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Abstract: We show that a general purpose graph rewriting system (GRS) nowadays is capable of simulating gene expression, the intra-cellular synthesis of proteins. The model organism we use is one of the best-studied prokaryotic life-forms in genetics, the *E. coli* bacteria. Our graph representation of the *E. coli* DNA consists of 23 million graph elements. In our case study we correctly synthesize the proteins of 30 consecutive genes. In this paper we describe our approach as well as our observations. Further on, we discuss some potential extensions to GRSs that would support more sophisticated simulations.

Keywords: Graph Rewriting, Biology, Genetics, Gene Expression

1 Introduction

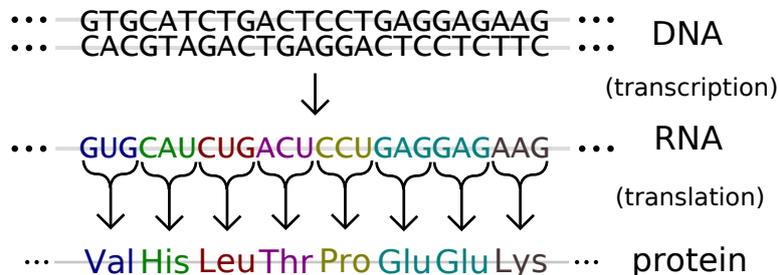
Current life science research aims at understanding the life cycle process of a human cell¹. Today, highly optimized string-based approaches exist to compute gene expression products. Since more sophisticated simulations considering chemical behavior can effectively not be done with string-based representations, several authors propose graph-based techniques. The recently increased performance of graph rewriting systems suggests the applicability of this alternative.

In this paper, we present a case study in which we investigated the simulation of gene expression using the general purpose graph rewriting system GrGen.NET [BG07]. Simulating the human genome is beyond the capabilities of graph rewriting on current PCs. Therefore, our subject was the (considerably smaller) DNA of *E. coli* – the simulation steps are similar. In comparison with the natural archetype, our simulation has several simplifications. In fact, our simulation does not outperform the string-based approaches in terms of ‘functionality’. However, the intent of this case study was to investigate whether and how such simulations are feasible using graph rewriting techniques. The results show that the graph-based approach is appropriate and performs sufficiently well.

2 Gene Expression Basics

All life forms are based on a large macro molecule called *deoxyribonucleic acid* (DNA). Gene expression is the biological process of creating proteins using the information stored in the DNA as a blueprint. The DNA itself is a concatenation of molecules called *nucleotides*. Four different types of nucleotides occur in the DNA: *Adenin* (A), *Cytosin* (C), *Guanin* (G) and *Thymin* (T). DNA is double stranded: The nucleotides of two strands are kept together by a chemical binding.

¹ For instance, the possibility to entirely simulate the genetic processes within a human cell is expected with the computing power of one ExaFLOP. Intel expects computers with this processing power around the year 2017 [Gel08].

Figure 1: Gene expression depicted⁵.

In opposing strands, Adenin binds to Thymin, while Cytosin binds to Guanin. Therefore, the nucleic sequence of one strand can easily be deduced from the other.

Another molecule that is involved in gene expression is the *ribonucleic acid* (RNA). It has the same structure as DNA, except that Thymin is replaced by the nucleotide *Uracil* (U). RNA is (usually) single-stranded.

A protein is built from molecules called *amino acids*. Twenty different types of amino acids exist, which can form to large strings. Such an amino acid sequence is called a protein's *primary structure*. For the protein to be fully functional, the three-dimensional layout of the amino acid chain matters. The three-dimensional layout of a protein is called its *secondary* (and *tertiary*) *structure*. In this study, we only simulate the primary structure².

To store the primary structure, each amino acid is encoded using three nucleotides (so called *triplets* of A, C, G, or T). Three consecutive nucleotides enable 64 different combinations, whereas only 20 amino acids exist: Several triplets represent the same amino acid, while others are reserved as special 'commands'³. In fact, the additional combinations are chosen in such a way that a) a possible mutation (i. e. alteration of a nucleotide) is likely to represent the original amino acid, or b) the mutation is at least likely to represent an amino acid with a similar chemical behavior, preserving the functionality of the encoded protein. A DNA subsequence that encodes a protein is called a *gene*⁴.

The process of actually creating a protein from a gene is called *gene expression* and is done in two steps, *transcription* and *translation*, as depicted in Figure 1. Both will be digested in the following sections.

