Exploring Life through Logic Programming:  
Logic Programming in Bioinformatics

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Abstract. This chapter provides a broad overview of how logic programming, and in particular Answer Set Programming, can be used to model and solve some popular and challenging classes of problems in the general domain of bioinformatics. In particular, the chapter explores the use of ASP in Genomics studies, such as Haplotype inference and Phylogenetic inference, in Structural studies, such as RNA secondary structure prediction and Protein structure prediction, and in Systems Biology. The chapter offers a brief introduction to biology and bioinformatics and working ASP code fragments for the various problems investigated.

1 Introduction

Recent advances in the field of Biology have uniquely built on the use of computational methods to model, simulate, and sift through dauntingly large data repositories. The synergistic interaction with biologists and other scientists in the life sciences has provided computer scientists with a broad range of challenging problems. Problems are often hidden or vaguely specified, and emerge only after long discussions with biologists, physicists, chemistry researchers, medical clinicians, etc. The effective resolution of these problems has the potential of having a profound impact on our ability to understand the basic mechanisms of life and to advance medical research. Bioinformatics can be broadly defined as the area of Computer Science that deals with modeling and solving problems from biology and related life sciences.

In very broad strokes, we can classify typical bioinformatics application domains in three general categories. The first class includes applications of computational methods as a support infrastructure for analysis and experiments; this is the case, for example, of automated environments for workflow management, description and annotation of experiments, minimal reporting requirements (e.g., MIAME), etc. The second class includes problems that are efficiently solvable (e.g., in polynomial time), but commonly applied to very large data sets (e.g., BLAST searches and other string matching problems [98, 67, 63]). Finally, the third class includes problems that are intractable (i.e., NP-complete or worse),
even under simplifying assumptions; this is the case, for example, of problems related to ab-initio protein structure prediction and to synthesis of gene regulatory networks.

Bioinformatics problems can also be classified according to the target of the study. Genomics studies focus on the investigation of genomes; genomic studies often rely on tractable algorithms applied to very large amounts of data, such as DNA and RNA sequences. Structural studies focus on the prediction and recognition of the spatial structure of biomolecules (e.g., proteins); problems in this class are often intractable and based on data sets that are relatively smaller than those used in genomic studies. Systems studies (a.k.a. systems biology) investigate the complex interactions among components (e.g., genes) of a biological system; these studies typically belong to high complexity classes. The interested reader can refer to [23, 81, 117] for an introduction to computational biology.

The literature has highlighted the potential for logic programming technology to address problems in all of these areas (see, e.g. [5]). Prolog and other logic programming paradigms have been employed in supporting workflow management—e.g., data formats interoperation, service composition (e.g., [91])—and querying tasks—e.g., querying phylogenetic databases [21]. Nevertheless, the capabilities of logic programming—especially Constraint Logic Programming (CLP) and Answer Set Programming (ASP)—to tackle complex combinatorial problems make this paradigm particularly appealing to address problems in the third class. Logic programming offers some distinct advantages; in particular:

- Models are rarely stable and static—logic programming provides the level of elaboration-tolerance to support model modifications and incremental addition of new knowledge.
- The use of more traditional techniques, such as linear programming, is often inadequate (e.g., incapable of capturing the complexity of realistic energy models).

In this manuscript, we will provide a broad overview of how ASP can be used to model and solve some popular and challenging classes of Bioinformatics problems. In particular, we will explore ASP in

- Genomics studies: we will investigate problems associated to Haplotype inference and Phylogenetic inference;
- Structural studies: we will investigate problems associated to RNA secondary structure prediction and Protein structure prediction;
- Systems studies: we will investigate problems associated to reasoning with biological networks.

The manuscript represents a coherent synthesis of previous research contributions (as found in the literature, including some the authors’ own work) as well as novel problem formalizations. The content of this paper also builds on our domain knowledge gained by organizing the Workshop on Constraint-based Methods in Bioinformatics yearly from 2005 (see, e.g., cp2013.a4cp.org/workshops/web), and related publications (e.g., [31, 36, 1]).
2 Biology in a nutshell

The well known central dogma of Biology was first introduced in 1958 by F. Crick [30] and describes how the biologically relevant information migrates from DNA sequences to RNA sequences and finally to proteins, whose function is determined by their 3D structure. The conversion DNA $\rightarrow$ RNA is called transcription, while the conversion RNA $\rightarrow$ Protein is called translation. Let us briefly review the above notions in order to understand the central dogma.

DNA (DeoxyriboNucleic Acid) is characterized by a sequence of nucleotides of 4 kinds: A, C, G, and T (Adenine, Cytosine, Guanine, Thymine). The nucleotides have a different atomic composition, omitted here; however they share a common substructure, that is used to connect sequences of nucleotides into polymeric strands of potentially unlimited length. Typically, DNA strands have a high propensity to pair: two strands can be aligned and facing nucleotides, one from each strand, can form a relatively stable binding. Some nucleotide matchings are more favorable than others; in particular, there is a notion of complementary string. Precisely, given a sequence $s \in \{A, C, G, T\}^*$, its complementary sequence $\bar{s}$ is obtained by reversing the sequence order and by substituting $A \leftrightarrow T$ and $C \leftrightarrow G$. A string $s$ and its complementary string $\bar{s}$ form bindings between each pair of corresponding nucleotides, and together they fold into the famous double helix (see, e.g., the DNA strands at both sides of Figure 1).

Observe DNA strings are huge ($10^6$–$10^{10}$ nucleotides). Differences between the DNAs of two members of the same specie are limited (e.g., 1 in 1,000 nucleotides for humans). Some fragments of the DNA encode proteins, as we show below. These regions encode information as described by the central dogma and they are called genes. Other regions perform other tasks that are beyond the...
focus of this manuscript. With the inception of the Human Genome Project,\(^1\) it became possible to estimate that the human genome contains anywhere between 20,000 to 25,000 protein-coding genes [73]—the most recent estimates indicate such number to be roughly 21,000 genes. Differences in some nucleotides within the same gene characterize properties which distinguish one individual from another. The set of all genes of an individual is called genome. Some important problems dealing with genome analysis are discussed in Section 3 and Section 4.

In order to be transcribed, the DNA double helix is locally detached by effect of specific molecules (enzymes)—see Figure 1 for an example. The two single strands are exposed to the surrounding environment and they can, under proper conditions, be used to initiate the transcription. Other enzymes regulate the process, which basically ensures that a complementary copy of the DNA fragment is generated. The new string of RNA (Ribonucleic Acid) is composed of four kinds of nucleotides: \(A, C, G,\) and \(U,\) where \(A, C, G,\) are the same as in DNA, while \(U\) (Uracil) can be seen as a “variant” of \(T.\) The RNA string is single stranded and it less stable and shorter than a DNA sequence. However, RNA can perform various tasks within the cell—while DNA sequences are confined in the cell’s nucleus, for eukaryotic organisms. After the transcription is over, the DNA double helix is formed again and the new RNA sequence acts as a messenger of the information drawn from the DNA. From the string manipulation point of view, a new RNA sequence \(r\) is obtained from a substring of \(s\) that is copied, complemented (\(T\) replaced by \(U\) nucleotides), and reversed.

![Fig. 2. Example of RNA secondary structure.](http://web.ornl.gov/sci/techresources/Human_Genome/)

The RNA string \(r\) folds in the space according to a series of favorable matches between pairs of nucleotides (base pairing). In this case, the presence of a single strand allows richer shapes w.r.t. the DNA helix. The so-called RNA secondary structure is the set of its base pairings (see, for example, Figure 2). In particular, \(A–U\) and \(C–G\) are the favored base pairings; these are also known as the Watson-Crick pairs. Note that it is also possible to find \(U–G\) pairs. The topology of the pairs influences and stabilizes the three dimensional functional shape of the strand. Predicting such three dimensional shape is extremely important, and

\(^1\) http://web.ornl.gov/sci/techresources/Human_Genome/
this is the topic of Section 5. It is interesting to note that, unlike the protein case, an accurate secondary structure prediction for RNA is sufficient to infer the overall three dimensional shape of the strand.

Let us focus now on the translation phase, namely how RNA sequences determine proteins sequences. In simple terms, triplets of consecutive nucleotides (codons, see the 3-letter boxes in Figure 1) identify a new object (monomer) called amino-acid. The association is defined by the so called universal code, and it is summarized in Figure 3. Even if the translation involves RNA nucleotides, it is a common practice to associate the corresponding DNA triplets, in order to easily interface to genomic data. There are $4^3 = 64$ possible distinct codons, whereas only 20 distinct amino acids are commonly generated; this is caused by the presence of redundant codons that encode the same amino acid. Notably, specific codons are used to signal to the dedicated machinery of the cell the start and the stop of a protein encoding sequence.

<table>
<thead>
<tr>
<th>Amino Acid</th>
<th>Codons</th>
<th>Amino Acid</th>
<th>Codons</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>GCT, GCC, GCA, GCG</td>
<td>E</td>
<td>TTA, TTG, CTT, CTC, CTA, CTG</td>
</tr>
<tr>
<td>R</td>
<td>CGT, CGC, CGA, CGG, AGA, AGG</td>
<td>K</td>
<td>AAA, AAG</td>
</tr>
<tr>
<td>N</td>
<td>AAT, AAC</td>
<td>M</td>
<td>ATG</td>
</tr>
<tr>
<td>D</td>
<td>GAT, GAC</td>
<td>F</td>
<td>TTT, TTC</td>
</tr>
<tr>
<td>C</td>
<td>TGT, TGC</td>
<td>P</td>
<td>CCT, CCC, CCA, CCG</td>
</tr>
<tr>
<td>Q</td>
<td>CAA, CAG</td>
<td>S</td>
<td>TCT, TCC, TCA, TCG, AGT, AGC</td>
</tr>
<tr>
<td>E</td>
<td>GAA, GAG</td>
<td>T</td>
<td>ACT, ACC, ACA, ACG</td>
</tr>
<tr>
<td>G</td>
<td>GGT, GGC, GGA, GGG</td>
<td>W</td>
<td>TGG</td>
</tr>
<tr>
<td>H</td>
<td>CAT, CAC</td>
<td>Y</td>
<td>TAT, TAC</td>
</tr>
<tr>
<td>I</td>
<td>ATT, ATC, ATA</td>
<td>V</td>
<td>GTT, GTC, GTA, GTG</td>
</tr>
<tr>
<td>START</td>
<td>ATG</td>
<td>STOP</td>
<td>TAA, TGA, TAG</td>
</tr>
</tbody>
</table>

Fig. 3. Universal genetic code: DNA triplets are associated to corresponding amino acid (represented by a single letter)

The process described so far can be summarized as follows: a gene contained in the DNA, through RNA transcription, uniquely determines a sequence of amino acids, called the primary sequence of a protein.

