Covariation-derived residue contacts in *ab initio* modelling and Molecular Replacement

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List of Abbreviations

CC Correlation Coefficient

LLG Log-Likelihood Gain

MR Molecular Replacement

MSA Multiple Sequence Alignment

PDB Protein Data Bank

RIO Residue-Independent Overlap RMSD Root-Mean-Square Deviation

TFZ Translation Function Z-score
TM-score Template-Modelling score

Introduction

Materials & Methods

Residue contacts predicted by evolutionary covariance extend the application of *ab initio* molecular replacement to larger and more challenging protein folds

Note: The majority of the work presented in this chapter was published in two independent pieces of work. All work relating to the globular targets was published by Simkovic et al. [1], and a great majority of work relating to the transmembrane targets by Thomas et al. [2]. As such, this chapter consists of extracts from both publications with additional information where appropriate. Text duplicated from either publication was written by Felix Simkovic, all other elements were adapted.

3.1 Introduction

The introduction of residue-residue contacts as distance restraints in *ab initio* protein structure prediction has proven to be a highly successful approach to limiting the conformation search space thereby enabling successful fold prediction of larger and more β -rich protein structures [e.g., 3–11]. In AMPLE, these two domains are the major limitation for a more successful approach [12]. This typically results in user success being limited to small globular and primarily α -helical folds, or time- and resource-demanding attempts most likely going to be unsuccessful for larger targets

With the advent of contact information, is has thus become essential to identify the extend to which this invaluable bit of information is going to help AMPLE users in the future.

3.2 Materials & Methods

3.2.1 Target selection

In this study, targets from the ORIGINAL and TRANSMEMBRANE datasets were used. This resulted in a final set of 21 globular and 17 transmembrain protein targets. For details in how the targets were selected refer to ????, and for details on each target refer to ????.

3.2.2 Contact prediction

For all globular targets, one contact map was predicted with the fully automated metapredictor PCONSC2 v1.0 [13]. In summary, four Multiple Sequence Alignment (MSA)s were generated with JACKHMMER v3.1b2 [14] against the uniref100 v2015-10 database and HHBLITS v2.0.15 [15] against the uniprot20 v2013-03 database [16] at E-value cutoffs of 10⁻⁴⁰, 10⁻¹⁰, 10⁻⁴ and 1. Each MSA was analysed with PSICOV v2.13b3 [17] and PLMDCA v2 [18] to produce 16 individual contact predictions. All 16 predictions and per-target PSIPRED v3 [19] secondary structure prediction, NET-

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SURFP v1.0 [20] solvent accessibility information and HHBLITS v2.0.15 [15] sequence profile were provided to the PCONSC2 deep learning algorithm [13] to identify protein-like contact patterns. The latter produced a final contact map for each target sequence.

An additional contact map for β-structure containing targets was predicted using CCMPRED v0.3 [21] and reduced to β-sheet contact pairs using the CCMPRED-specific filtering protocol BBCONTACTS v1.0 [22]. Each MSA for CCMPRED contact prediction was obtained using HHBLITS v2.0.15 [15]. This entailed two sequence search iterations with an E-value cutoff of 10⁻³ against the uniprot20 v2013-03 database [16] and filtering to 90% sequence identity using HHFILTER v2.0.15 [15] to reduce sequence redundancy in the MSA. Besides the contact matrix as input, BBCONTACTS requires a secondary structure prediction and an estimate of the MSA diversity. The secondary structure prediction was taken from the PCONSC2 step whilst the diversity factor was calculated using ??.

For each transmembrane protein target, a MSA was generated using HHBLITS v2.0.16 [15] against uniprot20 v2016-02 database [16]. Contact predictions for each transmembrane target were obtained using the metapredictor METAPSICOV v1.04 [23], which in turn used the contact prediction algorithms CCMPRED v0.3.2 [21], FREECONTACT v1.0.21 [24] and PSICOV v2.1b3 [17]. Additionally, a set of contacts was also generated using the MEMBRAIN server v2015-03-15 [25].

3.2.3 Contact-to-restraint conversion

For all targets, the predicted contact maps were converted to ROSETTA restraints to guide $ab\ initio$ structure prediction. The FADE energy function was used to introduce a restraint in ROSETTA's folding protocol. The implementation described by Michel et al. [4] was used, which defined a contact to be formed during folding if the participating C β atoms (C α in case of glycine) were within 9Å of one another. The top-L (L corresponds to the number of residues in the target sequence) contact pairs were converted to ROSETTA restraints, and if satisfied a "squared-well" bonus of -15.00 added to the energy function.

Additionally to above, all β-containing targets were subjected to a further conversion step in a separate condition. The approach of adding BBCONTACTS restraints to a previous prediction is outlined in ??.

3.2.4 Ab initio structure prediction

Fragments for all targets were selected using the make_fragments.pl script shipped with ROSETTA. To ensure no homologous fragments were included in the fragment libraries, the -nohoms flag was set. Each target's secondary structure prediction was

provided to the fragment picker using the -psipredfile argument. The fragment libraries, contact restraints and secondary structure prediction were subjected to the ROSETTA AbinitioRelax protocol [26] to predict 1,000 decoys per target. ROSETTA options were chosen according to the default protocol in AMPLE v1.0 [12]. ROSETTA v2015.05.57576 was used for globular targets and v2015.22.57859 for transmembrane ones for all ROSETTA-related protocols.

3.2.5 Molecular Replacement in AMPLE

All generated decoys were subjected to AMPLE v1.0 [12] for ensemble search model generation.

All transmembrane protein targets were processing using AMPLE's default parameters. Molecular Replacement (MR) trials were performed with software versions shipped in CCP4 v6.5.13 [27], with the exception of SHELXE v2014/14 [28] and ARP/wARP v7.5 [29].

All globular protein targets were subjected to AMPLE with two deviations from the default parameters. The -use_scwrl was set to subject all decoys to side-chain remodelling using SCWRL4 [30]. Furthermore, the number of clusters to trial was set increased from one to three via the -num_clustersparameter. All MR trials were performed with the version of software shipped with CCP4 v6.5.15 [27].

