Implications of AlphaFold2 for crystallographic phasing by molecular replacement

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The AlphaFold2 results in the 14th edition of Critical Assessment of Structure Prediction (CASP14) showed that accurate (low root-mean-square deviation) in silico models of protein structure domains are on the horizon, whether or not the protein is related to known structures through high-coverage sequence similarity. As highly accurate models become available, generated by harnessing the power of correlated mutations and deep learning, one of the aspects of structural biology to be impacted will be methods of phasing in crystallography. Here, the data from CASP14 are used to explore the prospects for changes in phasing methods, and in particular to explore the prospects for molecular-replacement phasing using in silico models.

1. Introduction

The quality of a model for phasing a crystal structure by molecular replacement, for a given diffraction resolution limit, depends on the root-mean-square deviation (r.m.s.d.) of the model to the target structure and the fraction of the total scattering ($f_m$) that it represents. Broadly, as the resolution of the experimental data decreases, the $f_m$ must increase (Fig. 1). Specific predictions based on r.m.s.d., $f_m$ and data resolution guide molecular-replacement phasing strategies in individual cases (Oeffner et al., 2018).

When the data resolution extends to better than ~1 Å, the model can be as tiny as a single atom. A single atom can be considered to be the perfect in silico substructure model, with an r.m.s.d. of zero to the target structure, although overall the structure factors calculated from this model have large errors because of the extremely low $f_m$. Following single-atom molecular replacement, log-likelihood gradient completion can rapidly locate the remaining ordered atoms and for these resolutions the phase problem is considered to be solved (McCoy et al., 2017).

Small secondary-structure elements of helix or $\beta$-sheet are viable models when the data extend to better than ~2.2 Å resolution, and in some cases this technique extends to resolutions of 2.5 Å. The atomic coordinates of the model can be extracted from known structures or generated in silico (Glykos & Kokkinidis, 2001). Success does not necessarily require a homologous model with a sequence identity of over 30%, despite this being a commonly quoted metric for the success of molecular replacement (Scapin, 2013). Density modification and model building, which are significantly more powerful at higher resolutions, are central to structure completion with this method. Accurate fragments are regularly used for molecular replacement in software such as ARCIMBOLDO (Sammito et al., 2014, 2015; Rodriguez et al.,...
2. CASP14

CASP14 has established a leap in protein structure prediction. The primary CASP metric for ranking models and modelling groups is the GDT_TS (Global Distance Test Total Score), a structure-similarity measure designed and developed for the structure-alignment program LGA (Local–Global Alignment) as an alternative to the r.m.s.d. (Zemla, 2003). The GDT measures the percentage of $C^\alpha$ atoms that are found within certain distance cutoffs of one another between the model and target (either dependent upon or independent of a sequence alignment): the cutoff distance(s) must be defined for each reported GDT value. The GDT_TS is the average of four cutoff distances (1, 2, 4 and 8 Å). Higher GDT values are achieved with better models, in contrast to the r.m.s.d., where lower values are better. In circumstances where a score that is sensitive to smaller differences is needed, the GTD_HA is used; this is a variant of the GDT_TS in which the cutoff distances are 0.5, 1, 2 and 4 Å.

Harnessing the power of correlated mutations, contact predictions and deep learning, the AlphaFold2 group (group 427 in the CASP14 numbering) from the commercial organization DeepMind (Service, 2018; Callaway, 2020) were ranked first on Z-score for GDT_TS, reaching values over twice, and up to three times, those of the second and subsequent ranked groups, depending
on the classification of targets considered. This was achieved on a background of major improvements from other groups, including the Baker group (BAKER group 473, ranked second, and BAKER-experimental group 403, ranked third; Hiranuma et al., 2021), who used similar methods in academic settings.

For the first time, structures submitted to CASP as targets were phased with the help of models provided during the assessment (Kryshtafovych et al., 2021). For target T1058, the structure was solved by MR-SAD with the AlphaFold2 models after attempting molecular replacement with homologous structures, domains thereof and server models. For T1089, the AlphaFold2 models gave a far higher molecular-replacement signal than using trimmed ensemble models. For T1100, several of the models submitted to CASP, including the AlphaFold2 models, gave a molecular-replacement solution where the NMR structures of individual domains failed. The AlphaFold2 model for T1064 has also been used to solve the SARS-CoV-2 ORF8 structure retrospectively by molecular replacement (Flower & Hurley, 2021).

3. AlphaFold and RoseTTAFold

While this paper was being reviewed and revised, two major developments occurred that have made models of the quality of those submitted to CASP14 readily available to all. Firstly, the AlphaFold algorithms have been published (Jumper et al., 2021), together with release of the source code, enabling the prediction of any protein structure of interest. In addition, predicted structures have been made available in a database hosted at the European Bioinformatics Institute for all proteins from a substantial number of proteomes; further releases are promised (Tunyasuvunakool et al., 2021). Secondly, RoseTTAFold has been developed and published by the Baker group (Baek et al., 2021); it has been made available as part of the Robetta server, and the source code has also been released. RoseTTAFold is a deep-learning algorithm with features inspired by what was revealed about AlphaFold2 at CASP14.

4. Molecular-replacement assessment

Suitability for molecular replacement is one of the metrics for high-accuracy assessment in CASP, including CASP14 (Pereira et al., 2021; Millán et al., 2021). The assessment uses a log-likelihood gain (LLG) calculated in Phaser (McCoy et al., 2007; Read & McCoy, 2016). The likelihood is the probability that the data would have been measured given the model, and the log-likelihood gain is the difference between the log-likelihood of the model and that calculated from a random distribution of the same atoms (Wilson, 1949).