2.1 Transcription

In order to create a protein from a gene, the gene's code has to be copied. This is performed by an enzyme called *polymerase*. The polymerase attaches to the start of the gene and slides along the

² The process of bringing proteins into their correct three-dimensional order is called *folding*. It is not yet thoroughly understood how this process works. A well-known research project in this area is Folding@home (<http://folding.stanford.edu/>).

³ Computer scientists would call them 'escape sequences'.

⁴ Not all parts of the DNA are genes.

⁵ Image from Wikipedia, http://en.wikipedia.org/w/index.php?title=Gene_expression&oldid=258825410.

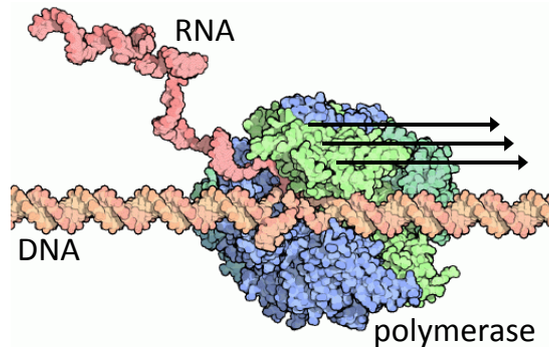


Figure 2: A polymerase, sliding along a DNA molecule and synthesizing RNA⁶.

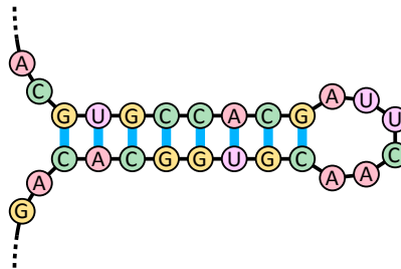


Figure 3: A hairpin structure with a stem-loop⁷.

DNA molecule towards the gene's end. In this process, the polymerase creates an RNA molecule that contains the same information as the gene (see Figure 2).

To enable a polymerase to find the exact starting location of a gene, each gene is preceded by a sequence called *promoter*. The promoter sequence has the chemical ability to attract polymerase enzymes and bind them to the DNA. In order to control which genes are transcribed in a cell, a multitude of proteins can attach to promoters: Some proteins enhance the binding behavior of a promoter while others disable a promoter. (However, such proteins are not part of our simulation.) The sequences of different promoters have a common structure, the most obvious one is a region called *TATA-Box*, as its sequence is Thymin-Adenin-Thymin-Adenin. Unfortunately, each promoter differs slightly. As we will show later, this fact is important for simulation with graph rewriting.

One option to terminate the transcription process is the *rho-independent termination*. Here, the last copied nucleotides have an 'inverted' palindromic order. Therefore, the corresponding nucleotides can form a chemical binding which leads to a *hairpin structure*. These nucleotides stick together to form a stem-loop as shown in Figure 3. This stem-loop can slide into the polymerase enzyme and detach it from the DNA. The newly created RNA molecule is then set free.

⁶ Image from Wikipedia, <http://de.wikipedia.org/w/index.php?title=RNA-Polymerase&oldid=52619006>.

⁷ Image from Wikipedia, <http://de.wikipedia.org/w/index.php?title=Haarnadelstruktur&oldid=39180278>.

2.2 Translation

The RNA molecule produced in the preceding transcription step serves as a blueprint for the protein. Like a polymerase attaches to DNA, a *ribosome* enzyme attaches to RNA during translation. As soon as the ribosome detects the triplet ‘AUG’, it starts to concatenate amino acids according to the triplet pattern found on the RNA strand (c. f. Section 2). The mapping from triplets to amino acids is shown in Figure 4. The translation process stops as soon as the ribosome detects one of the three stop-sequences: ‘UAG’, ‘UAA’, or ‘UGA’. Together with the start sequence AUG, these are all of the ‘special commands’ mentioned above.

Ala	Arg	Asn	Asp	Cys	Gln	Glu	Gly	His	Ile	Leu	Lys	Met	Phe	Pro	Ser	Thr	Trp	Tyr	Val
GCA	CGA	AAC	GAC	UGC	CAA	GAA	GGA	CAC	AUA	CUA	AAA	AUG	UUC	CCA	UCA	ACA	UGG	UAC	GUA
GCC	CGC	AAU	GAU	UGU	CAG	GAG	GGC	CAU	AUC	CUC	AAG		UUU	CCC	UCC	ACC		UAU	GUC
GCG	CGG						GGG		AUU	CUG			CCG	UCG	ACG				GUG
GCU	CGU						GGU			CUU			CCU	UCU	ACU				GUU
	AGA									UUA					AGC				
	AGG									UUG					AGU				

Figure 4: Mapping between amino acids and nucleotide triplets [Lew08]. Each triplet is mapped to the amino acid in the first cell of the corresponding column.

