

BioAutoML: automated feature engineering and metalearning to predict noncoding RNAs in bacteria

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Abstract

Recent technological advances have led to an exponential expansion of biological sequence data and extraction of meaningful information through Machine Learning (ML) algorithms. This knowledge has improved the understanding of mechanisms related to several fatal diseases, e.g. Cancer and coronavirus disease 2019, helping to develop innovative solutions, such as CRISPR-based gene editing, coronavirus vaccine and precision medicine. These advances benefit our society and economy, directly impacting people's lives in various areas, such as health care, drug discovery, forensic analysis and food processing. Nevertheless, ML-based approaches to biological data require representative, quantitative and informative features. Many ML algorithms can handle only numerical data, and therefore sequences need to be translated into a numerical feature vector. This process, known as feature extraction, is a fundamental step for developing high-quality ML-based models in bioinformatics, by allowing the feature engineering stage, with design and selection of suitable features. Feature engineering, ML algorithm selection and hyperparameter tuning are often manual and time-consuming processes, requiring extensive domain knowledge. To deal with this problem, we present a new package: BioAutoML. BioAutoML automatically runs an end-to-end ML pipeline, extracting numerical and informative features from biological sequence databases, using the MathFeature package, and automating the feature selection, ML algorithm(s) recommendation and tuning of the selected algorithm(s) hyperparameters, using Automated ML (AutoML). BioAutoML has two components, divided into four modules: (1) automated feature engineering (feature extraction and selection modules) and (2) Metalearning (algorithm recommendation and hyper-parameter tuning modules). We experimentally evaluate BioAutoML in two different scenarios: (i) prediction of the three main classes of noncoding RNAs (ncRNAs) and (ii) prediction of the eight categories of ncRNAs in bacteria, including housekeeping and regulatory types. To assess BioAutoML predictive performance, it is experimentally compared with two other AutoML tools (RECIPE and TPOT). According to the experimental results, BioAutoML can accelerate new studies, reducing the cost of feature engineering processing and either keeping or improving predictive performance. BioAutoML is freely available at <https://github.com/Bonidia/BioAutoML>.

Background

Considering advances in sequencing, an increasing number of biological sequences have been generated

[1, 2]. With the expansion in volume and complexity of biological data, machine learning (ML) algorithms have been successfully applied to their analysis [3–5].

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Received: March 2, 2022. **Revised:** May 6, 2022. **Accepted:** May 9, 2022

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ML algorithms can extract new and useful knowledge from biological data [6], allowing complex analyses, speeding up new findings and reducing research costs [7]. These advances bring important social and economical benefits, such as improving diagnosis, treatment and the design of new medications [7–9], e.g. coronavirus disease 2019 [8, 10], cancer diagnosis [11] and CRISPR/Cas9-based gene-editing technology [12, 13].

Moreover, with the advancement of next-generation sequencing technologies and multi-omics analysis [14], studies have focused on discovering and characterizing small noncoding RNAs (sRNAs) in bacteria and archaea, expanding the understanding of gene regulation and elucidating new biological mechanisms [15]. Moreover, Noncoding RNAs (ncRNAs) have distinct classes with specific functions, depending on their spatial structure, sequence composition and length [16]. Regarding genome annotation, the identification of protein-coding and non-protein-coding sequences is the first and most crucial step [17]. In addition, sRNAs control gene expression in prokaryotes, regulating processes, e.g. stress responses, nutrient acquisition, virulence and biofilm formation [18]. According to [19], there is a large number of regulatory ncRNAs, highlighting their potential links to bacterial pathogenesis.

Nevertheless, one of the main difficulties for applying ML algorithms to ncRNAs and other sequences is the categorical and unstructured nature of biological sequence data. A frequent alternative to deal with this problem is to apply, in a feature engineering process, feature extraction techniques (e.g. DNA sequences: A, C, T, G), to transform biological sequences into numerical data (e.g. GC content and k-mers) with a structured format. Feature extraction techniques based on various aspects have been proposed to extract numbers from these sequences, including physicochemical, biological and mathematical features [6, 9]. No matter the aspect, the features must capture the relevant information present in the biological sequence, as the predictive performance of the model induced by an ML algorithm strongly depends on the representativeness of the input feature vector [20]. A common approach to increase the representativeness of the features is to select, among the extracted features, the subset that leads to the best predictive performance of a model induced by an ML algorithm. This approach, known as feature selection using wrappers, is also a part of the feature engineering process.

