

Recent advances on the interval distance geometry problem

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- Abstract We discuss a discretization-based solution approach for a classic problem in global optimization, namely the distance geometry problem (DGP). We focus our attention on a par-2 ticular class of the DGP which is concerned with the identification of the conformation of з biological molecules. Among the many relevant ideas for the discretization of the DGP in Δ the literature, we identify the most promising ones and address their inherent limitations 5 to application to this class of problems. The result is an improved method for estimating 6 3D structures of small proteins based only on the knowledge of some distance restraints 7 between pairs of atoms. We present computational results showcasing the usefulness of the 8 new proposed approach. Proteins act on living cells according to their geometric and chem-9 ical properties: finding protein conformations can be very useful within the pharmaceutical 10 industry in order to synthesize new drugs. 11
- 12 Keywords Distance geometry · Discretization · Molecular conformation

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13 1 Introduction

Given a positive integer K and a simple weighted undirected graph G = (V, E, d), where d maps edges $\{u, v\} \in E$ to positive interval weights $[\underline{d}(\{u, v\}), \overline{d}(\{u, v\})]$, the Distance Geometry Problem (DGP) [38] is the problem of finding a realization of the graph G in a K-dimensional Euclidean space. In other words, the DGP requires the identification of a map $x : V \to \mathbb{R}^K$, satisfying the distance constraints:

$$\underline{d}(\{u,v\}) \le \|x(u) - x(v)\| \le \bar{d}(\{u,v\}), \quad \forall \{u,v\} \in E,$$
(1)

where $\|\cdot\|$ denotes the Euclidean norm.

A solution for (1) is called a *realization* or an *embedding*. In order to simplify the notation, we will use $x_u := x(u)$ and $d_{uv} := d(u, v) := d(\{u, v\})$ hereafter.

In structural biology, the problem of identifying molecular conformations from a given list of distance restraints between atom pairs is a DGP in dimension K = 3. This problem is also known in the scientific literature as the Molecular Distance Geometry Problem (MDGP). In this particular application, the distances may be exact (i.e. $\underline{d}_{uv} = \overline{d}_{uv}$) or represented by a positive real-valued interval (i.e. $\overline{d}_{uv} > \underline{d}_{uv} > 0$).

Exact distances are related to the chemical bonds whereas interval ones can be provided
by experimental techniques. Such techniques include Nuclear Magnetic Resonance (NMR)
[3], Förster resonance energy transfer (FRET) [7] and mass spectrometry (MS) cross-linking
[10].

The DGP is NP-hard [56] and there exist several approaches to this problem (see [38, 52] and Sect. 1.1), where the DGP is reformulated as a global optimization problem on a continuous search domain, whose objective function is generally a penalty function of the distance constraints. More recently, a discrete approach to the DGP was proposed [39], where the continuous domain of the optimization problem is transformed into a discrete domain.

37 **1.1 Literature review**

Distance Geometry (DG) has played a prominent part in Global Optimization (GO) inso-38 far as it has important applications to science (e.g. protein conformation) and engineering 39 (localization of sensor networks, structural rigidity, control of unmanned underwater vehi-40 cles and robotic arms), and it is naturally cast as a system of nonconvex constraints (Eq. (1)) 41 in terms of continuous decision variables. In general, DGPs are reformulated as a mini-42 mization of constraint violations. Such reformulations have the property that the optimal 43 objective function value is zero for feasible instances, and strictly positive for infeasi-44 ble ones. Various approaches have been proposed in this journal for the general case 45 [15,16,26,27,34,37,47,64,68], and many others on the application to protein conforma-46 tion [11,17,21,43,46,54]. In this paper we focus on the case where the input graph is rigid, 47 which implies that the search process has an inherently combinatorial side. 48

⁴⁹ Over the years, the solution to MDGPs (DGPs arising in structural biology) have been ⁵⁰ typically attempted by employing tools such as ARIA [42], CYANA [23] and UNIO [22], ⁵¹ which are all based on the Simulated Annealing (SA) meta-heuristic [28].

52 While molecular conformations are generally obtained by the above methods and succes-

sively stored in databases such as the Protein Data Bank (PDB) [4], a second class of methods

⁵⁴ based on Nonlinear Programming (NLP) solution techniques has emerged in the last decades.
 ⁵⁵ A well-known example is the DGSOL algorithm [48], which employs a homotopy method

⁵⁵ A well-known example is the DGSOL algorithm [48], which employs a homotopy method ⁵⁶ based on locally solving progressively finer Gaussian smoothings of the original problem.

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A third class of methods is based on Euclidean Distance Matrix Completion [1, 14]. This is 57 the case for the EMBED algorithm [12], which aims to fill in the missing distance bounds by 58 constraint propagation of triangle and tetrangle inequalities. Thereafter, a candidate distance 59 matrix (named dissimilarity matrix) is sampled from the completed interval distance matrix, 60 and atom coordinates are obtained by matrix decomposition [13, 58]. Since the dissimilarity 61 matrix is not guaranteed to be a Euclidean Distance matrix, some of the original constraints 62 might be violated. The last phase therefore consists in minimizing the constraint violation 63 by local minimization, using the obtained embedding as an initial point. 64

A fourth class is centered around the so-called Build-Up algorithm [15,64]. These methods 65 are based on the ancient idea of triangulation, used by humankind ever since navigation 66 existed. In the context of distance geometry, where a point position is determined by the 67 distances to it rather than the angles they subtend, this is known as "trilateration". Build-Up 68 algorithms in dimension K = 3 attempt to place an unknown point x_i (for some i < n) by 69 identifying at least four other points with known positions, and having known distances to 70 x_i . When dealing with proteins and experimental data, the assumption of having four known 71 exact distances to any given point may be excessively strong [44]. We point out, however, 72 that some variants of the Build-Up algorithm overcome this limitation. For example, in order 73 to address the uncertainty of the given distance values, the extension presented in [60] takes 74 into account atomic coordinates and an unknown radius representing the uncertainty. Another 75 variant [65] partly addresses the requirement of unknown vertices having at least four adjacent 76 vertices with known positions. This variant can find multiple valid realizations, but appears 77 to lack the ability to finding *all* possible incongruent realizations. 78

A fifth and very important class of methods is based on solving a Semidefinite Programming (SDP) relaxation of the DGP [6,29,45]. In particular, [29] exploits the cliques in the graph to reduce the size of the SDP formulation (also see [2]). This method was shown to be able to solve NMR instances containing real data and to reconstruct conformation models that are very close to the ones available on the PDB.