3 Implementation

In this section we describe our model of the gene expression process using the GrGen.NET graph rewriting system [BG07]. This section can also serve as a guide for implementations on other graph rewriting systems. Even though our simulation considerably simplifies the biological processes, it still produces the proteins found in the natural archetype.

In our graph model, nodes represent molecules or sets of molecules like enzymes. We use the hierarchical node type system of GrGen.NET to express similarities between nodes. For example, Adenin, Guanin, Cytosin, Thymin, and Uracil are represented by node types that extend the abstract node type **Nucleotide**.

Edges represent chemical bindings between molecules or the attachment of enzymes. In our simulation, we simplify complex chemical reactions by introducing special nodes that mark locations of pending reactions. For example, we connect promoters found during our simulation with a marker node in order to abstract and simplify the process of polymerase binding. Furthermore, these promoter marker nodes contain a field that stores the associated gene’s length.

Graph rewriting rules represent chemical reactions (processing uses the SPO-approach). The rewriting rules as well as the model can be obtained at <http://www.grgen.net/gxb/>. The rules we use in our simulation fall in three categories: promoter-identification, RNA-synthesis, and protein-synthesis.

Promoter-Identification. We created a rule for each promoter: These rules add marker nodes at each occurrence of the promoter they search for. A rule matches a promoter’s nucleic sequence against the DNA and places a marker node at the first nucleotide of the associated gene. As the

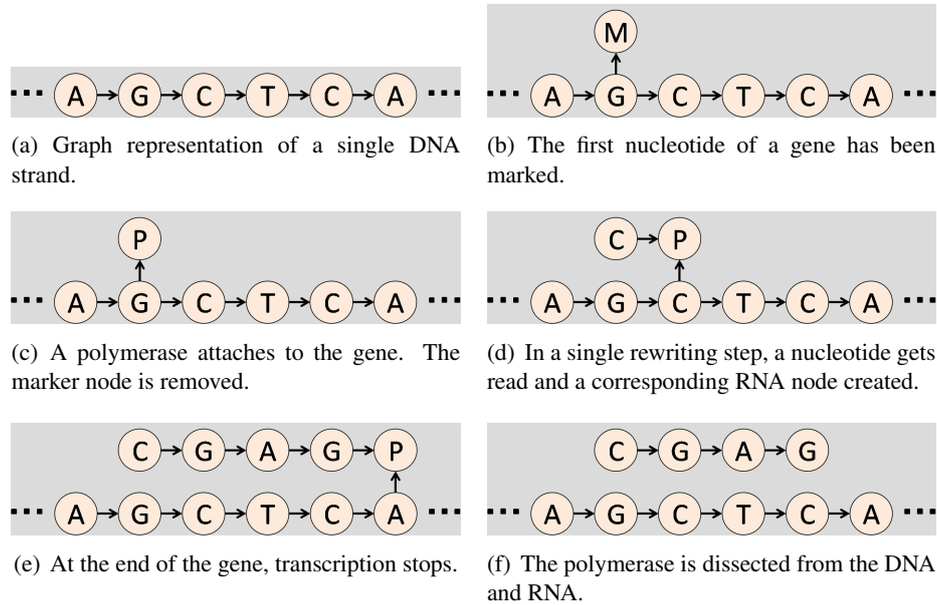


Figure 5: Simulation steps for transcription.

promoter sequences are rather long, we developed a tool to generate these graph rewrite rules. The tool can be used to prepare promoter rules for arbitrary genes of arbitrary species.

RNA-Synthesis (Transcription). After inserting the marker nodes, the RNA synthesis process starts and attaches a free polymerase enzyme (i. e. an isolated polymerase node) at a marked nucleotide. Furthermore, the polymerase copies and stores the gene's length from the marker node for the termination of the transcription process. Further rewriting rules move the attached polymerase node along the DNA strand while they create enchainned nucleotide nodes that represent the synthesized RNA strand. Each transcription step increments an internal counter of the polymerase node. The transcription process stops as soon as this counter equals the gene's length. The polymerase node detaches from the DNA and the newly created RNA strand. Afterwards, the polymerase node may initiate further transcription processes. This implies, that the total number of available polymerase nodes determines the maximal number of genes that can be transcribed concurrently in our simulation. It is also possible, that several polymerase nodes transcribe a single gene at the same time. Multiple transcription corresponds perfectly to the biological process. The RNA-synthesis is depicted in Figure 5. Listing 1 shows the GrGen.NET Code to detach the polymerase from a DNA strand.