The process of translation assembles amino acids as connected beads, linked together by peptidic bonds. All amino acid types contain a common part, called backbone, which is formed by a set of atoms that allows high flexibility, many degrees of freedom and yet robustness of the chain. One of these atoms is a Carbon atom called Cα and it shows amenable properties when modeling protein structures. In particular, given any protein structure in any spatial arrangement, the Cα’s belonging to two consecutive amino acids in a sequence are roughly 3.81Å apart, with a very low variance. Each amino acid is characterized by a type-dependent variable group of atoms that influences its specific physical and chemical properties. This group, called side chain, contains anywhere from 1 to 18 atoms and is connected to the Cα atom of each amino acid.

Another fundamental result by Anfinsen [2] states that the same protein sequence immersed in the same environment is capable of folding to a specific, and
often stable, three-dimensional shape, called the native state or native conformation. The process is spontaneous and arranges the molecule according to its minimal free energy. The spatial arrangement of the protein, thanks to the chemical properties of amino acids on the surface, determines the function and how the protein interacts and binds to other molecules (ligands). Thus, the combination of the previously discussed central dogma and the properties of proteins provides the fundamental explanation of how a gene expressed by DNA can encode a protein that is capable of performing specific functions.

Proteins can contain from as few as 10 amino acids all the way to 1,000 amino acids. An average globular protein is about 300 amino acids long. Each amino acid contains 7–24 atoms; therefore, the number possible arrangements of atoms in the three-dimensional space is well beyond any computational power. Given the atomic size of proteins and the difficulties to experimentally determine their native state, the prediction of their structure plays a crucial role. This problem is discussed in Section 6.

When studying a living cell and/or organism, we need to contend with the existence of a complex network of (partially known) interactions among the various active components. It is often the case that these interactions are not studied at the lowest level of atomic-level processes, but the are instead abstracted using higher level perspectives. This is often necessary to address the high computational complexity underlying these systems and the relatively limited knowledge about the interaction models. Depending of the level of simplification adopted, it is possible to investigate different properties of a system of cells and/or organisms. For example, metabolism, some signaling pathways of hormones and cell cycles can be modeled by fusing experimental data and stochastic analyses.

Systems biology describes the relationships among components through graphs and provides some computational models that can be used to reproduce particular behaviors. The typical objects modeled are genes and DNA/RNA fragments, proteins and enzymes, metabolites and external stimuli. The goals of systems biology are to (a) analyze the network of interactions, (b) predict the behavior of the system under specific conditions, and (c) integrate knowledge by merging diverse experimental data. The representation in use can vary, depending on the scope and granularity of the modeled system: a specific reaction chain may involve a few molecule types, a larger scale system may describe a cell behavior that involves genes expressions (e.g., quantity of RNA transcribed from a specific gene in a time unit), while a very large system may describe a complete organism with a more coarse description of single cell activities. Some techniques developed within the logic programming community to deal with problems from systems biology are presented in Section 7.

3 Phylogenetics

Phylogenies are artifacts used to describe the relationships among entities (e.g., biological entities like proteins or genomes) derived from a process of evolution.
We often refer to the entities studies in a phylogeny as *taxonomic units (TUs)* or, simply, as *Taxas*.

The field of *Phylogenetics* developed from the domain of biology, as a powerful instrument to investigate similarities and differences among entities as a result of an evolutionary process. Evolutionary theory provides a powerful framework for comparative biology, by converting similarities and differences into events reflecting causal processes. As such, evolutionary-based methods provide more reliable answers than the traditional similarity-based methods, as they employ a theory (of evolution) to describe changers instead of relying on simple pattern matching. Indeed, evolutionary analyses have become the norm in a variety of areas of biological analysis. Evolutionary methods have proved successful, not merely in addressing issues of interest to evolutionary biologists, but in regard to practical problems of structural and functional inference [107]. Evolutionary inference of pairing interactions determining ribosomal RNA structure [116] is a clear case in which progress was made by the preferential use of an evolutionary inference method, even when direct (but expensive and imprecise) experimental alternatives were available. Eisen and others [101, 82] have shown how an explicitly evolutionary approach to protein “function” assignment eliminates certain categories of error that arise from gene duplication and loss, unequal rates of evolution, and inadequate sampling. Other inference problems that have been addressed through evolutionary methods include studies of implications of SNPs in the human population [101], identification of specificity-determining sites [66], inference of interactions between sites in proteins [109], interactions between proteins [108], and inferences of categories of sets of genes that have undergone adaptive evolution in recent history [83].

Phylogenetic analysis has also found applications in domains that are outside of the realm of biology; for example, a rich literature has explored the evolution of languages (e.g., [62, 94, 102, 42]).

### 3.1 Modeling

A *phylogenetic tree* (or simply a *phylogeny*) is typically a labeled binary tree $(V, E, L, T, L)$ where:

- The leaves $L$ represent the taxonomic units being compared;
- The internal nodes $V \setminus L$ represent the (hypothetical) ancestral units; in rare cases, the internal nodes correspond to concrete entities (e.g., fossils);
- The edges $E$ of the tree describe evolutionary relationships; the structure of the edges describe the processes that hypothetically led to the evolution of the TUs, e.g., biological processes of speciation, gene duplication, and gene loss;
- Commonly, each TU is described by a collection of finite domain properties, referred to as *characters*. In the formalization, $T = (C, D, f)$ is the description of such properties, where
  - $C = \{c_1, \ldots, c_k\}$ is a finite set of characters;
− $D = (D_{c_1}, \ldots, D_{c_k})$ associates a finite domain to each character;\(^2\)
− $f : L \times C \rightarrow \bigcup_{c \in C} D_c$ is a function that provides the value of each character for each TU being studied.

- We are often interested in the length of the branches of a phylogeny and/or the assignment of dates to the internal nodes of the phylogeny; if this feature is present, then we will describe it as a function $L : E \rightarrow \mathbb{R}^+$. Whenever we do not have information about length of branches, we omit the component $L$ from the description of the phylogeny.

For example, Fig. 4 (top) shows a phylogenetic tree for the TUs $L = \{\textit{Mollusca}, \textit{Annelida}, \textit{Arthropoda}, \textit{Echinodermata}, \textit{Chordata}\}$. In this example, the set of characters is $C = \{\textit{Coelom}, \textit{Dark}\}$—Coelom denotes the presence/absence of coelom (a body cavity between the intestine and the body walls), while Dark denotes the phenotypical character of having dark color. In this example, these are both binary characters, i.e., $D_{\text{Coelom}} = D_{\text{Dark}} = \{0, 1\}$. The function $f$ describing the five TUs is given by the table underneath each TU in the figure—e.g., $f(\textit{Annelida}, \text{Coelom}) = 0$ and $f(\textit{Annelida}, \text{Dark}) = 0$.

While a significant part of the literature focused on binary, rooted trees for the description of phylogenies, there are several instances in which this model has to be relaxed.

While true evolutionary histories are indeed rooted trees, the actual location of the root is often challenging, especially because of the lack of appropriate historical references. As a result, it is not uncommon to encounter phylogenetic analyses that make use of unrooted trees (e.g., see Figure 5).

Further relaxations of the binary rooted tree view of phylogenies can also be found. The binary nature of the tree can be abandoned (allowing the creation of polytomies, i.e., nodes with more than two children) when a fully-resolved bifurcating tree cannot be determined with sufficient reliability.

Typically, we assume that each node in a phylogeny has at most one immediate ancestor; this reflects the assumption that evolutionary lineages, once separate, do not fuse; this assumption follows from the “biological species concept” based on reproductive isolation [85]. Even the restriction of single parentage may be abandoned, for strictly biological reasons, in the case of lateral transfer (a partial mixing of lineages due to transfer of some genes) or reticulate evolution (a genome-level mixing of lineages due to interbreeding between previously separate species) [75, 86]. This leads us to consider more general structures, e.g., phylogenetic networks.

**Phylogenetic Reconstruction** The most fundamental problem in phylogenetics is the phylogenetic tree reconstruction problem: given a set of data characterizing the entities being studied (e.g., species, genes, languages), identify a phylogeny that accurately describes the evolutionary lineages among the given entities. The key problem is how to define what does it mean to “accurately

\(^2\) Recent proposals have also started exploring the use of continuous characters, e.g., [95].
Fig. 4. A Sample Phylogeny (top); compatible (middle) and incompatible (bottom) characters
describe,” i.e., what measure of accuracy is used to compare plausible trees. A
variety of measures have been proposed, and various phylogenetic reconstruction
methods have been proposed based on the specific measure being used to assess
quality of the phylogeny.

A common method used in deriving phylogenies is based on the idea of character compatibility—a principle derived from Le Quesne’s idea of uniquely de-

derived characters [79, 80].

The intuitive idea of compatibility is as follows: a character $c$ is compatible
with a phylogeny if the TUs that present the same value for such character are
connected by a subtree within the phylogeny. More formally, given a phylogeny $P = (V, E, L, T, L)$, with $T = (C, D, f)$, a character $c \in C$ is compatible with $P$
if there is a mapping $h_c : V \to D_c$ such that:

- For each $t \in L$, we have that $h_c(t) = f(t, c)$;
- For each $i \in D_c$, the projection of the graph $(V, E)$ on the set of nodes $V_i^c = \{t \in V \mid h_c(t) = i\}$ has a subgraph that has $V_i^c$ as nodes and it is a
rooted tree.

A character that is not compatible with a phylogeny $P$ is said to be incompatible. The above (sub-tree) requirement implicitly states that when a character changes (during evolution) it never goes back to the previous value. This is referred to
as the Camin-Sokal requirement; moreover, it also accounts for the requirement
that the “change” occurs in a unique place, known as the Dollo requirement.

In the example of Fig. 4, the character Coelom is compatible with the given phylogeny—as shown in Fig. 4(middle). On the other hand, the character Dark is not compatible with this phylogeny (as shown in Fig. 4(bottom)).

The goal, in phylogeny reconstruction, is to determine a phylogeny that maxi-

mizes the number of characters that are compatible with it. This problem has
been often referred to as the $k$-incompatibility problem [53]. Formally, the $k$-
incompatibility problem is the problem of deciding, given a set $L$ of TUs, a
character description $T = (C, D, f)$ of $L$, and an integer $n \in \mathbb{N}$, whether there
is a phylogeny $(V, E, L, T)$ that has at most $k$ incompatible characters.
Remark 1. We report on some combinatorial considerations. Given a set of \( n \) TUs, the number of unrooted trees can be computed by the formula (using the so-called double-factorial) \((2n-5)!! = 1 \cdot 3 \cdot 5 \cdots 2n-5\), while the number of rooted trees is \((2n-3)!! = 1 \cdot 3 \cdot 5 \cdots 2n-3\). As such, the number of possible phylogenies is exponential in the number of TUs given.