All MR solutions were assessed for success using the criteria described in ??.

3.3 Results

In this study, the application of residue-residue contact predictions to *ab initio* protein structure prediction and subsequently MR was investigated. This proof-of-concept work is based on two datasets covering a range of globular and transmembrane protein targets. At the time of conducting this study, state-of-the-art contact prediction algorithms were applied to obtain the best possible contact predictions to identify the extend of pushing the boundaries previously incurred in AMPLE studies [12].

3.3.1 Residue-residue contact prediction

Accurate coevolution-based residue-residue contact prediction is highly dependent on the availability of many divergent homologous sequences. As such, it is important to validate that the selected targets in this study satisfy such requirement.

The depth of MSAs obtained for each target sequence suggests that sufficient num-

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bers of divergent homologous sequences are available. Across all globular targets, the minimum alignment depth is obtained for Galectin-3 domain (Protein Data Bank (PDB) ID: 1kjl) with 679 effective sequences and the maximum for G-protein Arf6-GDP (PDB ID: 1e0s) with 1,897 effective sequences (Fig. 3.1a). The median alignment depth for all globular targets is over 1,000, which is beyond the often suggested threshold of 200 sequences [31]. The MSAs for all transmembrane protein targets also surpass this threshold comfortably. The median alignment depth is much higher than for globular targets with 1,878 sequences (Fig. 3.1b). The minimum, which was obtained for Sensory rhodopsin II (PDB ID: 1gu8), is 692 sequences and the maximum for the sequence of Rhomboid protease GLPG (PDB ID: 2xov) is 6,583.

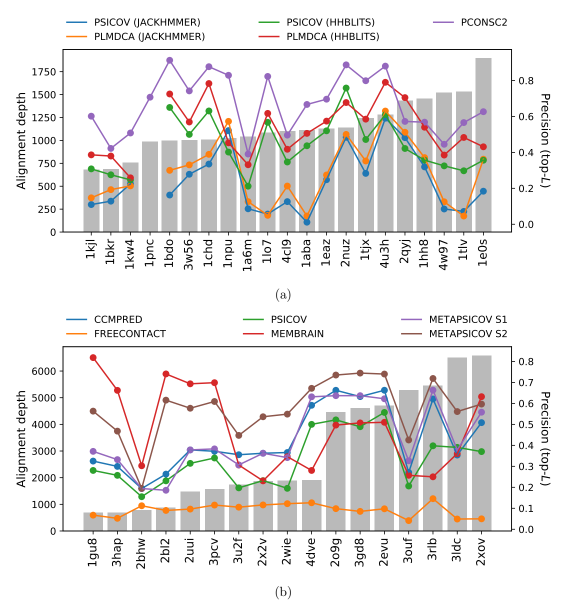


Figure 3.1: Alignment depth and contact precision analysis of (a) globular and (b) transmembrane protein targets. Contact predictions were obtained with several contact prediction algorithms. Precision scores were calculated for the top-L contact pairs. JACKHMMER and HHBLITS alignments for PSICOV and PLMDCA contact predictions in (a) were obtained with E-value $1e^{-4}$.

In coevolution-based contact prediction, the precision depends on alignment depth. Despite sufficient number of effective sequences across all targets, findings in this study suggest that some (meta-)predictors cannot fully utelise greater alignment depths to correct contact pairs (Fig. 3.1).

PCONSC2 — a metapredictor using numerous starting alignments and two contact predictors — outperforms its individual parts for almost all globular targets (Fig. 3.1a). Despite only a fraction of all generated contact predictions illustrated in Fig. 3.1a, the pattern translates across all 16 predictions per target. Such results suggest that precision greatly depends on the tool used to identify and select homologous sequences for the MSA. A closer inspection of mean precision scores resulting from HHBLITS-and JACKHMMER-based alignments shows higher precision scores for top-L contact pairs based on the former alignments (Table 3.1). Nevertheless, the Machine Learning approach in PCONSC2 to combine more and less precise individual predictions results in superior precision in the output (Table 3.1). No correlation could be established between alignment depth and precision for either individual predictors or the metapredictor PCONSC2 (Fig. 3.1a).

Table 3.1: Summary of mean PCONSC2 raw contact prediction precision based on JACKHMMER and HHBLITS alignments and PSICOV, PLMDCA and PCONSC2 coevolution-based contact prediction.

Contac	Alignment E-value cutoff				
		$1e^0$	$1e^{-4}$	$1e^{-10}$	$1e^{-40}$
PSICOV	JACKHMMER	0.240	0.239	0.213	0.167
rsicov	HHBLITS	0.439	0.435	0.354	0.209
PLMDCA	JACKHMMER	0.293	0.288	0.252	0.140
PLMDCA	HHBLITS	0.545	0.530	0.447	0.224
PCONSC2			(0.667	

Contacts for transmembrane protein targets in this study were predicted with the metapredictor METAPSICOV and the transmembrane-specific predictor MEMBRAIN. METAPSICOV STAGE 1 and STAGE 2 predictions outperform MEMBRAIN in nine and ten cases, respectively, whilst MEMBRAIN outperforms METAPSICOV for the rest (Fig. 3.1b). The METAPSICOV algorithm utelises the raw predictions by CCM-PRED, FREECONTACT and PSICOV to generate its STAGE 1 and STAGE 2 predictions. METAPSICOV STAGE 1 predictions are near identical to CCMPRED, whereby 15 of 17 targets show an absolute $\Delta_{TMscore}$ of less than 0.05 (Fig. 3.1b). This similarity does not propagate to METAPSICOV STAGE 2 predictions with only a single target showing such similar Template-Modelling score (TM-score) values (Fig. 3.1b). Amongst the three raw predictors used by METAPSICOV, FREECONTACT performs by far the worst with a mean TM-score of 0.09 across all transmembrane targets. PSICOV shows similar trend to CCMPRED when assessed by target, which results in a

mean absolute $\Delta_{TMscore}$ of 0.10.