An important component of the scoring of models with the LLG is the incorporation of the estimated error in coordinates. As part of the modelling, groups are encouraged to estimate the error (Δ) in each atomic position and to record this estimate in the B-factor column of the deposited PDB file. This error estimate can be converted to a B-factor for each atomic position through the relationship $B = 8\pi^2\Delta^2/3$ and thereby used to weight each atom in the LLG calculation appropriately. Accurate estimates of the error improve the LLG (Bunkóczi et al., 2015; Croll et al., 2019), and in practice will add value to the models by increasing the signal in molecular-replacement searches.

If, during molecular-replacement searches, a pose of a model has an LLG over a certain space-group-dependent value (60 in nonpolar space groups, 50 in polar space groups and 30 in P1), the pose is probably correct. However, achieving this LLG is not sufficient to determine whether the full structure can be traced and refined to a point suitable for interpretation, publication and deposition; this also depends on the accuracy and completeness of the model and the resolution of the data. Testing the ability of a model to phase includes validation of these steps downstream of nominally successful model replacement.

4.1. Model parameters

Before structure solution, the LLG that will be achieved for a correct pose of a model can be estimated using the ‘expected LLG’ (eLLG; McCoy et al., 2017). We have previously shown that the eLLG for each reflection $hkl$ can be calculated from the $\sigma_A$ parameter (Read, 1986):

$$e\text{LLG}_{hkl} \sim \frac{1}{2} \sigma_A^2.$$  \hspace{1cm} (1)

The total eLLG is the sum of this value over all reflections $hkl$. The resolution-dependent $\sigma_A$ term is approximated with a four-parameter curve which includes the r.m.s.d., the $f_m$ and also two solvent parameters which affect the $\sigma_A$ value at resolutions lower than approximately 8 Å (Murshudov et al., 1997). For reflections higher than 8 Å resolution, this curve is dominated by the dependence on the square root of $f_m$ and an exponential dependence on r.m.s.d.$^2$. For each reflection $hkl$ (resolution $d_{hkl}$),

$$\sigma_{A,hkl} \sim (f_m)^{1/2} \exp\left(-\frac{2\pi^2}{3d_{hkl}^2} r.m.s.d.^2\right).$$  \hspace{1cm} (2)

Since the eLLG is a good estimate of the LLG when there are no pathologies in the data (Oeffner et al., 2018), the above equation also shows the relationship between r.m.s.d., $f_m$ and the LLG. The dependence of the LLG on r.m.s.d. and $f_m$ is shown in Fig. 2.

Any parameter that correlates strongly with r.m.s.d. (such as the GDT_HA, when the GDT_HA is high) will show the same relationship to the LLG.

The r.m.s.d. is only useful for predicting the LLG when the same r.m.s.d. value describes the difference between the model and target both locally and globally. If it varies between regions, the r.m.s.d. is dominated by the regions where the r.m.s.d. is large, whereas the LLG score will be dominated by the regions where the r.m.s.d. is low. In practice, there will be regions that are better modelled than others, and the LLG obtained from a model will be higher when the estimated errors in each coordinate are good and are incorporated into the electron-density calculation via the $B$ factor, as described above.
4.2. Targets

There were 33 crystal structures provided as modelling targets for CASP14. Of these, 31 have a single protein sequence (sometimes in multiple copies) in the asymmetric unit and two have two protein sequences. The relationship between the naming of the targets and the crystal structures is not straightforward. In 30 of the 31 cases with a single sequence there is a one-to-one correspondence between a sequence and a CASP target number (for example T1032). The exception is the case of the structure with PDB code 6vr4, where there are nine separate targets (T1031, T1033, T1035, T1037, T1039, T1040, T1041, T1042 and T1043), each representing between 95 and 404 residues of the full polypeptide sequence of 2194 residues, two copies of which are present in the asymmetric unit. In 11 of the 30 cases the full sequence is deemed to be ‘multidom’ (multi-domain) and is also divided into two, three or four domains; these are treated as additional, separate targets, with the suffix ‘-D1’, ‘-D2’, ‘-D3’ or ‘-D4’ added to the target number for the whole structure (for example, T1024 is divided into T1024-D1 and T1024-D2). In eight of the 11 ‘multidom’ targets the whole sequence target is referenced suffixed with ‘-D0’ (for example T1038-D0) and not simply by the target number alone, as in the other three of the 11 targets (for example T1024). The 19 of the 30 cases that are not ‘multidom’ have a single domain defined within the full sequence, and the target is given the suffix ‘-D1’ (for example T1032-D1). In the two of the 33 cases where there are two sequences in the asymmetric unit, the corresponding two targets are named with the same target number with the suffix ‘s1’ or ‘s2’ added (T1046s1 and T1046s2; T1065s1 and T1065s2). Neither of these divide their constituents into domains, and the single targets are given the ‘-D1’ suffix.

We also considered one other target, PDB entry 6un9, corresponding to target T1048, which was cancelled from CASP14 because of a lack of tertiary structure; it is a single sequence that folds into a single helix and forms a coiled coil. A model for this structure was also prepared by the AlphaFold2 group before it was cancelled.

In total, we considered 72 CASP14 targets from the 34 crystal structures when the domains of ‘multidom’ targets are included in the total (Table 1). Crystallographic details of relevance to the difficulty of structure determination, such as the resolution limit and the number of copies in the asymmetric unit, are discussed below in the context of the molecular-replacement trials.

4.3. Classification of targets

CASP classifies the targets by modelling difficulty in four categories: free modelling (FM), template-based modelling (TBM-easy and TBM-hard) and structures on the boundary between free modelling and template-based modelling (FM/TBM). In the set of crystal structures, there was a good representation of all four classes (Fig. 3). All but two of the structures were from lower organisms (viruses, bacteria, archaeabacteria and tetrahymena), and these two structures were classed as TBM, which reflects the high coverage of fold space that has now been achieved in higher organisms (Supplementary Table S1).

4.4. Metrics for target quality

For the purposes of judging models for molecular replacement, the model r.m.s.d. and \( f_m \) are the important metrics.