The problem considered above, based on \(k\)-incompatibility, has been studied in the literature and determined to be NP-complete [39]. The proof relies on a reduction of the \(k\)-clique problem to the \(k\)-incompatibility problem. This result is not surprising, and similar complexity results can be found for other measures of phylogeny accuracy—e.g., the popular maximum parsimony method for the construction of phylogenies is shown to be NP-hard in [38].

3.2 ASP Encoding

The \(k\)-incompatibility problem admits an elegant encoding in ASP, thanks to its natural formulation as a generate-and-test problem.

Let us assume that the TUs for the problem at hand are represented in ASP as a collection of facts of the form \(\text{taxa}(i)\) where \(i\) is the name of a TU. For the sake of simplicity, let us assume that the TUs have been numbered and we use the index of a TU as its name—thus, we will use the ASP declaration \(\text{taxa}(1..N)\).

where \(N\) is the number of TUs. The character description \(\mathcal{T}\) will be provided extensionally using facts of the form:

\[
\begin{align*}
\text{character}(c). & \quad \text{%%% For each } c \in C \\
\text{domain}(c, s). & \quad \text{%%% For each } s \in D_c \\
\text{f}(i, c, v). & \quad \text{%%% For each } i \in L, c \in C \text{ such that } f(i, c) = v
\end{align*}
\]

The code is divided in two parts: the tree construction and the computation of \(h_c\). Let us focus on the tree construction first:

1. \(\text{num_of_taxas}(N) :- \text{N} = \#\text{count(taxa(L))}\).
2. \(\text{node}(1..2*\text{N}-1) :- \text{num_of_taxas}(\text{N}).\)
   \%
   Each internal node has exactly two children
3. \(2 \{ \text{edge}(I, J): \text{node}(J) \} 2 :- \text{node}(I), \text{not taxa}(I).\)
   \%
   Children are of smaller index
4. \(:- \text{node}(I; J), \text{edge}(I, J), I < J.\)
   \%
   Each node but the root has 1 fathers
5. \(1 \{ \text{edge}(I, J): \text{node}(I) \} 1 :- \text{node}(J), \text{num_of_taxas}(\text{N}), J < 2*\text{N}-1.\)

Nodes 1 to \(N\) are used for the TUs and they represent leaves of the tree being constructed. The tree has \(2N-1\) nodes and node \(2N-1\) is the root. Lines (1) and (2) compute the number of TUs and define the nodes of the trees. Line (3) is used to enforce the binary structure of the tree (each internal node has exactly two children) and to ensure that the TUs are appearing as leaves of the tree. Line (4) is used to ensure that edges point from bigger to smaller nodes, allowing
us to remove symmetries. Finally, line (5) allows us to ensure the creation of a tree, by verifying that each node, except for the root, has exactly one parent.

The second part of the code is used to assess the compatibility of the phylogeny with respect to the characters.

\[
\begin{align*}
\% & \text{h}_c \quad \text{(i.e., } h(_,C,\_)) \text{ is a total function} \\
\% & \text{h}_c \text{ is an extension of } f \\
\% & \text{Project the tree on each character} \\
\% & \text{A node can be either a root or not after the projection} \\
\% & \text{If there are two roots, it is not partially compatible} \\
\% & \text{If for all symbols is partially compatible, it is compatible} \\
\% & \text{Maximize the compatible characters} \\
\end{align*}
\]

Lines (6) and (7) provide the definition of the function \( h_c \), ensuring that it represents an extension of \( f \). The code in line (8) extracts the edges of the phylogeny that link nodes with the same value for a given character. This creates a new tree, composed of edges labeled by the pair \((c,v)\) of character and corresponding value. The projected trees are analyzed to determine whether they represent individual trees or forests (lines (11)-(13)). If there are multiple trees for the same \((c,v)\), then the phylogeny is incompatible for such character value. Lines (15) and (16) allow us to classify the phylogeny with respect to each character—as being compatible or not. Finally, Line (18) forces the search for phylogenies that maximize the number of compatible characters.

**Remark 2.** In the encoding above and in all encodings in the paper we modeled the optimization version of the problem. In general, for encoding a decision version, i.e. is there a solution of size at least (at most) \( k \) one have to replace the last two lines of the encodings with a line of the kind:

\[
\begin{align*}
\% & \text{Maximize the number of compatible characters} \\
\% & \text{At most k compatible characters} \\
\end{align*}
\]

where \( k \) can be given as input (e.g., \(-c k=2\) — \( k \) should be on the right for requiring at most.
3.3 Related work

The use of logic programming in the context of phylogenetic inference is a relatively new domain.

Erdem and her collaborators have extensively studied several variants of the phylogeny reconstruction problem, both using traditional Answer Set Programming, as well as more ad-hoc logic programming systems. A nice survey of their contributions can be found in [48]. The ASP modeling of phylogenetic reconstruction using the k-incompatibility problem was originally proposed in [11, 10], later expanded to address issues of polymorphic characters. This line of work was later extended to address the problem of formulating criteria of diversity or similarity in the generation of phylogenies, enabling the creation of pools of phylogenies that are sufficiently similar/dissimilar [14]; similarly, the problem of dealing with large numbers of plausible solutions has been addressed in [15] using mechanisms to weight each phylogeny.

The proposed solution described in the previous section is not suitable to deal with very large data sets; techniques based on divide-and-conquer have been successfully adopted to the problem of phylogeny reconstruction [10, 13]; similar ideas have been used in other areas associated to manipulation of phylogenetic knowledge (e.g., for supertree construction [100]).

The study of phylogenies that include time calibrations has been first considered in [50, 51], with special considerations for how to deal with real-valued time stamps and even larger data sets.

Another line of research focused on the application of logic programming technology to phylogenetic analysis and, more in general, to phyloinformatics, can be found in the context of the CDAOStore project [78, 21]. The CDAOStore relies on a triple-based encoding of phylogenetic artifacts (predominantly phylogenies and associated character data matrices), encoded using the Comparative Data Analysis Ontology (CDAO) [92]. Logic programming is employed, in CDAOStore, to implement a variety of queries to a phylogenetic repository, including structural queries (to retrieve phylogenies that meet certain structural constraints) and synthesis queries (e.g., to develop supertrees of sets of collected phylogenies).

4 Haplotype Inference

The DNA of diploid organisms, such as humans, is organized in pairs of not completely identical copies of chromosomes. The sequence of nucleotides from a single copy is called haplotype, while the conflation of the two copies constitutes a genotype. Each person inherits one of the two haplotypes from each parent. The most common variation between two haplotypes is a difference in a single nucleotide. Using statistical analysis within a population, it is possible to describe and analyze the typical points where these mutations occur. Each of such differences is called a Single Nucleotide Polymorphism (SNP). In other words, a SNP is a single nucleotide site, in the DNA sequence, where more than one type
of nucleotide (usually two) occur with a non-negligible population frequency. We refer to such sites as alleles.

Considering a specific genotype, a SNP site where the two haplotypes have the same nucleotide is called an homozygous site, while it is heterozygous otherwise. Research has confirmed that SNPs are the most common and predominant form of genetic variation in DNA. Moreover, SNPs can be linked to specific traits of individuals and with their phenotypic variations within their population. Consequently, haplotype information in general, and SNPs in particular, are relevant in several contexts, such as, for instance, in the study and diagnosis of genetic diseases, in forensic applications, etc. This makes the identification of the haplotype structure of individuals, as well as the common part within a population, of crucial importance. However, in practice, biological experiments are used to collect directly genotype data instead of haplotype data, mainly due to cost or technological limitations. To overcome such limitations, efficient and accurate computational methods for inferring haplotype information from genotype data have been developed during the last decades (for a review, the reader is referred to [72, 69, 70]).

4.1 Modeling

The haplotype inference problem can be formulated as follows. First, we apply an abstraction and represent genotypes and haplotypes by focusing on the collection of ambiguous SNPs sites in a population. Moreover, let us denote, for each site, the two possible alleles using 0 and 1, respectively. Hence, an haplotype will be represented by a sequence of \( n \) components taken from \( \{0, 1\} \). Each genotype \( g \), being a conflation of two (partially) different haplotypes \( h_1 \) and \( h_2 \), will be represented as a sequence of \( n \) elements taken from \( \{0, 1, 2\} \), such that 0 and 1 are used for its homozygous sites, while 2 is used for the heterozygous sites.

More specifically, following [77], let us define the conflation operation \( g = h_1 \oplus h_2 \) as follows:

\[
g[i] = \begin{cases} 
    h_1[i] & \text{if } h_1[i] = h_2[i] \\
    2 & \text{otherwise}
\end{cases}
\]

where \( g[i] \) denotes the \( i^{th} \) element of the sequence \( g \), for \( i = 1, \ldots, n \).

We say that a genotype \( g \) is resolved by a pair of haplotypes \( h_1 \) and \( h_2 \) if \( g = h_1 \oplus h_2 \). A set \( H \) of haplotypes explains a given set \( G \) of genotypes, if for each \( g \in G \) there exists a pair of haplotypes \( h_1, h_2 \in H \) such that \( g = h_1 \oplus h_2 \).

Given a set \( G \) of \( m \) genotypes, the haplotype inference problem consists of determining a set \( H \) of haplotypes that explains \( G \). The cardinality of \( H \) is bound by \( 2^m \) but, in principle, each genotype having \( k \leq n \) ambiguous sites, can be explained by \( 2^{k-1} \) different pairs of haplotypes. For instance, the singleton \( G = \{212\} \) (i.e., \( k = 2 \)) can be explained in two ways, namely by choosing \( H = \{011, 110\} \) or \( H = \{010, 111\} \) (see also Figure 6).

Hence, in general, there might be an exponential number of explanations for a given set \( G \). All of them are, from the combinatorial point of view, “equivalent” and a blind algorithm—not exploiting any biological insight—may result
in inaccurate, i.e., biologically improbable, solutions. What is needed is a genetic model of haplotype evolution to guide the algorithm in identifying the “right” solution(s).

Several approaches have been proposed, relying on the implicit or explicit adoption of some kind of assumptions reflecting general properties of an underlying genetic model. We focus on one of such formulations, namely parsimony. The main underlying idea is the application of a variant of Ockham’s principle of parsimony: the minimum-cardinality possible set of haplotypes is the one to be chosen as explanation for a given set of genotypes \( G \). For instance, the set \( G \) in Figure 6 admits two explanations. The one at the bottom, i.e., \( \{010, 111, 101\} \), is preferable by the parsimony principle. In this formulation, the haplotype inference problem has been shown in [77] to be APX-hard, through a reduction from the node-covering problem.

