Benchmarking Current RNA Folding Software and Improvements to the Current Regime

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Abstract

Finding the secondary structure of RNA is important for understanding how RNA will interact in a cell. Frequently computational algorithms are used to determine structure due to difficulties extracting good in vivo data of RNA structures. Many of the algorithms for RNA folding are computationally complex. In this paper we establish benchmarks for two commonly used RNA folding packages, mfold, and RNAfold Vienna, and compare them to an improved Four-Russians folding algorithm. We show ... These results will help software maintainers to understand the benefit of updating their algorithms. The results will also help guide users when choosing RNA folding software when looking for the most computationally optimal package.

1 Introduction

RNA is an essential macromolecule used in protein formation and performs other essential functions within the body(4). RNA does not stay in single stranded form and instead folds on itself to create the lowest energy conformation possible to ensure thermodynamic stability(6). When folding, RNA forms a 2D secondary structure(5) with A matching to U and G to C (figure 1). Using this data we can find a 3D tertiary structure(5).

Our paper focuses on benchmarking 2D secondary structure prediction software based off the Nussinov dynamic programming algorithm(3) for RNA folding. This algorithm is $O(n^3)$ time complexity. There have been multiple attempts to parallelize the Nussinov algorithm(15; 16) which have resulted in large speed increases, however, CPU intensive algorithms only slightly lowered the bound up until 2010(12; 13). In 2010 the Nussinov bound that was significantly improved by the Frid-Gusfield Four Russians method which established you could perform the DP method in $O(\frac{n^3}{loq(n)})$

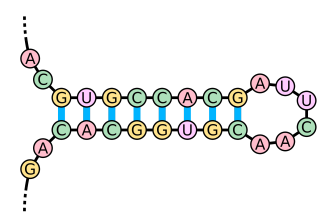


Figure 1: Example of an RNA molecule folding

time(1). Later a parallel method of the Four Russians algorithm presented proof that you can lower this bound to $O(\frac{n^2}{\log(n)})$ inside an NVIDIA CUDA environment(2).

There are two major RNA folding software packages, mfold(8; 9), and the Vienna RNA Package(7). These both utilize the Nussimov method to return results of the RNA secondary structure by finding the lowest possible thermodynamic conformations of the RNA(9; 7). In order to tell which was faster we performed application level benchmarks(14) to see which of these two applications could more quickly fold arbitrary length RNA of the circular and linear variety. The way mfold is written allows linear RNA to be treated as exceptional variants of circular ones(17). RNAfold Vienna was initially optimized to only handle linear RNA(17) but later improvements enabled it to speed the folding of circular ones(17). As a result we also wish to determine what kind of speed difference still exists between mfold and Vienna when performing analysis on circular RNA.

We then performed micro-benchmarks (10) of the folding algorithms utilized in mfold and Vienna for both circular and linear RNA, and then compared that to equivalent benchmarks for the Four Russians and parallel Four Russians algorithms. The first thing we found when comparing mfold and Vienna was When we compared mfold and Vienna to the Four Russians algorithms we found that without a graphics card and parallel processing capabilities a machine can see speed ups of ... in mfold and ... in Vienna for linear RNA. For circular RNA speed increases change to ... in mfold and ... in Vienna. With a graphics card and parallel processing folding clusters can see increases of up to ... for mfold and ... in Vienna. Circular RNA also sees speed bumps of ... in mfold and ... in Vienna.

These results show us that ... for natively differentiating between mfold and Vienna. They also help explain that both pieces of software can experience significant speed increases if they implemented the Frid-Gusfield method. Furthermore the authors of this paper would recommend both software packages to support GPU hardware to achieve even greater speed gains when inside a parallel capable environment.