The addition of BBCONTACTS contact pairs to improve structure prediction accuracy for β -structure containing targets is a novel aspect introduced in this study. The initial step of the addition of BBCONTACTS contact pairs included the filtering of one- and two-pair strand contacts from the original BBCONTACTS list, since those contained high numbers of false positives (Jessica Andreani, personal communication). The findings in this study confirm this for all β -structure containing targets. Precision values improved for all targets with changes ranging from 0.01 to 0.14 whilst retaining on average 80% of all contacts. Filtered BBCONTACTS predictions were combined with other contact maps, i.e. PCONSC2, to either upweight or add contact pairs. Findings in this study highlight that upweighted contact pairs are more precise than ones to be added. The minimum precision score for a set of upweighted contacts is 0.72 for 29 contact pairs and the maximum of 1.00 for up to 27 contact pairs. In comparison, novel BBCONTACTS contact pairs not present in the base contact map range in precision scores from 0.22 (nine contacts) to 0.76 (21 contacts).

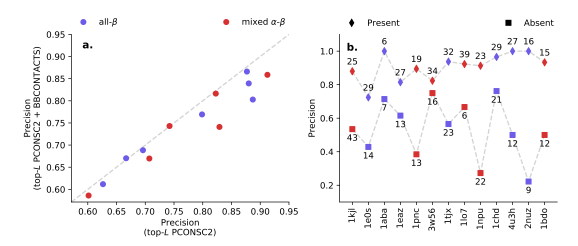


Figure 3.2: Evaluation of BBCONTACTS contact pairs. (a) Precision evaluation of PCONSC2 contact map with and without BBCONTACTS. (b) Precision evaluation of BBCONTACTS contact pairs split by status of presence or absence in the base PCONSC2 contact list. Numbers besides each marker indicate the number of contacts. The order scatter points is identical between both subplots and based on the PCONSC2 precision values (x-axis) in (a).

Despite the high precision of BBCONTACTS contact pairs, the merge of such pairs with top-L PCONSC2 contact pairs results in an expected loss in precision for the resulting contact set (Fig. 3.2). True positive contacts, which dominate the BBCONTACTS contact set, are usually also predicted by PCONSC2, and thus upweighted (Fig. 3.2). Since upweighting does not affect the precision, the value remains unaffected after this procedure. However, contact pairs unique to BBCONTACTS contact more false positives. Once added to the base PCONSC2 contact list, these contacts therefore reduce the precision value (Fig. 3.2). Either subset of BBCONTACTS con-

tacts does not show any correlation between the number it contains and its precision. The fold of the target does not show any clear distinction between better and worse sets of contacts either (Fig. 3.2).

3.3.2 Protein structure prediction

Predicted contact information is particularly useful to limit the conformation search space in *ab initio* protein structure prediction. Since such predictions are the basis for AMPLE studies presented in this thesis, it is important to analyse the improvement in decoy quality.

Globular protein targets benefit greatly from the addition of PCONSC2 residue contacts. All but one target see median TM-score improvements of at least 0.05 when comparing contact-guided PCONSC2 decoys with simple ROSETTA decoys (Fig. 3.3). The greatest improvement over 1,000 decoys was achieved for Oxy-myoglobin (PDB ID: 1a6m) with an improvement in median TM-score of 0.42. The decoys for Ankyrin (PDB ID: 2qyj) show a minor decrease in median TM-score of 0.04; however, the median TM-score for ROSETTA decoys is 0.78, and thus a minor decrease may be negligible.

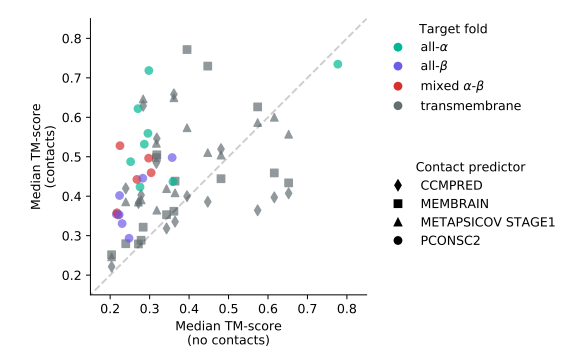


Figure 3.3: Effect of contact distance restraints on *ab initio* decoy quality by comparison of unrestrained (*no contacts*) and contact-restrained (*contacts*) median TM-scores for 1,000 decoys per target. Colours indicate the target fold and symbols the contact prediction algorithm.

Previously, *ab initio* protein structure prediction for globular targets was greatly limited by target fold and chain length. The addition of residue-residue contacts en-

hances decoy quality primarily for α -helical and mixed α - β protein targets (Fig. 3.4). Whilst only one all- α target has more than 50% native-like decoys in its ROSETTA decoy set, five targets surpass this threshold when PCONSC2 contact data is used to restrain the folding procedure. Similarly, the median TM-score of no mixed α - β target decoy set surpasses the TM-score threshold of 0.5 with ROSETTA decoys compared to one for PCONSC2 decoys with three further ones greater than 0.4. All- β targets also benefit from the addition of contact restraints, although decoy sets do not surpass the native-like threshold by median TM-score (Fig. 3.4). Larger targets do not benefit any more than smaller targets from the addition of residue contacts to the structure prediction protocol. The only real exception to this are the decoys for the CheB methylesterase domain (PDB ID: 1chd), for which the majority of ROSETTA decoys are almost random-like whilst PCONSC2 decoys are native-like (Fig. 3.4).

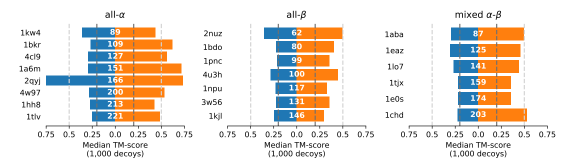


Figure 3.4: TM-score comparison for globular targets separated by fold and ordered by target chain length. Median TM-scores for 1,000 decoys generated with simple ROSETTA (orange) or contact-guided ROSETTA (blue) runs. White numbers in each row correspond to the target chain length. Bars surpassing the dark gray line indicate that the majority of structures are better than random, whilst the light gray line indicates that the majority of structures are native-like [32].