The authors of this paper are among the researchers who proposed and worked on a sixth 84 class of methods based on a combinatorial algorithm called Branch & Prune (BP) [36]. 85 Protein graphs share some common properties: for example, they can be decomposed into a 86 backbone subgraph and many side chains subgraphs [57] (these can be realized separately 87 and then put together [55]). The backbone subgraph is larger than the subgraphs related to 88 side chains, and hence most difficult to realize. However, it also defines an order on the atoms 89 with certain topological properties, which we formally discuss below (informally, we can say 90 that every atom in this order has at least three predecessors which are also adjacent in the 91 graph structure). Under this assumption, the search domain of the underlying optimization 92 problem can be reduced to a discrete set with a tree structure [32,51], which can be searched 93 by the BP algorithm [36]. If the distances are exact, BP can find all realizations of a given 94 protein backbone graph. Although an exhaustive search in the conformation tree is worst 95 case exponential [32], numerical experiments have shown that BP behaves polynomially in 96 protein-like instances [40]. In fact, it can be proved that, in such cases, the problem is Fixed 97 Parameter Tractable (FPT) [41]. In computational experiments, the parameter value could 98 always be fixed at a single constant, which explains the polytime behaviour. For protein 99 backbone instances with exact distances, BP is one of the fastest available methods, one of 100 the most reliable, and the only one which can certifiably find all incongruent realizations. 101

An adaptation of the BP to the interval distance setting was proposed in [34], where intervals are replaced with a finite set of discrete points. We refer to this BP adaptation as the *interval* BP (*i*BP). This algorithm was tested on real protein instances in [8]. Although this BP variant shows promise, its practical applicability is currently limited by the choice
 of discretization points.

Several other approaches for solving DGPs can be found in the scientific literature. The
 interested reader can refer to [33,38,52].

1.2 Aim of this paper

Our main motivation in this work is to improve the *i*BP algorithm proposed in [34] for solving MDGPs with interval data. For this purpose, we identify the main limitations of this discrete approach in presence of interval distances and propose a new variant of *i*BP to find approximate solutions for interval MDGPs.

The identification of the barriers against the successful application of iBP in real settings represents an important step towards a combinatorial methodology with the following properties:

- it is specifically suitable for solving the protein conformation problem from distance
 restraint data;
- it can work with uncertain data, specified as interval distances provided by experimental
 techniques;
- 121 it can potentially find all incongruent realizations of a given instance.

The paper is organized as follows. In Sect. 2, we define the subclass of DGP instances 122 describing protein backbone graphs: we discuss assumptions, discretization orders, the iBP123 algorithm variant, pruning devices, and the parameterization of the coordinates. Section 3 124 presents a method, based on some interval distance constraints, which is able to reduce the 125 set of candidate positions for certain vertices before the *i*BP branching phase. Section 4 126 addresses the main limitations in handling larger molecules in presence of interval data and 127 presents a heuristic for finding approximate realizations. The computational results in Sect. 5 128 illustrate the improvements due to the proposed approaches. 129

130 2 A combinatorial approach

Let G = (V, E, d) be a simple weighted undirected graph representing an instance of the MDGP. In the following, vertices of V will represent atoms of the given molecule and $\{u, v\} \in$ *E* if the distance between the atoms *u* and *v* is available. The map *d* relates each edge $\{u, v\} \in$ *E* to a positive interval weight $[\underline{d}(u, v), \overline{d}(u, v)]$. The MDGP asks to find a realization $x : V \to \mathbb{R}^3$ (see Introduction), i.e. a molecular conformation in three-dimensional space such that:

$$\underline{d}(u,v) \le \|x_u - x_v\| \le \overline{d}(u,v), \quad \forall \{u,v\} \in E.$$

$$(2)$$

Recall that $\underline{d}(u, v)$ and $\overline{d}(u, v)$ denote, respectively, the lower and upper bounds for the distance d(u, v) (with $\underline{d}(u, v) = \overline{d}(u, v)$ if d(u, v) is exact). We also suppose that the given set of distances is realizable in \mathbb{R}^3 .

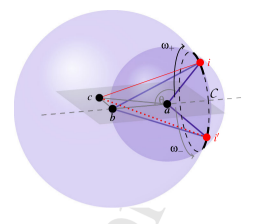
In order to discretize the search domain, MDGP instances need to satisfy some particular assumptions. The main requirement is that the atoms need to be sorted in a way that there are at least three *reference atoms* for each of them (aside, obviously, from the first three). We say that an atom u is a reference for another atom v when u precedes v in the given atomic order, and the distance d(u, v) is known. In such a case, candidate positions for v belong to the sphere centered in u and having radius d(u, v). When the *reference distance* d(u, v) is given through an interval, the sphere becomes a spherical shell, namely, the region between

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Fig. 1 The two feasible arcs obtained by intersecting *two* spheres and one spherical shell



an inner sphere of radius d and an outer sphere of radius \overline{d} with the same center. If three 148 reference atoms are available for v, then candidate positions (for v) belong to the intersection 149 of three spherical shells. The easiest situation is the one where the three available distances 150 are exact and the intersection gives, in general, two possible positions for v [32]. However, 151 if only one of the three distances is allowed to take values into a certain interval, then the 152 intersection gives two arcs of a circle, generally disjoint, where sample points can be chosen 153 [34]. In both last situations, the discretization can be performed. More details are given in 154 the next section. 155

156 **2.1** *i*BP algorithm and discretization orders

Let G = (V, E, d) be an instance of the MDGP and let us suppose that there exist an order for the atoms $v \in V$, so that we can assign a numerical label $i \in \{1, 2, ..., |V|\}$ to each of them. At each recursive call of the *i*BP algorithm, candidate positions for the current atom *i* are computed using the positions of the previously placed reference atoms and their distances to the atom *i*.

When the distances between *i* and its references are exact, the intersection of three spheres needs to be computed. If the reference atoms $\{a, b, c\}$ are not collinear, then such an intersection results in at most two points. When this situation is verified for all atoms i > 3, then the search domain has the structure of a binary tree [32].

However, if one of the three reference distances, say d_{ci} , is an interval, then the two spheres centered at x_a and x_b need to be intersected with a spherical shell centered at x_c . As a result, the intersection gives two candidate arcs (see Fig. 1). These arcs are over the dashed circle C defined by the intersection of the two spheres. When the intersection consists of two arcs, a finite number D of sample positions should be selected from each of them [34]. This way, we still have a discrete set of possible positions for the atom i.