Protein-Synthesis (Translation). The protein-synthesis starts with the application of a rule that accepts as input an RNA strand and a ribosome node that does not yet participate in a translation process. It attaches the ribosome node at the starting sequence (AUG) of an RNA molecule. RNA processing rules match an attached ribosome together with the next three nodes of the RNA strand. They emit and enchain the corresponding amino acids. These rules are

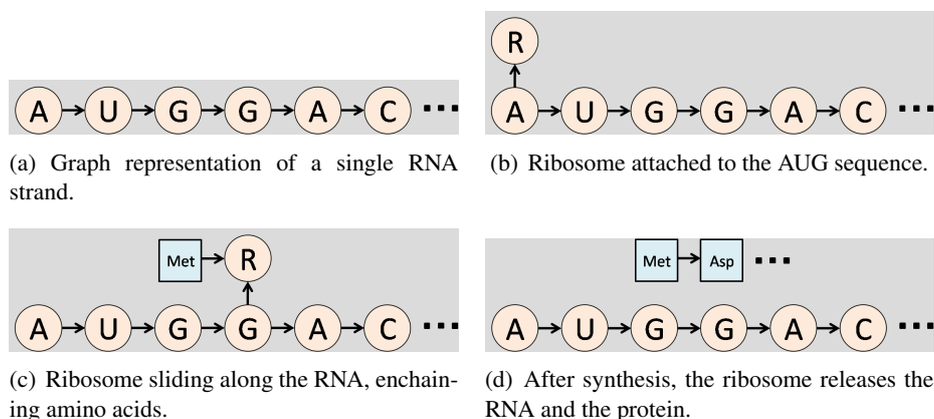


Figure 6: Simulation steps for translation.

sensitive for the stop sequences ‘UAG’, ‘UAA’, and ‘UGA’. If an RNA processing rule finds a stop sequence, the process terminates and the ribosome node gets disconnected from the RNA strand and the amino acid chain (i. e. the protein). The protein-synthesis is depicted in Figure 6. A sample screenshot of a simulation result can be found in Figure 7.

3.1 The Sample Organism

As sample data we used the genome of Escherichia Coli (E. coli), a bacterium that is well understood and frequently used as model organism. The genome of E. coli has a length of 4.6 million base pairs. As each base pair is modeled as two nodes that are connected by an edge, the graph representation takes 13.8 million graph objects. The interconnections of the two strands add two more edges per base pair, summing up to a total of 23 million graph objects. Other life forms may have much larger genomes. Table 1 lists some genome lengths of other organisms. The E. coli genome is available at [KRS⁺02].

To identify genes on the E. coli genome, we used 30 consecutive promoter sequences. The promoter sequences have been extracted using the gene data available at [KRS⁺02]. They have a length of 50 to 200 nucleotides.

4 Runtime Observations

For a long time, graph rewriting systems have not been an option to simulate complex biological processes, though the benefits of the formal representation of biochemical reactions are widely accepted. Performance is one of the main reasons for the lack of simulation implementations. In this section we present our experiences with GrGen.NET.

As described in Section 3, promoter search is one important part of the gene expression simulation. Our promoter search rules seek large nucleotide node sequences in the graph representation of a DNA molecule. It is obvious, that string based implementations can perform such search tasks faster. A simple string implementation using regular expression takes 0.3 seconds on a

Table 1: Genomes of several viruses, bacteria and organisms.

Name	Basepairs (bp)	Genes
Simian virus 40	5243	6
M13 phage	6407	10
Lambda phage	48502	ca. 50
Helicobacter pylori	1667867	1590
Mycobacterium tuberculosis	4411529	3924
Escherichia coli	4639211	4288
Yeast	12 million	6240
Pinworm	97 million	18240
Fly	180 million	13600
Mouse	3000 million	25000
Human	3000 million	25000
Corn	2400 million	30 - 40000
Rice	440 million	30 - 40000

common PC. GrGen.NET took about 8 seconds on the same PC. The execution time shrinks to 3.5 seconds on an 8-core PC using a preliminary multi threaded implementation of GrGen.NET. Although being slower, this is still fast enough to use graph rewriting approaches.

