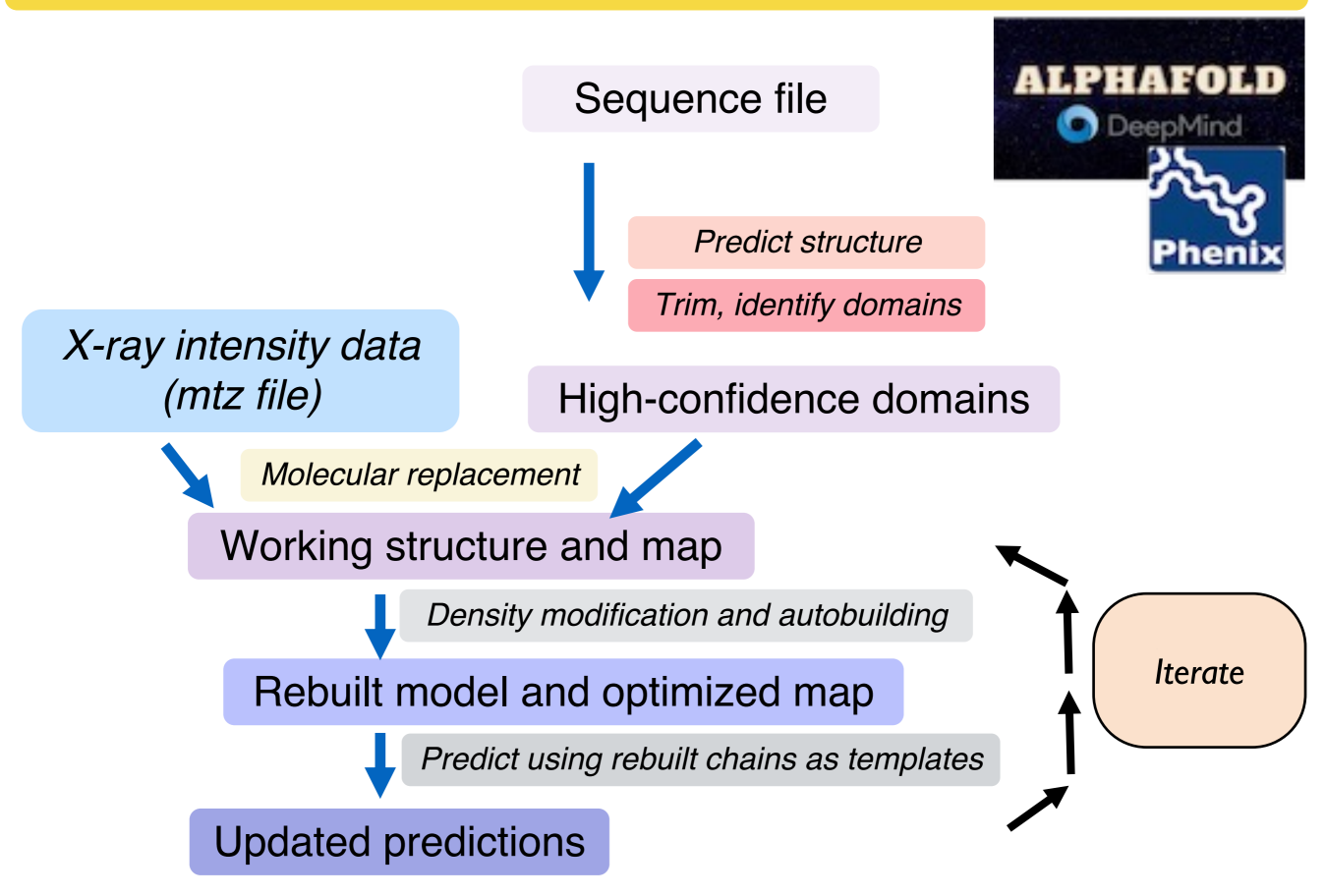
Compact high-confidence regions

Groupings of residues with low PAE values



DeepMind

#### X-ray structure determination with AlphaFold



#### Input and output from structure determination with AlphaFold

Experimental data (maps or X-ray data)

