Protein structure prediction guided by cross-linking restraints — A systematic evaluation of the impact of the cross-linking spacer length

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Recent development of high-resolution mass spectrometry (MS) instruments enables chemical cross-linking (XL) to become a high-throughput method for obtaining structural information about proteins. Restraints derived from XL-MS experiments have been used successfully for structure refinement and protein-protein docking. However, one formidable question is under which circumstances XL-MS data might be sufficient to determine a protein’s tertiary structure \textit{de novo}? Answering this question will not only include understanding the impact of XL-MS data on sampling and scoring within a \textit{de novo} protein structure prediction algorithm, it must also determine an optimal cross-linker type and length for protein structure determination. Whereas a longer cross-linker will yield more restraints, the value of each restraint for protein structure prediction decreases as the restraint is consistent with a larger conformational space.

In this study, the number of cross-links and their discriminative power was systematically analyzed \textit{in silico} on a set of 2055 non-redundant protein folds considering Lys-Lys, Lys-Asp, Lys-Glu, Cys-Cys, and Arg-Arg reactive cross-linkers between 1 Å and 60 Å. Depending on the protein size a heuristic was developed that determines the optimal cross-linker length. Next, simulated restraints of variable length were used to \textit{de novo} predict the tertiary structure of fifteen proteins using the BCL::Fold algorithm. The results demonstrate that a distinct cross-linker length exists for which information content for \textit{de novo} protein structure prediction is maximized. The sampling accuracy improves on average by 1.0 Å and up to 2.2 Å in the most prominent example. XL-MS restraints enable consistently an improved selection of native-like models with an average enrichment of 2.1.
1 Introduction

“Structural Genomics” — the determination of the structure of all human proteins — would have profound impact on biochemical and biomedical research with direct implication to functional annotation, interpretation of mutations, development of small molecule binders, enzyme design, or prediction of protein-protein interaction. Although significant progress towards this goal has been made through X-ray crystallography and nuclear magnetic resonance (NMR) spectroscopy, tertiary structure determination continues to be a challenge for many important human proteins. At present, high-resolution structures exist for about 5% of all human proteins in the Protein Data Bank (PDB). For many uncharacterized human proteins, construction of a comparative model is possible starting from the experimental structure of a related protein. Nevertheless, for about 60% (≈7800) of known protein families in the Pfam database not a single structure is deposited. Many of these proteins will continue to evade high-resolution protein structure determination.

Accordingly, researchers strive to develop alternative approaches. The most extreme approach includes computational methods that predict the tertiary structure of proteins from their sequence alone. Although computational methods are sometimes successful at the predicting the tertiary structure of small proteins with up to one hundred residues, for larger proteins the size of the conformational space to be searched as well as the discrimination of incorrectly folded models hinder structure prediction.

However, recent studies demonstrate that combining de novo protein structure prediction with limited experimental data, i.e. experimental data that alone is insufficient to unambiguously determine the fold of the protein, can yield accurate models for larger proteins. The structural restraints in those studies were acquired using electron paramagnetic resonance (EPR) spectroscopy, electron microscopy (EM), or NMR spectroscopy.

As an alternative technique, XL in combination with MS can be applied to obtain distance restraints, which can be used to guide protein structure prediction. Using bifunctional reagents with a defined length, functional groups within the protein can be covalently bridged in a native-like environment. Thus, it is possible to determine an upper limit for the distance between those residues after enzymatic proteolysis and identification of cross-linked peptides.

This method allows for a fast analysis of protein structures in a native-like environment at a low concentration and can even be applied to high molecular weight proteins or highly flexible proteins. If combined with affinity purification it becomes possible to study proteins inside the cell. Currently, the XL-MS technology is rapidly gaining importance driven by the liquid chromatography (LC)-MS instrument development, the generation of advanced analysis software and the direct integration in protein structure prediction workflows. Furthermore, hundreds of different cross-linking reagents with different spacer lengths, reactivities, and features for specific enrichment and improved detectability are now commercially available.

However, whereas the potential to combine XL-MS and computational modeling has been frequently demonstrated and many technical problems of XL-MS have been solved, several central questions have not yet been evaluated systematically.

(i) Cross-linking reagents are available with a spacer length ranging from 0 Å to more than 35 Å. Whereas longer reagents are likely to provide more distance restraints, shorter cross-links have higher information content in de novo structure prediction as the conformational search space is more restricted. Thus, the question arises, which cross-linker spacer length supports structure prediction best?

(ii) Cross-linking results are often used to confirm already existing structures. However, what is
the average gain in model accuracy and selection of correct models when using cross-linking data in conjunction with de novo protein structure prediction?

(iii) Cross-linking reagents vary in reactivity towards different functional groups present in different amino acids. For de novo protein structure prediction, what is the gain of using additionally cross-linkers with different reactivities?

In this study, we simulated cross-linking experiments on more than 2000 non-redundant protein structures to determine the number of possible and structurally relevant cross-links depending on the size of the protein as well as on the length and reactivity of the applied cross-linking reagents. We then tested the impact of cross-linking restraints on de novo protein structure prediction for fifteen selected proteins.

2 Materials and methods

2.1 Software and databases

A subset of the PDB containing 2055 non-redundant protein structures was downloaded from the PISCES server (Version 08.2012). This PDB subset was created by filtering all available structures with a resolution of at least 1.6 Å, a maximum sequence identity of 20%, and an R-factor cutoff of 0.25. Euclidean distances and shortest solvent accessible surface (SAS) path lengths between Cβ-Cβ, Lys-Nz-Lys-Nz, Lys-Nz-Asp-Cγ, and Lys-Nz-Glu-Cδ, as well as Arg-NH2-Arg-NH2 and Cys-SG-Cys-SG atom pairs with a maximum intramolecular distance of 60 Å were determined through the command line version of Xwalk.

2.2 Generation of sequence dependent distance functions

Tables containing the Euclidean distances and the sequence separation between cross-linking target amino acids (i) Lys-Lys, (ii) Lys-Asp, (iii) Lys-Glu, (iv) Arg-Arg, and (v) Cys-Cys were generated. Amino acid pair distances were sorted into 2.5 Å bins. The total number of observed pairs for each sequence and Euclidean distance was counted. Based on the result an approximation of the distance distribution for every sequence distance was created. The median of the distribution was determined. A logarithmic function was calculated as a regression curve in the form $E_{med} = a \times ln(S) + b$ to correlate the sequence separation $S$ to the median Euclidean distances $E_{med}$.

2.3 Calculation of the amino acid side-chain length

Based on the structure of calmodulin (PDB entry 2KSZ) the average Cβ-Nz, Cβ-Cγ, Cβ-Cδ, Cβ-NH2, and Cβ-SG distances of the side-chains of lysine, aspartic acid, glutamic acid, arginine, and cysteine were determined to be 4.5 Å, 2.3 Å, 3.6 Å, 5.1 Å, and 1.8 Å, respectively.