The feature engineering process often requires extensive domain knowledge, performed manually by a human expert, and is one of the most time-consuming steps in the ML pipeline [20]. Furthermore, according to [21], ncRNAs are divided into categories based on their cellular functionality and their sequential, thermodynamic and structural properties, assuming that their sequence can provide robust discriminative features. However, the same sequences can act as more than one type of ncRNAs, e.g. mature micro RNA (miRNA) can also be transfer RNA (tRNA) fragments. Consequently, most

computational approaches can predict only the presence of ncRNAs. Even those designed to classify more than one type (class) of ncRNAs do not work well with more than three types [21, 22].

These limitations motivated the development of a novel open-source software package, called BioAutoML, that can extract features based on different aspects, and automate the feature selection, algorithm(s) recommendation and algorithm(s) tuning steps for multi-class classification of biological data. BioAutoML is an end-to-end Automated Machine Learning (AutoML) tool for experiments using biological sequences; BioAutoML is able to deal with different categories of ncRNA in bacteria, such as small RNA (sRNA), tRNA, ribosomal RNA (rRNA), precursor-microRNA (pre-miRNA), miRNA, small nucleolar RNA (snRNA), small nuclear RNA (snRNA) and transfer-messenger RNA (tmRNA) [21, 23]. According to [24, 25], AutoML has a proposal similar to the area of hyper-heuristics, automatically recommending pipelines, algorithms or hyper-parameters for specific tasks, reducing dependence on user knowledge. These tasks can include different ways of preprocessing or feature engineering, as well as algorithms and optimization of its parameters (hyper-parameter tuning) [24–26]. In this study, BioAutoML calls the MathFeature package [27, 28] to extract feature descriptors representing relevant numerical information from ncRNA sequences (**Feature Extraction module**). After receiving the feature values, BioAutoML, automatically recommends, using Bayesian Optimization [29], the best pair of selected features and predictive model.

To select the features (**Feature Selection module**), BioAutoML follows the wrapper approach, using a predictive model to assess how good a feature set is. The feature extraction and feature selection model are part of the Feature Engineering process. To recommend the best model, BioAutoML recommends the best number of predictive models, and the ML algorithm to be used for the induction of each model (Algorithm Recommendation module). These two tasks are carried simultaneously, whereby one feeds information to the other in the Bayesian optimization process, until the predictive performance obtained by the pair is not further improved. It is important to point out that the predictive model used in the wrapper and induced by the recommended algorithm can be an ensemble of predictive models.

This occurs when the Algorithm recommendation model recommends more than one ML algorithm. In this case, each recommended algorithm induces a predictive model, and the induced models are combined in an ensemble. As an ensemble of predictive models is, by itself, a predictive model, for the sake of simplicity, we will name the ensemble also a predictive model. Having finished the **Feature Engineering module** and the **Algorithm Recommendation module**, BioAutoML goes to the fourth module, which uses AutoML to fine tune the hyper-parameters of the recommended ML

algorithms [24–26], aiming to improve the predictive performance of the model (**Tuning and Combination module**). If the predictive model is an ensemble, the algorithm that induces each model in the ensemble has its hyper-parameters tuned. Afterwards, the predictive models induced by the tuned algorithms are combined in the ensemble.

To make the role clearer of each module in the whole process, the part of the pipeline with the **Feature Extraction module** and the **Feature Selection module**, as they mainly work at the feature level, is named here **Feature Engineering**. The other part of the pipeline, with the **Algorithm Recommendation module** and the **Tuning and Combination module**, as their work at the ML level, is called henceforth **Metalearning**. Thus, BioAutoML creates an automated pipeline working at the data feature and algorithm level. In this research, we have investigated several insights to support our hypothesis, as follows:

- **Hypothesis:** Automated feature engineering and metalearning provides an efficient mechanism to extract features based on different aspects, and automates the feature selection, algorithm(s) recommendation and tuning steps, and hence, a high-quality prediction of categories of ncRNAs in bacteria.