4.2 ASP Encoding

Following [52] let us start by introducing the decision version of the haplotype inference by pure parsimony problem:

**HIPP-DEC** Given a set \( G = \{g_1, \ldots, g_m\} \) of \( m \) genotypes, each with \( n \) sites, and a positive integer \( k \), decide whether there is a collection \( H \) of at most \( k \) distinct haplotypes such that \( H \) explains \( G \).

As before, we denote genotypes and haplotypes by sequences of elements from \( \{0, 1, 2\} \) and \( \{0, 1\} \), respectively. To simplify the description, we consider \( H \) as made of \( 2m \) (not necessarily distinct) haplotypes \( h_1, \ldots, h_{2m} \) and, consequently, every \( g_i \in G \) is explained by the haplotypes \( h_{2i} \) and \( h_{2i-1} \).

Our encoding is close to the problem definition—differently from what used in [52], where efficiency considerations are also taken into account. We assume that \( G \) is given extensionally using a predicate \( g/3 \), where \( g(i, j, k) \) means that the \( i \)-th genotype \( (i \in \{1, \ldots, m\}) \) in position \( j \) \( (j \in \{1, \ldots, n\}) \) has the value \( k \) \( (k \in \{0, 1, 2\}) \). For example, the instance of Figure 6 is represented by:
Moreover, the domain predicates `geno(1..m)`, `site(1..n)`, `haplo(1..2m)` are also part of the input. In the current example we have:

`geno(1..2)`. `site(1..3)`. `haplo(1..4)`.

We propose the following ASP encoding; we define predicates `h(i,j)` where `h(i,j)` belongs to the model if and only if the corresponding haplotype `h_i` is such that `h_i[j] = 1`—for `i ∈ {1,...,2m}` and `j ∈ {1,...,n}`.

% If g(i,j)=0 or 1 assign deterministically h(2i) and h(2i-1).
(1)  h(2*I-1,J) :- g(I,J,1).
(2)  h(2*I,J) :- g(I,J,1).
(3)  :- h(2*I-1,J), g(I,J,0).
(4)  :- h(2*I,J), g(I,J,0).
% If g(i,j)=1 then exactly one between h(2i) and h(2i-1) is 1
(5)  h(2*I-1,J) :- g(I,J,2), not h(2*I,J).
(6)  h(2*I,J) :- g(I,J,2), not h(2*I-1,J).
% Check if an amino acid is a representative
(7)  representative_haplo(A) :- haplo(A), not cover_someone(A).
(8)  cover_someone(B) :- haplo(A;B), A < B, samehaplo(A,B).
% Verify if two haplotypes are the same
(9)  { samehaplo(A,B) } :- haplo(A), haplo(B), A < B.
(10) :- samehaplo(A,B), haplo(A;B), A < B, site(S), h(A,S), not h(B,S).
(11) :- samehaplo(A,B), haplo(A;B), A < B, site(S), not h(A,S), h(B,S).
% Count the number of representative and minimize it
(12) min_haplo(N) :- N = #count{ representative_haplo(A) }.
(13) #minimize[min_haplo(N)=N].

In lines (1)–(4), we deterministically require `h(2i, j)` and `h(2i−1, j)` to be in a model if `g(i, j) = 1`, and to be out of a model if `g(i, j) = 0`. In lines (5)–(6), we introduce the non-deterministic choice for `h(2i, j)` and `h(2i−1, j)`. In lines (7)–(8), we define a predicate to determine whether the haplotype `A` is a representative, namely, moving from 1 to `2m`, it is the first occurrence of the same haplotype. This definition makes use of the auxiliary predicate `same_haplotype` defined in lines (9)–(11). A minimization of the number of representative haplotypes is requested in lines (12)–(13).

### 4.3 Related Work

In [52, 24, 49], the authors present a number of heuristics aimed at establishing lower and upper bounds for the value of `k` w.r.t. the set of genotypes at hand. Experimental results (see [52, 24]) reveal that the ASP-based approach can successfully compete with approaches based on integer linear programming and constraint programming.

A number of techniques (e.g., greedy and branch-and-bound algorithms, reduction to SAT, integer programming, stochastic local search) have been applied to address the problem of haplotype inference by pure parsimony, in its
initial formulation as well as in a number of variants and refinements, such as [12, 18, 24, 40, 64, 65, 99, 115, 118]. Among the various proposals appearing in literature, we mention the rule-based method proposed by Clark [22], which in its general form suffers from the NP-hardness of the problem.

Several alternative principles, different from parsimony, have also been explored. One of them is called the infinite sites model. It postulates that, during the evolution of a population, a mutation that results in a SNP occurs only once in the history; this reflects the Dollo and Camin-Sokal principles discussed in the previous section. Hence, all individuals sharing a specific allele must be descendants of a single ancestor.

Methods relying on \textit{maximum likelihood} criteria assume that the probability of observing a genotype is strictly related to the probability of observing its constituent haplotypes. Hence, these methods estimate the probability of (each specific set $H$ of) haplotypes from observed genotypes frequencies in the real population. These methods seek the most probable solution.

Other methods make hypotheses on the historical evolution of the characters in a population. The notion of \textit{perfect phylogeny} is introduced to represent such evolution as a rooted tree (coalescent model, see [68]). The tree-structure not only imposes constraints on the candidate solutions, but also reduces the computational complexity of the problem.

For a much detailed treatment of these and other methods, we refer the interested reader to [72, 70] and to the references therein.

5 RNA secondary structure prediction

In this chapter, we focus on the prediction of the secondary structure arrangement for RNA strands. We provide a model for a simplified version of the problem and we show an elegant ASP encoding.

As anticipated in Section 2, an RNA sequence $S = s_1 \cdots s_n$ is a string of symbols from $\{A,C,G,U\}$. The sequence can fold in the space, so that pairs of bases can physically interact. The secondary structure can be described by a set of pairs of interacting bases. In Figure 7, we show an example of an RNA sequence and a set of pairs of bases (denoted by solid lines). In general, the energetic evaluation of these interactions can be computed in a precise way, regardless of the overall three dimensional arrangement of the sequence. Therefore, it is possible to compute the best secondary structure while ignoring the actual folding in space. This gives rise to specific prediction algorithms that are much more reliable when compared to the prediction tools for amino acid sequences (see also Section 6).

In order to identify the most probable secondary structure, various energetic models have been studied and efficient methods for its estimation have been proposed. In the following, we present two simple approximations of energy functions. The first one is based of the simplest approximation and maximizes the number of favorable pairings (i.e., $A-U$, $C-G$, and $U-G$). A more refined energy function scores the frequency of the pairs.
Fig. 7. An example of pairings on a sequence of length 14. Solid lines between $i$ and $j$ represent a pair between corresponding $s_i$ and $s_j$. The addition of the dashed line introduces a pseudo-knot.

Fig. 8. Example of 2D hypothetical arrangement of secondary structure

Figure 8 shows an hypothetical 2D pairing of secondary structure associated to Figure 7 without the dashed pair. The first section (bases 1–7) forms a ladder with a bulge, in this case containing only one base. This structure is referred to as an *hairpin* or *stem* loop, which is rather stable and it can roll into a double helix shape in the actual three dimensional space, just as for a DNA paired double strand.  

5.1 Modeling

Given $S = s_1 \cdots s_n$ with $s_i \in A, C, G, U$ and a partial injective function $P : \{1, \ldots, n\} \to \{1, \ldots, n\}$ such that

- $P(i) = j \iff P(j) = i$ and
- $P(i) = j \Rightarrow (s_i, s_j) \in \{(A, U), (U, A), (C, G), (G, C), (U, G), (G, U)\}$

For the sake of simplicity, we will represent $P$ as a the set of pairs $\{(i, j) \mid P(i) = j\}$. $P$ represents a possible secondary structure. The RNA secondary structure

\[\text{Figure 8 was produced using the Varna Applet } \text{http://varna.lri.fr/}.\]
prediction problem is that of finding, among all such functions $P$, the one that minimizes a given energy function. We experiment with two simple energy functions:

$$
E_1 = -|P|
$$

$$
E_2 = c_1(n - |P|) + c_2|AU - 0.35|P| + c_3|CG - 0.53|P|
$$

where $AU$ ($CG$) is the number of contacts of bases $A-U$ ($C-G$, respectively) and $c_1, c_2, c_3$ are some suitable weights.

The energy function $E_1$ was proposed by Nussinov [90] and represents the simplest approximation of the problem. It maximizes the number of admissible pairs, and thus promotes the tightest packing of the structure.

The energy function $E_2$ is an adaptation from [76, 9] and promotes a distribution of pair types as similar as possible to a typical statistical distribution retrieved from experimental observations.

5.2 ASP Encoding

Let us explore the encoding of this problem in ASP. The sequence $S = s_1 \cdots s_n$ is provided as a set of $n$ facts of the form $\text{seq}(i, s_i)$; for example:

```
seq(1,a). seq(2,g). seq(3,u). seq(4,c). seq(5,c). seq(6,a).
```

The main program models the predicate $\text{pairing}$, which represents the partial function $P$ and contains pairs of indices corresponding to interacting bases.

```
%% domain predicates
(1) sequence_index(X) :- seq(X,_).
(2) sequence_base(B) :- seq(_,B).

%% Definition of the pairing function
(3) 0 \{pairing(X,Y):sequence_index(Y)} 1 :- sequence_index(X).

%% the pairing is injective and symmetric
(4) :-sequence_index(X1;X2;Y),X1<X2,pairing(X1,Y),pairing(X2,Y).

(5) pairing(B,A):- sequence_index(A;B),pairing(A,B).

%% wrong associations
(6) wrong(X,X):- sequence_base(X).
(7) wrong(a,c). wrong(a,g). wrong(c,u).
(8) :-wrong(B1,B2),seq(X1,B1),seq(X2,B2),pairing(X1,X2).

%% each position can have at most one pairing
%% a base cannot match with itself and the successive
(9) :-sequence_index(X1), pairing(X1,X1).
(10) :-sequence_index(X1;X2), X1=X2+1, pairing(X1,X2).

%% Optional constraint: no pseudo-knots
(11) :- sequence_index(X1;X2;X3;X4), X1<X3,X3<X2,X2<X4,
    pairing(X1,X2),pairing(X3,X4).

%% Nussinov Energy E1
(12) contacts(C):- C = #count\{ pairing(A,B) \}.
(13) #maximize[contacts(C)=C].
```
The clauses in lines (1)–(2) extract domain knowledge from the input. Line (3) defines the partial function \texttt{pairing}, which is forced to be injective, by the constraint in line (4), and symmetric, by the clause in line (5). Lines (6)–(8) discard non-admissible pairs, according to the definitions in the modeling section. The constraints in line (9) and (10) forbid the pairing of a base with itself and its neighbors. The constraint in line (11) enforces the absence of pseudo-knots (see Sect. 5.3). If this constraint is removed, then the program will model the NP-complete version of the problem. The clauses in lines (12) and (13) collect and minimize the number of contacts, respectively.