The enhancement of β -structure specific contact pairs is an important part of this study. Previously, the precision of the added BBCONTACTS contact pairs has been demonstrated. The next essential step is to explore how the BBCONTACTS supplement enhances or degrades decoy quality after ROSETTA ab initio protein structure prediction. Given 13 β -structure containing targets, eight targets achieve better overall decoy quality with added BBCONTACTS (Fig. 3.5a). The smallest improvement is observed for target 1e0s with 0.01 TM-score units, whilst the largest for target 1eaz with 0.05 units. The remaining five targets — PDB IDs 1chd, 1bdo, 1npu, 4u3h and 1tjx — saw decreases in median TM-score up to 0.03 when BBCONTACTS contact pairs were added as restraints (Fig. 3.5a). No clear difference between fold classes, i.e. mixed α - β or all- β targets, could be observed, although mixed α - β targets do show slightly greater extremes (Fig. 3.5a).

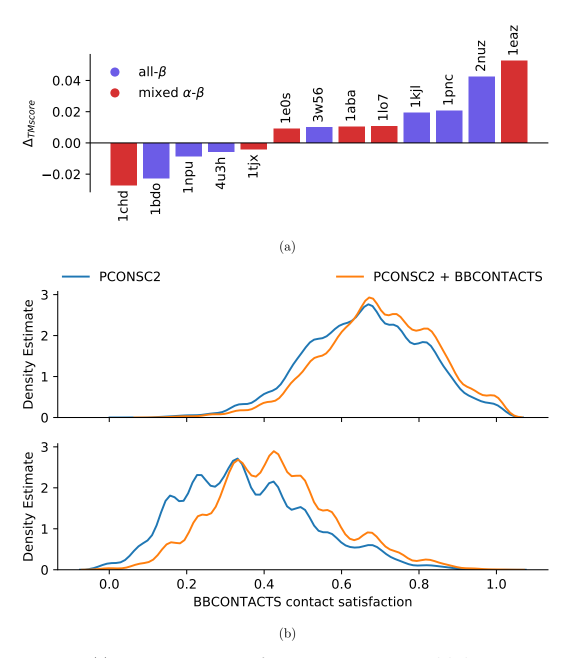


Figure 3.5: (a) TM-score comparison for β -structure containing globular targets separated by fold and ordered by the difference in median TM-score between PCONSC2 and PCONSC2+BBCONTACTS decoys. Positive values indicate a better median TM-score in favour of PCONSC2+BBCONTACTS decoys, whilst negative values those for PCONSC2. PDB IDs are provided alongside each bar. (b) Satisfaction of BBCONTACTS contact pairs in decoys with added β -structure contact restraints (PCONSC2+BBCONTACTS) and those without (PCONSC2).

An analysis of the satisfaction of BBCONTACTS contact pairs in decoys where extra β -sheet contact pairs were used as restraints compared those where not highlights a greater satisfaction in the former. This indicates that the added and upweighted BBCONTACTS β -structure contact restraints enhance the formation of β -sheets in the resulting decoys, which would explain the overall improved decoy quality for more than half the targets outlined previously. A separation of contact satisfaction by upweighted

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and added BBCONTACTS contact pairs indicates that the upweighting has less effect compared to the addition (Fig. 3.5b). Although the former shows a marginal improvement in BBCONTACTS contact satisfaction of decoys without upweighted restraints, the difference is minimal. In comparison, PCONSC2 decoys without the added BB-CONTACTS restraints show less satisfaction for such contacts, indicating that they did not form as often compared to PCONSC2+BBCONTACTS. In combination with the upweighting, this explains that β-rich regions are predicted more accurately when BBCONTACTS contact pairs supplement PCONSC2 contacts.

Transmembrane protein targets were modelled using residue-residue contact predictions derived with CCMPRED, MEMBRAIN and METAPSICOV STAGE1. A ROSETTA benchmark was also run to compare contact-guided decoys to the current norm. Findings in this study highlight the much improved decoy quality for almost all targets when contact information is used to reduce the conformational sampling space (Fig. 3.3). Across all methods, only two targets suffered from the addition of contact restraints during *ab initio* protein structure prediction, namely the domains of ATP synthase C chain (PDB ID: 2wie) and ATP synthase subunit C (PDB ID: 3u2f). In both cases, ROSETTA generates decoys with median TM-score of greater than 0.6 when no contact restraints are used. This contrasts strongly with contact-guided decoy sets, for which only METAPSICOV STAGE1 guide the modelling sufficiently to obtain overall native-like decoys, i.e. median TM-score of greater than 0.5.

A split for decoy quality comparison between no-contact and contact-guided decoy sets by contact prediction algorithm shows that CCMPRED contacts are not sufficiently precise to always improve decoy quality. Six out of 16 targets are predicted more accurately without CCMPRED contact information (Fig. 3.6). In comparison, MEMBRAIN and METAPSICOV STAGE1 contact predictions result in enhanced decoy quality to the extend that only three and two decoy sets are worse than their no-contact counterparts (Fig. 3.6). Most notably, either of the three contact predictions per target performs better for certain targets. The most extreme example may be the decoy sets for Bacteriorhodopsin (PDB ID: 3hap) for which CCMPRED contacts result in decoy quality degradation of 0.06, METAPSICOV STAGE1 in a slight improvement of 0.06 and MEMBRAIN in an improvement of 0.28 TM-score units. This translates into absolute decoy counts with native-like fold — i.e., TM-score > 0.5 of the follwing: 274 for decoys without contact guidance, 289 for CCMPRED contact guidance, 538 for METAPSICOV STAGE1 contact guidance, and 996 for MEMBRAIN contact guidance. Similar examples exist (e.g., PDB IDs 1gu8, 3rlb or 4dve in Fig. 3.6) and highlight that no single method yields the best decoys in all circumstances.