The generated graph rewriting rules for promoter search are rather large. GrGen.NET translates each rewriting rule into C# source code. The generated C# file for all 30 promoter patterns has a size of about 700 megabytes. This is unwieldy, even for generated code. A solution to this problem could be a generation mode that optimizes code size.

A problem that still persists is memory consumption. Nodes as well as edges are represented as .NET objects. A node requires 28 bytes, while an edge requires 44 bytes. Therefore, the graph representation of the E. coli genome requires 824 megabytes. This is acceptable on current workstation PCs. For a similar graph representation of the human genome 525 gigabytes of memory would be required⁸. As the current representation is very coarse grained the memory consumption for advanced genetic applications will be even higher.

5 Improvements of the Simulation

Even though our simulation produces the correct proteins from the real genes, our simulation relies on several simplifications. In this section, we discuss these simplifications in an attempt to give an outline for the next steps required to reach a more realistic simulation.

A major simplification is the omission of spacial relations. In a real environment, an unbound enzyme and the availability of reactants is not sufficient for most reactions to happen: All reactants need to be physically close to each other and in a certain angle for the reaction to happen. Unfortunately, graph elements do not provide a notion of locality. Locality could be simulated by connecting the nodes of our graph to a three-dimensional grid of nodes representing posi-

⁸ Current workstation PCs have about 8 gigabytes of main memory.

tions in an euclidean space. The resulting effort would be immense. However, if the GRS itself had a notion of spatial positioning, internal data representation could possibly benefit from optimizing data structures such as octrees [YKFT84]. An octree is a tree data structure that stores hierarchical three-dimensional data in a compact way.

Another simplification concerns timely relations. No known graph rewriting system implements time constraints, although theoretical approaches exist [GHV02]. The synthesis of an RNA strand in *E. coli* proceeds with 50 to 100 polymerisation steps per second. Molecules located on the DNA may slow down the process. The speed of current graph rewriting systems depends on the processing speed of software and hardware. For a realtime simulation graph rewriting rules have to be extended by timely relations.

During implementation of the study, we realized that not all details of the gene expression process can be simulated using GrGen.NET. As stated in Section 3, a graph rewriting system needs to know the specific sequence of a promoter in order to attach a polymerase to the corresponding gene. Unfortunately, in reality, this pattern is only imprecisely known and its nucleic sequence differs slightly from instance to instance. Current graph rewriting systems are not able to detect such fuzzy patterns. A fuzzy matching mode would be a helpful extension. For instance, if a search pattern consists of fifty nodes and a match candidate has 48 correctly identified nodes, it could be considered as a valid match.

The inability to fuzzy-match also prevents a more realistic termination of the RNA synthesis: As stated in Section 2, a stem-loop in the newly created RNA-strand initiates the rho-independent transcription termination. The rho-independent transcription termination requires a palindromic sequence. Current graph rewriting systems are in fact able to detect *perfect* palindromic sequences, but nature is a little more tolerant in this concern.

To circumvent the following simplification, additional genetic and software engineering research is required. Our simulation only uses known promoters, so only known genes can be found. In real cells, polymerases react with promoters because of the chemical properties of the promoters' nucleic sequences. Simulating such chemical reactions requires a DNA representation at molecular level and an accurate description (in terms of graph rewriting rules) of organic chemistry. This requires in-depth knowledge of chemistry and is subject to further studies. In [YKS04] a first approach is presented. Nevertheless, rewriting rules could be able to identify new promoters during runtime, as the sequence of a promoter follows a certain pattern.

6 Related Work

Simulating gene expression using graph rewriting systems has already been proposed by McCaskill and Niemann [MN01]. While in their work a graph rewriting system has been created solely for the purpose of genetic simulation, we use a standard multipurpose graph rewriting system. McCaskill and Niemann implemented a specialized mechanism to differentiate between molecules of different types, whereas we can use typed attributed graphs, for example. We see the future of genetic graph rewriting in multi purpose systems, as we can easily benefit from further improvements in graph rewriting technology. Yadav, Kelley, and Silverman propose to simulate chemical reactions at the molecular level [YKS04], too. They provide hints on extending graph rewriting systems for that purpose but their work lacks an implementation. Rosselló

and Valiente propose the computation of signal transduction in cells using graph transformation [RV04]. One can sum up approaches like these to the goal of simulating the inner workings of biological cells as a whole. Rosselló and Valiente give an overview on more approaches to simulate different parts of these fundamentals in [RV05]. We propose to sum up all these approaches into an integrated simulation based on graph rewriting systems. Currently available tools focus on static computation or data storage rather than simulation. Gene expression products, as they are generated in the application presented in this paper, are stored in the ecocyc database [KRS⁺02]. However no tool tries to simulate the genetic processes as a whole. We propose that graph rewriting may lead to such a software at some point in the future.