2.4 Distinguishing impossible, possible and structurally valuable cross-links

Cross-linker spacer lengths between 1 Å and 60 Å distances were evaluated and classified in either (i) impossible cross-links, meaning that the distance between the Cβ-atoms of the cross-linked amino acids exceeds the sum of the spacer lengths and the side-chain lengths, or (ii) possible cross-links, meaning that the Cβ-Cβ distance is below the sum of the spacer lengths and side-chain lengths. The latter group was subdivided into cross-links potentially useful for structure
2.5 Estimation of optimal spacer lengths for a given protein molecular weight

Over all proteins in each molecular weight (MW) bin, the total number of possible distance pairs (#possible) as well as the number of distance pairs useful for structure determination (#valuable) were computed for each cross-linker spacer length. Furthermore, the maximum number of valuable cross-links observed for all spacer lengths (#valuable_max) was determined. For each MW bin the ratios (#valuable/#possible) and (#valuable/#valuable_max) were plotted as a function of the cross-linker spacer length. The optimal cross-linker length for each MW bin was approximated as intersection points of the two functions using a local regression (Figure 2 on page 6). The estimated values for the optimal cross-linker spacer length were plotted as a function of the MW and were fitted using a cubic regression curve. The script used for the calculation is available at http://www.ufz.de/index.php?en=19910.

2.6 Simulation of cross-linking restraints

Seventeen proteins with known tertiary structure determined via X-ray crystallography (resolution of <1.9 Å) were selected from the data set of structures as test cases to evaluate the influence of cross-linking restraints on de novo protein structure prediction. To thoroughly benchmark the
Table 1: Proteins used for the cross-link spacer length benchmark. The fifteen proteins for the benchmark set were selected from high-resolution structures deposited in the PDB with varying content of lysines. The structures were selected to cover a wide range of the structural features sequence length as well as percentage of residues within α-helices and β-strands.

| Protein | Uniprot | resolution | weight | length | Lysine | α-helix | β-strand |
|---------|---------|------------|--------|--------|--------|---------|---------|
| 1HRC    | P00004  | 1.9 Å      | 12 368 Da | 105    | 18 %   | 40 %    | 1 %     |
| 3IV4    | Q7A6S3  | 1.5 Å      | 13 235 Da | 112    | 6 %    | 49 %    | 25 %    |
| 1BGF    | P42228  | 1.5 Å      | 14 504 Da | 124    | 5 %    | 79 %    | 1 %     |
| 1T3Y    | Q14019  | 1.2 Å      | 15 835 Da | 141    | 9 %    | 35 %    | 29 %    |
| 3M1X    | C4LXT9  | 1.2 Å      | 15 882 Da | 138    | 7 %    | 25 %    | 28 %    |
| 1X91    | Q9LNF2  | 1.5 Å      | 16 419 Da | 153    | 7 %    | 76 %    | 0 %     |
| 1JL1    | P0AY74  | 1.3 Å      | 17 483 Da | 155    | 7 %    | 34 %    | 30 %    |
| 1MBO    | P02185  | 1.6 Å      | 17 980 Da | 153    | 12 %   | 77 %    | 0 %     |
| 2QNL    | Q11XA0  | 1.5 Å      | 19 218 Da | 162    | 5 %    | 70 %    | 2 %     |
| 2AP3    | Q8NX77  | 1.6 Å      | 23 190 Da | 199    | 23 %   | 81 %    | 0 %     |
| 1J77    | Q9RGD9  | 1.5 Å      | 24 226 Da | 209    | 8 %    | 62 %    | 1 %     |
| 1ES9    | Q29460  | 1.3 Å      | 25 876 Da | 232    | 3 %    | 41 %    | 11 %    |
| 3B5O    | D0VWS1  | 1.4 Å      | 27 506 Da | 244    | 3 %    | 71 %    | 0 %     |
| 1QX0    | P0A2Y6  | 2.3 Å      | 32 821 Da | 293    | 7 %    | 38 %    | 20 %    |
| 2XM    | Q15257  | 1.5 Å      | 34 798 Da | 303    | 7 %    | 60 %    | 3 %     |
| FGF2    | P09038  | 1.5 Å      | 17 859 Da | 145    | 10 %   | 9 %     | 34 %    |
| P11     | P60903  | 2.0 Å      | 11 071 Da | 95     | 13 %   | 63 %    | 3 %     |

algorithm, the benchmark set covers a wide range of protein topologies and structural features. The sequence lengths of the proteins range from 105 to 303 residues, the number of secondary structure elements (SSEs) ranges from 5 to 19 with varying α-helical and β-strand content (Table 1). For these proteins, all solvent accessible surface Cβ-Cβ distances between target amino acids in the structure which were within the range of either homobifunctional Lys-reactive cross-linkers or heterobifunctional Lys-Asp/Glu reactive cross-linkers were determined through Xwalk. For the predicted optimal cross-linker length (read above) and spacer lengths of 2.5 Å, 7.5 Å, 17.5 Å and 30.0 Å lists of structurally possible cross-links were generated.

For the two proteins horse heart cytochrome c (PDB entry 1HRC) and oxymyoglobin (PDB entry 1MBO) restraints were also derived from published cross-linking MS experiments deposited in the XL database. Experimental cross-linking data of FGF2 (PDB entry 1FGA) and P11 (PDB entry 4HRE) were derived from Young et al. and Schulz et al. respectively.

2.7 Translating cross-linking data into structural restraints

Explicitly rebuilding coordinates for a cross-link is comparable to solving the loop closure problem. During de novo, protein structure prediction the cross-link would have to be reconstructed each time the conformation of the protein changes. In a typical Monte Carlo (MC) simulation with a maximum of 12 000 MC steps per model and 5000 models for each protein this would result in a maximum number of 60 million attempts to build the cross-link, which is too resource demanding for usage in de novo protein structure prediction. Therefore, we developed a fast approach to estimate the chance that a particular model fulfills a XL-MS restraint. The surface path of a cross-link is approximated by laying a sphere around the protein structure and computing the arc length between the cross-linked residues (Figure on page 20). The geometrical center of the protein structure is used as the center of the sphere. If takeoff and landing point have different distances to the center of the sphere, the longer distance is used as
the radius. During the protein structure prediction process, the side-chains of the residues are not modeled explicitly but represented on a simplified way through a 'super atom'. While this simplification vastly reduces the computational demand of the algorithm, it also adds additional uncertainty due to the unknown side-chain conformations. The agreement of the model with the cross-linking data is quantified by comparing the distance between the cross-linker lengths \((l_{XS} + l_{SS1} + l_{SS2})\) with the computed arc lengths \((d_{arc})\), with \(-1\) being the best agreement and \(0\) being the worst agreement. To account for the uncertainty of side-chain conformations a cosine-transition region of \(7\ \text{Å}\) was introduced (Figure 6 on page 20).