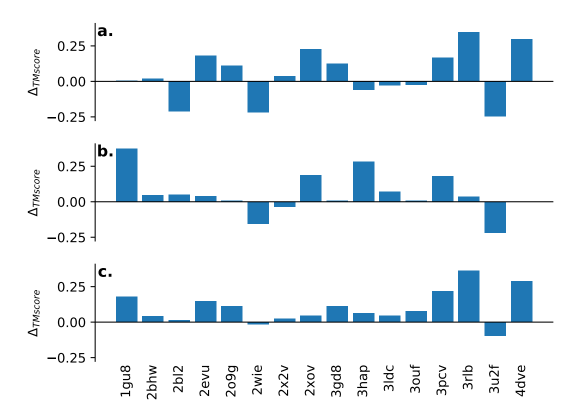


Figure 3.6: TM-score difference between contact-guided and simple ROSETTA decoys for transmembrane protein targets. Positive $\Delta_{TMscore}$ values indicate more accurate contact-guided decoys, whilst negative values better decoys without the addition of contacts. $\Delta_{TMscore}$ values were computed by median TM-score. Contact restraints were obtained with (a.) CCMPRED, (b.) MEMBRAIN, and (c.) METAPSICOV STAGE1.

3.3.3 Molecular Replacement

The most important aspect of this study is the impact of contact-guided decoys in AMPLE-MR. The most limiting factors in AMPLE runs without contact information are target chain length and fold, which typically cannot exceed 150 residues and performs poorly for β -rich folds [12]. Findings presented in previous sections has shown great in overall decoy quality when contact information was used as distance restraints in *ab initio* protein structure prediction. However, it is yet to be shown how such improved decoy quality translates to MR structure solutions.

Structure solutions were attempted for a total of 21 globular targets. Simple ROSETTA decoys — those without contact restraints and AMPLE's current default — resulted in nine structure solutions (Fig. 3.7). The addition of PCONSC2 contact-restraints to the structure prediction procedure improved decoy quality to achieve four additional structure solutions. However, the structure solution of N-terminal region of P67Phox (PDB ID: 1hh8) was lost when PCONSC2-restrained decoys compared to simple ROSETTA ones. The addition of BBCONTACTS distance restraints to up-

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weight and supplement PCONSC2 contacts enabled a further unique solution for the Phosphoinositol (3,4)-bisphosphate PH domain (PDB ID: 1eaz) (Fig. 3.7).

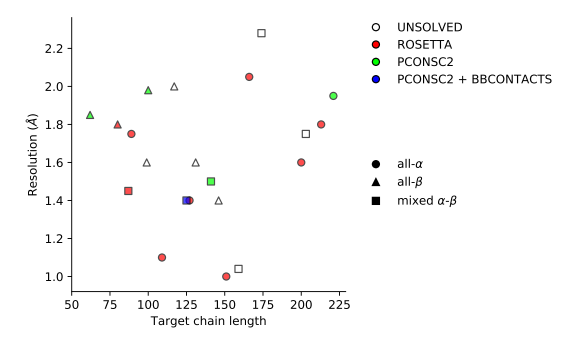


Figure 3.7: Summary of structure solutions obtained with AMPLE using no-contact ROSETTA, PCONSC2-contact-restraint-guided ROSETTA and PCONSC2 + BBCONTACTS-guided ROSETTA decoy sets. Empty markers indicate unsolved targets. Filled markers highlight the minimum decoy set by complexity of the prediction procedure to obtain MR structure solution. Marker shape distinguishes the fold class.

The majority of structure solutions were obtained for all- α targets in the dataset, with a total of eight structure solutions (Fig. 3.7). Seven of those eight structure solutions were achieved with unrestrained ROSETTA decoys, with target chain lengths up to 213 residues. The largest target in the globular dataset, and the only all- α target that required residue contacts, totals 221 residues in target chain length, which exceeds AMPLE's previously benchmarked limits for globular targets greatly [12]. In comparison to all- α targets, β -structure containing proteins require the contact restraints to result in sufficiently accurate decoys for MR. Across all- β and mixed α - β targets, only two structure solutions were obtained with unrestrained ROSETTA decoys. This contrasts to an additional three targets when PCONSC2 restraints were used during *ab initio* structure prediction. Furthermore, the addition of BBCONTACTS contact restraints enabled an additional structure solution, yielding much greater success for β -structure containing protein targets compared to the previous default. Structure solutions for β -containing targets were obtained for target chain lengths up to 141 residues (Fig. 3.7).

In this study, three exceptional cases exemplify the application of contact predictions and their benefit to MR. The first, the structure solutions for 4-hydroxybenzoyl CoA thioesterase (PDB ID: 1lo7) based on AMPLE ensemble search models derived from the PCONSC2 and PCONSC2+BBCONTACTS decoy sets. Without contact re-

straints, AMPLE search models do not accurately represent the target fold (Fig. 3.8a). In comparison, precise residue-residue contacts primarily restraining the large β-sheet yield decoys of sufficient quality to achieve MR structure solution with both PCONSC2 (Fig. 3.8b) and PCONSC2+BBCONTACTS (Fig. 3.8c) decoy sets.

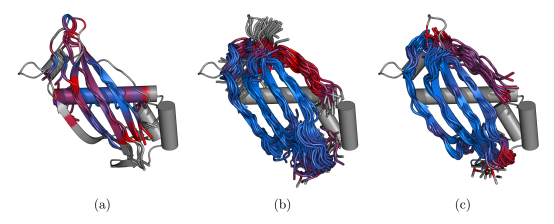


Figure 3.8: Structural superposition of the (a) ROSETTA (C α Root-Mean-Square Deviation (RMSD) 2.814 Å; ensemble contains two structures), (b) PCONSC2 (C α RMSD 1.748 Å; 30 members) and (c) PCONSC2+BBCONTACTS (C α RMSD 1.760 Å; 15 members) search-model ensembles for 4-hydroxybenzoyl CoA thioesterase (PDB ID: 1lo7). Examples are the highest scoring search models based on SHELXE Correlation Coefficient (CC) score, with only (b) and (c) leading to successful MR structure solutions. Search models are shown as tubes and crystal structures as cartoons. (a) and (c) are 50% of the target sequence, while (b) is 55%. The colour scale illustrates the pairwise C α RMSD between each search-model ensemble (represented by its first member) and the crystal structure, with blue representing the minimum C α RMSD and red the maximum. Unaligned residues are coloured grey.