Of the metrics reported by CASP, the sequence-independent LGA (4 Å) parameters RMSD and LGA_S are most closely allied with the r.m.s.d. and \( f_m \). The RMSD is the...
root-mean-square deviation for the subset of Cα atoms from the model that correspond to the residues from the target structure in the sequence-independent LGA superposition. LGA_S is the sequence-similarity score, which is a combination of the GDT score and the LCS score, where the LCS is the longest continuous segment (as a percentage of the total sequence) that can fitted under a r.m.s.d. of a given cutoff. The LGA_S scores are similar to the GDT_TS scores for closely aligned structures. LGA_S is not sensitive to out-of-sequence register errors, in the same way as the \( f_m \) of a model for molecular-replacement phasing is not directly sensitive to any registration errors in the model.

A histogram of the metrics RMSD and LGA_S for the AlphaFold2 models across the 44 CASP crystallographic targets of interest is shown in Figs. 3(b) and 3(c). The LGA_S is skewed towards almost full sequence coverage, with an average of 87%, and the RMSD is clustered around the average RMSD of 1.27 Å. The RMSD and LGA_S also show the superiority of the AlphaFold2 models over the models submitted by other groups. Table 2 shows the metrics for the AlphaFold2 models with the best LGA_S and the best RMSD (of the five submitted) for the 44 CASP crystallographic targets of interest, compared with the best models by the same metrics overall. In only two cases (T1073 and T1085) was an AlphaFold2 model not the best as scored by LGA_S. In the case of T1085, the difference in the LGA_S was negligible (less than half a percent), and the AlphaFold2 model had a much lower RMSD (0.85 versus 1.39 Å). In the case of T1073, the differences between the models were mostly confined to a short region of the N-terminal helix that extended from the body of the globular fold. In 15 cases a non-AlphaFold2 model had a lower RMSD; however, this was exclusively at the expense of a lower (usually a far lower) LGA_S.

These two metrics [LGA_S and RMSD from LGA (4 Å)] are not ideal metrics for representing r.m.s.d. and \( f_m \). Calculation of r.m.s.d and \( f_m \) is critically dependent on the alignment of the electron density rather than coordinates, a problem that will be addressed elsewhere.

### Table 1

| Crystal No. | Target No. | CASP target | Residues | CASP domain | Residues | 'Multidom' domain(s) | Residues |
|-------------|------------|-------------|----------|-------------|----------|---------------------|----------|
| 1           | 1          | T1024       | 408      | T1024-D0    | (273)    | T1024-D1            | 193      |
| 2           | 2          | T1030       | 273      | T1030-D0    | (273)    | T1030-D2            | 204      |
| 3           | 3          | T1031       | 95       | T1031-D1    | (95)     | T1031-D1            | 154      |
| 4           | 4          | T1033       | 100      | T1033-D1    | (100)    | T1033-D1            | 119      |
| 5           | 5          | T1035       | 102      | T1035-D1    | (102)    | T1035-D1            |          |
| 6           | 6          | T1037       | 404      | T1037-D1    | (404)    | T1037-D1            |          |
| 7           | 7          | T1039       | 161      | T1039-D1    | (161)    | T1039-D1            |          |
| 8           | 8          | T1040       | 130      | T1040-D1    | (130)    | T1040-D1            |          |
| 9           | 9          | T1041       | 242      | T1041-D1    | (242)    | T1041-D1            |          |
| 10          | 10         | T1042       | 289      | T1042-D1    | 276      | T1042-D1            |          |
| 11          | 11         | T1043       | 148      | T1043-D1    | (148)    | T1043-D1            |          |
| 12          | 12         | T1032       | 284      | T1032-D1    | 169      | T1032-D1            |          |
| 13          | 13         | T1034       | 156      | T1034-D1    | (156)    | T1034-D1            |          |
| 14          | 14         | T1038       | 199      | T1038-D0    | 190      | T1038-D1            | 114      |
| 15          | 15         | T1046s1     | 216      | T1046s1-D1  | (72)     | T1046s1-D1          |          |
| 16          | 16         | T1046s2     | 216      | T1046s2-D1  | 141      | T1046s2-D1          |          |
| 17          | 17         | T1048       |          | T1048-D1    |          | T1048-D1            |          |
| 18          | 18         | T1049       | 141      | T1049-D1    | 134      | T1049-D1            |          |
| 19          | 19         | T1050       | 779      | T1050-D1    |          | T1050-D1            |          |
| 20          | 20         | T1052       | 832      | T1052-D0    | (832)    | T1052-D1            |          |
| 21          | 21         | T1053       | 580      | T1053-D0    | 576      | T1053-D1            | 171      |
| 22          | 22         | T1054       | 190      | T1054-D1    | 143      | T1054-D1            |          |
| 23          | 23         | T1056       | 186      | T1056-D1    | 169      | T1056-D1            |          |
| 24          | 24         | T1058       | 382      | T1058-D0    | (382)    | T1058-D1            |          |
| 25          | 25         | T1064       | 106      | T1064-D1    | 92       | T1064-D1            |          |
| 26          | 26         | T1065s1     | 225      | T1065s1-D1  | 11       | T1065s1-D1          |          |
| 27          | 27         | T1065s2     | 225      | T1065s2-D1  | 98       | T1065s2-D1          |          |
| 28          | 28         | T1067       | 220      | T1067-D1    | (221)    | T1067-D1            |          |
| 29          | 29         | T1070       | 335      | T1070-D1    |          | T1070-D1            |          |
| 30          | 30         | T1073       | 58       | T1073-D1    | (59)     | T1073-D1            |          |
| 31          | 31         | T1074       | 131      | T1074-D1    | (132)    | T1074-D1            |          |
| 32          | 32         | T1079       | 483      | T1079-D1    | 451      | T1079-D1            |          |
| 33          | 33         | T1080       | 137      | T1080-D1    | 133      | T1080-D1            |          |
| 34          | 34         | T1082       | 97       | T1082-D1    | 75       | T1082-D1            |          |
| 35          | 35         | T1083       | 196      | T1083-D1    | 92       | T1083-D1            |          |
| 36          | 36         | T1084       | 146      | T1084-D1    | 71       | T1084-D1            |          |
| 37          | 37         | T1085       | 588      | T1085-D0    | 406      | T1085-D1            |          |
| 38          | 38         | T1086       | 408      | T1086-D0    | 381      | T1086-D1            |          |
| 39          | 39         | T1087       | 186      | T1087-D1    | 93       | T1087-D1            |          |
| 40          | 40         | T1089       | 404      | T1089-D1    | 377      | T1089-D1            |          |
| 41          | 41         | T1090       | 193      | T1090-D1    | 191      | T1090-D1            |          |
| 42          | 42         | T1091       | 863      | T1091-D1    |          | T1091-D1            |          |
| 43          | 43         | T1100       | 338      | T1100-D0    |          | T1100-D1            |          |
| 44          | 44         | T1101       | 318      | T1101-D0    | 307      | T1101-D1            |          |