The approach will help us to answer our Research Question (RQ), and consequently be used to confirm or deny the hypothesis, described as follows:

- **RQ:** Is it possible to predict different categories of bacterial ncRNAs using automated feature engineering and metalearning pipelines?

Finally, to support our hypothesis and research question, we have validated BioAutoML into three different case studies using seven bacterial phyla.

Feature engineering

According to Chou’s five-step rule [3, 30], numerically representing biological sequences with an efficient and adequate mathematical expression is one of the most relevant steps to establish an effective statistical predictor for a biological system. In ML, biological sequences must be represented by a fixed number of features (e.g. binary, categorical or continuous), transforming originally unstructured data into a structured format. Feature extraction or feature encoding is a key step in the construction of high-quality ML-based models, determining the effectiveness of trained models in bioinformatics applications, such as biological sequence classification [5, 31, 32]. Nevertheless, the feature engineering process is a time-intensive step and requires domain knowledge of experts [20, 33, 34], which is a complex exercise [34]. Therefore, to develop our proposal and answer our research question, we define the automated feature engineering task formally explained as follows:

- Given a set of sequence data, D , divided into train (D_{train}) and test (D_{test}), a set of feature descriptors, F_d , where $F_d = [f_{d1}, f_{d2}, \dots, f_{dn}]$, our aim is to select the best numerical representation, that is, the feature vector (V_f), combining different feature descriptors in the training set (D_{train}), using an objective function that considers the most important feature descriptor (I_{fd}) to evaluate the best V_f .

Metalearning

One of the main difficulties in applying ML algorithms to a new dataset is selecting the most adequate algorithm for this dataset. Each ML algorithm has an inductive bias, which can be defined by the way it searches for a robust model, e.g. starting with simple models and gradually increasing the complexity of the models, until a robust model is found, and the format adopted to represent the models, e.g. a model represented by a decision tree. Although it can be seen as a limitation, the bias is necessary for learning to occur. As a consequence, each algorithm fits better datasets with particular conformations. Thus, there is no champion ML algorithm that performs better than all the others in every situation, but each ML algorithm performs better than the others on some datasets, which are not known beforehand [35]. A good alternative to select the best ML algorithm for a new dataset is to use previous knowledge regarding the performance of a set of algorithms in previous learning experiences. This is the idea behind a particular approach for metalearning, defined in [36] as learning to learn. According to the authors, metalearning is a research area that investigates how to recommend the most suitable algorithm, or set of algorithms, for a new task. In this study, we use metalearning to do the following:

- Given a set of selected features, recommend the ML algorithm(s) able to induce the best predictive model, which can be a set of algorithms, each one inducing a model, and combine these models into an ensemble (P_{mi}), recommending the best algorithm. Ensemble methods can boost the performance of simple classifiers (e.g. using multiple prediction models for solving the same problem) and has proven its effectiveness in bioinformatics [37–39].

Related works

After a carrying out systematic literature review, we found 14 related studies proposing packages that use feature engineering (feature extraction and selection) and ML algorithms for biological sequence classification: PseAAC [40], propy [41], PseKNC-General [42], SPiCE [43], Pse-in-One [3], repDNA [44], RcpI [45], BioSeq-Analysis [46], PyFeat [31], iLearn [5], iLearnPlus [6], BioSeq-BLM [47], autoBioSeqpy [48] and AutoGenome [49]. For each package, we checked if it uses AutoML for feature engineering, ML algorithms and, when they use these algorithms, tune their hyper-parameters. Table 1 summarizes our findings.