2.8 Structure prediction protocol for the benchmark set

The protein structure prediction protocol is based on the BCL::Fold protocol for soluble proteins. In a preparatory step, the glsplsse are predicted using the SSE prediction methods PsiPred and Jufo9D and an SSE pool is created. Subsequently a Monte Carlo Metropolis (MCM) energy minimization algorithm draws random glsplsse from the predicted SSE pool and places them in the three-dimensional space (Figure 3 on the following page). Random transformations like translation, rotation or shuffling of glsplsse are applied. After each MC step the energy of the resulting model is evaluated using knowledge-based potentials which, among others, evaluate the packing of glsplsse, exposure of residues, radius of gyration, pairwise amino acid interactions, loop closure geometry and amino acid clashes. Based on the energy difference to the previous step and the simulated temperature a Metropolis criterion decides whether to accept or reject the most recent change.

The protein structure prediction protocol is broken into multiple stages, which differ regarding the granularity of the transformations applied, and the emphasis of different scoring terms. The first five stages apply large structural perturbations, which can alter the topology of the protein. Each of the five stages lasts for a maximum of 2000 MC steps. If an energetically improved structure has not been generated within the previous 400 MC steps, the stage terminates. Over the course of the five assembly stages, the weight of clashing penalties in the total score is ramped up as 0, 125, 250, 375 and 500.

The five protein assembly stages are followed by a stage of structural refinement. This stage lasts for a maximum number of 2000 MC steps and terminates if no energetically improved model is sampled for 400 MC steps in a row. Unlike the assembly stages, the refinement stage only consists of small structural perturbations, which will not drastically alter the topology of the protein model.

Through multiple prediction runs with different score weights, the optimal contribution of the cross-linking score to the total score was determined to be 40\% to 50\%. Consequently, the
weight for the scoring term evaluating the agreement of the model with the cross-linking data was set to 300 over all six stages, which ensures that the cross-linking score contributes between 40% and 50% to the total score.

2.9 De novo folding simulations without and with cross-linking restraints

To evaluate the influence of cross-linking restraints on protein structure prediction accuracy, each protein was folded in the absence and in the presence of Lys-Lys, Lys-Glu, and Lys-Asp cross-linking restraints. Independent structure prediction experiments were performed for the predicted optimal as well as two shorter and two longer cross-linker spacer lengths each of the five spacer lengths (Table 2 on page 21). Additionally, predictions were performed using combination of all spacer lengths as well as using restraints obtained by the optimal spacer length of all three cross-linker reactivities. For the two proteins of which experimentally determined cross-linking data were available, protein structure prediction was additionally performed for the experimentally determined restraints. For each protein and cross-linker length used, 5000 models were sampled in independent MCM trajectories. Due to the randomness of the employed MC algorithm, ten sets of 5000 models were sampled for each protein without restraints. Improvements in prediction accuracy can be compared to the standard deviations to identify statistically significant improvements (Table 3 on page 21).

2.10 Metrics for comparing calculating model accuracy and enrichment

The prediction results were evaluated using the protein-size-normalized root-mean-square-deviation (RMSD100) and enrichment metrics. The RMSD100 metric was used to quantify the sampling accuracy by computing the normalized root-mean-square-deviation between the backbone atoms of the superimposed model and native structure. The enrichment metric was used to quantify the discrimination power of the scoring function by computing which percentage of the most accurate models can be selected by the scoring function. The enrichment metric is used to assess the influence of the cross-linking restraints to discriminate among the sampled models. First, the models of a given set $S$ are sorted by their RMSD100 relative to the native structure. The 10% of the models in $S$ with the lowest RMSD100 are assigned to subset $P$ (positives) and the remaining 90% of the models are assigned to subset $N$ (negatives). Second, the models in $S$ are sorted by their BCL score. The 10% of the models in $S$ with the best score are assigned to subset $FS$ (favorable score). The intersection $TP = FS \cap P$ contains the
most accurate models which the scoring function can select (true positives). The enrichment 
\[ e = \frac{\text{TP}}{\text{FP}} \times \frac{\text{TP} + \text{TN}}{\text{FP} + \text{TN}} \]
of the most accurate models the scoring function can select. In order to reduce the influence of the sampling accuracy on the enrichment values, the positive models are considered the 10% of the models with the lowest RMSD100 and \( \frac{\text{TP} + \text{TN}}{\text{FP} + \text{TN}} \) is fixed at a value of 10.0. Therefore, the enrichment ranges from 0.0 to 10.0, with a score of 1.0 indicating random selection and a value above 1.0 indicating that the scoring function enriches for native-like models.

3 Results

3.1 Creation of an in silico cross-linking database

We performed in silico cross-linking experiments on 2055 non-redundant proteins. Covering an MW range from 1.4 kDa to 139 kDa, 59% of the proteins have an MW below 25 kDa. For each of those proteins all Lys-Lys, Lys-Asp, and Lys-Glu sequence and Euclidean distances as well as the SAS distance between the \( C_\beta \)-atoms were determined. Thus, the resulting database contained information on 391902 Lys-Lys, 395815 Lys-Glu, and 360101 Lys-Asp pairs which built the basis for the determination of the number of possible cross-links, cross-links useful for structure prediction, and finally for the prediction of the optimal cross-linker length for studying a selected protein (Figure 3 on the previous page).

3.2 Estimation of the possible cross-links per protein

Next we estimated how many and which of the distances could be cross-linked with a cross-linker of a given length and specificity. We considered cross-links possible if the sum of the spacer length and the length of the two connected side-chains (\( C_\beta-C_\beta \), Lys-Nz-Lys-Nz, Lys-Nz-Asp-C\( _\gamma \), or Lys-Nz-Glu-C\( _\delta \)) is longer than the \( C_\beta-C_\beta \)-SAS-distance between the amino acids. As the lengths of the side-chains of Lys (\( C_\beta-Nz \)), Asp (\( C_\beta-Oz \)), and Glu (\( C_\beta-Oz \)) 4.5 Å, 2.4 Å, and 3.6 Å were used, which were determined as average values from the crystal structure of calmodulin (PDB entry 1CLL). In silico cross-linking experiments were conducted for all of the 2055 proteins using homobifunctional Lys-Lys-reactive, as well as heterobifunctional (Lys-Asp- and Lys-Glu-reactive) cross-linking reagents with lengths from 1 Å to 60 Å (step size 1 Å).

