

# TargetRNA3



## **User Manual**

TargetRNA3 uses machine learning to predict targets of small regulatory RNAs (sRNAs) throughout a genome. At its core, TargetRNA3 employs a gradient boosting classification algorithm that has been trained on thousands of evinced interactions between sRNAs and their regulatory targets in various prokaryotes. When making target predictions, the machine learning algorithm uses a variety of features indicative of regulatory interactions, including the thermodynamics of the interaction and potential homologous interactions in related organisms, if available.

The quickest and easiest way to use TargetRNA3 is via the webserver: https://cs.welleslev.edu/~btjaden/TargetRNA3

However, some users may prefer to use TargetRNA3 on their own machine. This user manual provides details on downloading TargetRNA3 and using it locally.

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#### **REQUIREMENTS**

Source code and data files for TargetRNA3 are available from GitHub (shown in Table 1): https://github.com/btjaden/TargetRNA3

TargetRNA3 is written in Python and requires at least version 3 of Python. Beyond the Python standard library, TargetRNA3 uses three libraries that are available from the Python Package Index:

- numpy
- pandas
- scipy

TargetRNA3 uses three applications from the <u>BLAST+ suite provided by NCBI</u>:

- blastn
- blastp
- blastdb aliastool

TargetRNA3 uses two applications from the <u>ViennaRNA Package</u>:

- RNAplfold
- RNAplex

The BLAST+ and ViennaRNA applications should be available to TargetRNA3 for execution in the current path, or else the executables for these applications should be in the same directory as the TargetRNA3 source code.

The contents of two further data files are necessary. Because of their larger size (8 GBs and 4 GBs), they are not provided on GitHub. The two files are located at:

- https://cs.wellesley.edu/~btjaden/TargetRNA3/DB FNA.tar
- <a href="https://cs.wellesley.edu/~btjaden/TargetRNA3/DB">https://cs.wellesley.edu/~btjaden/TargetRNA3/DB</a> FAA.tar

These two files should be placed in the DataFiles/ directory and their contents extracted:

Each of these two files contains 15 files within it. Once extracted, the DataFiles/ directory should contain 45 files (15 from each of these two files and 15 from GitHub as shown in Table 1).

Table 1. Source code and data files from GitHub

File	Description		
TargetRNA3.py	Source code		
AddGenome.py	Source code for adding a new genome		
SgrS.fa	FASTA file containing an example sRNA sequence		
Prrf1.fa	FASTA file containing an example sRNA sequence		
DataFiles/model.pickle	Trained machine learning model		
DataFiles/gene_genome.pickle	Mapping genes to genomes		
DataFiles/genome_IDs.pickle	Mapping genomes to unique identifiers		
DataFiles/assembly_summary.txt.archaea.gz	Used for adding a new archaeal genome (optional)		
DataFiles/assembly_summary.txt.bacteria.gz	Used for adding a new bacterial genome (optional)		
DataFiles/16S.fna.ndb	BLAST database file		
DataFiles/16S.fna.nhr	BLAST database file		
DataFiles/16S.fna.nin	BLAST database file		
DataFiles/16S.fna.njs	BLAST database file		
DataFiles/16S.fna.nog	BLAST database file		
DataFiles/16S.fna.nos	BLAST database file		
DataFiles/16S.fna.not	BLAST database file		
DataFiles/16S.fna.nsq	BLAST database file		
DataFiles/16S.fna.ntf	BLAST database file		
DataFiles/16S.fna.nto	BLAST database file		
Genomes/GCF_000005845.2/ closest_relatives.txt	E. coli genome files (optional)		
Genomes/GCF_000005845.2/			
GCF_000005845.2_ASM584v2_feature_table.txt.gz	E. coli genome files (optional)		
Genomes/GCF_000005845.2/ GCF_000005845.2_ASM584v2_genomic.fna.gz	E. coli genome files (optional)		
Genomes/GCF_000005845.2/ GCF_000005845.2_ASM584v2_genomic.gff.gz	E. coli genome files (optional)		
Genomes/GCF_000005845.2/ GCF_000005845.2_ASM584v2_protein.faa.gz	E. coli genome files (optional)		
Genomes/GCF_000005845.2/	E. coli genome files (optional)		
GCF_000005845.2_ASM584v2_rna_from_genomic.fna.gz	L. con genome mes (optional)		
Genomes/GCF_000005845.2/ mRNA.homologs	E. coli genome files (optional)		
Genomes/GCF_000005845.2/ RNAplfold results/*	Many E. coli genome files (optional)		
Genomes/GCF_000006765.1/	P. aeruginosa genome files (optional)		
closest_relatives.txt Genomes/GCF 000006765.1/			
GCF_000006765.1_ASM676v1_feature_table.txt.gz	P. aeruginosa genome files (optional)		
Genomes/GCF_000006765.1/ GCF_000006765.1_ ASM676v1_genomic.fna.gz	P. aeruginosa genome files (optional)		
Genomes/GCF_000006765.1/	P. aeruginosa genome files (optional)		
GCF_000006765.1_ ASM676v1_genomic.gff.gz Genomes/GCF 000006765.1/			
GCF_000006765.1_ ASM676v1_protein.faa.gz	P. aeruginosa genome files (optional)		
Genomes/GCF_000006765.1/ GCF_000006765.1_ ASM676v1_rna_from_genomic.fna.gz	P. aeruginosa genome files (optional)		
Genomes/GCF_000006765.1/ mRNA.homologs	P. aeruginosa genome files (optional)		
-			
Genomes/GCF_000006765.1/ RNAplfold_results/*	Many P. aeruginosa genome files (optional)		

### **USAGE: TargetRNA3**

As input, TargetRNA3 requires genome information and a sRNA sequence. Two example sets of genome files (Genomes/GCF\_000005845.2/ for *E. coli* and Genomes/GCF\_000006765.1/ for *P. aeruginosa*) and two example sRNA sequences (SgrS.fa and Prrf1.fa) have been provided.