Therefore, the discretization strongly depends on an *order* for the vertices (atoms) of *G* satisfying specific properties. Definition 1 formalizes the assumptions mentioned above.

174 **Definition 1** The *interval* Discretizable DGP in dimension 3 (*i*DDGP₃)

Given a simple weighted undirected graph G = (V, E, d), where $E' \subset E$ is the subset of

edges for which their weights are exact distances, we say that G represents an instance of the

 $iDDGP_3$ if there exists a total order on the vertices of V verifying the following conditions:

(a) $G_C = (V_C, E_C) \equiv G[\{1, 2, 3\}]$ is a clique and $E_C \subset E'$;

(b) $\forall i \in \{4, \dots, |V|\}$, there exists $\{a, b, c\}$ such that

Algorithm 1 The *i* BP algorithm.

1: iBP(i, n, d, D)2: **if** (i > n) **then** 3. *// one solution is found* 4: print current conformation; 5: else 6. // coordinate computation 7: if $(d_{ci}$ is an interval) then 8: compute the two candidate arcs and add them to the list L 9: else 10: compute the two candidate positions and add them to the list L 11: end if 12: for j = 1, ..., |L| do 13: if (L(j) is an arc) then 14: take D samples from the arc; set N = D; 15: else 16: set N = 1; 17: end if 18: // verifying the feasibility of the computed positions 19: for k = 1, ..., N do if $(x_i^{j,k}$ is feasible) then 20: $i \dot{B} P(i + 1, n, d, D);$ 21. 22: end if 23. end for 24: end for 25: end if

- 180 1. a < i, b < i, c < i;
- 181 2. $\{\{b, i\}, \{c, i\}\} \subset E' \text{ and } \{a, i\} \in E;$

182 3. $\Delta_S(a, b, c) > 0$,

where $\Delta_S(a, b, c)$ stands for the area of the triangle formed by $\{a, b, c\}$. Assumption (a) allows us to place the first 3 atoms uniquely and fixes the realization with respect to rotation and translations. Assumptions (b.1) ensures the existence of three reference atoms for every i > 3, and assumption (b.2) ensures that at most one of the three reference distances may be represented by an interval. Finally, assumption (b.3) requires that the area $\Delta_S(a, b, c)$ is strictly positive, which prevents the references from being collinear. Under these assumptions, the MDGP can be discretized.

Algorithm 1 is a sketch of the *i*BP algorithm for solving *i*DDGP₃ instances. In the algorithm call, *i* is the current atom for which the candidate positions are searched, *n* is the total number of atoms forming the considered molecule, *d* is the list of available distances (exact or interval distances), and *D* is the discretization factor, i.e. the number of sample points that are taken from the arcs in case the distance d_{ci} is represented by an interval. In the algorithm (see lines 8 and 10), we make use of a list *L* of positions and arcs, from which candidate positions are extracted.

Given an order for the vertices in V satisfying the assumptions in Definition 1, the algo-197 rithm calls itself recursively in order to explore the tree of candidate positions. Every time 198 a new atomic position is computed, it defines a new branch of the tree. This phase in iBP199 is named *branching phase*. For every computed atomic position, its feasibility is verified by 200 checking the constraints (2), up to the current tree layer, or other additional feasibility criteria 201 based on properties of the molecule, e.g., van der Waals' separation distance (VdW), chirality 202 constraints, and others [8, 53]. This phase in *i*BP is named *pruning phase*, and the criteria 203 are called *pruning devices* (see line 20 of Algorithm 1). 204

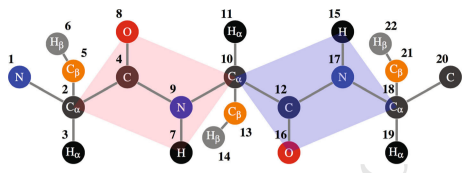


Fig. 2 A model for the protein backbone and a possible discretization order

Even if the tree grows exponentially (in the worst case scenario), the pruning devices allow *i*BP to focus the search on the feasible parts of the tree. The easiest and most efficient pruning device is the Direct Distance Feasibility (DDF) criterion [32], which consists in verifying the ε -feasibility of constraints involving distances between the current atom *i* and previously placed atoms:

 $\underline{d}(h,i) - \epsilon \le ||x_h - x_i|| \le \overline{d}(h,i) + \epsilon, \quad \forall \{h,i\} \in E, \text{ with } h < i \text{ and } h \notin \{a,b,c\}.$ (3)

²¹¹ The distances involved in the above constraints are called *pruning distances*.

212 2.2 Protein backbone model: discretization orders and pruning devices

A necessary preprocessing step for solving MDGPs by this discrete approach consists in 213 finding suitable atomic orders allowing each atom v to have at least three reference atoms. 214 We name such orders *discretization orders* [9]. In previous works, discretization orders have 215 been either handcrafted [34], or obtained by looking for paths on *pseudo* de Bruijn graphs 216 consisting of cliques of G [50], or even automatically detected by a greedy algorithm [31,49]. 217 In fact, if we consider distances defined by bond lengths and bond angles as exact, along with 218 the peptide plane geometry, it is possible to find orders for the protein backbone (and also for 219 side chains [11]) satisfying the assumptions required for the discretization. This preprocessing 220 step can be performed efficiently, in polynomial time [49], so that the necessary assumptions 221 can be fulfilled by graphs related to proteins. However, we point out that when some additional 222 assumptions are imposed to the searched orders, such as the *consecutivity* of the reference 223 vertices, this problem becomes NP-hard [9]. In this work, we consider a model for the protein 224 backbone as depicted in Fig. 2. 225

With the backbone atoms N, C_{α} , and C, we also considered the attached H and H_{α} , and we have included only the C_{β} and H_{β} atoms to represent the side chain of each amino acid. There are two exceptions for this amino acid model: the glycine, where C_{β} and H_{β} are replaced by one H, and the proline, which has a missing hydrogen. The backbone model in Fig. 2 only considers 3 amino-acids, but it can be repeated for all amino-acids in a longer protein sequence, because of the regular pattern defining the protein backbone.

The order depicted in Fig. 2 is the one used in our numerical experiments.