2 Methods

2.1 Standardizing the Testing Environment

Benchmarking is renown as a difficult thing to perform effectively (10; 14). There are many processes that can be executing on a computer at any one moment that it is possible that a benchmark can give inaccurate information due to conflicting processes running in the background (10). As a result we used a machine solely dedicated for benchmarking and no other tasks. We also standardized on the following specifications for our runs (11):

Architecture	Operating	Compiler	
	\mathbf{System}		
8 core Intel i7 CPU 4.00	Linux 4.2.5-201	GCC	
GHz 16G RAM GeForce	Fedora 22	5.1.1 - 4.fc22	
GTX 960			
"	"	gcc-gfortran	
		5.1.1 - 4.fc 22	
"	"	NVIDIA CUDA	
		version 5.5	

We used the following applications with corresponding versions and requirements in our test runs:

Application	Version	Requirements
mfold	3.6	GCC, Fortran
RNAfold Vienna	2.1.9	GCC
Frid-Gusfield Four Russians	N/A	GCC
Parallel Four Russians	N/A	GCC, NVIDIA CUDA

Our testing architecture was laid out where we would SSH into the benchmarking machine and then execute tests. Test results would then be reported back to the user's central machine where they could be stored in a database for later analysis (figure 2). Our testing required no internet connectivity besides the ssh access required to initiate our testing so all calculations were performed locally. Also there were no IO operations except for post processing of mfold and Vienna results.

2.2 Data Inputs

For input data we give inputs of RNA as strings in a file. An example of this would be the 10 character RNA string AUGCCAUGGA. This same RNA sequence can be treated as circular by providing parameters to the mfold and RNAfold Vienna programs that tell it the RNA is circular(18; 19).

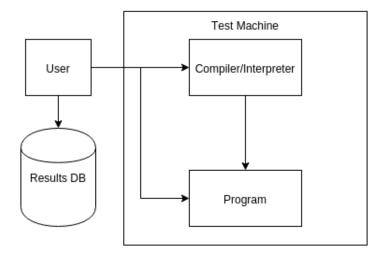


Figure 2: Test Architecture

2.3 Application Benchmarks

The first type of benchmark we perform is the application level benchmark. An application benchmark is designed to measure the performance of an entire application and the resources it consumes on an individual machine(20). In our case we wish to evaluate the amount of time that mfold and RNAfold Vienna take to return RNA secondary structures given different length RNA strands varying on linear and circular variety. Since a single run of an application may vary in time even for identical inputs. Because of this we evaluate each input of RNA 30 times and report the mean μ , standard deviation, σ^2 , of the runs corresponding to each sequence length.

2.4 Micro-benchmarks

The most basic type of benchmark to perform is the micro-benchmark. The micro-benchmark is a single piece of code executed many times in serial so that we can get a profile of its run characteristics(14; 10). Once these characteristics are observed we can then make inferences about its performance and ways that it can be improved. Micro-benchmarks have the downside of losing generality of performance across the entire application(14; 10). A good example of this is if an IO heavy function made many consecutive calls to the read function on the OS while the rest of the application made no calls to read whatsoever. If we tried to generalize this one function to the rest of the application we would misguidedly attempt to optimize disk IO across our entire system.

We avoid this trap in our paper by benchmarking only parts of the code that execute the Nussinov algorithm in mfold and the Vienna package. We then report these results back to our test results database for later analysis. After this we compare these results to runs of the serial Frid-Gusfield algorithm and parallel Frid-Gusfield algorithm.

2.5 Frid-Gusfield Four-Russians Algorithm

The Four-Russians Algorithm (1) is an algorithm to improve the above-mentioned Nussinov $O(n^3)$ Algorithm by Four-Russian method. The Frid and Gusfield is an $O(\frac{n^3}{\log n})$ algorithm. The Four Russian algorithm achieves this speed up by understanding that we can make certain optimizations to the matrix of matching base pairs required by the Nussinov algorithm. Particularly, the values along a column from bottom to top and along a row from left to right are monotonically non-decreasing. Consecutive cells differ at most by 1(1). As a result we can perform pre-processing of specific operations that the Nussinov algorithm must compute manually.