The second exceptional case, PDB ID 1e0s, does not yield any MR structure solution with either decoy set according to the stringent criteria for MR success applied in this study (see ??). However, a Residue-Independent Overlap (RIO) analysis of PHASER solutions, i.e. after MR, indicates that some PCONSC2 and PCONSC2+BBCONTACTS AMPLE search models were placed partially correctly (Fig. 3.9a). For the top PCONSC2 search model, 40% (12 residues) of the search model residues are correctly superimposed, albeit out of register on the target structure (PHASER Translation Function Z-score (TFZ)=4.7, PHASER Log-Likelihood Gain (LLG)=16). For the top PCONSC2+BBCONTACTS search model, 77% (30 residues) of the search model were superimposed in an in-register fashion (PHASER TFZ=5.3, PHASER LLG=17) (Fig. 3.9b). For the latter, expert manual intervention might allow structure determination, but in this case the correct solution was not prominent in the list of MR placements. Nevertheless, it is clear that even when overall structure solution was not automatically achieved the PCONSC2+BBCONTACTS search model provided better results which might be recoverable as successes in the future as MR and post-MR software improves still further.

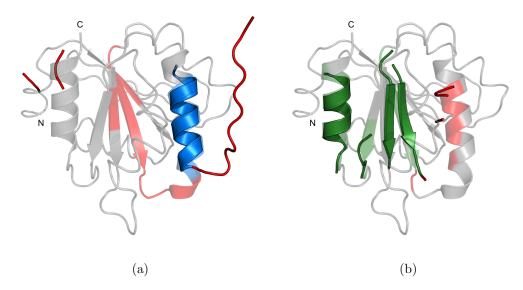


Figure 3.9: Top-PHASER solutions for PDB ID 1e0s based on RIO scores for (a) PCONSC2 (RIO score 12) and (b) PCONSC2+BBCONTACTS (RIO score 30) search models. Search-model colour coding indicates useful superposition of residues by insequence (green) or out-of-sequence register (blue) residues as well as misplaced (red) residues. The addition of BBCONTACTS restraints produced a more accurate model with correctly placed β -strands that was placed correctly. Both structures are shown in cartoon representation with the crystal structure shown as a transparent cartoon. Unaligned reference crystal structure residues are coloured grey.

With much improved decoy quality deriving from the use of predicted residue restraints to guide ab initio structure prediction, the question arises whether AMPLEs existing cluster-and-truncate approach remains the most suitable for obtaining a conserved, native-like core from the decoys found in the largest clusters. For globular targets solved using simple ROSETTA decoys, certain features throughout AMPLEs cluster-and-truncate approach typically correlated with eventual success in structure solution [12]. In general, the greater the number of decoys in the largest cluster the more likely the success was with derived search models. Truncation removed structurally variant parts leading to smaller more accurate ensemble subsets of the cluster decoys. Although successful search models were found at every truncation interval, the majority were derived with search models containing around 30 residues. Lastly, each of the potential nine search models derived at each truncation level (three subclustering radii with three side chain treatments each) can lead to non-redundant structure solutions. Similar observations, particularly with respect to the most successful search model size range were made for other target classes [2, 33] and for ab initio models made with QUARK [34].

A size comparison of the largest clusters of ROSETTA and PCONSC2+BBCONTACTS (or PCONSC2 for all-α) decoys indicated a median increase of 122 decoys per cluster in the latter. All cluster sizes increased except for target 2qyj. More accurate ab initio models are directly linked to larger cluster sizes because of the associated increase in

convergence [35]. Here, as expected, the largest cluster contains better than average quality decoys but the size of the largest cluster does not link to the total number of successful search models.

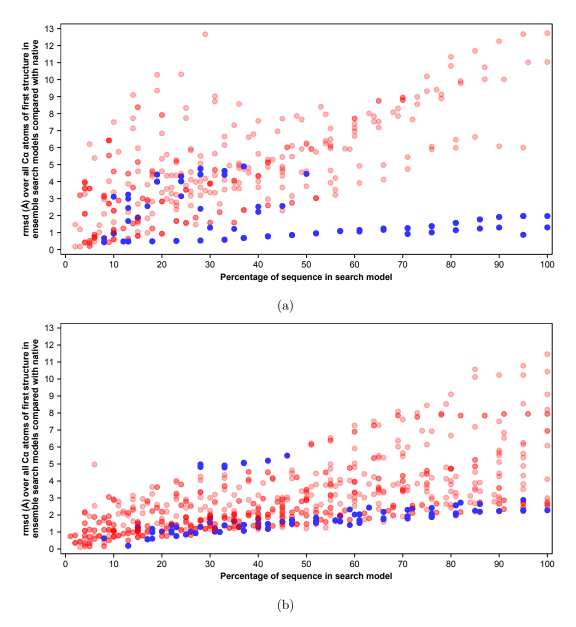


Figure 3.10: The percentage of sequence in the search model is mapped against the RMSD over all $C\alpha$ atoms of the first representative of each search-model ensemble derived from the largest cluster against the native structure. Successful structure solutions of individual search models are highlighted in blue and unsuccessful solutions in red. Progressively darker shades of either colour correspond to increasing numbers of overlapping points. Progressive truncation is shown for (a) ROSETTA and (b) PCONSC2+BBCONTACTS decoys (or PCONSC2 decoys for all- α targets).