The first three atoms, $N-C_{\alpha}-H_{\alpha}$, of the first amino-acid can be used as initial clique (see assumption (a) in Definition 1) for the discretization order because the involved distances are defined by bond lengths and angles, that can be considered as exact [12]. Analogously, taking into account the peptide plane distances and the distances between hydrogens provided by NMR, it is not hard to verify that assumptions (b.1)–(b.2) of Definition 1 are satisfied by the
 order given in Fig. 2.

On the basis of the model in Fig. 2 for the protein backbone geometry, it is possible to conceive other pruning devices [8,53] to be integrated with DDF (see, Eq. 3), based on the following considerations:

- Helices in proteins can be either right or left-handed. The former situation is statistically more common, because of side chains steric constraints. In this work, we do not consider side chains explicitly, but we suppose that it is possible to understand, from an analysis of the protein sequence, whether right-handed or left-handed helices are expected to be present. We call this pruning device as the *chirality-based device*: in some situations, it can allow for placing uniquely some atoms during the execution of the search. For the carbon C, in fact, we can get only one (instead of two) possible positions by using $N-C_{\alpha}-H_{\alpha}$ as reference atoms. An analogous reasoning can be applied to C_{β} . The chirality defines the orientation of the tetrahedron formed by C, C_{β} , N, C_{α} , H_{α} , where C_{α} is the chiral center, and it can be used to avoid unnecessary branching;

²⁵² - The tetrahedron around C_{α} forms a clique as well as the peptide planes [2]. Such local ²⁵³ structures define rigid regions of the protein backbone. Using the peptide plane clique, ²⁵⁴ it is possible to find a unique position for C_{α} . It is also possible to place N uniquely, ²⁵⁵ because its relative orientation with respect to H, C and C_{α} of the same peptide plane can ²⁵⁶ be computed by taking into account the van der Waals minimum distance;

The oxygen atoms in Fig. 2 are included in the model because they participate in hydrogen bonds. Each oxygen can be placed uniquely by using the exact distances with the other atoms of the peptide plane.

260 2.3 Computing coordinates for candidate positions

The method employed to compute the candidate positions at each recursive call of Algorithm 1 has a fundamental importance. While looking for candidate atomic positions for the atom *i*, it is supposed that the reference atoms $\{a, b, c\}$ are already positioned. These reference atoms define a local coordinate system centered at *a* [19,62]. This coordinate system is illustrated in Fig. 3.

Let v_1 be the vector from b to a and v_2 be the vector from b to c. The x-axis for the system in a can be defined by v_1 , and the unit vector in this direction is $\hat{x} = v_1/||v_1||$. Moreover, the vectorial product $v_1 \times v_2$ gives another vector that defines the z-axis, whose corresponding unit vector is \hat{z} . Finally, the vectorial product $\hat{x} \times \hat{z}$ provides the vector that defines the y-axis (let the unit vector be \hat{y}).

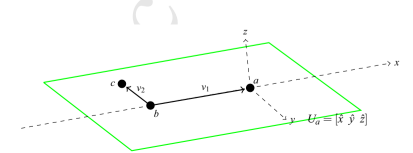


Fig. 3 The reference vertices a, b and c induce a system of coordinates

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These three unit vectors are the columns of a matrix $U_a = \begin{bmatrix} \hat{x} & \hat{y} & \hat{z} \end{bmatrix}$, whose role is to convert directly vertex positions from the coordinate system defined in *a* to the canonical system.

Once the matrix U_a has been computed, the canonical Cartesian coordinates for a candidate position for the vertex *i* can be obtained by:

$$x_{i}(\omega_{i}) = x_{a} + U_{a} \begin{bmatrix} -d_{ai} \cos \theta_{i} \\ d_{ai} \sin \theta_{i} \cos \omega_{i} \\ d_{ai} \sin \theta_{i} \sin \omega_{i} \end{bmatrix},$$
(4)

where θ_i and ω_i are the angles related to the spherical coordinates of vertex *i*.

We will use the symbol θ_i in order to refer to the angle formed by the two segments (i, a)and (a, b), and we will use the symbol ω_i to refer to the angle formed by the two planes defined by the triplets (a, b, c) and (b, a, i) (see Fig. 1). The cosine of the angles θ_i and ω_i can be computed by exploiting the positions of the reference vertices a, b and c, as well as the available distances d_{ai}, d_{bi} and d_{ci} . Thus,

$$\cos \omega_i = \frac{\cos \theta_{c,b,i} - \cos \theta_{a,b,i} \cos \theta_{a,b,c}}{\sin \theta_{a,b,i} \sin \theta_{a,b,c}}$$

where we consider the positive values for the sines, and

$$\cos \theta_i = \cos \theta_{b,a,i} = \frac{d_{ab}^2 + d_{ai}^2 - d_{bi}^2}{2 \, d_{ab} \, d_{ai}}.$$

Recall from Sect. 2.1 that if the three reference distances are all exact, then the three 286 spherical shells are in fact three spheres, whose intersection gives 2 points, with probability 287 1 [32]. The two points x_i^+ and x_i^- correspond to two possible opposite values, ω_i^+ and ω_i^- , 288 for the angle ω_i . When one of the three distances is instead represented by an interval (see 289 Definition 1), the third sphere becomes a spherical shell, and the intersection provides two 290 curves (see Fig. 1). These two curves correspond to two intervals, $[\underline{\omega}_i^+, \overline{\omega}_i^+]$ and $[\underline{\omega}_i^-, \overline{\omega}_i^-]$, 291 for the angle ω_i . In order to discretize these intervals, a certain number of points, say D, can 292 be chosen from the two curves. 293

As shown in [19], the generalized procedure for the computation of atomic coordinates in Algorithm 1, based on equation (4), is very stable when working on MDGP instances related to real proteins. Moreover, equation (4) is also at the basis of an important technique that can be used to reduce the feasible arcs obtained by sphere intersection. This technique for arc reduction was firstly proposed in [20]. Another approach for arc reduction, based on Clifford algebra, is presented in [30].

300 3 Pruning distances and arc reduction

When candidate atomic positions, at each recursive call of the *i*BP algorithm (see Algorithm 1), are computed by intersecting two spheres with one spherical shell, a continuous set of positions is obtained, which generally corresponds to two disjoint arcs, related to two intervals for the corresponding torsion angle values.