Let us briefly show how to encode the energy model $E_2$. Line (13') replaces the optimization instruction (13) by asking for a minimal energy. The clauses in lines (14)–(16) define the count of the total number of AU and CG pairs in the sequence, respectively. The energy predicate, in line (18), defines the energy function $E_2$. We assign here $c_1 = c_2 = c_3 = 1$. Note that the first coefficient and lines (22)–(23) are multiplied by 100 in order to scale from floating point to integer numbers.

\begin{verbatim}
(13') #maximize[energy(E)=E].
(14) total(N) :- N=#count{ seq(X,Y)}.  
(15) au(N) :- N=#count{pairing(A,B):seq(A,a):seq(B,u)}.  
(16) cg(N) :- N=#count{pairing(A,B):seq(A,c):seq(B,g)}.  
(18) energy(E) :- C1=1, C2=1, C3=1,  
    total(N), contacts(C), au(AU), cg(CG),  
    E = C1 * (N-C/2) + C2 * #abs(100*AU - 35*C) +  
    C3 * #abs(100*CG - 53*C).
\end{verbatim}

Let us observe that the latter version of the solution encounters significant grounding problems, even for average size inputs.

### 5.3 Related work

In the literature, RNA secondary structure prediction has been addressed by a large number of proposals, presenting different algorithms and various energy functions. Starting from the seminal work by Tinoco [110], where propensity of forming helices was addressed, other popular methods include the one by Jacobson and Nussinov [90], introduced more than 30 years ago. It promotes the maximal number of admissible pairs in the sequence. This approach can be addressed by a polynomial dynamic programming approach for a restricted class of solutions (in absence of pseudo-knots, see the complexity notes below).

More refined energy functions have been proposed, with the specific goal to better estimate hairpin loops. The Zucker’s algorithm [119] proposes a more precise estimate of stacking pairs forming a hairpin loop, as well as contributions for opening and closing a loop. The computational complexity of the solution is polynomial in absence of pseudo-knots, and non-logic programming techniques, i.e., dynamic programming, have been proposed. The original algorithm proposes an energy minimization approach.
Other energy functions have been proposed as the result of probabilistic analyses. In particular, the mean distribution of base pairs found in actual RNA secondary structures has been used as the target of the secondary structure prediction. The most favorable structure is the one that best approximates such distribution [76, 9].

In order to improve the structural results, Sankoff proposed a simultaneous alignment and folding approach, which goes beyond the scope of this section [96].

In the general case, the decisional version of the problem is NP-complete—see [84] for the original proof. The main idea is to reduce a 3-SAT formula into a particular sequence such that there exists a specific number of contacts if and only if the original formula is satisfiable. An interesting polynomial class of problems has been characterized by the absence of pseudo-knots (see dashed pair drawn in Figure 7). A pseudo-knot is encountered when the following property holds: for two pairings \((i, j)\) and \((\ell, k)\) in \(P\),

\[
i < \ell < j \Rightarrow j < k \lor k < i
\]

The work in [8] provides examples of energy functions for general structures with pseudo-knots. Moreover, a constraint propagation approach for alignment and folding in the presence of pseudo-knots has been presented in [37]. For pseudo-knot free structures, dynamic programming algorithms can be used with a time complexity ranging from \(O(n^2)\) to \(O(n^4)\), depending on the energy function being considered, while space complexity ranges from \(O(n)\) to \(O(n^2)\).

6 Protein Structure Prediction

The problem we introduce here is a simplification of the protein structure prediction (PSP) problem. In particular, we consider two levels of simplification:

- **Protein model:** we consider only one atom per amino acid. Moreover, we partition the 20 amino acids in two families: \(h\) (hydropic) and \(p\) (polar);
- **Spatial model:** we focus exclusively on a simple 2D discrete lattice representation of the space.

We provide some more details on the full version of the problem and to (some) approaches for dealing with it in Subsection 6.3.

6.1 Modeling

We refer to such simplified version of the problem as the 2DHP-PSP. The problem can be formulated in a simple way as follows. The input is a list of amino acids \(S = s_1 \cdots s_n\), where \(s_i \in \{h, p\}\). Imagine \(S\) as a pearl necklace, where the \(s_i\)s are the pearls, while the distance between \(s_i\) and \(s_{i+1}\) is constant. This string of pearls lays down on a large 2D matrix, where the size of the square is exactly the distance among pearls—for the sake of simplicity, we will assume such distance to be 1—with the further constraints that \(s_i\) must occupy the intersection
of two lines, and that no two distinct $s_i$ and $s_j$ occupy the same point. We call this necklace placement a folding. In a given folding, if a pair of amino acids $s_i$ and $s_j$ are at distance 1, we say that they are in contact. We are interested in contacts between pairs of amino acids of type $h$ ($h$-contacts). The 2DHP-PSP is the problem of finding a folding that maximizes the number of $h$-contacts.

The decision version of the problem:

Given a sequence $S$ and a number $k$, is there a folding with at least $k$ $h$-contacts?

has been shown to be NP-complete in [29].

We model the 2DHP-PSP in a mathematical way as follows. Given a list $S = s_1 \cdots s_n$, with $s_i \in \{h, p\}$, a folding is a function $\omega : \{1, \ldots, n\} \rightarrow \mathbb{N}^2$ such that

1. $\forall i \in \{1, \ldots, n - 1\}$ $\text{next}(\omega(i), \omega(i + 1))$
2. $\forall i, j \in \{1, \ldots, n\}$ ($i \neq j \rightarrow \omega(i) \neq \omega(j)$)

where $\text{next}(\langle X_1, Y_1 \rangle, \langle X_2, Y_2 \rangle)$ holds if and only if $|X_1 - X_2| + |Y_1 - Y_2| = 1$. The 2DHP-PSP is the problem of finding the folding that minimizes the following energy function:

$$E(S, \omega) = \sum_{1 \leq i < j \leq n} \text{Pot}(s_i, s_j) \cdot \text{next}(\omega(i), \omega(j))$$

where $\text{Pot}(p, p) = \text{Pot}(h, p) = \text{Pot}(p, h) = 0$ and $\text{Pot}(h, h) = -1$.\(^4\)

Without loss of generality, we can consider only foldings within $[1..2n] \times [1..2n]$; moreover, to remove some symmetries in the solution space, we can set $\omega(1) = \langle n, n \rangle$ and $\omega(2) = \langle n, n + 1 \rangle$.

\(^4\) With a slight abuse of notation, we assume $\text{next}$ to return 1 when it holds and 0 when it does not.
Contiguous occurrences of \( h \) in the input sequence \( S \) (namely, when \( s_i = s_{i+1} = h \)) contribute in the same way to the energy associated to each folding and, thus, they are not considered in the objective function.

An easy observation (that unfortunately cannot be “lifted” to the general PSP formulation) is that \( s_i \) and \( s_j \) can be in contact only if \( j - i \) is odd.

6.2 ASP Encoding

The essential part of the ASP encoding of this problem is presented in [43]. A specific instance \( S = s_1, \ldots, s_n \) of the problem is represented by a set of \( n \) facts of the kind \( \text{prot}(i, s_i) \). For instance, the protein \( \\text{phphpphp} \) of Figure 9 is described as:

\[
\begin{align*}
\text{prot}(1, \text{p}). & \quad \text{prot}(2, \text{h}). & \quad \text{prot}(3, \text{p}). & \quad \text{prot}(4, \text{h}). \\
\text{prot}(5, \text{p}). & \quad \text{prot}(6, \text{p}). & \quad \text{prot}(7, \text{h}). & \quad \text{prot}(8, \text{p}).
\end{align*}
\]

The ASP code used in the encoding is reported below:

\[
\begin{align*}
% & \text{domains} \\
(1) & \text{size}(N) :- N = \#\text{count} \{ \text{prot}(I,J) \}. \\
(2) & \text{board}(1..2*N) :- \text{size}(N). \\
(3) & \text{range}(X,Y) :- \text{size}(N), \text{board}(X;Y), \#\text{abs}(X-N)+\#\text{abs}(Y-N) < N. \\
% & \text{set the first two positions} \\
(4) & \text{sol}(1,N,N) :- \text{size}(N). \\
(5) & \text{sol}(2,N,N+1) :- \text{size}(N). \\
(6) & 1 \{ \text{sol}(I,X,Y) : \text{range}(X,Y) \} 1 :- \text{prot}(I, \text{Amino}). \\
% & \text{add geometrical constraints} \\
(7) & :- \text{prot}(I1,A1), \text{prot}(I2,A2), I1< I2, \\
& \quad \text{sol}(I1,X,Y), \text{sol}(I2,X,Y), \text{range}(X,Y). \\
(8) & :- \text{prot}(I1,A1), \text{prot}(I2,A2), I2>1, \\
& \quad I1==I2-1, \text{not next}(I1,I2). \\
(9) & \text{next}(I1,I2) :- \text{prot}(I1,A1), \text{prot}(I2,A2), I1< I2, \\
& \quad \text{sol}(I1,X1,Y1), \text{sol}(I2,X2,Y2), \\
& \quad \text{range}(X1,Y1), \text{range}(X2,Y2), \\
& \quad I1==\#\text{abs}(Y1-Y2)+\#\text{abs}(X2- X1). \\
% & \text{Count the number of h-h energy_pairs and minimize it} \\
(10) & \text{energy_pair}(I1,I2) :- \text{prot}(I1,h), \text{prot}(I2,h), \\
& \quad \text{next}(I1,I2), I1+2<I2, \text{I}=(I2-I1) \#\text{mod} 2. \\
(11) & \text{energy_value}(N) :- N = \#\text{count} \{ \text{energy_pair}(I1,I2) \}. \\
(12) & \text{#maximize}[\text{energy_value}(N)=N].
\end{align*}
\]

Rules (1)–(3), together with the predicate \text{prot}, define the domains of the variables to be used later. Precisely, rule (2) and (3) state the admissible values for \(x\) and \(y\) coordinates: it is easy to see that the Manhattan distance w.r.t. the initial point (set in rule (4)) is less than \(N\). We aim at defining the solution \(\text{sol}\) predicate that states, for each amino acid \(1, \ldots, n\) its spatial position. Rules (4) and (5) fix the positions of the two initial amino acids. Rule (6) states that each amino acid occupies exactly one position. The ASP-constraints (7) and (8) state that there are no self-loops and that two contiguous amino acids must satisfy
the next property. Rule (9) defines the next relation, also including the odd property of the lattice. The objective function is defined by Rule (10) and (11), which determines the energy contribution of the amino acids, and rule (12), that searches for answer sets maximizing the energy.