† Cancelled.
5. Molecular-replacement methods

The high LGA_S and low RMSD scores of the best AlphaFold2 models indicated that these models were good prospects for achieving phasing by molecular replacement.

Initially, the 31 targets with a one-to-one correspondence between a sequence and a CASP crystal structure, the two targets each for the two heterodimeric structures and the nine targets for PDB entry 6vr4 were used for molecular replacement (a total of 44 CASP crystallographic targets to be placed in 34 crystal structures).

If molecular replacement with the target representing the full sequence failed, and the target was one of the 11 classified as ‘multidom’, then molecular replacement was attempted with the domains.

The AlphaFold2 models for any given target almost exclusively superimposed with very little coordinate variability, and thus creating an ensemble structure did not indicate where poorly modelled regions could be trimmed (by divergence between models) unless exceptionally small divergence distance thresholds were used (for example 0.1 Å). Rather than using a tiny deviation threshold, trimming was performed using a threshold for the predicted error per residue supplied as part of the AlphaFold2 structure prediction.

The AlphaFold2 models (full and domain targets, un-trimmed and trimmed) were used for molecular replacement in Phaser.voyager (manuscript in preparation). Phaser.voyager uses the phasertng codebase (McCoy et al., 2021). The initial VRMS (effective r.m.s.d.) was set to 1.2 Å and then refined for posed models. The five submitted AlphaFold2 models were used as an ensemble. The pose was checked to see whether it was a match for the target coordinates with phenix.famos (Oefner et al., 2012). To confirm the solution, we used phenix.autobuild for the initial R value and R_free (Terwilliger et al., 2008). Structures were considered to be solved if the correlation between σ_A-weighted density maps computed with the AutoBuild model and the final structure was over 0.3. If the R_free was high, model improvement was attempted with phenix.morph_model. Further manual model building and refinement was not pursued.

In one case (described below) molecular replacement with Phaser.voyager failed, and molecular replacement was performed with ARCIMBOLDO_LITE for coiled coils (Caballero et al., 2018).

6. Molecular-replacement results

Of the 34 crystal structures, 31 could be solved by molecular replacement with the AlphaFold2 models, two could be partially solved and one could not be solved with the AlphaFold2 model.

The case that could not be solved with the AlphaFold2 model was crystal 8, the coiled-coiled structure with PDB code 6un9, target T1048. Although not solvable with the full AlphaFold2 model, this structure could be solved with a generic 20-residue polyalanine helix using ARCIMBOLDO_LITE for coiled coils (Caballero et al., 2018).

A partial solution was achieved for crystal 3, the polymerase structure with PDB code 6vr4, for which six of the nine constituent CASP targets (in two copies each) could be placed. The full structure was not designated as a CASP target. Should the whole structure, or even larger pieces, have been a target, and had an AlphaFold2 model been available, it is possible that such a model would have also succeeded in molecular replacement.

A partial solution was also achieved for crystal 2, the all-helical structure with PDB code 6poo. The full structure was designated as a ‘multidom’ CASP target with two domains. The second domain T1030-D1 could be placed unambiguously. The first domain T1030-D2 could be placed by molecular replacement but gave a very high final R_free.

Of the 31 solved structures, 28 were solved straightforwardly with the default Phaser.voyager protocol (Table 3).

Details are given below for crystal 8 (no solution), crystals 2 and 3 (partial solutions) and crystals 20, 23 and 33, for which
the structure solution was successful but proved to be more problematic.

6.1. Crystal 2 (PDB entry 6poo, target T1030)

T1030 is a helical bundle classified as ‘multidom’ with two domains. D2 could be placed unambiguously by molecular replacement, but the best pose for D1 was only able to superimpose a portion of the fragment, and the $R_{free}$ after molecular replacement was greater than 0.50.

The overall C-atom r.m.s.d. of the first ranked AlphaFold2 model to the target for D1 was 2.8 Å over 154 residues and that for D2 was 1.2 Å over 119 residues.

The high $R_{free}$ of the molecular-replacement solution can be attributed to model/target differences in the D1 binding angles and the angular disposition of the six constituent helices. Since the helices are long (residue lengths of 18, 15, 35, 15, 38 and 22) these differences result in systematic deviation of the coordinates, so that an overall r.m.s.d. does not give a complete picture of coordinate divergence.

Analysis with the HELANAL-Plus server (Kumar & Bansal, 2012) showed that the six AlphaFold2 helices were classified as ‘linear’, ‘curved’, ‘linear’, ‘linear’, ‘unassigned’ and ‘curved’, while the target helices were classified as ‘curved’, ‘curved’, ‘kinked’, ‘linear’, ‘kinked’ and ‘curved’, respectively, based on the average and maximum bending angles. Analysis with helixang from CCP4 (Winn et al., 2011) gave angles between the helical axes of helix 1 and helices 2–6 for the AlphaFold2 model of 173°, 7°, 194°, 19° and −154°, respectively, and for the target of 172°, 5°, 171°, −23° and −153°, respectively; most notable were the differences in the dispositions of helices 1 and 4 (a difference of 23°) and helices 1 and 5 (a difference of 42°).