Table 1. Use of AutoML for feature engineering, recommendation of ML algorithm and hyper-parameter tuning

Study	Feature Engineering	ML algorithm	Tuning
PseAAC	–	–	–
propy	–	–	–
PseKNC-General	–	–	–
SPiCE	–	–	–
Pse-in-One	–	–	–
repDNA	–	–	–
Rcpi	–	–	–
BioSeq-Analysis	–	–	–
PyFeat	–	–	–
iLearn	–	✓	✓
iLearnPlus	–	✓	✓
BioSeq-BLM	–	–	–
autoBioSeqpy	–	✓	✓
AutoGenome	–	✓	✓
BioAutoML	✓	✓	✓

The most similar packages to BioAutoML are iLearn, iLearnPlus, autoBioSeqpy and AutoGenome, which apply AutoML to recommend ML algorithms, but they do not use automated feature engineering. The most similar package to our proposal, iLearn, requires an initial configuration file (choosing descriptors and classifiers), which needs domain knowledge from a human expert. Even in its most sophisticated version, iLearnPlus, a file needs to be inserted with the extracted features, instead of automatic feature engineering. The autoBioSeqpy and AutoGenome packages focus on recommending the best deep learning architecture. Thus, to the best of our knowledge, BioAutoML automates the longest pipeline for biological sequence analysis, encompassing feature engineering, ML algorithm recommendation and hyper-parameter tuning. Furthermore, BioAutoML is a user-friendly tool for non-experts.

Looking at more general applications of AutoML, we can cite RECIPE [24] and TPOT [50]. Tree-based Pipeline Optimization Tool (TPOT) is an AutoML tool that optimizes ML pipelines using genetic programming. REsilient Classification Pipeline Evolution (RECIPE) is an AutoML framework with grammar-based genetic programming. One of the most notable methods of RECIPE is how it uses grammar to organize the knowledge acquired from the literature [24]. RECIPE can also be an alternative to TPOT, as TPOT can create ML pipelines that are arbitrary, failing to solve a classification problem, therefore leading to a waste of computational resources [24]. The major difference compared with BioAutoML is the lack of a feature extraction module for biological sequences. These two packages are for any application domain, requiring a previously selected feature vector.

Prediction techniques of ncRNAs in bacteria

Many ML-based techniques have been proposed to identify ncRNAs in bacteria [51–56]. In [55], the authors compare the predictive performance of different techniques for RNAs classes, such as tRNAs, rRNAs and mRNAs. For such, they use normalized minimum free

energy of folding, motif frequency and several RNA-folding parameters, such as base-pairing propensity, Shannon entropy and base-pair distance. The model induced by the Random Forest algorithm presented 89.5% of predictive accuracy. Another related study [51] constructed ML models to discriminate bona fide sRNAs applying five ML algorithms to random genomic sequences from five bacterial species. Seven features were used, including secondary structure. In [53] the support vector machine (SVM) algorithm was applied to a Non-Coding DNA (ncDNA) benchmark dataset, collected from *Saccharomyces cerevisiae*. SVM was also used in [57] to identify sRNAs in bacteria, particularly *Salmonella Typhimurium* LT2, *Escherichia coli* (*E. coli*) K-12 and *Salmonella Typhi* (*S. Typhi*). Some features are combined to achieve better results with accuracy of 81.25% and 88.82% for *E. coli* K-12 and *S. Typhi* Ty2. Unlike BioAutoML, these approaches did not apply an end-to-end ML pipeline.

BioAutoML package

BioAutoML is a user-friendly multi-class and binary classification package that allows the use of automated feature engineering and metalearning, as illustrated by Figure 1. Its use does not require specialized human assistance. BioAutoML only needs a training dataset of biological sequences (FASTA files) to perform an end-to-end ML experiment, from the feature engineering to generating of the predictive model induced by tuned ML algorithms. Nevertheless, the modules implemented in the BioAutoML package can be run independently, i.e. users can just generate the best numerical representation and send it to another ML model generation package, or they can use features extracted from other packages to generate a predictive model. For such, BioAutoML has two components with two modules each: (1) automated feature engineering (feature extraction and selection) and (2) Metalearning (algorithm recommendation and hyperparameters tuning). In the next sections, we briefly describe each component and module.

Feature extraction

This module, which is the first feature engineering stage, extracts feature descriptors using the MathFeature package [28], e.g. Mathematical descriptors (Fourier, Shannon, Tsallis, among others) and Conventional descriptors [Nucleic Acid Composition (NAC), dinucleotide composition (DNC), trinucleotide composition (TNC), ORF Features, Xmer k-Spaced Ymer composition frequency (kGap), Fickett score, among others]. As a result, more than 15 feature extraction techniques can numerically represent information found in biological sequences.