During a typical run of Algorithm 1, every time the reference distance d_{ci} is represented by an interval, D equidistant samples are taken from each arc [34]. As a consequence, 2Datomic positions are generated in total, and 2D new branches are added to the tree, at the current layer, for every branch at the upper level. After their computation, the feasibility of each atomic position is verified. On the one hand, too large D values can drastically

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increase the width of the tree; on the other hand, too small values can generate trees where
no solutions can be found (all branches are pruned, because all positions, at a certain layer,
are not compatible with pruning distances).

In [20], an adaptive scheme was proposed for tailoring the branching phase of the *i*BP algorithm so that all computed candidate positions are feasible at the current layer. The idea is to identify, before the branching phase of the algorithm, the subset of positions on the two candidate arcs that is feasible with respect to all pruning distances to be verified on the current layer.

Let us suppose that, at the current layer *i*, the distance d_{ci} is represented by the interval $[\underline{d}_{ci}, \overline{d}_{ci}]$. By using Equation (4), two intervals for the angle ω_i can be identified: $[\underline{\omega}_i^+, \overline{\omega}_i^+] \subset [0, \pi]$ and $[\underline{\omega}_i^-, \overline{\omega}_i^-] \subset [\pi, 2\pi]$, such that the distance constraints

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$$\|x_a - x_i(\omega_i)\| = d_{ai},$$

$$\|x_b - x_i(\omega_i)\| = d_{bi},$$

$$\underline{d}_{ci} \le \|x_c - x_i(\omega_i)\| \le \overline{d}_{ci},$$
(5)

322 are satisfied.

However, there may be pruning distances, at layer *i*, that could be exploited for tightening these two arcs. Let us suppose there is an $h \in \{j < i \mid j \notin \{a, b, c\}\}$, such that the distance d_{hi} is known and lies in the interval $[\underline{d}_{hi}, \overline{d}_{hi}]$. The solution set of the inequalities

$$\underline{d}_{hi} \le \|x_h - x_i(\omega_i)\| \le d_{hi} \tag{6}$$

consists of intervals for ω_i that are compatible with the distance d_{hi} .

A discussion about how to solve the inequalities (6), by using Eq. (4), is presented in details in [20].

The feasible positions for the atom *i* can be therefore obtained by intersecting the two previously computed arcs (in bold in Fig. 1), and several spherical shells, each of them defined by considering one pruning distance between *i* and h < i. For each available pruning distance, other inequalities (6) can be defined and new arcs on the circle *C* may be identified. The final subset of *C* which is compatible with all available distances can be found by intersecting the arcs obtained for each pruning distance with the two initial disjoint arcs, given by Eq. (5).

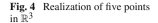
After considering all pruning distances, i.e., after performing all intersections, the final 336 result provides a list of arcs on C that are feasible with all the distances that can be verified 337 at the current layer. All positions that can be taken from these arcs are feasible at the current 338 layer: all of them generate a new branch and may serve as a reference for computing new 339 candidate positions on deeper layers of the tree. In order to integrate the iBP algorithm with 340 this adaptive scheme, there are two main changes to be performed on Algorithm 1. On line 8, 341 the adaptive scheme needs to be invoked for taking into consideration the information about 342 the pruning distances. Moreover, the use of the DDF pruning device has become unnecessary, 343 and it should not be considered at line 20 of Algorithm 1. 344

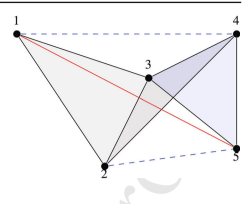
It is important to remark that this adaptive scheme is not supposed to speed up the execution 345 of the search, but rather to help in defining search trees that can actually contain solutions. 346 Without the use of this adaptive scheme, all sample positions selected from the two arcs 347 obtained with the discretization may be discovered to be infeasible as soon as the DDF 348 pruning device is invoked. The other extreme situation is instead the one where the adaptive 349 scheme can allow us to select a subset of sample positions that all bring to the definition of a 350 solution. Naturally, the second situation is desirable, even if, in terms of complexity, it tends 351 352 to increase the total computational cost.

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4 Limitations of the current approach: finding approximate realizations

For DGP instances where all available distances are exact, the presented discrete approach is extremely efficient, allowing for example to realize graphs having thousands of vertices in few seconds with a standard computer [32].

However, for *i*DDGP₃ instances, there are some difficulties encountered by the *i*BP algorithm [34], even for finding one solution. Such limitations, related to the presence of interval data in both discretization and pruning distances, are discussed in this section and a heuristic to overcome such barriers is proposed.

361 4.1 Sampled distances and embeddability

Recent computational experiments have shown that taking equidistant sample points on the feasible arcs (or equidistant samples from the interval distance, see Algorithm 1 in Sect. 2.1), even after the intersection with the available pruning distances (see Sect. 3), is not enough to allow the *i*BP algorithm to solve some MDGPs within a predefined precision. The sampled distances are taken independently in each layer of the tree and, in particular for small *D* values, it is *not* likely that they are compatible with each other and with other pruning distances available at deeper layers.

The underlying issue is related to the embeddability of a given set of distances. Suppose that we are positioning the atom *i* and that the interval distance $[\underline{d}_{ci}, \overline{d}_{ci}]$ is used in the discretization. Even if we assume that there exists a distance value d_{ci}^* which is compatible with the other distances in *E* leading to a solution, we cannot ensure that, with a finite number *D* of samples taken from $[\underline{d}_{ci}, \overline{d}_{ci}]$, the compatible distance d_{ci}^* is actually sampled.

In order to illustrate this fact, consider the following example where five points in \mathbb{R}^3 are embedded (Fig. 4).

Suppose that the straight lines represent exact distances, and let the black lines be the exact distances used in the discretization. The dashed blue lines are the interval distances (used to compute the possible positions of atoms 4 and 5) and the red straight line represents one pruning distance (that can be used to validate the possible positions for the atom 5). The associated distances are the following: $d_{12} = d_{23} = d_{24} = d_{34} = d_{35} = d_{45} = 1$, $d_{13} = \sqrt{2}$, $d_{14} = \sqrt{x} \in [0.5, 2]$, $d_{15} = \sqrt{3}$, $d_{25} = \sqrt{y} \in [0.5, 2]$.