To draw conclusions from the correlation of this in silico cross-linking experiments to the MW of the studied proteins the proteins were grouped into 45 bins with a step size of 2.5 kDa. For example, a protein with a MW in the range of 25 kDa to 27.5 kDa contains on average 15.1 Lys, 14.4 Asp, and 16.7 Glu. On average, 182 Lys-Lys, 173 Lys-Glu, and 144 Lys-Asp cross-links exist per protein within this specific MW bin. Theoretically, all of those could be cross-linked with a cross-linker of 60 Å. In contrast by utilization of cross-linkers of 13 Å (as e.g. BS3) only about 33% of the cross-links are formed in silico. When going to a cross-linker of length of 1 Å (e.g. close to EDC), only 10% of all possible amino acid pairs are linked.

3.3 Estimation of structurally relevant cross-links

In protein structure prediction approaches, the enrichment of low RMSD structures among thousands of generated models is crucial. Therefore, we hypothesized that restraints that are valuable for structure prediction will reduce the conformational search space substantially. For the present study, we classify a cross-linking restraint as useful for structure prediction if it discriminate at least 50% of all possible conformations. Thus, in a second step each of the
possible cross-links was evaluated in terms of its potential to discriminate at least 50% of incorrect structures (useful for structure determination) or whether the cross-linked amino acids are so close in sequence that it can be derived from sequence separation that the distance can be bridged by the cross-linker independently of the protein’s structure (not useful for structure determination).

In order to develop a stringent measure for usefulness we did not simply assume the maximum distance that can be bridged by an amino acid chain of a certain length. Instead, the Euclidean distance distributions for Lys-Lys, Lys-Glu, and Lys-Asp were computed for the sequence separations ranging from 1 to 60 amino acids within our database of protein structures. For example, in the more than 2000 analyzed structures there are 3132 Lys-Lys pairs, which are separated by ten amino acids. For this sequence distance Euclidean distances bins of 2.5 Å were defined in which the occurrences of residue pairs were counted. The pairs were present in bins ranging from 2.5 Å to 35.0 Å. As the median distance, we found 15.5 Å. For the same sequence distance the distribution of Lys-Glu (3336 pairs) and Lys-Asp (3010 pairs) are quite similar and the median values were 15.6 Å and 15.3 Å.

Similarly, for sequence separations of 15 amino acids we observed 3024 Lys-Lys pairs, 3200 Lys-Glu pairs, and 2835 Lys-Asp pairs. The median values are 20.8 Å, 20.9 Å, and 20.7 Å, respectively. For sequence separations of 60 amino acids, we observed 2167 Lys-Lys pairs, 2212 Lys-Glu pairs, and 2167 Lys-Asp pairs. The median values are 23.0 Å, 23.0 Å, and 23.0 Å, respectively (Figure 1 on page 4).

Approximating the proteins structures as spheres, we applied a logarithmic model to fit the relationship between the sequence separation S and the median Euclidean distance \( E_{\text{med}} \). We find (i) \( E_{\text{Lys-Lys}} = 5.46 \times \ln(S_{\text{Lys-Lys}}) + 2.2 \), (ii) \( E_{\text{Lys-Glu}} = 5.37 \times \ln(S_{\text{Lys-Glu}}) + 2.36 \), and (iii) \( E_{\text{Lys-Asp}} = 5.19 \times \ln(S_{\text{Lys-Asp}}) + 2.36 \) for Lys-Lys, Lys-Glu, and Lys-Asp distances, respectively.

Secondly, using our derived functions constituting the \( S/E \) relationships, we considered every cross-link as of reasonable discriminative power, i.e. which fulfills the criterion that the sum of the cross-linker spacer length and the average length of both contributing side-chains is shorter than the median of the sequence/Euclidean-distance distribution. If we examine the 25 kDa MW bins of Lys-Lys targets with a 1 Å spacer cross-link 1167 of the possible 22398 target pairs fulfilled this criterion and were considered as of sufficient discriminative power (Figure 7 on page 22). These cross-links, which represent 4% of all Lys-Lys distances we defined therefore as useful for protein structure prediction. Application of a 13 Å spacer length results in 2935 valuable target pairs (12% of all Lys-Lys distances, see Figure 7 on page 22). In contrast, a cross-linker with a spacer length of 60 Å would allow to cross-link all distances. However, none of the cross-links would have discriminative power for native-like models (Figure 7 on page 22). For the proteins of the 25 kDa MW bins the number of valuable cross-links as a function of the cross-linker length has a log-normal character never exceeding a roughly 25 Å spacer. The intermediate length of 13 Å resulted in an almost equal contribution of valuable and structurally invaluable cross-linking pairs. Whereas 29% of all possible reactive amino acid pairs are linked, 12% are considered valuable according for structure prediction (Figure 7 on page 22).

### 3.4 Prediction of molecular weight dependent optimal cross-linker spacer lengths

Whereas usage of a short cross-linker will result in only a few but mostly structurally valuable restraints, a longer cross-linker will yield more restraints but a lower ratio of valuable restraints. Furthermore, the ratio of valuable restraints as well as the number of possible restraints depends on the size of the protein. In agreement with prior studies regarding structural modeling driven
by sparse distance restraints, we hypothesize that a compromise between maximizing the portion of valuable cross-links compared to all cross-links which can be formed with a given cross-linker length ($\frac{\text{#valuable}}{\text{#possible}}$) and maximizing the relative number of achievable cross-links with any spacer length ($\frac{\text{#valuable}}{\text{#possible}_{\text{max}}}$) might yield the best results.

Following our hypothesis, for each MW bin we derived the optimal spacer length as the intersection point of the two functions as it is shown exemplarily for MW 25 kDa in figure 2 on page 6.

The derived optimal spacer lengths for Lys-Lys, Lys-Asp, and Lys-Glu were plotted as function of the MW (Figure 4 on the following page). The relationship was fitted using a cube root function. For our observable MW sample space for Lys-Lys cross-links, all spacer lengths reached dimensions between 5.0 Å and 18.6 Å. No optimal spacer length was further than 2.5 Å separated from the regression curve. The average distance from the modeled spacer lengths was 0.7 Å. The MW term as well as the side-chain term has been modeled as an exponential fraction in respect to the relation between volume and distances in spherical objects.

Additionally, the optimal spacer lengths were also predicted for homobifunctional arginine and for homobifunctional cysteine cross-linking reagents analogously to the procedure being described for the homo- and heterobifunctional lysine-containing cross-links. Consistently, the optimal spacer lengths depend on the MW as well as the lengths of the cross-linked side-chains SS1 and SS2 and could be calculated by $l_{opt}[\text{Å}] = k \times \sqrt[3]{\text{MW}} + \sqrt[3]{\text{SS1} + \text{SS2}}$. $k$ was determined to be 0.32, 0.31, 0.34, 0.34 and 0.35 for Lys-Lys, Lys-Asp, Lys-Glu, Arg-Arg, and Cys-Cys, respectively.