BioAutoML - selection and recommendation

The second module carries out automated feature engineering, selecting the best feature vector and ML algorithm to induce predictive models, which can be an

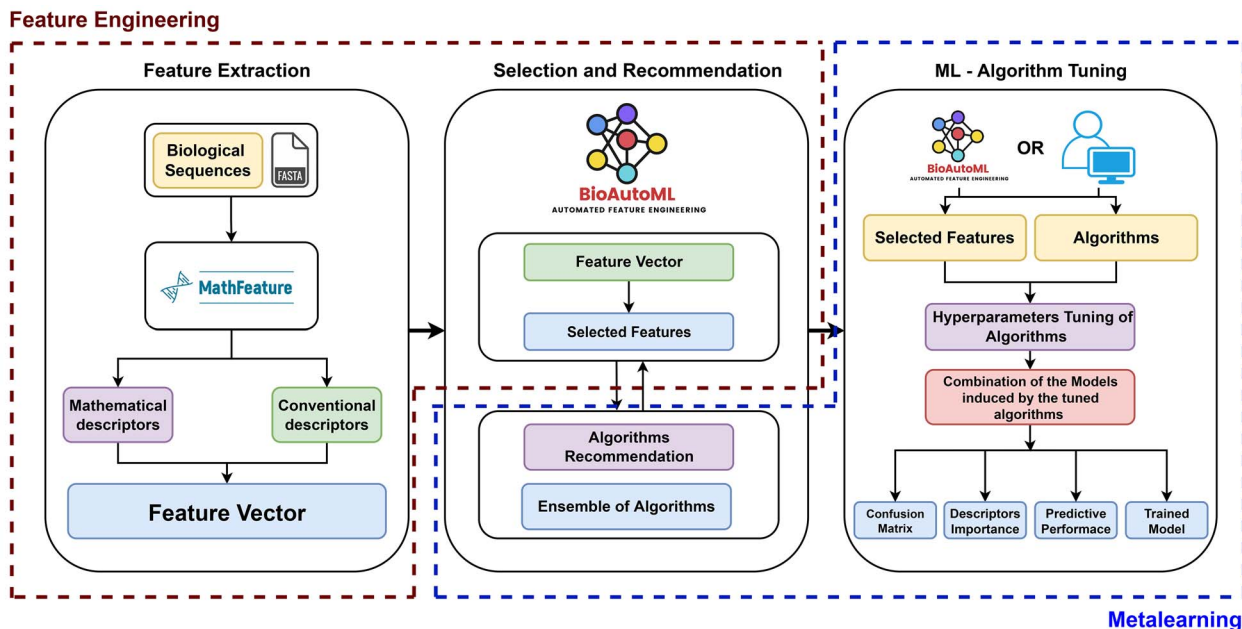


Figure 1. Components implemented in the BioAutoML package: (1) Automated Feature Engineering (feature extraction and selection) and (2) Metalearning (algorithm recommendation and hyper-parameters tuning).

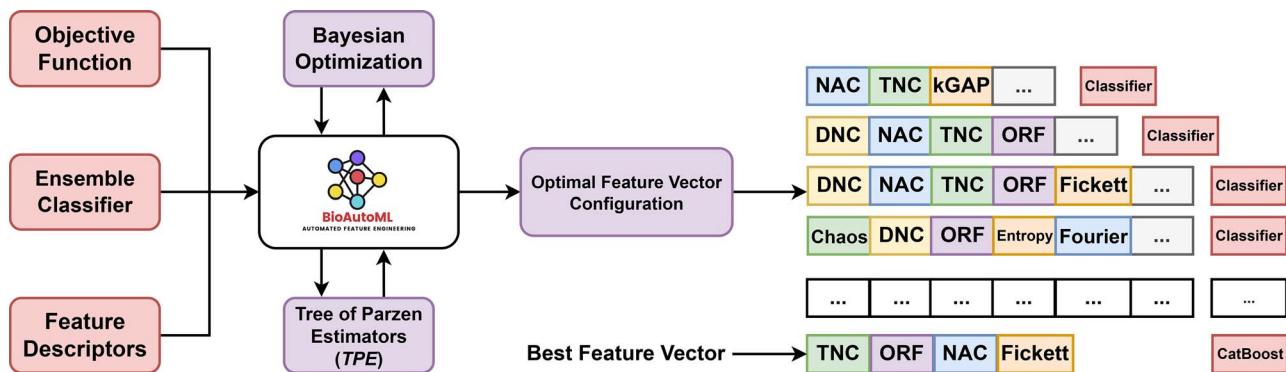


Figure 2. Illustration of how BioAutoML works: selection and recommendation module.