According to the Cayley–Menger conditions [38,59], for this set of distances to be realizable in \mathbb{R}^3 , it is necessary that

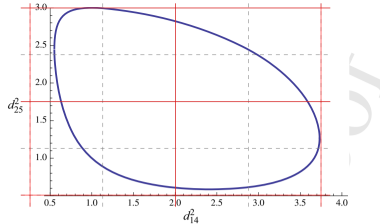


Fig. 5 Solution set for the five point Cayley–Menger determinant with d_{14}^2 and d_{25}^2 as missing distances

| 0 1 2 x 3 1 | |
|----------------------------|-----|
| 1011y1 | |
| 210111 | - 0 |
| x 1 1 0 1 1 | =0, |
| 3 y 1 1 0 1 1 1 1 1 1 0 | |
| 111110 | |

where the above matrix is a bordered distance matrix and $|\cdot|$ denotes its determinant. The solution set of this equation (the values for the missing (interval) squared distances $x = d_{14}^2$ and $y = d_{25}^2$) is represented by the blue curve in Fig. 5.

It is easy to see that, unless the grid is sufficient refined (number of samples *D* is sufficient large), a valid pair of distances (d_{14}^2, d_{25}^2) can be sampled with probability zero.

391 4.2 Long-range distance restraints

Long-range distance restraints are related to atoms that are at least four amino-acids apart in the protein sequence. Even if far in the protein sequence, some atom pairs may be in condition to be detected by an experimental technique. For example, if we consider NMR, it is typical to detect distances between atoms that are very far in the sequence, but quite close in space $(\leq 5 \text{ Å})$.

In case of all available distances are exact, the pruning distances efficiently guide the search in the binary tree corresponding to the discretized search space [32,40]. However, when interval distances are present, the search tree is no longer binary, because *D* samples are taken from each feasible arc. Moreover, the DDF pruning criterion (3) becomes much less effective when the bounds [$\underline{d}, \overline{d}$] are loose, resulting in a large number of active nodes in the tree, which increases exponentially the cost of exploring a whole subtree.

Furthermore, since other interval distances are also employed in the discretization, the sampled positions in the feasible arcs for previous atoms are only approximations for their true positions, and such a sequence of approximate positions may lead to an infeasibility at a further layer. For this reason, the longest-range pruning distances may fail to be verified (even if they are represented by an interval).

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| | - | 1.0 - | | |
|--------|-------------|--------------|------------|---------|
| | _ | 0.5 | 1.0 | 1.5 |
| Fig. 5 | Solution se | t for the fi | ve point C | ayley–M |

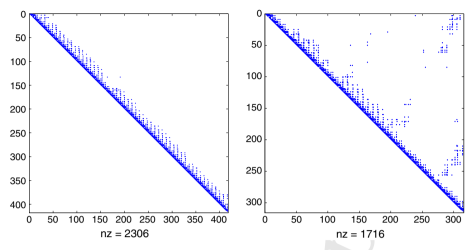


Fig. 6 Available distances for the instances 1FJK (left) and 2E2F (right)

To illustrate this fact, we depict in Fig. 6 the available entries of the upper triangu-408 lar part of the distance matrices for two instances belonging to our set: 1FJK and 2E2F. 409 Notice that the almost-band structure close to the main diagonal is a consequence of the 410 assumptions concerning the discretization. In fact, the distances between pairs (i - 1, i)411 and (i-2, i) are generally derived from the bond lengths and angles, while the distances 412 (i - 3, i) can be generally obtained from the analysis of the torsion angle among the 413 quadruplet of atoms (i - 3, i - 2, i - 1, i). Moreover, the distance (i - 3, i) may be also 414 estimated by applying an experimental technique such as NMR. Other distances that are 415 far from the main diagonal of the matrix can be obtained applying an experimental tech-416 nique. 417

Although 1FJK has more atoms than 2E2F, the former instance can be easily solved 418 by *i*BP in a few seconds, whereas the latter cannot be solved in less than one minute 419 (using $D \leq 20$). The difficulty in solving 2E2F is related to the presence of long-range 420 pruning distances: there are several entries in its distance matrix that are far from the diag-421 onal. 422

4.3 Approximate realizations 423

The presence of interval distances implies uncertainty on the atomic positions obtained by 424 sampling points in the intersection between spheres and spherical shells: even a small error 425 introduced at tree layer i can have a relevant propagation until the layer j >> i and, when 426 the pruning distance is finally tested, it is likely that the propagated error leads to infeasible 427 positions for atom i. 428

Thus, an error introduced during the intersection discretization in a certain tree layer, 429 might make every sampled candidate position infeasible with pruning distances in a further 430 layer. This phenomenon is more evident when considering long-range distance restraints. One 431 possibility to avoid pruning out all branches of the search tree, in order to obtain approximate 432 solutions to the problem, is to relax the distance constraints related to long-range distances. 433 We define the set 434 435

$$\mathcal{L} = \{\{i, j\} \in E \mid |i - j| \ge M\},\tag{7}$$

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where *M* is a positive integer used to identify long-range distance restraints. Our relaxation consists in avoiding the application of the DDF feasibility test (Eq. 3), as well as the intersection scheme (Sect. 3), to pruning distances in \mathcal{L} .

Naturally, when such pruning distances are neglected, some information is lost and this can have an impact on the found solutions. In fact, long-range distance restraints are the main responsible for the global fold. Thus, in order to mitigate this effect, we introduce another pruning criterion based on the partial Mean Distance Error (MDE) at the current layer k:

$$PMDE_{k}(X) = \frac{1}{|J_{k}|} \sum_{\{i,j\}\in J_{k}} \left[\frac{\max\left\{ \underline{d}_{i,j} - \|x_{i} - x_{j}\|, 0 \right\}}{\underline{d}_{i,j}} + \frac{\max\left\{ \|x_{i} - x_{j}\| - \overline{d}_{i,j}, 0 \right\}}{\overline{d}_{i,j}} \right],$$
(8)

443

444 where

445

 $J_k = \{\{i, j\} \in E \mid i \le k \land j \le k\}.$

Let n = |V| and note that $J_n = E$. It is common to measure the quality of a realization by the Mean Distance Error measure:

$$MDE(X) = PMDE_n(X).$$

Thus, by monitoring the $PMDE_k(X)$ for k < n, we can control the quality of partial realizations. This suggests the *PMDE pruning device*: if at layer k, $PMDE_k(X) > \hat{\varepsilon}$, then the candidate partial realization should be pruned. We set $\hat{\varepsilon} > \varepsilon$, where ε is the tolerance used in DDF (Eq. 3).