6.2. Crystal 3 (PDB entry 6v4r, targets T1031, T1033, T1035, T1037 and T1039–T1043)

For CASP14, the single polypeptide chain of the virion-packaged DNA-dependent RNA polymerase of crAss-like phage phi4:2 was divided into nine assessment domains, which we refer to here by numbering them 1–9 (Fig. 4). Eight domains were classed as FM and one was classed as FM/TBM.

There were two copies of the monomer in the asymmetric unit (PDB entry 6v4r; Drobysheva et al., 2021) related by a noncrystallographic twofold. The assessment domains were used as models for molecular replacement. In total, 12 of the 18 domains could be placed, giving a 2/3 complete solution, which was insufficient for phasing the remaining fragments of the structure given the limited resolution of 3.5 Å. The partial solution was achieved by running Phaser from the command line. Domains were placed sequentially and the order of placement of the 12 domains was 4, 4, 7, 7, 2, 3, 2, 3, 8, 5, 5. The second copies of domains 2, 3 and 8 were not placed by molecular replacement, but by applying the noncrystallographic symmetry operator to the already placed copies and performing rigid-body refinement. After the placement of the first domain 2, 40 cycles of REFMAC (Murshudov et al., 2011)
Table 3
Summary of phasing of the 34 crystal structures of interest in CASP14 with AlphaFold2 models.

Crystals and targets listed in bold are discussed in the text.

| Crystal code | Target | PDB code | $d_{min}$ (Å) | No. of reflections | $Z^+$ | Domains Filter† | TFZ‡ | $R_{free}$ after morphing |
|--------------|--------|----------|---------------|-------------------|------|----------------|------|--------------------------|
| T1024        |        | 6liz     | 2.9           | 12686             | 1    | 1              | No   | 24.7 0.52 0.42           |
| T1030        |        | 6pp0     | 3.0           | 7525              | 1    | 1              | No   | 20.4 0.53                |
| T1031        |        | 6nr4     | 3.5           | 92907             | 2    |                |      |                          |
| T1033        |        |          |               |                   |      |                |      |                          |
| T1035        |        |          |               |                   |      |                |      |                          |
| T1037        |        |          |               |                   |      |                |      |                          |
| T1039        |        |          |               |                   |      |                |      |                          |
| T1040        |        |          |               |                   |      |                |      |                          |
| T1041        |        |          |               |                   |      |                |      |                          |
| T1042        |        |          |               |                   |      |                |      |                          |
| T1043        |        |          |               |                   |      |                |      |                          |
| T1048        |        | 6un9     | 2.8           | 19203             | 4    |                |      |                          |
| T1049        |        | 6yf4     | 1.8           | 12228             | 1    | 1              | No   | 23.7 0.34                |
| T1050        |        | 6tmd     | 2.7           | 97731             | 3    | 1              | No   | 39.0 0.30                |
| T1052        |        | 6ya2     | 2.0           | 88914             | 2    | 1              | No   | 48.6 0.43                |
| T1053        |        | 7n7a     | 3.2           | 49627             | 4    | 1              | No   | 51.0 0.35                |
| T1054        |        | 6v4v     | 1.7           | 25547             | 1    | 1              | No   | 33.2 0.34                |
| T1056        |        | 6jy1     | 2.3           | 17863             | 2    | 1              | No   | 19.7 0.37                |
| T1058        |        | 7abw     | 3.1           | 20228             | 2    | 1              | No   | 25.5 0.44                |
| T1064        |        | 7jkl     | 2.0           | 16787             | 2    | 1              | No   | 19.1 0.41                |
| T1065s1      |        | 7m5f     | 1.6           | 35695             | 1    | 2              | No   | 48.7 0.22                |
| T1065s2      |        |          |               |                   |      |                |      |                          |
| T1067        |        | 6liz     | 1.4           | 51025             | 1    | 1              | No   | 57.2 0.26                |
| T1070        |        | 6liz     | 2.5           | 25412             | 1    | 1              | No   | 6.8 0.49 0.42           |
| T1073        |        | 6liz     | 1.9           | 27326             | 4    | 1              | No   | 24.1 0.39                |
| T1074        |        | 7oe9     | 1.5           | 25800             | 1    | 1              | No   | 21.1 0.29                |
| T1079        |        | 6liz     | 3.2           | 47985             | 4    | 1              | No   | 33.6 0.38                |
| T1080        |        | 6liz     | 1.7           | 106570            | 6    | 1              | No   | 20.9 0.39                |
| T1082        |        | 6liz     | 6.0           | 97762             | 2    | 1              | No   | 33.6 0.44 0.31          |
| T1083        |        | 6liz     | 1.3           | 81236             | 4    | 1              | No   | 30.0 0.48 0.35          |
| T1084        |        | 6liz     | 1.9           | 23901             | 3    | 1              | No   | 32.1 0.38                |
| T1085        |        | 6liz     | 2.5           | 10578             | 1    | 3              | No   | 22.0 0.40                |
| T1086        |        | 6liz     | 2.3           | 21887             | 1    | 1              | No   | 18.7 0.41                |
| T1087        |        | 6liz     | 1.4           | 69617             | 4    | 1              | No   | 43.2 0.25                |
| T1089        |        | 6liz     | 2.2           | 55192             | 2    | 1              | No   | 63.5 0.29                |
| T1090        |        | 6liz     | 1.8           | 22947             | 1    | 1              | No   | 27.2 0.29                |
| T1091        |        | 6liz     | 2.2           | 62789             | 1    | 1              | No   | 22.0 0.44                |
| T1090        |        | 6liz     | 3.1           | 36829             | 4    | 1              | No   | 31.1 0.48 0.45          |
| T1091        |        | 6liz     | 1.4           | 58030             | 1    | 1              | No   | 16.2 0.34                |

† Number of copies in the asymmetric unit. †† Threshold for predicted RMSD below which residues were pruned from the model. †§ Translation-function Z-score, a measure of significance of the MR solution. †† Partial solution. †‡ No solution.

refinement were performed to improve the partial structure before continuing. This procedure was repeated after placing the second domain 3. Domains 1, 5 and 9 could not be placed; these domains had a very high r.m.s.d. to the target of over 2.5 Å (Table 4).

