1) Gathering information	
Prior models	h3 comparative model, template 1TZY:G
	h4 comparative model, template 1TZY:H
Physical principles and statistical preferences	Excluded volume
	Sequence connectivity
Experimental data	170 pE-MAP derived distance restraints
2) Representing the system	10 1
Composition (number of copies)	h3: 1
Atomic (atmesterned) common ente	h4: 1 h3: 39-136
Atomic (structured) components	h4: 20-103
Un atmentioned accompany and	None
Unstructured components	
Resolution of structured components	1 [R1] residue per bead
Resolution of unstructured components	None
Structural coverage	100.0 %
Rigid body (RB) definitions	RB1: h3 ₃₉₋₁₃₆
Contint motorints and latints are in found in	RB2: h4 ₂₀₋₁₀₃
Spatial restraints encoded into scoring function	Excluded volume; applied to the R1 representation Sequence connectivity; applied to the R1 representation
	1 7, 11
	pE-MAP MIC pair-restraints; applied to the R1 representation
3) Structural Sampling	
Sampling method	Replica Exchange Gibbs sampling, based on Metropolis Monte
- •	Carlo
Replica exchange temperature range	1.0 - 2.5
Number of replicas	4
Number of runs	60
Number of structures generated	2500000
Movers for rigid bodies	Random translation up to 0.3 Å
	Random rotation up to 0.1 radians
Movers for flexible string of bead	Random translation up to 4.0 Å
CPU time	4 hours on 20 processors
	•
4) Validating the model	
Models selected for validation	0,0000
Number of models after equilibration	2500000
Number of models that satisfy the input information	1162507
Number of structures in samples A/B	639091/523416
p-value of non-parametric Kolmogorov-Smirnov two-sample	0.006 (threshold p-value > 0.05)
test	0.00
Kolmogorov-Smirnov two-sample test statistic, D	0.96
Thoroughness of the structural sampling	
Sampling precision	2.03 Å
Homogeneity of proportions χ^2 test (p-value)/Cramers V value	0.679/0.006 (thresholds: p-value>0.05 OR Cramer's V<0.1)
Number of clusters	1 1 00 6 07
Cluster populations	cluster 1 : 98.6 %
Cluster precisions	cluster 1 : 1.04 Å
Average cross-correlation between localization probability den-	cluster 1: 1.0
sities of samples A and B	
Validation by information used for modeling	99 %
Percent of sequence connectivity restraints satisfied per struc-	33 /0
Persont nF MAP restraints satisfied non structure	97 07
Percent pE-MAP restraints satisfied per structure Percent of confuded violume restraints satisfied per structure	87 % 99 %
Percent of excluded volume restraints satisfied per structure	99 /0
5) Benchmark	
Structural accuracy (95 % CI)	3.8 (2.6-5.1) Å
6) Software and data availability	
Software	1
Software Modeling programs	IMP PMI module version devolor 20e22e602
Modeling programs	IMP PMI module, version develop-39c22a602 Integrative Modeling Platform (IMP), version develop-

Modeling scripts	https://github.com/salilab/pemap
Homology detection and structure prediction	HHPred, version 2.0.16
Visualization and plotting	UCSF Chimera, version 1.10
	Matplotlib, version 3.0.3
Data	
PDB-dev accesion code	TBD
pE-MAP data deposition	https://github.com/salilab/pemap

MODELLER, version 9.21