When this new pruning device is introduced, a solution found by Algorithm 1 is actually an approximate solution in the sense that it satisfies all distances in $E \ \mathcal{L}$ (with tolerance ε), while some distances in \mathcal{L} can be violated. However, the total MDE value for such a solution remains relatively small, because of the new pruning device based on (8). By applying this scheme, together with the chirality and peptide plane constraints (see Sect. 2.2), we expect that the fold of the obtained conformation mimics the fold of the true protein. This is the case for the set of instances used in the computational experiments.

5 Computational experiments

In this section we present some computational results on a set of artificially generated MDGP instances. Our aim is to assess the improvements on *i*BP (Algorithm 1) due to the integration of a set of recently proposed techniques: the pruning devices based on chirality and peptide plane geometry, described in Sect. 2.2; the arc reduction technique presented in Sect. 3; and the partial MDE pruning device introduced in Sect. 4.3.

The instances that we consider in our experiments were generated as it follows. The 466 protein conformations were extracted from the PDB: by using the coordinates of a known 467 conformation, all pairwise distances between atom pairs of the backbone were computed. 468 Then, only a small subset of all distance pairs is kept for defining an instance. The distances 469 related to bond lengths and those that can be obtained from bond angles are considered as 470 exact, as well as distances between atoms belonging to the same peptide plane (see Fig 2). 471 Torsion angles on the protein backbones give rise to the definition of interval distances, related 472 to the minimal and maximal extension of such torsion angles. Distances between pairs of 473 hydrogens are also included, as specified in the next subsection. 474

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During the generation of our instances, we rely on the premise that experimental techniques,
such as NMR spectroscopy, are able to give information about all distances between hydrogen
atoms that are close in space [35]. Moreover, these distances can be supposed to be more
precise than other ones. Statistics on such distances [5,66], with the geometry of consecutive
peptide planes, validate this assumption.

We will consider therefore that all distances between hydrogens belonging to the same or to two consecutive amino-acids are available, and we suppose that they lie in an interval having width 0.1Å. Besides these distances, responsible for the local geometry, we also consider distances between hydrogens that belong to amino acids that are far in the protein sequence, but close in space. These distances are responsible for the global fold. We include these distances in our generated instances whenever they are smaller than 5Å and consider that imprecisions lead to an interval of width 1Å.

Hydrogen bonds H-O, responsible for stabilizing α -helices and β -strands, are also considered. If the distance between H and O of distinct amino-acids is greater than 1.3Å and less than 3.5Å, such a distance is included in our instances as an interval of width 1Å. All intervals have a predefined width and their extremes are randomly generated in a way that the interval contains the true distance.

493 **5.2 Numerical results**

Let us refer to the algorithm presented in [34] as *i*BP, while we will name "New *i*BP" the algorithm integrated with the new method for the computation of candidate positions (see Sect. 2.3), with the technique for arc reduction (see Sect. 3), with the chirality and peptide plane pruning devices (see Sect. 2.2), and with the pruning device introduced in Sect. 4.3.

In both *i*BP variants, the tolerance used in the experiments for the DDF criterion (Eq. 3) is $\varepsilon = 0.001$. In new *i*BP, for the PMDE-based pruning device, we used $\hat{\varepsilon} = 0.01$ and set M = 40 in definition of \mathcal{L} (see Eq. 7). We gradually increased the number of samples *D* taken from the feasible arcs until the first solution is found in less than 60 s (timeout).

The numerical experiments were run in a Intel MacBook Pro, 2Ghz, 2GB RAM, and the Algorithm 1 was implemented in C programming language, compiled using GNU GCC with flag -O3.

Table 1 shows a comparison between *i*BP and New *i*BP. The number of amino acids (*aa*), atoms (|V|) and available distances (|E|) are given for each instance. For the two versions of *i*BP, the performance is evaluated by the minimum number of samples *D* (taken from interval arcs) necessary to find one solution, the number of recursive calls and the CPU time in seconds. The quality of the realization is assessed through the MDE. The character "*" means that the instance could not be solved in less than one minute for any value of $D \le 20$.

We can notice that the arc reduction technique presented in Sect. 3 is very effective in 511 reducing the number of sample positions that we need to extract from the arcs in order to 512 513 obtain at least one solution. This is an important improvement because it is not known a priori how many samples are sufficient to allow *i*BP to find a conformation. We can observe 514 that the number of calls and CPU time were reduced in 9 out of 11 instances. It is also worth 515 to mention that the pruning criteria based on peptide plane geometry and chirality helped 516 the new version of *i*BP in reducing the number of calls in some instances and improving the 517 global fold as well. 518

⁵¹⁹ Concerning the MDE, the original iBP seems to be more stable, although it fails to ⁵²⁰ solve four of the instances (within the specified timeout). On the other hand, since the New

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| PDB ID | D Instance | | | <i>i</i> BP from [34] | | | New <i>i</i> BP | | | | |
|--------|------------|-----|------|-----------------------|-----------|-------|-----------------|---|-----------|------|-------|
| | aa | V | E | D | Calls | Time | MDE | D | Calls | Time | MDE |
| 2JMY | 15 | 120 | 660 | 13 | 37,658 | 0.13 | 3e-06 | 5 | 2983 | 0.01 | 1e-16 |
| 2KXA | 24 | 177 | 973 | 10 | 215,669 | 0.92 | 5e-06 | 3 | 5064 | 0.01 | 6e-03 |
| 1DSK | 28 | 222 | 1210 | 14 | 31,309 | 0.13 | 4e-06 | 4 | 53,890 | 0.14 | 1e-06 |
| 2PPZ | 36 | 287 | 1522 | 9 | 2,372,242 | 11.34 | 2e-06 | 3 | 442,965 | 1.87 | 4e-08 |
| 1AQR | 40 | 310 | 1596 | * | * | * | * | 4 | 114,671 | 0.20 | 6e-03 |
| 2ERL | 40 | 324 | 1792 | 14 | 1,495,282 | 6.14 | 4e-06 | 3 | 10,410 | 0.03 | 1e-03 |
| 2E2F | 41 | 315 | 1716 | * | * | * | * | 3 | 19916 | 0.06 | 9e-03 |
| 1FJK | 52 | 417 | 2306 | 12 | 115,426 | 0.73 | 4e-06 | 4 | 925,090 | 3.07 | 2e-06 |
| 2JWU | 56 | 448 | 2416 | * | * | * | * | 4 | 226870 | 0.81 | 1e-02 |
| 2KIQ | 57 | 455 | 2452 | 20 | 1,217,945 | 12.79 | 6e-06 | 4 | 317,136 | 1.12 | 7e-04 |
| 2LOW | 64 | 497 | 2650 | * | * | * | * | 3 | 3,738,152 | 8.79 | 2e-07 |