3.5 Generation of in silico and experimental cross-linking data for testing the effect of different spacer length for de novo modeling

To evaluate the effect of cross-linking data derived from experiments with different spacer length we folded seventeen proteins de novo with BCL::fold (Figure 3 on page 7). Thirteen proteins were part of our data set while for four proteins experimental cross-linking data were available (1MBO, 1HRC, 1FGA, and 4HRE) (Table 1 on page 5). All proteins have a MW in the range from 13 kDa to 27 kDa. Most structures were mainly α-helical with fewer β-strand SSEs. The β-strand content ranged from 0% to 51%. The α-helical content ranges from 2% to 81%. The highest β-strand content showed 1LMI with also the fewest α-helices. The portion of lysines was between 3% and 23%, which resulted in minimal 4 and maximal 46 lysine residues per structure. For the two structures 1MBO and 1HRC, which were studied experimentally, we used the published experimental data, which were obtained using DSG, DSS/BS3, and DEST. For 1MBO, there were 8 cross-links in total with the 11.4 Å reagent BS3 four of them confirmed with the 7.7 Å reagent DSG. For 1HRC, 48 cross-links were reported. 9 DSS, 31 BS3, 6 DSG, and 9 with DEST (11 Å). Six cross-links had been identified with different cross-linking reagents. 18 BS3 cross-links were published for 1FGA, whereas 3 intramolecular BS3 cross-links were available for 1HRE. For the thirteen proteins as well as for 1MBO and 1HRC, we predicted all cross-links, which are possible with the predicted optimal cross-linker length as well as with two shorter and two longer cross-linking reagents (Table 2 on page 21) and used these data as restraints during modeling (Figure 6 on page 20).

3.6 Cross-linking restraints improve the sampling accuracy of de novo protein structure prediction
XL-MS data provides structural restraints that reduce the sampling space in *de novo* structure determination. Thereby a fraction of incorrect conformations is excluded and the sampling density in all other areas of the conformational space is increased. To evaluate the power of cross-linking restraints to guide *de novo* protein structure determination we computed the RMSD100 values of the most accurate models for each protein for structure prediction with and without cross-linking restraints. Using cross-linking restraints not only increases the frequency with which accurate models are sampled, but the best models achieve an accuracy not sampled in the absence of cross-linking data (Table 3 on page 21). Across all benchmark proteins, the accuracy of the best models was, on average, 6.6 Å when no cross-linking data was used. By using cross-linking, data for the spacer length deemed optimal the average RMSD100 value was improved to 5.6 Å, which corresponds to two standard deviations. By using restraints obtained for all five spacer lengths, the average accuracy of the best model improved to 5.2 Å. For the proteins 1XQ0, 2IXM, and 3B50, even with cross-linking data, it was not possible to sample a native-like conformation. We attribute this to limitations in the sampling algorithm resulting in the native conformation not being part of the sampling space. For other proteins, significant improvements could be observed. While the accuracy of the best models for 1ES9 and 1J77 was 7.3 Å and 6.8 Å, cross-linking restraints improved the accuracy to 5.7 Å and 4.5 Å, respectively. For 1MBO, the accuracy could be improved from 7.1 Å to 4.2 Å by using a combination of Lys-Glu/Asp reactive cross-linkers (Figure 8 on page 23).

### 3.7 Cross-linking restraints improve the discriminative power of the scoring function

The ability of the scoring function to identify the most accurate models among the sampled ones was quantified using the enrichment metric (see section 2 on page 3). Enrichments were computed for proteins predicted without cross-linking data, for each spacer length and for all spacer lengths combined. For protein structure prediction without cross-linking restraints an average enrichment of 1.1 was measured, which is barely better than random selection. The scoring function has almost no discriminative power. Using cross-linking restraints yielded by the
optimal spacer length improved the enrichment to 2.1 (Table 3 on page 21), which corresponds to three standard deviations. Using all five spacer lengths to obtain additional restraints further improves the enrichment to 2.4. The most significant improvement could be observed for 1J77, for which the enrichment could be improved from 0.5 to 2.4.

3.8 The cross-linker length determines improvements in sampling accuracy and discrimination power

The length of the cross-linker determines the number of obtainable restraints as well as their information content. While a longer cross-linker is able to form more cross-links and therefore yields a larger number of restraints, the longer cross-linker length can be fulfilled by a larger number of conformations, reducing the discriminative power of the restraint. In order to assess the influence of the cross-linker length, and therefore the number of restraints and restraint distances, on the sampling accuracy and discrimination power, the protein structure prediction protocol was conducted with restraints derived from different cross-linker lengths.

The cross-linker lengths were separated into five groups: optimal, which is the predicted optimal cross-linker length, short1 and short2, which are shorter cross-linker lengths, and long1 and long2, which are longer cross-linker lengths. The cross-linker length predicted to be optimal yielded the most useful restraints for protein structure prediction in terms of sampling accuracy and discriminative power. Across all proteins the average RMSD100 values of the most accurate models for the optimal cross-linker length were 5.6 Å — an improvement by 15% — while...
they were 6.3 Å, 6.2 Å, 5.9 Å and 6.1 Å — improvements by 5%, 6%, 11% and 8% — for the shorter and longer cross-linker lengths, respectively (Figure 5 on the previous page). The longest cross-linkers have a less significant impact on the sampling accuracy due to their ambiguity, whereas the shortest cross-linkers failed to yield a sufficient number of distance restraints to impact prediction accuracy. The discriminative power, quantified through the enrichment metric, for the optimal cross-linker length was 2.1, whereas it was 1.4, 1.5, 1.9 and 1.7 for the shorter and longer cross-linkers, respectively (Figure 5 on the preceding page). For the proteins 1X91 and 3M1X, the optimal cross-linker length did not yield any cross-links with a sequence separation of at least ten and therefore did not provide relevant structural information. In those cases protein structure prediction with longer cross-linker lengths provided better results. By combining restraints obtained for all five cross-linker lengths, the average enrichment value could be improved to 2.4.

3.9 Combination of cross-linkers with different reactivities results in improvements larger than seen when varying the spacer lengths

In order to obtain valuable restraints for de novo protein structure prediction a maximum number of SSE pairs needs to be cross-linked. The availability of Lys-Asp/Glu reactive cross-linkers allows for a better sequence coverage and therefore a wider coverage of SSE pairs. Cross-links with different spacer lengths were simulated for the proteins in the benchmark set using Xwalk. To assess the influence of Lys-Asp/Glu reactive cross-linkers on protein structure prediction the same protocol was applied as for the Lys-Lys reactive cross-linkers. For the Lys-Glu reactive cross-linkers a prediction accuracy of 5.7 Å and enrichment of 2.2 on average could be achieved, which is comparable to the results for the Lys-Lys reactive cross-linkers.