However, a model for this structure was generated by AlphaFold2 before the target was cancelled.

This structure is a 61-residue coiled coil. Coiled-coil structures are notoriously difficult to solve by molecular replacement because of the modulation of the data due to the helical repeats (Caballero et al., 2018; Thomas et al., 2015).

The structure had four copies of the target in the asymmetric unit. The overall C-alpha r.m.s.d. of the AlphaFold2 model to chain A in the target structure was 2.1 Å for 67 residues. Analysis with the HELANAL-Plus server (Kumar & Bansal, 2012) showed that the maximum bending angle was markedly different (7.4° versus 25.7°); the model helix was classified as ‘curved’, while the target was classified as ‘kinked’. The structure could not be solved with the AlphaFold2 model in Phaser.voyager, even with the model

6.4. Crystal 8 (PDB entry 6un9, target T1048)

T1048 is a single helix and was cancelled from CASP14 on 20 October 2020 for a ‘lack of tertiary structure’. However, a model for this structure was generated by AlphaFold2 before the target was cancelled.

This structure is a 61-residue coiled coil. Coiled-coil structures are notoriously difficult to solve by molecular replacement because of the modulation of the data due to the helical repeats (Caballero et al., 2018; Thomas et al., 2015).

The structure had four copies of the target in the asymmetric unit. The overall C-alpha r.m.s.d. of the AlphaFold2 model to chain A in the target structure was 2.1 Å for 67 residues. Analysis with the HELANAL-Plus server (Kumar & Bansal, 2012) showed that the maximum bending angle was markedly different (7.4° versus 25.7°); the model helix was classified as ‘curved’, while the target was classified as ‘kinked’. The structure could not be solved with the AlphaFold2 model in Phaser.voyager, even with the model...
optimized by trimming to reduce the r.m.s.d. to the deposited structure (at the expense of a lower fraction scattering).

ARCIMBOLDO_LITE in coiled-coil mode was able to solve the structure using a generic 20-residue polyalanine helix. The advantage of using short generic helices for coiled-coil structures is that they are able to superimpose with multiple short sections of the coiled-coil helices with low r.m.s.d.. Structure solution required the ‘verification’ step, which is a powerful method for distinguishing the true solution from the abundance of false solutions that arise merely from helical placements that satisfy the helical modulations of the data (Caballero et al., 2018).

6.5. Crystal 20 (target T1073)

All groups modelled the 12-residue N-terminal helix of T1073 with high r.m.s.d. to the target. This helix extends from the compact body of the fold. Structure solution was achieved by removing all sections of the AlphaFold2 model with a predicted error over 1 Å in a standard model-preparation protocol with Phaser.voyager.

The challenge in this case was with data preparation not model preparation; the model preparation was unremarkable, but we found that this case also required some additional attention be given to the crystallographic data. A number of different data sets were available in the file provided. Molecular replacement was achieved, after phenix.xtriage (Zwart et al., 2005) analysis, with one of the data sets and with the resolution restricted to 2.8 Å.

6.6. Crystal 23 (target T1080)

T1080 was classified as FM/TBM. There were six copies of the target in the asymmetric unit in two trimers.

This was the only case where the five submitted AlphaFold2 models showed significant deviation. Model 3 differed from the consensus fold of the other four in the first 40 N-terminal residues; these 40 residues took a very different conformation in model 3. In analysis, the consensus fold of the four models was correct, and model 3 was incorrect, although the incorrect conformation could be considered as a ‘trimer swap’ error, with the chain partly following the fold of a neighbouring monomer in the trimer. The molecular-replacement model trimmed these residues and residues with a predicted deviation of more than 1.2 Å, leaving 78 of 133 residues. The molecular-replacement model was therefore 60% of the target structure. After structure solution, the full AlphaFold2 consensus fold was superimposed on the solution and used for refinement.

6.7. Crystal 33 (target T1100)

T1100 was classified as ‘multidom’ with two domains. There were four copies in the asymmetric unit in two dimers with translational noncrystallographic symmetry between the two dimers. D2 is a compact globular structure. D1 is a helical bundle structure with four helices of 52, 11, 64 and 28 residues. Within the dimer, the D1 helices formed a coiled coil.

Table 4

| Model       | Z   | Type    | Residues | R.m.s.d. (Å) | Chain |
|-------------|-----|---------|----------|--------------|-------|
| T1031TS427_1-D1 | 2   | FM      | 95       | 2.91         |       |
| T1033TS427_1-D1 | 2   | FM      | 100      | 1.58         | EG    |
| T1035TS427_1-D1 | 2   | FM/TBM  | 102      | 0.81         | FH    |
| T1037TS427_1-D1 | 2   | FM      | 404      | 1.25         | AB    |
| T1039TS427_1-D1 | 2   | FM      | 161      | 2.50         | KL    |
| T1040TS427_1-D1 | 2   | FM      | 130      | 2.76         |       |
| T1041TS427_1-D1 | 2   | FM      | 242      | 1.70         | CD    |
| T1042TS427_1-D1 | 2   | FM      | 276      | 1.79         | IJ    |
| T1043TS427_1-D1 | 2   | FM      | 148      | 2.46         |       |

Figure 4

Crystal 3, PDB entry 6vr4, targets T1031 (1, red), T1033 (2, orange), T1035 (3, yellow), T1037 (4, khaki), T1039 (5, green), T1040 (6, blue), T1041 (7, purple), T1042 (8, magenta) and T1043 (9, violet). Two targets are discontinuous in the primary sequence. (a) Structure with targets highlighted; regions not corresponding to a target are shown in grey. The figure was created with Mol* (Sehnal et al., 2021). (b) Sequence with targets highlighted; regions that are not highlighted were not included in targets.
D2 could be placed unambiguously by molecular replacement.