ensemble of predictive models, as shown in Figure 2. For such, it uses the Bayesian optimization technique [29]. We use this technique because there is a large number of alternatives for the types and number of feature descriptors, characterizing an NP-hard problem. This module receives the following as input:

- 1) All feature descriptors extracted by the first module;
- 2) An objective function, e.g. in our case, balanced accuracy for binary problems and F1-score (weighted) for multi-class problems;
- 3) ML algorithms (CatBoost [58], AdaBoost [59], Random Forest [60] and LightGBM [61]). These classifiers are responsible for analyzing the potential of the selected features. These algorithms are used for the wrapper-based feature selection, using different feature subsets as input. We chose these ML algorithms because they have good predictive performance and induce interpretable predictive models, allowing the understanding of the internal decision-making process [62]. The algorithms are widely adopted in the bioinformatics literature [37–39].

To represent the search space (for selecting feature descriptors and recommending ML algorithms), we use a partially binary input vector, e.g. $[1, 0, 1, 0, 0, 1, [2]]$, when the last position can be a value from the set $0, 1, 2, 3$, representing each of the four ML algorithms. In the other position, value 0 means that the feature descriptor was not selected for the subset to be evaluated, and value 1 that was selected. Next, using Bayesian optimization (Hyperopt library - Tree of Parzen Estimators [63]), BioAutoML selects a quasi-optimal feature vector, regarding the predictive performance of the model used in the wrapper. We chose Bayesian optimization based on studies in the literature [29, 64, 65], which demonstrate that it saves time and improves performance, presenting benefits over random search [66]. As can be seen in Figure 2, BioAutoML generates combinations of features and ML algorithm(s) until it finds the best pair (selected feature set, recommended ML algorithm) to send to the fourth module, hyper-parameter tuning. We adopted as stopping criterion for the optimization when the predictive performance reaches a plateau or after

assessing 50 pairs (this number can be changed by the user). Let us remember that this module can recommend one ML algorithm or a set of ML algorithms (when an ensemble model induced by ML algorithms is recommended).

ML algorithm(s) hyper-parameter tuning

The last module is tuning, where users can generate a predictive model using the recommendation of the feature vector and ML algorithms (among those whose implementation is available at BioAutoML). These algorithms will use the feature vector to induce a set of classification models, whose output will be combined using an ensemble-based approach. The quality of the classification models will be affected by the hyper-parameter values used for the recommended ML algorithms. In this work, Bayesian optimization is used to tune their hyper-parameters. For such, we separate part of the training set. The hyper-parameters tuned for each algorithm are defined by their official documentation, e.g. Random Forest (`n_estimators`, `max_features`, `criterion`, `max_depth`, `min_samples_split`, `min_samples_leaf` and `bootstrap`). The optimization stops when the predictive performance reaches a plateau or after assessing 100 possible sets of values. In addition, this module generates important performance analysis files as outputs, e.g. best features, performance results, trained model and feature importance, among others.

Experimental results

The main purpose of this article was to provide a user-friendly and open-access package that allows automated feature engineering and metalearning for the analysis of biological sequences. To assess the relevance of the proposed package, we evaluate its predictive performance in three case studies, described in the following sections: (1) Case Study I - Genomic Pipeline, (2) Case Study II - Pipeline with Annotated Bacterial Sequences and (3) Case Study III - BioAutoML versus other proposed packages for automated experiments.