6.3 Related work

The literature on the protein structure prediction problem is extensive (e.g., see some reviews [87, 57, 34]). We just focus on some approaches based on logic and/or constraint programming. The 2DHP version of the problem is just a toy model that, however, can be used to understand the difficulty of combining a self-avoiding-walk with a non linear energy function (see running times in the previous section). The HP function was introduced by Dill [41] to model the fact that polar (p) amino-acids tend to stay outside (in contact with water) and, as a consequence, hydrophobic (h) amino-acids tend to stay in contact inside the protein.

A 3D lattice model for the HP problem (FCC) was introduced by Backofen and Will [3, 4]; using clever symmetry breaking results and a geometric notion of allowed cores, they are able to find the best folding for proteins of length 200 and more. In [32] the authors generalizes their approach using a Prolog encoding with a more precise contact energy function. An ad-hoc constraint solver for that modeling was then developed [35]. In [33] the same authors consider an orthogonal view of the same problem, namely the approach is that of assembly admissible fragments of the protein. The notion of a discrete lattice is no longer needed and results are more realistic, even remaining in a discrete world (the number of allowed fragments for each subset of a protein is finite). An ad-hoc constraint solver for that modeling has been described in [17].

7 Systems Biology

A cell can be considered as a complex system composed of different interacting components. Such components can be small molecules (e.g., water, amino acids, nucleotides, etc), DNA, RNA, and proteins. The interactions among these elements determine the diverse cellular functions and can be situated and analyzed at different levels/stages of the process that, starting from DNA replication, ends with protein synthesis. Each of these stages are usually modeled by means of some kind of network. At the DNA level, transcription factors control the transcription of genes in mRNA synthesis (transcriptional regulatory network). This affects the activities of genes through a so-called gene regulatory network (notice that genes, in turn, control the transcription factors). At the protein level several proteins can interact and affect the products of the transcription step and form protein complexes. These interactions are modeled by protein interaction network. Biochemical reactions occurring in the cell and involving metabolites (such as enzymes, substrates, etc.) constitute the metabolic network. Signaling networks are introduced to model the complex processes that took place in a
cell in order to receive different signals from the cellular environment and, possibly, from other cells. Figure 10 intuitively illustrates these networks and their relationships.

![Fig. 10. Various kinds of network models used to represent, at different granularity levels, the interactions between molecules in a cellular system.](image)

A typical problem in system biology consists of developing a network model by combining existing biological knowledge. This model has to be validated with respect to experimental data. Often, because of incompleteness in the biological knowledge and partial unreliability of experiments' outcome, the model has to be checked for consistency, modified, repaired, or extended. On the other hand, analysis of the model may give hints on how to design new experiments.

In this section we describe some of the declarative approaches aimed at automating specific steps of the process of model definition and refinement.

7.1 Gene Regulatory Networks

7.1.1 Modeling

A gene regulatory network is a directed graph having genes as nodes and their relationships as edges. The most commonly used model to relate the concentrations of network constituents (genes, metabolites, proteins, etc.) and their evolution is derived from (systems of) ordinary differential equations. In many cases, the incompleteness of information about biochemical reactions makes the task of defining such equations hard. Moreover, in a simplified setting, one may be interested in abstracting from quantitative aspects and introducing a qualitative model. Such model should focus on the controlling relationships among
genes/nodes of the network, namely, activations and inhibitions of genes’ expression as effect of other genes’ expressions.

Fig. 11. An influence graph modeling lactose operon in Escherichia coli

One of the simpler proposed models makes use of the so-called influence graph. In such network model, the edges are labeled either by + or by −, to represent the positive, resp. negative, influence between constituents. Formally, an influence graph is a directed graph $G = (N, E, \sigma)$, where $N$ is a set of nodes, $E$ is a set of edges, and $\sigma : E \to \{+,-\}$ is a (possibly partial) labeling of the edges.

Graphically, the edges labeled + are represented as arrows while those labeled − have a tee head. Figure 11 describes a simplified influence graph for the lactose operon in Escherichia coli [61]. In this graph, for example, the node $L_e$ is not regulated by other constituents, hence it is considered as an input of the system. By increasing or decreasing the levels of input constituents one can observe how the system evolves as result of reciprocal activations and inhibitions of nodes. This represents a (simplified) simulation of the modeled biological system. Since each node may be either active or inhibited, there exists only a finite number of possible states of the network. Thus, such simulation must end in a steady state or, otherwise, enter a cycle of states. Viewed from the qualitative perspective, at a higher abstraction level, steady states can be considered as counterparts for the solutions of the system of differential equations.

In reasoning on biological networks represented as influence graphs, one often adopts the so-called Sign Consistency Model. After a model has been designed and its steady state(s) are determined (corresponding to different input configurations), we may want to verify if its behavior reflects the outcome of real experiments. At this abstraction level, such outcome is specified in terms of a node labeling $\mu$ that labels constituents by + (resp., −), depending on the vari-
ations of their concentrations, as exhibited by experimental data. Consistency of the model with respect of the experimental data is formalized as follows.

Let $G = (N, E, \sigma)$ be an influence graph and $\mu : N \rightarrow \{+, -\}$ a (partial) node labeling. Then, $G$ and $\mu$ are consistent if there are extensions $\sigma'$ of $\sigma$ and $\mu'$ of $\mu$ such that, for each non-input node $n \in N$, there is an edge $e = (m, n) \in E$ such that $\sigma'(e) \circ \mu'(m) = \mu'(n)$, where the usual rule for signs, namely $+ \circ + = - \circ - = +$ and $- \circ + = + \circ - = -$, is adopted.

Even in this simplified setting, the problem of establishing consistency of a model w.r.t. a data set is not trivial. In [114], for instance, it is shown that in presence of incomplete information the decision problem is NP-hard.

In case inconsistency is detected, this can be caused by missing or inaccurate knowledge involved in the design of the network and by the presence of partially unreliable data. In such a situation, one may ask whether it is possible to reconcile model and data. A possibility consists in applying some changes in the network model or in the data (or both). Changing the data is an attempt in singling out those portions of the experimental outcome to be considered as unreliable. A modification of the network should yield a more refined biological model. Finding a set of such repairing operations consists in solving the repair problem. This is usually solved with respect to a collection of experimental profiles—observations or experimental evidences—represented as a collection of different node labeling.

Considering a set of experimental profiles, a typical repair operation consists of: 1) introducing a new edge in the network; 2) flipping the label of an edge; 3) considering a node to be an input node in all experimental profiles; 4) considering a node to be an input node in a specific profile; 5) flipping the label for a node in a specific profile. 1) and 2) correct incompleteness and inaccuracy of the model; 3) indicates that for that node some form of regulation is missing in the model; 4) and 5) revise the experimental observations.

We are interested in applying a minimal number of repair operations. Thus we are looking for repairs that are minimal with respect to some criteria. Two possibilities explored in literature [59], are subset-minimality and cardinality-minimality. As noted in [59], both choices are not computationally trivial, since evaluating subset-minimality and cardinality-minimality involves solving problems that are, in general, in $\Sigma^P_2$ and in $\Delta^P_2$, respectively.

It is possible that many repairs exist for reconciling the same inconsistency between a network and a set of profiles. We may be interested in detecting those parts in the edge/node labeling that are common to all these solutions. In other words, we are interested in determining the consequences of each minimal repair. This consists of solving the prediction problem.

7.1.2 ASP Encoding

The following encoding is inspired from [61, 59] but it avoids the use of disjunctive heads and it deals with consistency detection and with a simple notion of repairing.
The program aims to compute the predicates *label_vertex* and *label_edges* (although typically edges are already labeled). The input of the problem is given through a set of facts of the following kind: *vertex*(Vertex Name), *edges*(From Vertex, To Vertex), *input*(vertex Name)–for input vertices, and a set of observation for nodes and edges of the form *observed*(Vertex name, Sign) (for nodes) and *observed*(From Vertex, To Vertex, Sign) (for edges), where Sign is either plus or minus.

% domain predicates
(1) sign(minus). sign(plus).
(2) opposite(minus,plus). opposite(plus,minus).
% Non deterministic labels for nodes and edges
(3) 1 {label_vertex(V,S): sign(S)} 1 :- vertex(V).
(4) 1 {label_edge(U,V,S): sign(S)} 1 :- edge(U,V).
% choice for reversing an edge
(5) {wrong(U,V)} :- edge(U,V).
% labeling nodes consistent with observation
(6) label_vertex(V,S) :- observed(V,S).
% labeling edges consistent with observation, if possible
(7) label_edge(U,V,S) :- wrong(U,V), observed(U,V,T), opposite(S,T).
(8) label_edge(U,V,S) :- not wrong(U,V), observed(U,V,S).
% Rules of signs
(9) receive(V, plus) :- edge(U,V), sign(S),
    label_edge(U, V, S), label_vertex(U, S).
(10) receive(V, minus) :- edge(U,V), opposite(S,T),
     label_edge(U, V, S), label_vertex(U, T).
% All vertices (but inputs) must be labeled in justified way
(11) :- label_vertex(V, S), not receive(V, S), not input(V).
% Minimize edge reversing to guarantee consistency
(12) edges_reversed(N) :- N = #count{ wrong(U,V) }.
(13) #minimize [edges_reversed(N)=N].

After defining the domain predicate *sign* in line (1), and the predicate opposite between signs, the mutual exclusion between plus and minus for vertices and edges is stated in lines (3) and (4). The choice for the predicate *wrong* stating, intuitively, that an edge should be reversed is added in line (5). The labeling we are looking for should be consistent with known observations. In particular, node labeling is required to be consistent in line (6) and the edge labeling is consistent if the edge is not wrong, otherwise the complementary value is chosen (lines (7) and (8)). Rules of signs propagation are stated in lines (9) and (10), using the auxiliary predicate *receive*. Then, in line (11) it is stated that the labeling of the non-input vertex must be justified by the predicate *receive*, namely by the rule of signs. Lines (12) and (13) are inserted for looking for the answer set with a minimum number of wrong edges. If *edges_reversed(0)* is in the result, then the network is already consistent.
7.2 Metabolic Networks

7.2.1 Modeling

Large amounts of data are made available through experiments, making possible the study of the dynamic metabolic behavior of living cells (or systems of cells) in response to perturbations and stimuli coming from their surrounding environment. This is usually done by analyzing metabolic networks. In such networks, the nodes represent the metabolites while the edges represent the reactions. A metabolic network describes a collection of metabolite components together with a functional readout of the cellular state. The modeling and the reconstruction of biochemical reaction pathways/networks are far from being easy tasks, because of the complexity of the molecules and the reactions that may take place. This fact calls for sophisticated and refined computational techniques for reconstructing and simulating metabolic networks.