7 Conclusion

Even though the processing of string representations is considerably faster, we see the future of genetic simulations in graph representation and graph rewriting. String representations are inflexible in handling multidimensional structures: A polymerase binding simulation at a detailed level involves hundreds of RNA and protein molecules and needs to consider their three dimensional layout and position. Graphs can easily represent multi-dimensional relations containing large numbers of objects.

All genetics simulation approaches aim at simulating the whole life cycle process of a cell at some point. As the size of the human genome does not grow, we are confronted with a fixed problem size. Therefore, we may concentrate on simulation approaches offering a good abstraction level such as graph rewriting. The loss of performance will be overcome by better computing devices in the future. The trend towards parallel computing systems can be used to advantage, as genetic processes offer ample parallelism by nature.

Listing 1: Rule to detach a polymerase from DNA

```
1 rule RNA_Pol_Finish {
2 pattern {
3   pol:Polymerase -cur_conn:PolymeraseToDna->cur_nuc:Nucleotide
4     -:DNA_Backbone-> next_nuc:Nucleotide;
5   pol -rna_conn:PolymeraseToRna->cur_rna_nuc:Nucleotide;
6   if { pol.MaxNucleotide > 0 &&
7     pol.CurrentNucleotide == pol.MaxNucleotide;
8     }
9   }
10 modify {
11   delete(cur_conn); delete(rna_conn);
12   eval { pol.MaxNucleotide = 0; pol.CurrentNucleotide = 0; }
13   }
14 }
```

Table 2: Simulation summary.

Sample organism	Escherichia coli
Total number of DNA base pairs	4,639,211
Total number of DNA base pairs in simulation	4,639,211
Total number of promoters/genes	4,288
Average size of promoters	50-200 nucleotides
Genes used in benchmark	30
Graph rewriting system	GrGen.NET (version 1.4)
Max. number of nodes in simulation	9.3 million
Max. number of edges in simulation	13.9 million
Number of rules	69
Total size of rewriting-rules source code	700 megabytes
Total size of rewriting-rules object code (MSIL-DLL)	13 megabytes
Memory consumption of graph elements	824 megabytes
Runtime (only promoter search), string-based implementation	0.3 seconds
Runtime (only promoter search), GrGen.NET	7.7 seconds
Runtime (only promoter search), parallel GrGen.NET	3.5 seconds
Total runtime of the simulation, GrGen.NET	7.9 seconds

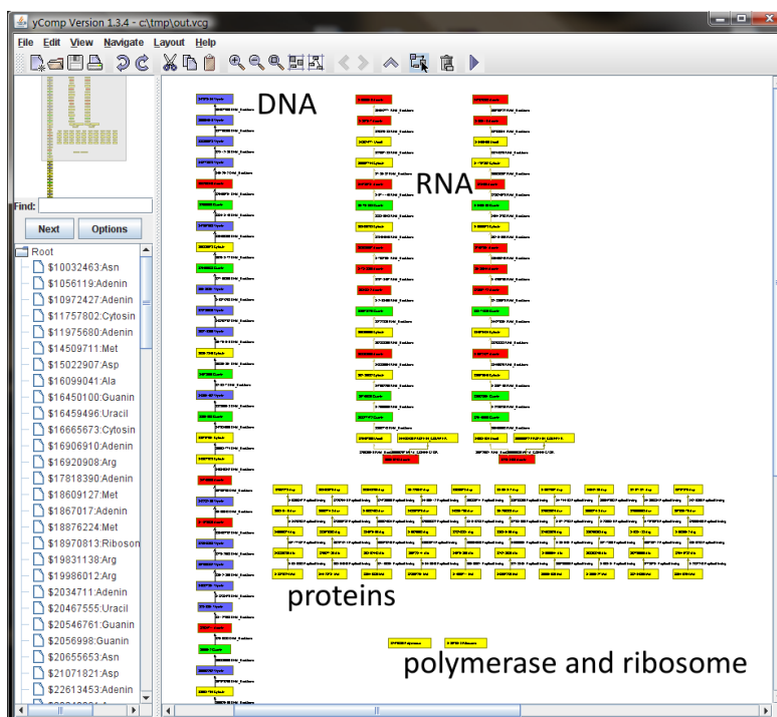


Figure 7: A yComp-visualization of the result graph of our simulation process.

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