While Lys-Asp reactive cross-linkers also achieve improvements in prediction accuracy and enrichment when compared to protein structure prediction without restraints, the results are slightly worse than for Lys-Lys reactive cross-linkers with an average prediction accuracy of 6.0 Å versus 5.6 Å and an average enrichment of 1.7 versus 2.1 (Table 3 on page 21). To a large part, the difference in the overall results is caused by the results for the proteins 1ES9, 1T3Y, and 3M1X for which Lys-Asp reactive cross-linkers failed to yield restraints between SSE pairs and therefore failed to reduce the conformational space significantly. Besides deviations regarding the average improvements over all proteins, the spacer length deemed optimal also provides the best results for Lys-Asp/Glu reactive cross-linking. Combining the restraints yielded for the optimal spacer lengths with Lys-Lys/Asp/Glu reactive cross-links improves the sampling average sampling accuracy to 5.1 Å and the average enrichment to 2.6. Combining the restraints yielded by all spacer lengths and cross-linker reactivities failed to further improve prediction results.

4 Discussion

4.1 Prediction of the optimal cross-linker spacer length

It has been shown frequently that chemical cross-linking data can be used to guide de novo structure prediction and selection of native-like models. Surely, the sensitivity, the broad applicability to almost all proteins, the nearly physiological experimental condition during the chemical cross-linking reaction, and the potential of automation are the main advantages for using XL-MS to generate such restraints. However, the small number and high uncertainty of
restraints from chemical cross-links limit impact on de novo proteins structure prediction, in particular when compared to more data rich methods such as NMR spectroscopy. One major limitation is the fact that distances between the functional groups in long and flexible amino acid side-chains are measured. Therefore, a significant uncertainty is added to the cross-linker length when converting XL-MS data into Cβ-Cβ restraints. A second challenge of chemical cross-links is that only the maximum distance is restricted, but no information is obtained on the minimum distance or the favored distance distribution. In result, even a “zero length” cross-linker restricts the Cβ-Cβ distance to the sum of the lengths of the two connected side-chains (e.g. for Lys-Lys cross-links 9.1 Å).

In most of the cross-linking experiments, lysine residues are targeted. Lysines are excellent targets because of their overrepresentation on protein surfaces and the clean chemistry of amine modification. Nevertheless, their frequency is on average only about 7%. As a consequence the number of cross-links, which can be formed for example in a 25 kDa protein with a standard homobifunctional Lys-Lys-reactive cross-linking reagents with a spacer length of 6.4 Å (length of DST) are in the range of about 20. Only a small fraction of these restraints will substantially limit the conformational space for the protein. This number is usually too small to restrict the conformational space to an unambiguous single protein fold. To increase the number of restraints it is possible to use cross-linkers with longer spacer length or target amino acids such as Asp, Glu, Tyr, Ser, Thr, Arg, or Cys.

Restraints obtained with longer cross-linking reagents are less restrictive to the conformational space. To evaluate the value of cross-links for protein structure prediction we determined for each sequence distance (0 to 60 amino acids) how long a cross-linker has to be to link the target amino acids. For example two lysines, which are separated by eight amino acids in sequence were found to be linkable in all 3488 cases by a homobifunctional cross-linker with a length of >30 Å (as it is in BS(PEG)9). In our study, we stated the hypothesis that it would be desirable if two target amino acids can only be linked in 50% of all models created meaning that 50% of all structures could be discarded based on a single cross-link. For example, for two lysines separated by 10 amino acids this would be the case for cross-linker lengths of 14.8 Å (distance distributions for other amino acids distances are shown in figure [ ] on page [ ]). Cross-links, which could only be formed in less than 50% for the corresponding sequence distance, were considered as being valuable. Based on this definition for all 2055 structures of the applied non-redundant protein structure database the optimal spacer length was calculated. With this optimal spacer length, the number of structural valuable cross-links has been maximized taking into account that in general for modeling approaches few distance restraints of highly discriminative character are less favorable than a higher number with a smaller discriminative power.

Since the optimal cross-linker length should depend on the protein size in a cubic root fashion to convert volume into distance, it is not unexpected that this was also observed for the dependency on the MW (Figure [ ] on page [ ]). However, one has to keep in mind that the formula might not be applicable to non-globular proteins and multi-domain proteins. However, in case of multidomain proteins this formula should be applicable to the separated domains. Remarkably, based on our simulation for proteins with MWs of 10 kDa, 25 kDa, 50 kDa and 100 kDa the recommended spacer lengths are 9.0 Å, 11.5 Å, 13.9 Å and 17.0 Å, respectively, which is quite close to the homobifunctional amine-reactive commercially available cross-linkers DSG (7.7 Å), BS3 (11.4 Å), and EGS (16.1 Å), which are currently the preferred choice to study small (<20 kDa), medium (20 kDa to 50 kDa), and large proteins (>50 kDa), respectively.

Addressing different functional groups is a second approach to increase the total number of distance restraints. The consequence is that the cross-linking reaction is either less effective or specific (Asp, Glu, Tyr, Ser, Thr) creating challenges in data interpretation or the target amino
acids are less frequent (Arg and Cys) limiting the number of restraints observed. However, using the same theoretical approach revealed that optimal spacer length for heterobifunctional Lys-Asp and Lys-Glu cross-linker (Figure 8 on page 23) as well as homobifunctional Cys-Cys and Arg-Arg cross-linker can be predicted with the same equation: 

\[ l_{opt}[\text{Å}] = k \times \sqrt[3]{MW} + \sqrt[3]{SS1} + SS2 \]

with \( k \approx \frac{1}{3} \) in which \( SS1 \) and \( SS2 \) are the average lengths of the cross-linked side-chains.

Comparing the two approaches to increase the number of valuable cross-links, it should be pointed out that using several cross-linking reagents with different reactivities results in significantly higher improvement of the model quality than using only lysine reactive cross-linking reagent but with different spacer length.

4.2 Challenges in using cross-linking data to guide \textit{de novo} modeling

To test whether the cross-linker with the predicted optimal spacer length indeed perform best in modeling we have chosen a \textit{de novo} structure prediction approach BCL::Fold for testing. Even though comparative modeling using known protein structures as a template usually performs better then \textit{de novo} modeling, our goal was to maximize impact of XL-MS restraints.