Several formal techniques and approaches have been adopted to model and reason about metabolic networks. Among them, we find Petri nets, Flux balance analysis, and Process calculi (see [20], and the references therein, for further details).

In the rest of this section we describe a qualitative approach based on ASP to study this class of problems. We rely on the notion of metabolic network borrowed from [89, 97, 25]. Intuitively speaking, a metabolic network models situations where a reaction can only occur if all of its substrates are available as nutrients or can be produced by other reactions. Starting from some initially given nutrients, called seeds, the network is expanded by adding all of the enabled reactions together with their products. Such a process iterates until no further reactions can occur. The set of metabolites in the expanded resulting network is called the scope of the given seeds. It represents all the metabolites that can be, potentially, synthesized from the seeds by the given network.

We adopt the following definition: a metabolic network is a directed bipartite graph \( G = (R \cup M, E) \) where \( R \) and \( M \) are sets of nodes representing reactions and metabolites, respectively, and \( E \subseteq (R \times M) \cup (M \times R) \).

Given an edge \( (m, r) \in E \), the metabolite \( m \in M \) is called reactant of the reaction \( r \in R \). Similarly, for an edge \( (r, m) \in E \), the metabolite \( m \in M \) is called product of the reaction \( r \). We say that a reaction \( r \) is reachable from a set of metabolites \( S \) if \( S \) contains all the reactants of \( r \). Moreover, we say that a metabolite \( m \) is reachable from a set of metabolites \( S \) if either \( m \in S \) or \( m \) is a product of some reaction \( r \) reachable from \( S \). We define the scope of a set \( S \) of metabolites in the network \( G \) (briefly, \( \Sigma_G(S) \)) to be the set of metabolites that are (transitively) reachable from \( S \).

The authors of [97, 25] propose ASP-based solutions for two specific reasoning tasks on metabolic networks: metabolic network completion and inverse scope problem.

The metabolic network completion problem can be stated as follows. Let us consider a metabolic network \( G = (R \cup M, E) \), two sets \( S, T \subseteq M \) of seed and target metabolites, and a reference metabolic network \( (R', M', E') \). We wish to
find a set of reactions $R'' \subseteq R' \setminus R$ such that $T \subseteq \Sigma_G(S)$, where:

$$G = ((R \cup R'') \cup (M \cup M''), E \cup E'')$$

$M'' = \{ m \in M' \mid r \in R'' \text{ and } m \text{ is a reactant or a product of } r \}$

$E'' = \{ (m,r) \in E' \mid r \in R'' \text{ and } m \text{ is a reactant of } r \} \cup$

$\{ (r,m) \in E' \mid r \in R'' \text{ and } m \text{ is a product of } r \}$

The set $R''$ is the completion of $(R \cup M, E)$ from $(R' \cup M', E')$ w.r.t. $S, T$.

For instance, let us consider the network in Figure 12. It represents the reference network $(R' \cup M', E')$, while $G$ is its subgraph where $R = \{r_3,r_4,r_6\}$ (the dark nodes), $M = M'$ and $E'$ is the subset of edges that concerns nodes in $R$. If $S = \{m_1,m_2\}$ and $T = \{m_5,m_7\}$, then further reactions should be used (i.e., inserted in $R''$). A solution with a minimal size of new reactions is $R'' = \{r_1\}$.

Variants of this problem impose minimality requirements on the set $R''$, typically asking for subset-minimal or cardinality-minimal solutions. Other variants minimize the production of metabolites occurring in a set of “forbidden” products. Moreover optimality can be imposed on the distance (i.e., path length) between seeds and scope.

The inverse scope problem can be formulated as follows. Given a metabolic network $G = (R \cup M, E)$ and a set $T \subseteq M$ of target metabolites, find a set of seed metabolites $S \subseteq M$ such that $T \subseteq \Sigma_G(S)$. Also in this case minimality requirements can be imposed to obtain optimized versions of the problem, as well as specification of “forbidden” products.

As regards computational complexity, it can be shown that, in general, the problem of metabolic network completion, as well as the inverse scope problem and other related problems, are NP-hard [89, 88].

7.2.2 ASP Encoding

We report here a simplified encoding of the problem of metabolic network com-
pletion. The interested reader can refer to [97] for a description of variants of this encoding as well as for the ASP formulation of the inverse scope problem.

The instance of the problem is described by the specification of two networks. This is done using facts of the forms:

- **draft(Net)** to identify the network named Net as the network $G$.
- **reaction(R,Net)** states that the reaction node is in $G$;
- Other reactions of $G'$ are defined by **reaction(R,a)** (a stands for “all”; w.l.o.g. let us assume that the network name Net is not a, and that it is different from the symbol x, as well).
- Edges are represented by facts of the form **reactant(M,R)** and **product(M,R)** to specify the topology of the network.
- **seed(S)** and **target(T)** specify seeds and targets.

For instance, considering again the network in Figure 12, one can state **draft(d)** to identify $G$, along with the facts

$$\text{reaction(r1, a). reaction(r2, a). reaction(r3, d). ...}$$

The graph is given by facts of the form:

$$\text{reactant(m1, r1). reactant(m2, r2). ...}$$
$$\text{product(m3, r1). product(m3, r2). ...}$$

Finally, seeds and target are defined as

$$\text{seed(m1). seed(m2). target(m5). target(m7).}$$

The following code implements the solution to the the problem of metabolic network completion.

```prolog
%%% Type of reaction. d: graph G, a: all, x: reactions added
(1) type(Net) :- draft(Net).
(2) type(a). type(x).
%%% Extended graph.
(3) reaction(R,x) :- reaction(R,Net), draft(Net).
(4) { reaction(R,x) } :- reaction(R,a).
%%% reachability predicate
(5) scope(M, T) :- seed(M), type(T).
(6) scope(M, T) :- type(T), product(M,R), reaction(R,T), scope(M2,T) :- reactant(M2,R).
(7) :- target(M), not scope(M,x).
%%% new reactions and their minimization
(8) new(R) :- reaction(R,x), draft(Net), not reaction(R,Net).
(9) reactions(S) :- S = #count{ new(R) }.
(10) #minimize[reactions(S)=S].
```

In the above code, lines (1)–(2) define the three types of reactions, d stands for the network $G$, a stands for the reference network, and x will denote the reactions of the intermediate network $G''$ we are looking for. Lines (3)–(4) define nondeterministically which reaction can be considered by the intermediate network $G''$. All reactions of $G$ are included, while those in $G' \setminus G$ can be considered or
not. Lines (5) and (6) define the reachability relation. In particular, let us observe the use of the conditional literal \( \text{scope}(M_2, T) : \text{reactant}(M_2, R) \) in the body of the rule of line (6). It is a shorthand for \( \text{scope}(m_1, T), \ldots, \text{scope}(m_n, T) \) where \( m_1, \ldots, m_n \) are all the metabolites that are reactants of the reaction \( R \) active in the network identified by \( T \). This kind of syntactic extension to ASP is replaced by the proper list of usual literals by the grounding stage. The constraint in line (7) imposes that all targets must be obtained in the extended network. Finally, lines (8)–(10) ask for a cardinality-minimal set of new reactions.

### 7.3 Related work

It is practically impossible to compile here a comprehensive bibliography on Systems Biology. We refer the reader to [44], which provides a general overview of the field and an extensive bibliography, and to [20] for a network-oriented perspective. Let us focus on the declarative approaches developed in the subfields of Systems Biology treated in this chapter.

In this section, we described the approach relying on the notion of Influence Graph and the Sign Consistency Model for gene regulatory networks. Other methods have been studied; for instance, [28, 27, 19, 26, 58] adopt, and in some cases extend, the logical formalism introduced in [104, 105, 106]. Notions of network, such as regulatory graphs and interaction graphs [54], are defined similarly to influence graphs, but by admitting numerical labeling or activation/inhibition thresholds. Implementations have also been developed, such as SysBiOX [27] and PINT [58], exploiting computational logic solvers, mainly based on SAT, ASP, or CSP.

Boolean networks have been also used, for instance in [71], to declaratively tackle the learning problem for biological network models. The proposal exploits an ASP-solver as reasoning engine. Similarly, [54] relies on ASP to model and reason about protein interactions, while [93] integrates numerical weights in Boolean networks in order to introduce a notion of preference among solutions. Another extension of Boolean networks is used in [55, 56] to develop an ASP-based system to model regulatory networks with meta-interactions. The implementations of a collection of deductive tasks for reasoning on biological networks, developed by the authors of [71, 45, 59, 61] (including those presented in this section), are described in [60].

The use of Action Description Languages to solve different computationally hard problems in Systems Biology has been explored in [112, 111, 6, 7, 113]. The authors also realized a tool, named BioSigNet-RR, exploiting such ideas. Similar approaches to signaling, metabolic, and regulation networks have been proposed in [46, 47], among others, by introducing the ADL $C_{TAID}$ and the system BioC.

In [16] a software environment for modeling biochemical systems (BIOCHAM) has been described. It allows the analysis and simulation of Boolean, kinetic, and stochastic models and the formalization of biological properties in temporal logic. Parts of the tool are implemented in logic programming.
8 Conclusions

This chapter reviews some classical problems that arise in Bioinformatics, namely Phylogenetics, Haplotype, RNA secondary structure prediction, protein structure prediction, and systems biology. The chapter shows how logic programming can be employed to model these problems by describing the properties of the system, rather than the process of computing the problem’s solutions. The problems have been modeled by means of Answer Set Programming, a logic programming language that allows concise and versatile programming style. The reader is guided into the modeling process and the intuitive encoding into ASP. The relatively short encodings are fully functional and they can be tested in conjunction with input reported in the Appendix.

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References


A Instances

For readers’ convenience we report here some instances on which run the ASP encodings presented in the paper (just cut & paste from the pdf—line numbers should be removed from ASP codes). Expected running time will depend on the machine of course, but instances are chosen to run in at most a couple of minutes.