 Table 1
 Numerical results on artificially generated instances from the PDB

iBP uses the relaxed pruning criterion PMDE, it cannot ensure a better MDE for all the 521 instances. For some of them we observe a better MDE which is a consequence of the other 522 considered pruning devices. Although we relaxed some distance constraints by using PMDE, 523 the chirality constraints helped in improving the local geometry, resulting in a better MDE. 524 For those instances with a worse MDE, like 2KXA, 2E2F or 2KIQ, the PMDE relaxation was, in some sense, the way to "pass-through" the long-range distance constraints and find a 526 realization in a affordable time. Additionally, we remark that an MDE value around 10^{-3} is 527 able to guarantee a sufficient detection of the global fold of the protein. In fact, the realizations 528 found by the New *i*BP are not so far from the true ones. The quality of such realizations is 529 discussed in the next subsection. 530

531 5.3 Quality of a realization and practical usage

While looking at Table 1, a natural question emerges: how good are the realizations X with $MDE(X) \approx 10^{-3}$ when compared to the "true" protein ?

Since we have relaxed the distance constraints related to long-range distances, in principle, we cannot ensure that the underlying molecule is recovered. However, we will illustrate that the realization found by the New *i*BP gives a very good approximation of the true conformation.

First, let us take a look at the instance 2KXA. Figure 7 shows the realizations found (first found solutions) by the original *i*BP and the New *i*BP. Although the MDE of the first is smaller than the second, 10^{-6} against 10^{-3} , we can see that the conformations are roughly the same, except by partial reflections. The New iBP produced a right-handed helix because it contains, in its list of pruning devices, the chirality-based device.

⁵⁴³ Now, let us consider the instance 2E2F. According to Table 1, the MDE for the realization ⁵⁴⁴ found by the New *i*BP is approximately 10^{-2} . By superimposing the realizations found with ⁵⁴⁵ the true one from the PDB (first model), see Fig. 8, we obtain a RMSD value equal to 0.7 Å ⁵⁴⁶ (according to TM-align [67]).

Therefore, although the realization found by New *i*BP does not fit perfectly with the true conformation, it is close enough to identify its global fold, and it also can be used as a smart

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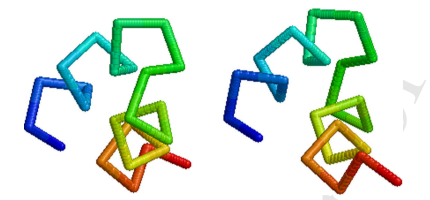
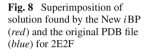
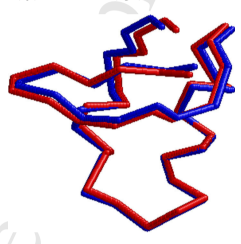


Fig. 7 Realization for 2KXA found by the original *i*BP (*left*) and the new *i*BP (*right*)





starting point for a local optimization intended to minimize the MDE and enforce VdW constraints [61].

Once the first solution X_1 is found by the "New *i*BP", a set of feasible exact distances for the distances that were originally represented by intervals can be selected. This set of distances defines a DGP instance with exact distances which contains X_1 in its finite solution set. Moreover, by solving such an instance with the basic BP algorithm (for exact distances), we can compute all other feasible conformations that can be obtained from X_1 by partial reflections [32,38]. This procedure gets rid of the flexings ¹ in the molecule, but only in this case the solution set is finite.

Applying this scheme to a modified 1AQR instance, where hydrogen distances between consecutive amino-acids were removed and the threshold was lowered to 4.5 Å, four incongruent conformations are obtained, as depicted in Fig. 9.

We claim that, even though interval distances pose some difficulties to the extension of this combinatorial approach, it is still possible to explore all the (discrete) conformational space obtained with discretization. Henceforth, we propose our New *i*BP as an exploratory tool to enumerate protein conformations that satisfy most of the given distance restraints, that can be further improved by local minimization procedures.

¹ Continuous motions of part of the structure preserving all distance restraints.

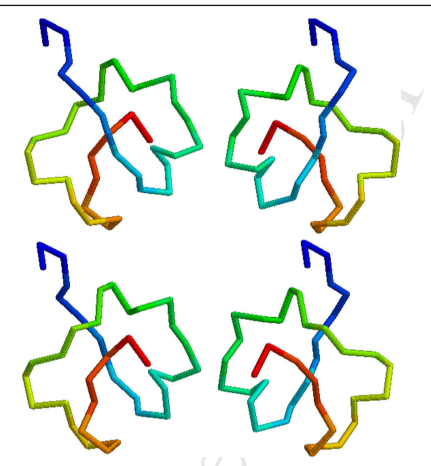


Fig. 9 Four incongruent realizations for 1AQR. The conformations in the *bottom* are reflections of the *top* ones. All four conformations differ by partial reflections

566 6 Conclusion and future work

We collected in this paper the most recent and promising advances in solving the MDGP with our combinatorial approach. The main contributions presented in this paper can be summarized as follows:

- identification of the main barriers against the successful application of the discrete
 approach to real MDGPs with interval data;
- another pruning devices based on chirality and peptide planes, whose easy implementa tion is allowed by the model and discretization order presented in Sect. 2.2;
- 3. a new pruning device that "relaxes" long-range distance constraints (Sect. 4.3) which
 allows us to obtain approximate realizations.

⁵⁷⁶ Computational experiments on artificially generated instances showed the effectiveness of all above mentioned points, when integrated in the *i*BP algorithm. We are in fact able to find

approximate realizations for protein backbones up to 64 amino acids in an affordable time,

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and with reasonable precision, that can be further improved by using our solutions as starting 570 points for a local minimization solver. 580

In the presented experiments, the two compared versions of the iBP algorithm were both used for identifying only one solution to the problem. However, as remarked before 582 and illustrated in Sect. 5.3, the *i*BP algorithm can potentially enumerate the entire solution 583 set of a discretizable MDGP. Research is currently focused on efficiently enumerating all 584 conformations belonging to the search tree. Due to the presence of interval distances, many 585 solutions may belong to the same cluster/ensemble of conformations. Hence, the next step is 586 to define a method to classify the solutions in equivalence classes and integrate *i*BP with an 587 scheme able to pick only one representative conformation from each incongruent ensemble. 588

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