A major limiting factor for \textit{de novo} protein structure prediction methods is the vast size of the conformational space. Cross-linking restraints can aid the computational prediction of a protein’s tertiary structure by drastically reducing the size of the sampling space. Cross-linking experiments yield a maximum Euclidean distance between the cross-linked residues, which increases the sampling density in the relevant part of the conformational space.

A major limitation of using cross-linking restraints to guide protein structure prediction when compared to restraints obtained from EPR and NMR spectroscopy is that the cross-linker length cannot be directly translated into a Euclidean distance between the cross-linked residues. While cross-link prediction and evaluation methods like Xwalk\cite{22} are successful at predicting if a certain cross-link can be formed in a given structure, explicit modeling approaches are computationally too expensive for usage in a rapid scoring function required for protein structure prediction. Approximations, such as the great circle on a sphere presented here, are fast to compute but associated with increased uncertainties. Most of the cross-linkers used can cover a long Euclidean distance and therefore the yielded restraints can be fulfilled by a wide variety of conformations. In spite of this, cross-linking restraints display some potential in limiting the size of the sampling space, resulting in a higher density of accurate models. Further, the geometrical restraints derived from XL-MS allow for the discrimination of a significant fraction of models representing incorrect topologies and therefore improve the discriminative power of the scoring function.

4.3 Abilities and limitations of protein structure prediction from limited experimental data

We showed that incorporation of cross-linking data into a \textit{de novo} protein structure prediction method improves the accuracy of the structure prediction. The two major challenges of \textit{de novo} predictions are the sampling of structures as well as the discrimination of inaccurate structures. In this study reduction of the conformational space was achieved through the assembly of predicted SSEs with limited flexibility and the incorporation of geometrical restraints derived from cross-linking data. The discrimination of inaccurate models is performed through a scoring function which approximates the free energy. Assuming that the native structure is in the global energy minimum, complete sampling and an accurate methods to measure free energy would lead to the correct identification of the native conformation. However, the conformational space is too large to be extensively sampled and the free energy needs to be approximated,
which results in ambiguity regarding the model which is most similar to the native structure. Incorporating cross-linking data provides geometrical restraints, which can be used as additional criteria to discriminate inaccurate models. While an average sampling accuracy of 5.1 Å, when using restraints yielded by XL-MS, is a significant improvement over the 6.6 Å, when not using cross-linking data at all, only for four proteins it was possible to sample models with an RMSD100 of less than 4 Å when compared to the crystal structure. Cross-linking data yields an upper boundary for the Euclidean distance of the cross-linked residues, which allows for the placement of the second residue within a sphere of volume \( \frac{4}{3} \pi r^3 \) around the first residue. Depending on the cross-link distribution, topologically different models can fulfill the same restraint set. Discrimination among those models is impossible with XL-MS restraints.

4.4 Comparison of experimental and in silico cross-links

In order to draw general conclusions based on the analysis of hundreds of different structures this study relies mainly on virtual cross-linking experiments. Unfortunately, although extensive XL-MS data sets have been published for several proteins, it proved difficult to obtain suitable experimental data sets for the present benchmark due to additional requirements: (i) the protein must be monomeric and small enough for de novo protein folding with BCL::Fold, (ii) an experimental atomic detail structure for comparison, and (iii) a large data set of intramolecular cross-links must be available. Results for the four cases P11, FGF2, cytochrome c, and oxymyoglobin that came closest are reported to demonstrate our efforts to work not only with simulated data. However, for P11 and FGF2 using experimentally determined restraints did not improve the prediction results in a statistically significant way. For P11, only three restraints were available with a maximum sequence separation of nine residues. Because of the small sequence separation, these restraints contain very limited structural information and no improvement in de novo folding can be expected. The tertiary structure of FGF2 contains twelve β-strands with several β-strands that are strongly bent. This protein is too large for de novo structure determination with BCL::Fold. As BCL::Fold is unable to sample the conformation of the protein in the first place, no significant improvement was expected or observed when XL-MS data were added. Nevertheless, the value of the predicted cross-links in comparison to experimental cross-links could be validated with the two proteins cytochrome c and oxymyoglobin for which experimental cross-links had been published in the XL database. For cytochrome c (PDB entry 1HRC), we indeed found that the cross-linker with predicted optimal spacer length of 10.2 Å performed best. However, for oxymyoglobin (PDB entry 1MBO) the longer spacers improved the accuracy slightly more than the cross-linker with the optimal spacer length. Interestingly, on the one hand for both proteins several cross-links, which should be possible, could not be detected, which might be due to experimental or analytical reasons. On the other hand, also several cross-links, which were experimentally identified which were not predicted. An examination of these data revealed that most of these cross-links are not present in the virtual data set because their \( C_\beta-C_\beta \) distances exceed the expected maximum length. This finding is in agreement with Merkley et al., who evaluated protein structures by molecular dynamics and reported that usually a high number of experimental approved cross-links exceed the theoretical maximal spatial distance due to structure flexibility. It was concluded for the investigation of Lys-Lys distances using a BS3/DSS cross-linking reagent an upper bound of 26 Å to 30 Å for Cα-atoms.

On the other hand, spacer conformations usually adapt lengths that are somehow distributed between their minimal and maximal lengths. In line it was also reported that many spacers in commercially available cross-link agents preferable adopt conformations, which are significantly
below the cited maximal spacer length. Thus, ideally cross-linking results should be evaluated based on experimentally derived or simulated ensembles of in-solution structures instead of using X-ray structures as reference. However, to address all degrees of flexibility during the de novo structure prediction is currently too resource intensive. Furthermore, there are many additional practical challenges, which may prevent the formation or identification of cross-links, and thus may result in more meaningful results using a cross-linker with a non-optimal length. Nevertheless, for both structures the sampling accuracies could also be improved by 0.7 Å based on the experimentally determined restraints, which is only slightly worse than the improvement of 1.0 Å observed based on in silico cross-links.

5 Conclusion

Recent development of high-resolution MS instruments enables the analysis of proteins not accessible to NMR spectroscopy and X-ray crystallography. Data obtained from those experiments can be translated into structural restraints to guide protein structure prediction. The information content of a geometrical restraint obtained from XL-MS experiments is directly dependent on the used spacer length. Thus, the choice of the spacer length is an important step.

Firstly, for amino acids pairs close in sequence only minimum structural information is obtained if the spacer is too long. Here we determine the optimal spacer length to gain structural information on lysines with a sequence separation of $S$, we estimated a length as $E = 5.5 \times \ln(S) + 2.2$. Secondly, we demonstrate that for de novo protein structure prediction the optimal spacer length depends on the MW of the protein of interest and the length of the cross-linked side-chains ($SS1$ and $SS2$) and can be predicted as $l_{opt} = k \times \sqrt{MW} + \sqrt{SS1 + SS2}$, with $k \approx \frac{1}{3}$.