In comparison to the clustering step, the progressive truncation of decoys in the largest cluster at 20 different intervals directly affects the number of successful search models. An analysis of the progressive truncation and the effects on search model accuracy revealed that all successful search model ensembles had a $C\alpha$ -RMSD better

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than 5.5Å compared to the native structure (Fig. 3.10). Although the latter cutoff is independent of whether contact information was provided during ab initio modelling, a clear difference between the ROSETTA and PCONSC2+BBCONTACTS (or PCONSC2 for all-α) ensemble search models for all targets can be observed. In total, ROSETTA decoys for all targets produced 1,314 ensemble search models based on the largest clusters. In comparison, PCONSC2+BBCONTACTS decoys generated for the same targets 2,469 search model ensembles from the largest clusters. This increase is the result of a more successful subclustering process due to the increased structural homogeneity across the decoys in the largest cluster. The most notable difference between the two sets is detected for the Small G-protein ARF6-GDP (PDB ID: 1e0s), which produced three ensemble search models based on ROSETTA decoys and 90 based on PCONSC2+BBCONTACTS decoys. Additionally, ensemble search models with structural fragments of 15-40 residues of the target sequence are more likely to succeed in MR phasing than larger or smaller search models [12]. Here we find that the same range is most successful for contact-guided decoys. Out of 246 successful search models for PCONSC2+BBCONTACTS decoys derived from the largest cluster (PCONSC2 for all- α), 101 successful search models contained 15-40 residues. Significantly, some cases like the PH domain of TAPP1 (PDB ID: 1eaz) and the N-terminal bromodomain of human BRD4 (PDB ID: 4cl9) only solved with truncated search models in this size range. Nevertheless, structure solutions were also achieved with larger or smaller search models. The smallest search model leading to a structure solution contained nine residues (8% of total sequence) and solved the Calponin homology (CH) domain from human Beta-spectrin (PDB ID: 1bkr). In comparison, the largest successful search model in terms of residues was found for the designed full consensus ankyrin (PDB ID: 2qyj) domain with 158 residues (95\% of total), and in terms of percentage of the total sequence the untruncated, 62 residue search model for -spectrin SH3 domain (PDB ID: 2nuz) was successful. Therefore, although truncating the ab initio models at different levels remains essential for contact-guided decoys, biasing sampling into the most successful size range may be advantageous in future runs.

The truncated decoys are further processed by subclustering at three different atomic radii, with the resulting subclusters previously found to be similarly successful [12]. Similar trends are seen here: 36% of structure solutions with ROSETTA decoys were achieved with a subclustering radius of 1Å, 36% at a radius of 2Å, and 28% at a radius of 3Å. For PCONSC2+BBCONTACTS (or PCONSC2 for all-α) decoy sets similar numbers were observed (35% at radius of 1Å; 40% at 2Å; 25% at 3Å). Nevertheless, in terms of number of targets solved all three subclustering radii were essential. Largest-cluster decoys for target 1eaz produced a total of 327 search models, but only one solved and this derived from a subclustering radius of 1Å. In comparison, contact-guided decoys from the largest cluster for target 4u3h achieved structure solutions solely with decoys subclustered at 2Å. A single search model with subclustering radius of 3Å solved the target 4cl9 with ROSETTA decoys. The final step in search model creation is the side-chain processing of each subclustered ensemble. Similarly to the subcluster-

ing, no difference was observed between ROSETTA and PCONSC2+BBCONTACTS decoys. For both the polyalanine treatment is most successful, covering 37% of successful search models for ROSETTA decoys and 44% for PCONSC2+BBCONTACTS decoys. For almost all targets, the polyalanine side-chain treatment would be enough to obtain a structure solution. However, some cases, like the target 1eaz, only solve with either or both of the remaining treatments. Thus, relying solely on polyalanine side-chain treatment may limit the overall success rate, although trialling polyalanine ensemble search models first might lead to structure solution faster.

Evaluation of ROSETTA distance-restraint energy functions on contact-guided ab initio structure prediction

Alternative *ab initio* structure prediction algorithms for AMPLE

Decoy subselection using contact information to enhance MR search model creation

Protein fragments as search models in Molecular Replacement

Conclusion & Outlook

Appendix A

Appendix

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Bibliography

- F. Simkovic, J. M. H. H. Thomas, R. M. Keegan, M. D. Winn, O. Mayans, D. J. Rigden, en, *IUCrJ* July 2016, 3, 259–270.
- [2] J. M. H. Thomas, F. Simkovic, R. M. Keegan, O. Mayans, Y. Zhang, D. J. Rigden, C. Zhang, Y. Zhang, D. J. Rigden, Acta Crystallographica Section D Structural Biology Dec. 2017, 73, 985–996.
- [3] D. S. Marks, L. J. Colwell, R. Sheridan, T. A. Hopf, A. Pagnani, R. Zecchina, C. Sander, en, PLoS One Dec. 2011, 6, e28766.
- [4] M. Michel, S. Hayat, M. J. Skwark, C. Sander, D. S. Marks, A. Elofsson, en, Bioinformatics Sept. 2014, 30, i482–8.
- [5] T. Kosciolek, D. T. Jones, en, *PLoS One* Mar. **2014**, *9*, e92197.
- [6] S. Ovchinnikov, L. Kinch, H. Park, Y. Liao, J. Pei, D. E. Kim, H. Kamisetty, N. V. Grishin, D. Baker, en, Elife Sept. 2015, 4, e09248.
- [7] S. Ovchinnikov, D. E. Kim, R. Y.-R. Wang, Y. Liu, F. DiMaio, D. Baker, en, *Proteins* Sept. 2016, 84 Suppl 1, 67–75.
- [8] M. Michel, D. Menéndez Hurtado, K. Uziela, A. Elofsson, Bioinformatics 2017, 33, i23–i29.
- [9] S. H. P. de Oliveira, E. C. Law, J. Shi, C. M. Deane, en, Bioinformatics Nov. 2017, DOI 10.1093/bioinformatics/btx722.
- [10] S. Ovchinnikov, H. Park, N. Varghese, P.-S. Huang, G. A. Pavlopoulos, D. E. Kim, H. Kamisetty, N. C. Kyrpides, D. Baker, en, Science Jan. 2017, 355, 294–298.
- [11] S. Wang, S. Sun, Z. Li, R. Zhang, J. Xu, en, PLoS Comput. Biol. Jan. 2017, 13, e1005324.
- [12] J. Bibby, R. M. Keegan, O. Mayans, M. D. Winn, D. J. Rigden, en, Acta Crystallogr. D Biol. Crystallogr. Dec. 2012, 68, 1622–1631.
- [13] M. J. Skwark, D. Raimondi, M. Michel, A. Elofsson, en, PLoS Comput. Biol. Nov. 2014, 10, e1003889.
- [14] L. S. Johnson, S. R. Eddy, E. Portugaly, en, BMC Bioinformatics Aug. 2010, 11, 431.
- [15] M. Remmert, A. Biegert, A. Hauser, J. Söding, en, Nat. Methods Dec. 2011, 9, 173–175.