D1 was more difficult to place. The problem can be attributed to model/target differences in helical bending angles. In general, the helices in the model were straighter than those in the target, with average bending angles of 4.5°, 7.1°, 4.6° and 3.6° versus 8.5°, 9.3°, 8.5° and 7.8°, respectively. When the differences in bending angles were compounded over the long helices, particularly helix 1 (~75 Å) and 3 (~90 Å), it was not possible to simultaneously superimpose both ends of the model and the target. Molecular replacement gave several closely related poses for D1, superimposing different portions of the model and target helix.

7. Survey of phasing methods

To discern the impact of high-accuracy in silico models on crystallographic phasing methods, we undertook a survey of crystallographic phasing methods since the turn of the millennium.

We can divide crystallographic phasing strategies into four broad categories: direct methods, experimental phasing, molecular replacement and difference Fourier methods (Fourier synthesis). The use of direct-methods phasing is negligible for macromolecular crystallography, in contrast to its supreme dominance in small-molecule crystallography (Sheldrick, 2008). Within the experimental phasing category are MAD (multi-wavelength anomalous dispersion), SAD (single-wavelength anomalous dispersion) and various IR (isomorphous replacement) methods, SIR (single isomorphous replacement), SIRAS (single isomorphous replacement with anomalous scattering), MIR (multiple isomorphous replacement) and MIRAS (multiple isomorphous replacement with anomalous scattering) (for a review, see Rupp, 2010).

The PDB mostly records macromolecular crystal structures that are published in peer-reviewed journals. The PDB is a record of novel crystal forms, if not novel structures. Our analysis only included those entries where protein was a component of the crystal.

The phasing method for each PDB entry is recorded in the ‘structure determination method’ field, which should allow a survey of phasing methods; however, analysis is not straightforward for a number of reasons listed below.

(i) Although the ‘structure determination method’ field has been compulsory for submissions commenced after 29th January 2019, a significant portion of the historical entries are null. Entries recorded before 2000 were regarded as too sparse for analysis. Null entries may be biased towards particular categories of phasing.

(ii) Although the ‘structure determination method’ field has been restricted to a few text strings for submissions commenced since 29th January 2019, historically it was ‘free format’ and highly variable. For this study, all historical text entries were scanned by eye to assign each to one of the new restricted values. If the field referenced a number of methods (for example SAD with molecular replacement or SIRAS/MAD) then the most senior phasing method was allocated, with the order of precedence being MIRAS, MIR, MAD, SIRAS, SIR, SAD, molecular replacement and Fourier synthesis.

(iii) Phasing by direct methods was not included in the study because a survey of entries with the ‘structure determination method’ field ab initio showed that although these entries included those phased with direct methods, in the majority of cases ab initio referred to fragment-based approaches to molecular replacement or the use of direct methods for anomalous substructure determination. Since very few entries were categorized as ab initio, removing these from consideration did not significantly bias the results.

(iv) Checking a small sample of the entries in the ‘structure determination method’ field against the method recorded in the corresponding publication showed that the field was not always accurate. Inaccuracies may be biased towards particular categories of phasing.

(v) Each entry has a deposition date, a release date and a revision date, therefore dating each entry is problematic. The deposition and release dates are commonly separated by a year, but can be three years apart or even more. Revision dates are commonly very recent, as they include PDB-wide changes to PDB nomenclature. Only entries with ‘entity ID’ with value ‘1’ were considered. In order to track the evolution of structure determination methods, we considered five-year intervals and restricted PDB identifiers for each interval such that both the deposition date and the release date were within the five-year interval. Hence, our analysis sampled a subset of the PDB entries.

(vi) The understanding of the definition of different phasing methods may vary between crystallographers. For example, there is a degree of overlap between Fourier synthesis and molecular-replacement methods, as the former can be considered to be the latter but without an initial wide-radius search strategy (a search strategy employing rotation and translation functions); if the pose is outside the radius of convergence of rigid-body refinement, then (nominally) isomorphous crystals cannot be phased by difference Fourier methods and molecular replacement with a local or global search is used. Similarly, crystallographers may not distinguish between various similar experimental phasing methods (for example SAD versus SIRAS).

Despite these caveats, the trend in the change of structure determination methods over the last 20 years is clear (Fig. 5a) and mirrors anecdotal experience. Molecular replacement now accounts for around 80% of phasing, increasing from around 50% of phasing in 2000, and molecular-replacement and Fourier synthesis (difference Fourier) methods combined account for 95% of phasing. It is possible that an even higher proportion of structures would have been amenable to phasing by molecular replacement had it been attempted.

Within the experimental phasing strategy, the method of experimental phasing has changed considerably, with MAD dominating in 2000 but SAD dominating today; SAD phasing, commonly using selenomethionine-substituted protein, now accounts for 82% of experimentally phased structures (Fig. 5b).
Of particular note is the decline in IR methods, which were originally the backbone of macromolecular phasing. To obtain an overview of the type of structures that currently require IR to phase, we examined the structures submitted and released in 2020 in more detail (Supplementary Table S2). 13 structures met this criterion, were phased by IR methods and had publications available at the time of writing. Of these 13, we found one was actually solved by Se-SAD, one by Os-SAD, one by Pt-MAD and one by MR. Of the nine confirmed examples of IR, only two structures used multiple derivatives.