We also demonstrate that restraints obtained from cross-linking experiments contribute moderately to solving the major challenges of de novo protein structure prediction — the vast size of the conformational space and discrimination of inaccurate models. Using restraints from cross-linking experiments significantly increases the sampling density of native-like models and contribute to the discrimination of incorrect models. By combining cross-linking restraints with knowledge-based scoring functions, the average accuracy of the sampled models could be improved by up to 2.2 Å and the average enrichment of accurate models could be improved from 11% to 24%.

Conclusively, we believe this study can help in the planing of XL-MS experiments as well as to evaluate how much information can be gained by XL-MS experiments and the ambiguity that remains.

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Parts of the data analysis were performed using the R package. The renderings of the models were created using Chimera. Part of the text was generated using the R package.
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7 Supplementary data

Figure 6: Implicit translation from cross-linking data into structural restraints. Explicit simulation of the cross-linker conformation is computationally expensive and prohibitive for use in a rapid scoring function required for protein structure prediction. Instead, the cross-linker conformation and the path crossed by the cross-linker were approximated through computing the arc length connecting the two cross-linked residues (A). The agreement of a model with cross-linking data was evaluated by computing the difference between the arc length ($d_{arc}$) and the cross-linker length ($d_{xl}$). The agreement of the model with the cross-linking data is quantified with a score between $-1$ and $0$, with $-1$ being the best agreement and $0$ being the worst agreement (B).
Table 2: Lys-Lys cross-links yielded by different spacer lengths. Cross-links obtained for the benchmark proteins. Simulated and experimentally determined cross-links were obtained for the fifteen benchmark proteins. For each protein, an optimal spacer length was determined (optimal). Additional cross-links were simulated for two shorter (short1 and short2) and two longer (long1 and long2) spacer lengths. The number of yielded cross-links (§ rest) is shown for each spacer length. For the two proteins 1HRC and 1MBO, experimentally determined cross-links were published.

| Protein | Protein | optimal | short1 | short2 | long1 | long2 |
|---------|---------|---------|--------|--------|-------|-------|
| 1HRC    | 10.2 Å  | 13 Å    | 2.5 Å  | 0 Å    | 7.5 Å | 17.5 Å | 27 Å | 30 Å |
| 3IV4    | 10.4 Å  | 5 Å     | 2.5 Å  | 2 Å    | 7.5 Å | 17.5 Å | 7 Å  | 30 Å |
| 1BGF    | 10.7 Å  | 6 Å     | 2.5 Å  | 3 Å    | 7.5 Å | 17.5 Å | 10 Å | 30 Å |
| 1T3Y    | 10.9 Å  | 35 Å    | 2.5 Å  | 9 Å    | 7.5 Å | 20 Å   | 17.5 Å | 42 Å | 30 Å |
| 3M1X    | 10.9 Å  | 1 Å     | 2.5 Å  | 0 Å    | 7.5 Å | 0 Å    | 17.5 Å | 5 Å  | 30 Å |
| 1X91    | 11.0 Å  | 2 Å     | 2.5 Å  | 0 Å    | 7.5 Å | 1 Å    | 17.5 Å | 8 Å  | 30 Å |
| 1JL1    | 11.2 Å  | 7 Å     | 2.5 Å  | 0 Å    | 7.5 Å | 3 Å    | 17.5 Å | 11 Å | 30 Å |
| 1MBO    | 11.3 Å  | 9 Å     | 2.5 Å  | 0 Å    | 7.5 Å | 3 Å    | 17.5 Å | 23 Å | 30 Å |
| 2QNL    | 11.5 Å  | 6 Å     | 2.5 Å  | 4 Å    | 7.5 Å | 14 Å   | 15 Å | 30 Å |
| 2AP3    | 12.1 Å  | 53 Å    | 2.5 Å  | 0 Å    | 7.5 Å | 19 Å   | 136 Å | 30 Å | 427 |
| 1J77    | 12.2 Å  | 29 Å    | 2.5 Å  | 7 Å    | 17.5 Å | 16 Å  | 36 Å | 30 Å |
| 1ES9    | 12.5 Å  | 8 Å     | 7.5 Å  | 0 Å    | 17.5 Å | 1 Å   | 37.5 Å | 17 Å | 45 Å |
| 3B5O    | 12.7 Å  | 15 Å    | 7.5 Å  | 2 Å    | 17.5 Å | 8 Å   | 37.5 Å | 21 Å | 45 Å |
| 1XQ0    | 13.3 Å  | 9 Å     | 7.5 Å  | 0 Å    | 17.5 Å | 4 Å   | 37.5 Å | 14 Å | 45 Å |
| 2IXM    | 13.5 Å  | 41 Å    | 7.5 Å  | 20 Å   | 17.5 Å | 41 Å  | 37.5 Å | 49 Å | 45 Å |

Table 3: Protein structure prediction results for different cross-linker reactivities. Comparison between structure prediction results with and without cross-linking restraints. By using geometrical restraints obtained from cross-linking experiments, the size of the sampling space can be reduced resulting in an improved sampling accuracy. This is shown by significant improvements in the RMSD100 value of the most accurate model (best). Furthermore, cross-linking restraints provide geometrical information, which improves the discrimination power of the scoring function, leading to an improvement in the enrichment (e). Without restraints, ten independent prediction trajectories were conducted and the standard deviation of the accuracy of the best model (σ_{best}) and the enrichment (σ_e) are reported.
Figure 7: Lys-Lys pair distributions. Distribution of all possible and valuable Lys-Lys pairs for a 25 kDa to 27.5 kDa weight bin. Gray bars show all theoretical pairs in their specific distance cluster of ± 2.5 Å. Red bars show pairs that could be connected in respect to their surface distance by a specific cross-link (here 1 Å, 13 Å and 60 Å) always including the side-chain contribution to the overall length. Green bars show pairs that are considered valuable by our proposed scoring function. Pie charts show the accumulated number of cross-links for every spacer length.
Figure 8: Selected prediction results from cross-linking data. Most accurate models sampled with and without using cross-linking restraints. The RMSD100 values of the most accurate models sampled for 1X91, 1J77, and 1MBO were 4.8 Å, 6.8 Å and 7.1 Å. By using restraints yielded by Lys-Lys/Asp/Glu reactive cross-linkers, the accuracy could be improved to 2.7 Å, 5.0 Å and 4.2 Å. Shown are the native structures of 1X91, 1J77, and 1MBO (A, D, G), the most accurate models sampled without cross-linking restraints (B, E, H), and the most accurate models sampled with cross-linking restraints (C, F, I). Selected restraints are shown that are not fulfilled in the model predicted without cross-linking data (red bars), but that are fulfilled in the model predicted with cross-linking data (black bars).