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A. Bateman, M. J. Martin, C. O'Donovan, M. Magrane, E. Alpi, R. Antunes, B. Bely, M. Bingley, C. Bonilla, R. Britto, B. Bursteinas, H. Bye-AJee, A. Cowley, A. Da Silva, M. De Giorgi, T. Dogan, F. Fazzini, L. G. Castro, L. Figueira, P. Garmiri, G. Georghiou, D. Gonzalez, E. Hatton-Ellis, W. Li, W. Liu, R. Lopez, J. Luo, Y. Lussi, A. MacDougall, A. Nightingale, B. Palka, K. Pichler, D. Poggioli, S. Pundir, L. Pureza, G. Qi, S. Rosanoff, R. Saidi, T. Sawford, A. Shypitsyna, E. Speretta, E. Turner, N. Tyagi, V. Volynkin, T. Wardell, K. Warner, X. Watkins, R. Zaru, H. Zellner, I. Xenarios, L. Bougueleret, A. Bridge, S. Poux, N. Redaschi, L. Aimo, G. ArgoudPuy, A. Auchincloss, K. Axelsen, P. Bansal, D. Baratin, M. C. Blatter, B. Boeckmann, J. Bolleman, E. Boutet, L. Breuza, C. Casal-Casas, E. De Castro, E. Coudert, B. Cuche, M. Doche, D. Dornevil, S. Duvaud, A. Estreicher, L. Famiglietti, M. Feuermann, E. Gasteiger, S. Gehant, V. Gerritsen, A. Gos, N. Gruaz-Gumowski, U. Hinz, C. Hulo, F. Jungo, G. Keller, V. Lara, P. Lemercier, D. Lieberherr, T. Lombardot, X. Martin, P. Masson, A. Morgat, T. Neto, N. Nouspikel, S. Paesano, I. Pedruzzi, S. Pilbout, M. Pozzato, M. Pruess, C. Rivoire, B. Roechert, M. Schneider, C. Sigrist, K. Sonesson, S. Staehli, A. Stutz, S. Sundaram, M. Tognolli, L. Verbregue, A. L. Veuthey, C. H. Wu, C. N. Arighi, L. Arminski, C. Chen, Y. Chen, J. S. Garavelli, H. Huang, K. Laiho, P. McGarvey, D. A. Natale, K. Ross, C. R. Vinayaka, Q. Wang, Y. Wang, L. S. Yeh, J. Zhang, en, Nucleic Acids Res. Jan. 2017, 45, D158–D169.

- [17] D. T. Jones, D. W. A. Buchan, D. Cozzetto, M. Pontil, Bioinformatics Jan. 2012, 28, 184–190.
- [18] M. Ekeberg, T. Hartonen, E. Aurell, J. Comput. Phys. Nov. 2014, 276, 341–356.
- [19] D. T. Jones, en, J. Mol. Biol. Sept. 1999, 292, 195–202.
- [20] B. Petersen, T. N. Petersen, P. Andersen, M. Nielsen, C. Lundegaard, en, BMC Struct. Biol. July 2009, 9, 51.
- [21] S. Seemayer, M. Gruber, J. Söding, en, *Bioinformatics* Nov. 2014, 30, 3128–3130.
- [22] J. Andreani, J. Söding, en, *Bioinformatics* June 2015, 31, 1729–1737.
- [23] D. T. Jones, T. Singh, T. Kosciolek, S. Tetchner, en, Bioinformatics Apr. 2015, 31, 999–1006.
- [24] L. Kaján, T. A. Hopf, M. Kalaš, D. S. Marks, B. Rost, en, BMC Bioinformatics Mar. 2014, 15, 85.
- [25] J. Yang, R. Jang, Y. Zhang, H. B. Shen, en, Bioinformatics Oct. 2013, 29, 2579–2587.
- [26] C. A. Rohl, C. E. M. Strauss, K. M. S. Misura, D. Baker, en, Methods Enzymol. 2004, 383, 66–93.
- [27] M. D. Winn, C. C. Ballard, K. D. Cowtan, E. J. Dodson, P. Emsley, P. R. Evans, R. M. Keegan, E. B. Krissinel, A. G. W. Leslie, A. McCoy, S. J. McNicholas, G. N. Murshudov, N. S. Pannu, E. A. Potterton, H. R. Powell, R. J. Read, A. Vagin, K. S. Wilson, en, Acta Crystallogr. D Biol. Crystallogr. Apr. 2011, 67, 235–242.
- [28] A. Thorn, G. M. Sheldrick, en, Acta Crystallogr. D Biol. Crystallogr. Nov. 2013, 69, 2251–2256.
- [29] S. X. Cohen, M. Ben Jelloul, F. Long, A. Vagin, P. Knipscheer, J. Lebbink, T. K. Sixma, V. S. Lamzin, G. N. Murshudov, A. Perrakis, en, Acta Crystallogr. D Biol. Crystallogr. Jan. 2007, 64, 49–60.
- [30] G. G. Krivov, M. V. Shapovalov, R. L. Dunbrack, Proteins: Struct. Funct. Bioinf. 2009, 77, 778–795.

Chapter A 37

[31] F. Simkovic, S. Ovchinnikov, D. Baker, D. J. Rigden, *IUCrJ* May 2017, 4, 291–300.

- [32] J. Xu, Y. Zhang, en, Bioinformatics Apr. 2010, 26, 889–895.
- [33] J. M. H. Thomas, R. M. Keegan, J. Bibby, M. D. Winn, O. Mayans, D. J. Rigden, en, *IUCrJ* Mar. 2015, 2, 198–206.
- [34] R. M. Keegan, J. Bibby, J. M. H. Thomas, D. Xu, Y. Zhang, O. Mayans, M. D. Winn, D. J. Rigden, en, Acta Crystallogr. D Biol. Crystallogr. Feb. 2015, 71, 338–343.
- [35] K. T. Simons, C. Kooperberg, E. Huang, D. Baker, J. Mol. Biol. 1997, 268, 209–225.