2.6 CUDA Parallel Implementation for F-G Method

Compute Unified Device Architecture (CUDA) is a parallel computing platform created by NVIDIA. By using CUDA API, Venkatachalam presented an $O(\frac{n^2}{logn})$ algorithm for RNA folding is presented (2). The CUDA implementation parallelizes the two-vector method so that achieve an enhancement of another factor of O(n).

3 Results

3.1 Application Benchmarks

In this part, we performed the benchmark for two packages using the Dynamic Programing (DP) paradigm with both linear and circular RNA. We report the number of RNA bases or size, the number of times the folding program was run with our specific input or N runs, the mean timing of the runs as μ , and the standard deviation of the runs denoted as σ^2 . For the benchmarking of both linear and circular types of RNA, we select the RNA of the size, ranging from 200 nucleobases to 1000 nucleobases. The time to fold these structures are compared in the following Table 1.

Timing of m -fold package for linear RNA (sec.)										
size	200	300	400	500	600	700	800	900	1000	
N runs	30	30	30	30	30	30	30	30	30	
μ	2.523	6.635	6.313	8.336	10.492	9.384	12.212	14.319	14.402	
σ^2	0.343	0.251	0.180	0.547	0.144	0.094	0.176	0.118	0.211	
	Timing of Vienna for linear RNA (sec.)									
size 200 300 400 500 600 700 800 900 100								1000		
N runs	30	30	30	30	30	30	30	30	30	
μ	0.029	0.072	0.112	0.163	0.508	0.491	1.934	11.723	8.296	
σ^2	0.006	0.001	0.001	0.001	0.002	0.002	0.006	0.336	0.011	

Table 1: The time taken by two packages to predict secondary structures of linear RNA of different length.

In a more vivid way, the plot of the bechmark time for linear RNA is shown in the Fig 3.

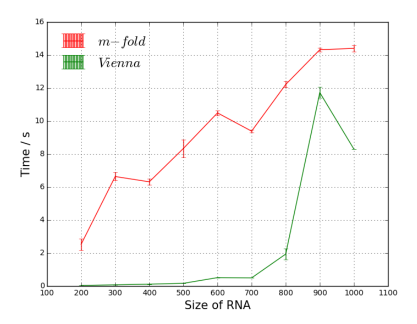


Figure 3: Benchmark of *Vienna* and *m-fold* packages with linear RNA of different sizes.

We can clearly see that m-fold takes the most time to complete. However as mentioned before, Vienna is optimized to handle linear RNA so these results are not necessarily surprising. We benchmark circular RNA. The time in seconds to determine their secondary structures are listed in the following Table 2.

Timing of m -fold package for circular RNA (sec.)										
Size	200	300	400	500	600	700	800	900	1000	
N runs	30	30	30	30	30	30	30	30	30	
average	2.409	6.633	6.269	8.889	11.274	9.074	11.358	14.678	16.712	
std (σ^2)	0.060	0.354	0.362	0.159	0.1176	0.161	0.364	0.1999	0.285	
	Timing of Vienna for circular RNA (sec.)									
Size	200	300	400	500	600	700	800	900	1000	
N runs	30	30	30	30	30	30	30	30	30	
average	0.027	0.084	0.129	0.153	0.664	0.351	1.943	7.915	50.15	
std (σ^2)	0.002	0.001	0.001	0.002	0.002	0.002	0.006	0.036	0.126	

Table 2: The time taken by two packages to predict secondary structures of circular RNA of different length.

The following Fig 4 is plotted to better visualize how the two packages compete with each other.

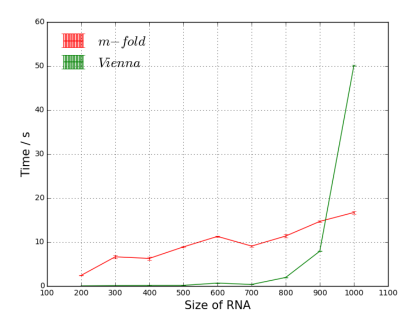


Figure 4: Benchmark of *Vienna* and *m-fold* with circular RNA of different sizes.