Case study I—genomic pipeline

We designed an experiment to classify ncRNA families in bacteria, using three known types of bacterial RNA: sRNA, tRNA and rRNA. These RNAs are often considered and studied to analyze ncRNA sequences, e.g. (1) tRNAs and rRNA can contaminate sRNA samples isolated from cytoplasmic total RNA extracts [67], and (2) sRNAs in bacteria, key actors in transcriptional and posttranscriptional regulation [55], emphasizing the importance of accurate prediction of these sequences. To further demonstrate the usefulness of our package, we generated our dataset using a standard bioinformatics pipeline, as shown in Figure 3, extracting sequences from genomes and then applying ML algorithms to predictive models. Our aim is to demonstrate that non-experts can easily connect their pipeline to BioAutoML.

Table 2. Number of sequences from *E. coli* K-12 used for training and testing

RNA type	Samples	Training	Testing
sRNA	166	133	33
tRNA	50	40	10
rRNA	40	32	8

Table 3. Results: *E. coli* K-12—case study I

	Precision	Recall	F1-Score
sRNA	1.00	0.97	0.98
tRNA	1.00	1.00	1.00
rRNA	0.89	1.00	0.94
Macro Average	0.96	0.99	0.98
Weighted Average	0.98	0.98	0.98

To collect the RNAs from genomes, we used the Infernal application [68]. First, in our genomic pipeline, we accessed the Rfam Public MySQL Database obtaining a list of families for each RNA type [69], using the Rfam database in its 14.7 version. Next, with the lists and the complete Rfam Covariance Model (CM), we generated three CM files using `cmfetch`, i.e. one for each RNA type. We use `cmsearch` considering the gathering cutoff (GA) selected by the Rfam curators to extract the sequences for the RNA types [70], given the CM files and a genome. Once the sequences are extracted, they are passed as input to BioAutoML. Thereby, we selected *Escherichia coli* K-12 genome for an initial experiment with the genomic pipeline. In Table 2, we show the sequences generated for training and testing (Hold-out 80% training and 20% test).

BioAutoML returned a combination of three feature descriptors that considered to better numerically represent this dataset, kGap, Fourier and Tsallis entropy. After automated feature engineering, our package selects a feature vector and recommended an ML algorithm to be finely tuned. The final results are shown in Table 3 and Figure 4-A. As our problem is multi-class, we report the main results using precision (Macro and Weighted), recall (Macro and Weighted), F1-score (Macro and Weighted) and confusion matrix [71].

BioAutoML was performed between 0.96 and 0.98 (macro and weighted average) in this initial experiment, showing a robust numerical representation for the input genome. Next, we used the recommended algorithm to induce a predictive model to classify new unknown sequences. To test the potential of our package, we did a more complex experiment using bacterial phyla, as shown in Table 4. We analyzed the generalization potential for the classification of new bacterial genomes as new organisms will not be in the training set, e.g. training with *Chloracidobacterium* and classifying a new genome as *Terriglobus roseus*. Moreover, according to [72], the different GC content skew patterns throughout bacterial phylogenetic groups could change relevant characteristics of the sequences' primary structure used for the generation of descriptors. The bacterial phyla