8. Discussion

Self-evidently, crystallography requires crystals; crystallization is a bottleneck, albeit one that has become far less constraining with advances in expression systems, fluidics, robotics and computer vision. Not only must there be crystals, but the crystals must diffract to better than 4 Å resolution to be useful for structural biology. With the data collected to the highest resolution that a crystal form (space group, unit cell, asymmetric unit contents) will allow, it is usual to regard the crystallographic data as the ‘fixed’ component of molecular-replacement phasing and to regard the model as the ‘dynamic’ component. Phasing pipelines are primarily designed for the automated exploration of many model structures prepared in different ways, with the hope that one will be accurate enough to be placed and allow model building and refinement to proceed with the single data set provided.

To some degree, the AlphaFold2 models upend this paradigm. The need for the extensive generation of models through different combinations of homologs in ensembles, different levels of trimming, the mining of domain databases and the use of small secondary-structure elements as models is likely to be greatly reduced as these high-accuracy models become available. In essence, the AlphaFold2 models distil the information from all of these methods, and more, in a single structure. The crystallographic problem may become one of finding a crystal form (for example, with fewer copies in the asymmetric unit) that is amenable to molecular replacement with the in silico model(s). As an example of this, successful molecular replacement with T1073 (crystal 20) was achieved after massage of the data, rather than the model.

For five of the 19 structures that were classified as ‘multidom’, we used the domains rather than the model of the whole structure for molecular replacement. In this approach, the whole structure is built up by addition as domains are placed sequentially in the asymmetric unit. It is necessary when the disposition of the domains in the target is largely determined by crystal packing or allosteric effects.

The structures that were the most challenging to solve with the AlphaFold2 models contained extended helices. The problem was twofold. Firstly, although helical secondary structure is very amenable to prediction, the subtle bends and kinks in the helices are more elusive, and these have long-range effects in the fit of the model to the target. Secondly, coiled coils induce modulations in the diffraction data that confound the maximum-likelihood targets in molecular replacement, a known issue and an active area of crystallographic methods development.

The statistics in Table 3 show that molecular replacement with the AlphaFold2 models, followed by simple refinement strategies, does not give structures that are suitable for immediate submission to the PDB. Investment in manual model building, informed by a degree of biological understanding of the structures, would have been required to obtain final structures, which was beyond the scope of this study.

The use of in silico models for molecular replacement will also impact downstream model building and refinement. Model building and refinement can already be assisted by techniques borrowed from ab initio modelling (Terwilliger et al., 2012). With models representing 100% of the polypeptide chain in approximately the correct conformation, model building is directed towards local minimization into electron density rather than de novo model building. In this study, we used phenix.morph_model to improve parts of the structure.
with an initially poor fit to the density. In regions where the electron density is weak or absent due to static disorder in the crystal, constraining the structure to the model may lower the R factors and improve the interpretation of the density. In extremis, the diffraction data may not need to be as good as they would need to be to refine, pass validation metrics and publish the structure in the absence of the model. In effect, the diffraction data need only verify the model.

There is some work to do to optimize the use of high-accuracy in silico models for the purposes of molecular replacement. The lack of conformational variability in the models is different from models drawn from homologs. Whereas homologs tend to vary most in the regions where they also deviate from the target structure, the AlphaFold2 models are very consistent (insistent) even in regions where they differ from the target structure. If taken purely at face value, this will lead to, for example, rejection of molecular-replacement solutions due to (false) packing clashes. We can also improve how we make use of the estimated error in the coordinates in model preparation. It is also likely that this will lead to, for example, rejection of molecular-replacement solutions due to (false) packing clashes. We can also improve how we make use of the estimated error in the coordinates in model preparation. It is also likely that improvements can be made in the estimation of $\sigma_A$ for these models, since optimization of $\sigma_A$ estimation has been calibrated for homologs rather than in silico models (Hatti et al., 2020).

The CASP14 crystal structures mostly represent a particular type of crystal structure: those that have a single protein sequence in the asymmetric unit and consist of one or few domains where the domain is unrelated, or poorly related, to known structures. These types of crystal structures are selected by CASP since they represent the more challenging structures for structure prediction. However, in support of structural biology, crystallography often focuses on protein complexes with peptide motifs, oligomeric associations and multi-domain structures, often with domains that already have homologous structures in the PDB. That these can already be solved by molecular replacement or Fourier synthesis in at least 95% of cases is evident in the statistics and does not diminish their scientific interest.

Our survey of phasing methods indicates that IR phasing is becoming a specialist method. Despite the undoubted power of IR to obtain spectacularly good phases, even with low resolution and poor data, there are other factors that mean that IR is avoided where possible. If using heavy metals, it requires handling highly toxic metal salts, which also bind to protein crystallographers, not just to proteins (Blundell & Johnson, 1976). Methods that incorporate noble gases such as xenon require specially designed high-pressure cells and appropriate training and support to use. We note that two of the nine structures phased by IR in our survey of 2020 were phased by SIRAS using iodine, which is a nontoxic and simple method.

Crystallographic phasing strategies have evolved continuously since 1913 (Ewald, 1962; Brooks-Bartlett & Garman, 2015), and the contribution of the high-accuracy models is continuing this evolution. It is already enabling crystallographers to concentrate their efforts even more keenly on the structural biology by making crystallographic phasing even more straightforward. We should look forward to the biological insights that this will bring.

9. Related literature

The following references are cited in the supporting information for this article: Bahat et al. (2020), Eckenroth et al. (2021), Filipčík et al. (2020), Gandini et al. (2020), Garcia-Doval et al. (2020), He et al. (2020), Jäger et al. (2020), Jensen et al. (2020), Micevski et al. (2020), Nie et al. (2021), Sakurai et al. (2020), Zhu et al. (2020) and Zimanyi et al. (2020).

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