Contents of asymmetric unit (sequence file)

Rebuilt model Optimized map

Map and model ready for next steps Docked predicted models

Useful as high-quality reference models



Output

Input

Phenix tools for structure determination with AlphaFold

**PredictModel** (Predict with AlphaFold)

**ProcessPredictedModel** (Trim and identify domains)

ResolveCryoEM, LocalAnisoSharpen (map improvement)

EMPlacement, DockInMap (Docking of single, multiple chains)

DockAndRebuild (Morphing and rebuilding)

RealSpaceRefine (Refinement)

Phaser-MR (Molecular replacement)

AutoBuild (Density modification and rebuilding)

Phenix.refine (Refinement)

PredictAndBuild (Prediction and structure determination)

X-ray

automation

AlphaFold

models

Cryo-EM

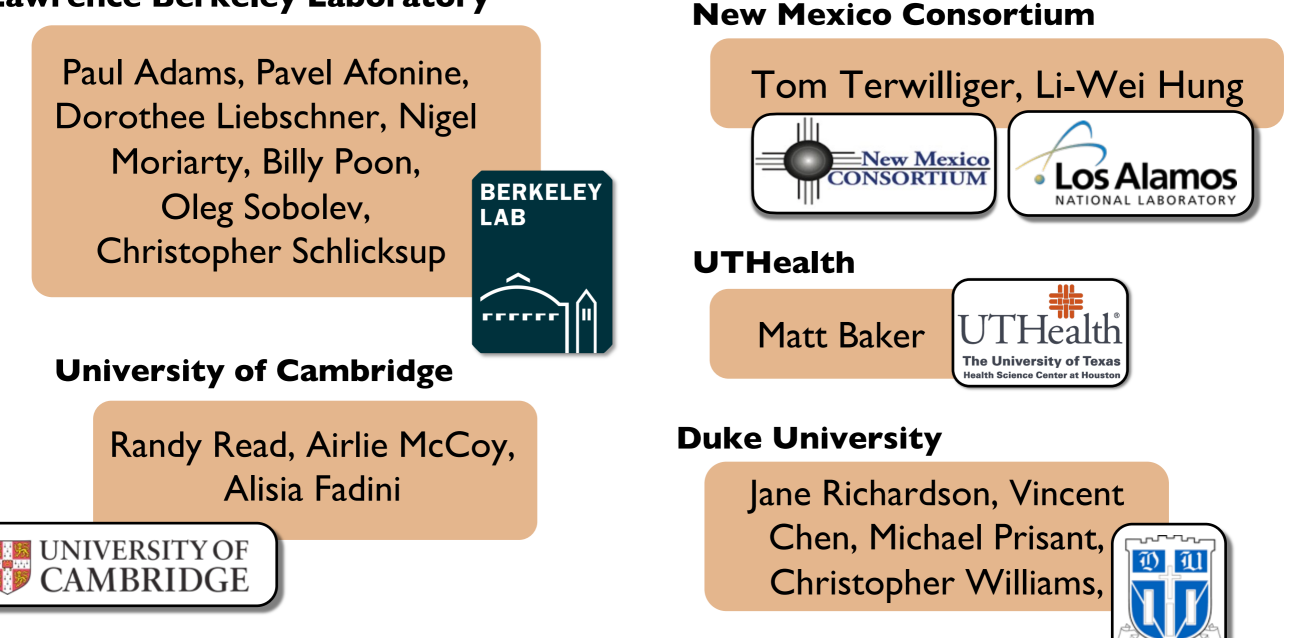






Los Alamos National Laboratory

#### Lawrence Berkeley Laboratory





An NIH/NIGMS funded Liebschner D, et al., Macromolecular structure determination using X-rays, neutrons and electrons: recent developments in Phenix. Acta Cryst. 2019 **D75**:861–877