As we can see Vienna initially outperforms m-fold but as RNA size hits 1000 bases the time to complete the folding increases dramatically. So it appears the optimizations made to Vienna by Hofacker and Stadler(17) break down around 1000 nucleobases.

3.2 Micro-benchmarks

The timing for Nussinov, Frid-Gusfield (FG), and CUDA Four-Russian algorithms are listed below with different sizes of RNA sequences 500 to 6000. The data is presented in Table 3.2.

Timing of Algorithms for RNA Folding (sec.)									
size	500 1000 2000 3000 4000 5000 6000								
Nussinov	0.2790	2.0751	16.7033	57.8146	145.2998	301.4874	519.6531		
F-G	0.0903	0.6092	5.5868	19.6117	49.3309	95.6461	162.9072		
CUDA	0.0088	0.1988	0.4690	1.1943	2.5817	4.8506	8.2735		

The time take by Nussinov, F-G and CUDA to construct the secondary structures of RNA of different size.

More clearly by plotting Figure 5, we observe the F-G method has a vast advantage over Nussinov when the size of RNA sequence is larger than 3000, and the CUDA method achieves approximately linear increase as the length of RNA sequences.

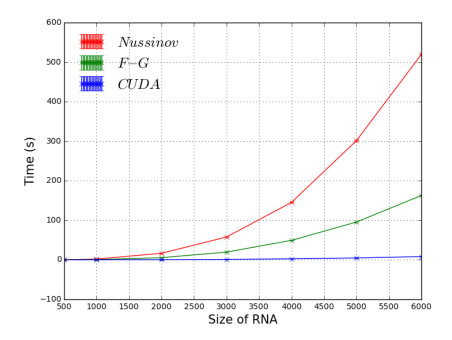


Figure 5: Timing of Algorithms for RNA Folding (sec.)

4 Discussion

In this paper, we performed both application and micro benchmark for current packages and some popular methods available for RNA secondary structure prediction problem, namely, RNA folding. For the application benchmark, RNA fold Vienna and m-fold have been applied to predict the secondary structure of both linear and circular RNA. In both linear and circular cases, the Vienna package is clearly optimal for chains of RNA smaller than 900 bases. While m-fold is time optimal in the prediction of circular RNA structures of 1000 nucleobases and likely for greater numbers of bases. However further amounts of testing is necessary this conclusion can be further confirmed by testing more RNA sequences at the same length. Although the two packages are all based on Dynamical Programming paradigm, their average performance may be different. The difference might come from the ways how they implement the DP algorithm. For example, the code used for packages and the I/O management are probably the main reasons for this difference.

When the size of RNA is within 1000 nucleobases, the Vienna packages is more efficient than the m-fold package. But the Vienna package increases faster than the m-fold package. Probably, the Vienna Package is not advantageous to predict the secondary structure of super large RNA.

The *Vienna* can do a more efficient prediction when the circular RNA has nucleobases smaller than 950. As the operations required for *Vienna* prediction increases

rapidly for RNA longer than 900 nucleobases, the Vienna package loses its advantages. And the m-fold package demonstrates better performance.

Through the above benchmark analysis, we have developed a clear insight into the performance of two packages, *Vienna* and *m-fold* packages. Linear and circular types of RNA have been well tested at several length. In both of the cases, the *Vienna* packages is time-optimal for small RNA no larger than 900 nucleobases. However, this package loses its advantages to *m-fold* after 1000 nucleobases.

When we come to the micro-benchmark, we found that the preliminary Nussinov DP method is significantly slowing down in the cases that the number of nucleotides in the RNA sequence is more than 3000. That's intrinsically caused by its $O(log(n^3))$ time complexity characteristic. While F-G method (1) has enhanced the performance by a factor of log(n), in some cases the speed is still not satisfactory for long RNA sequences. With the introduction of CUDA onto two-vector method invented by (2), we almost achieve a linearly increasing time complexity for different size of RNA sequences, which is quite impressive.

In the future if Vienna or m-fold etc. application packages employ F-G method or even the parallelized CUDA method, we could witness a great speed improvement in RNA design or visualization applications.

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