A.1 Phylogenetic Reconstruction

We report here the instance used in Figure 4 and other two simple instances:

- taxa(1..5).
  % characters and values on leaves (function f)
  f(1,coelom,0). f(2,coelom,0). f(3,coelom,0). f(4,coelom,1). f(5,coelom,1).
  f(1,dark,1). f(2,dark,0). f(3,dark,0). f(4,dark,0). f(5,dark,1).
- taxa(1..5).
  f(1,a,1). f(2,a,0). f(3,a,0). f(4,a,0). f(5,a,1).
  f(1,b,0). f(2,b,1). f(3,b,1). f(4,b,1). f(5,b,0).
  f(1,c,1). f(2,c,1). f(3,c,0). f(4,c,0). f(5,c,1).
  f(1,d,1). f(2,d,1). f(3,d,1). f(4,d,0). f(5,d,0).
- From http://evolution.berkeley.edu/evolibrary/article/phylogenetics_07

  taxa(1..7).
  % 1 = Sharks and Relatives
  f(1,vertebrate,1). f(1,skeleton,0). f(1,fourlimbs,0).
  f(1,amnioegg,0). f(1,hair,0). f(1,twopostorbital,0).
  % 2 = Ray-finned fishes
  f(2,vertebrate,1). f(2,skeleton,1). f(2,fourlimbs,0).
  f(2,amnioegg,0). f(2,hair,0). f(2,twopostorbital,0).
  % 3 = Amphibians
  f(3,vertebrate,1). f(3,skeleton,1). f(3,fourlimbs,1).
  f(3,amnioegg,0). f(3,hair,0). f(3,twopostorbital,0).
  % 4 = Primates
  f(4,vertebrate,1). f(4,skeleton,1). f(4,fourlimbs,1).
  f(4,amnioegg,1). f(4,hair,1). f(4,twopostorbital,0).
  % 5 = Rodents and rabbits
  f(5,vertebrate,1). f(5,skeleton,1). f(5,fourlimbs,1).
  f(5,amnioegg,1). f(5,hair,1). f(5,twopostorbital,0).
  % 6 = Crocodiles and relatives
  f(6,vertebrate,1). f(6,skeleton,1). f(6,fourlimbs,1).
  f(6,amnioegg,1). f(6,hair,0). f(6,twopostorbital,1).
  % 7 = Dinosaurs and birds
  f(7,vertebrate,1). f(7,skeleton,1). f(7,fourlimbs,1).
  f(7,amnioegg,1). f(7,hair,0). f(7,twopostorbital,1).
A.2 Haplotype Inference

We just report one instance with $|G| = 20$ and $n = 8$. To test other instances just commute turn 2 into 0 or 1 or viceversa.

\[
g(1,1,2). g(1,2,1). g(1,3,1). g(1,4,0). g(1,5,0). g(1,6,0). g(1,7,0). g(1,8,0).
g(2,1,2). g(2,2,0). g(2,3,1). g(2,4,1). g(2,5,2). g(2,6,2). g(2,7,2). g(2,8,0).
g(3,1,2). g(3,2,0). g(3,3,1). g(3,4,2). g(3,5,2). g(3,6,0). g(3,7,2). g(3,8,0).
g(4,1,2). g(4,2,2). g(4,3,2). g(4,4,2). g(4,5,1). g(4,6,2). g(4,7,2). g(4,8,0).
g(5,1,1). g(5,2,2). g(5,3,0). g(5,4,1). g(5,5,0). g(5,6,0). g(5,7,1). g(5,8,0).
g(6,1,1). g(6,2,2). g(6,3,0). g(6,4,2). g(6,5,2). g(6,6,0). g(6,7,1). g(6,8,2).
g(7,1,1). g(7,2,2). g(7,3,2). g(7,4,2). g(7,5,1). g(7,6,2). g(7,7,2). g(7,8,2).
g(8,1,1). g(8,2,1). g(8,3,2). g(8,4,2). g(8,5,1). g(8,6,2). g(8,7,2). g(8,8,2).
g(9,1,1). g(9,2,1). g(9,3,2). g(9,4,1). g(9,5,1). g(9,6,2). g(9,7,2). g(9,8,1).
g(10,1,1). g(10,2,0). g(10,3,2). g(10,4,2). g(10,5,1). g(10,6,0). g(10,7,1). g(10,8,1).
g(11,1,2). g(11,2,1). g(11,3,1). g(11,4,0). g(11,5,2). g(11,6,0). g(11,7,2). g(11,8,1).
g(12,1,2). g(12,2,0). g(12,3,0). g(12,4,2). g(12,5,1). g(12,6,1). g(12,7,0). g(12,8,1).
g(13,1,2). g(13,2,0). g(13,3,1). g(13,4,2). g(13,5,2). g(13,6,0). g(13,7,2). g(13,8,2).
g(14,1,2). g(14,2,2). g(14,3,2). g(14,4,2). g(14,5,1). g(14,6,2). g(14,7,2). g(14,8,0).
g(15,1,1). g(15,2,2). g(15,3,0). g(15,4,1). g(15,5,2). g(15,6,0). g(15,7,1). g(15,8,0).
g(16,1,1). g(16,2,2). g(16,3,0). g(16,4,2). g(16,5,2). g(16,6,0). g(16,7,1). g(16,8,0).
g(17,1,1). g(17,2,2). g(17,3,2). g(17,4,2). g(17,5,1). g(17,6,2). g(17,7,2). g(17,8,0).
g(18,1,1). g(18,2,2). g(18,3,2). g(18,4,2). g(18,5,1). g(18,6,2). g(18,7,2). g(18,8,0).
g(19,1,1). g(19,2,2). g(19,3,2). g(19,4,1). g(19,5,1). g(19,6,2). g(19,7,2). g(19,8,1).
g(20,1,1). g(20,2,0). g(20,3,0). g(20,4,2). g(20,5,1). g(20,6,0). g(20,7,1). g(20,8,1).
geno(1..20). site(1..8). haplo(1..40).
\]

A.3 RNA secondary prediction

Try these instances with the two different energy functions.

1. $E_1$: 18 contacts, $E_2$: 16 contacts.

\[
\text{seq(1,a). seq(2,c). seq(3,g). seq(4,a). seq(5,a). seq(6,a).}
\text{seq(7,u). seq(8,c). seq(9,g). seq(10,a). seq(11,a). seq(12,a).}
\text{seq(13,c). seq(14,g). seq(15,c). seq(16,c). seq(17,c). seq(18,a).}
\text{seq(19,u). seq(20,u). seq(21,u). seq(22,u). seq(23,g). seq(24,u).}
\]

2. $E_1$: 16 contacts, $E_2$: 14 contacts.

\[
\text{seq(1,c). seq(2,c). seq(3,a). seq(4,a). seq(5,g). seq(6,a).}
\text{seq(7,u). seq(8,g). seq(9,u). seq(10,g). seq(11,g). seq(12,a).}
\text{seq(13,g). seq(14,g). seq(15,c). seq(16,u). seq(17,g). seq(18,g).}
\text{seq(19,g). seq(20,g). seq(21,u). seq(22,c). seq(23,a). seq(24,g).}
\]

A.4 Protein Folding

1. The first instance has many equivalent solutions, but it is easy to understand that the first one cannot be improved, therefore the search is very fast

\[
% hppppppppppppppph
\text{prot(1,h). prot(2,p). prot(3,p). prot(4,p).}
\text{prot(5,p). prot(6,p). prot(7,p). prot(8,p).}
\text{prot(9,p). prot(10,p). prot(11,p). prot(12,p).}
\text{prot(13,p). prot(14,p). prot(15,p). prot(16,h).}
\]
2. Here the $h$ stay in a a central core. The search concludes in less than one minute with a 6-contact solutions. You can extend with $h(pph)^n$ obtaining nice results.

```prolog
% hpphppppppppppp
prot(9,p). prot(10,h). prot(11,p). prot(12,p).
```

3. This instance has several equivalent best solutions but it is not easy to understand that a best solution cannot be improved. Call it with length 12 (as stated), then increase.

```prolog
% hhhhhhhhhhhhh
prot(1,h). prot(2,h). prot(3,h). prot(4,h).
prot(5,h). prot(6,h). prot(7,h). prot(8,h).
prot(9,h). prot(10,h). prot(11,h). prot(12,h).
```

A.5 Inconsistencies in Networks

This is the instance used in [61].

```prolog
vertex(le). vertex(li). vertex(g). vertex(lacY).
vertex(lacZ). vertex(lacI). vertex(a). vertex(cAMPcRP).

% input(le). input(g).

% edge(le,li). edge(li,g). edge(lacY,le). edge(lacY,li).
edge(lacZ,a). edge(a,lacI). edge(g,cAMPcRP). edge(cAMPcRP,lacY).
edge(cAMPcRP,lacZ).

% observed(le,li,plus). observed(li,g,plus). observed(lacY,le,minus).
observed(lacY,li,plus). observed(lacI,lacY,minus). observed(lacI,lacZ,minus).
observed(lacZ,li,minus). observed(lacZ,g,minus). observed(lacZ,a,plus).
observed(a,lacI,minus). observed(g,cAMPcRP,minus). observed(cAMPcRP,lacY,plus).
observed(cAMPcRP,lacZ,plus).
```

Different results can be obtained by adding different observations on nodes. For instance with the following input, a minimal solution with two edge revered is obtained:

```prolog
observed(le,plus). observed(li,plus).
observed(g,plus). observed(lacY,plus).
observed(lacZ,plus). observed(lacI,plus).
observed(a,plus). observed(cAMPcRP,plus).
```

while giving the following partial observations the network is consistent and an answer set with 0 changes is found.

```prolog
observed(le,plus). observed(g,plus).
```
A.6 Metabolic Networks

This is the complete instance of the example in Figure 12.

draft(d).
reaction(r1, x). reaction(r2, x). reaction(r3, d).
reaction(r4, d). reaction(r5, x). reaction(r6, d).
reactant(m1, r1). reactant(m2, r2). reactant(m2, r3).
reactant(m3, r4). reactant(m4, r4). reactant(m1, r5).
reactant(m4, r5). reactant(m4, r6).
product(m3, r1). product(m3, r2). product(m4, r3).
product(m5, r4). product(m6, r5). product(m7, r6).
seed(m1). seed(m2). target(m5). target(m7).

The following example is from Sven Thiele PhD thesis [103, p. 42].

draft(d).
reaction(r1, d). reaction(r2, d). reaction(r3, d).
reaction(r4, d). reaction(r5, d). reaction(r6, d).
reaction(r7, a). reaction(r8, a). reaction(r9, a).
reaction(r10, a).
reactant(m2, r1). reactant(m7, r2). reactant(m8, r2).
reactant(m3, r3). reactant(m4, r3). reactant(m6, r4).
reactant(m9, r4). reactant(m9, r5). reactant(m12, r6).
reactant(m13, r7). reactant(m12, r8). reactant(m15, r9).
reactant(m1, r10).
product(m4, r1). product(m3, r1). product(m9, r2).
product(m5, r3). product(m6, r3). product(m10, r4).
product(m10, r5). product(m11, r5). product(m13, r6).
product(m5, r7). product(m15, r8). product(m10, r9).
product(m7, r10). product(m8, r10).
seed(m1). seed(m2). seed(m12).
target(m5). target(m10).