#### To search for targets of the sRNA SgrS in E. coli:

```
python TargetRNA3.py -s SgrS.fa -g Genomes/GCG 000005845.2
```

### To search for targets of the sRNA Prrf1 in *P. aeruginosa*:

```
python TargetRNA3.py -s Prrf1.fa -g Genomes/GCG 000006765.1
```

Command line arguments are shown in Table 2.

**Table 2. Command line arguments for TargetRNA3.py** 

		<u> </u>	
<u>Flag</u>	<u>Value</u>	<u>Description</u>	Required?
-S	String	File in FASTA format containing sRNA sequence	Required
		Path to directory containing genome information including files	
-g	String	<pre>genomic.fna, protein.faa, rna_from_genomic.fna, and</pre>	Required
		<pre>feature_table.txt (files may be gzipped or not)</pre>	
-0	String	File to which results should be output (default is standard out)	Optional
-prob	Decimal	Probability above which a gene is predicted as a target (default is 0.5)	Optional
-pval	Decimal	P-value below which a gene is predicted as a targe (default is 0.05)	Optional
-n_threads	Integer	Number of threads (default is based on self-identification of number of processors)	Optional
-db	String	Path to BLAST database (default is DataFiles/combined.fna)	Optional
-model	String	Path to ML model file (default is DataFiles/model.pickle)	Optional
-h		Print usage and description, ignore all other flags	Optional
-help		Print usage and description, ignore all other flags	Optional

As output, TargetRNA3 produces a ranked list of candidate regulatory targets that it predicts for a sRNA. The main measure of the likelihood that a sRNA interacts with a candidate target is the *probability*. Probabilities greater than 0.5 indicate that a candidate is more likely than not to be a target, as determined by the machine learning algorithm. Probabilities less than 0.5 indicate that a candidate is more likely *not* to be a target, as determined by the machine learning algorithm.

If TargetRNA3 reports no significant targets (or too few significant targets) for a sRNA, the sensitivity can be increased so as to predict more targets by lowering the probability threshold with the -prob command line argument.

For predicted targets of a sRNA, TargetRNA3 outputs a variety of information for each target, including the name and annotation of the target, a predicted structure and energy (based on thermodynamics of hybridization and structural accessibility of the sRNA and target) for the sRNA:mRNA target interaction, the location of the interaction within the sRNA and within the target (relative to the start of the target), the probability that there is an interaction between

the sRNA and target as determined by the machine learning algorithm, and a corresponding *p*-value for the interaction.

## **USAGE: Adding a New Genome**

Two example sets of genome files (Genomes/GCF\_000005845.2/ for *E. coli* and Genomes/GCF\_000006765.1/ for *P. aeruginosa*) have been provided. Using AddGenome.py, new genomes can be added for TargetRNA3 to search. AddGenomes.py provides two different ways to add a new genome: (1) with the assembly accession identifier or (2) with manual download.

#### (1) Adding a genome with the assembly accession identifier

First, determine the assembly accession identifier for the genome of interest. For *Escherichia coli* str. K-12 substr. MG1655, the identifier is GCF\_000005845.2. For *Pseudomonas aeruginosa* PAO1, the identifier is GCG\_000006765.1.

For bacteria, assembly accession identifiers can be found in the file DataFiles/assembly\_summary.txt.bacteria.gz or online at: <a href="https://ftp.ncbi.nlm.nih.gov/genomes/refseq/bacteria/assembly\_summary.txt">https://ftp.ncbi.nlm.nih.gov/genomes/refseq/bacteria/assembly\_summary.txt</a>

For archaea, assembly accession identifiers can be found in the file DataFiles/assembly\_summary.txt.archaea.gz or online at: https://ftp.ncbi.nlm.nih.gov/genomes/refseq/archaea/assembly\_summary.txt

To add a new genome (e.g., *Bacillus subtilis* subsp. subtilis str. 168 with assembly accession identifier GCF 000009045.1) using the assembly accession identifier:

```
python AddGenome.py -a GCF_000009045.1
```

#### (2) Adding a genome with manual download

In case a genome cannot be added with its assembly accession identifier, e.g., because the genome assembly is not available from RefSeq, a genome can be added manually. Four genome files are needed: the genome sequence in FASTA format (genomic.fna), protein sequences in FASTA format (protein.faa), RNA sequences in FASTA format (rna\_from\_genomic.fna), and the feature table (feature\_table.txt). These four files can be gzipped or not.

To add a new genome (e.g., assuming the four genome files are located in a directory Genomes/MyNewGenome) based on the provided genome files:

```
python AddGenome.py -q Genomes/MyNewGenome
```