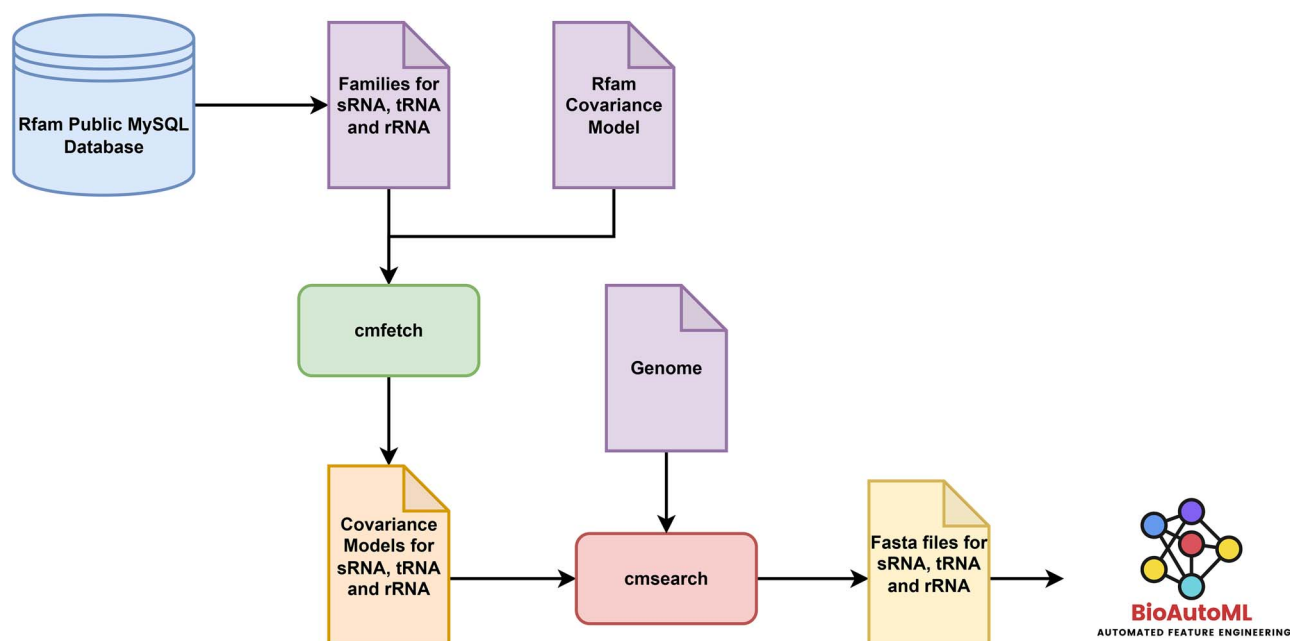


Figure 3. Case Study I - Genomic Pipeline.

Table 4. Number of sequences for bacterial phyla by RNA type

Phylum	Bacteria	sRNA	tRNA	rRNA	Gram	NCBI Taxonomy ID	Used as
Acidobacteria	<i>Chloracidobacterium</i>	21	50	7	Negative	2821542	Train
	<i>Terriglobus roseus</i>	4	47	11	Negative	926566	Test
Actinobacteria	<i>Corynebacterium diphtheriae</i>	–	43	14	Positive	1450520	Train
	<i>Mycobacterium tuberculosis</i>	11	45	8	Positive	83332	Test
Bacteroidota	<i>Flavobacterium sediminis</i>	–	41	19	Negative	2201181	Train
	<i>Mucilaginibacter gossypii</i>	–	46	20	Negative	551996	Test
Cyanobacteria	<i>Oscillatoria acuminata</i>	9	65	19	Negative	56110	Train
	<i>Prochlorococcus marinus</i>	2	39	7	Negative	167539	Test
Firmicutes	<i>Staphylococcus aureus</i>	84	35	31	Positive	93061	Train
	<i>Staphylococcus epidermidis</i>	44	38	11	Positive	1282	Test
Proteobacteria	<i>Escherichia coli</i>	166	50	40	Negative	83333	Train
	<i>Salmonella enterica</i>	118	52	39	Negative	99287	Test
Verrucomicrobia	<i>Akkermansia glycaniphila</i>	–	44	7	Negative	1679444	Test
	<i>Luteolibacter ambystomatis</i>	29	46	14	Negative	2824561	Train

Table 5. Results: Bacterial phyla—case study I

	Precision	Recall	F1-Score
sRNA	0.97	0.97	0.97
tRNA	0.98	1.00	0.99
rRNA	0.99	0.95	0.97
Macro Average	0.98	0.97	0.98
Weighted Average	0.98	0.98	0.98

used for this experiment are shown in Table 4 and Figure 4-B.

We randomly selected one type of bacteria from each phylum for a fair split, as shown in Table 4. We used seven bacteria for training and testing. The number of sequences generated by RNA type is also presented. The sequences were extracted using the same pipeline exposed in Figure 3. The performance metrics can be seen in Table 5.

Our package recommended six feature descriptors that best represent this new scenario: NAC, TNC, kGap, Fourier and ORF. Two of these descriptors were in the initial experiment (kGap and Fourier). Again, BioAutoML showed good predictive results, between 0.97 and 0.98 (macro and weighted average). However, we were classifying new bacterial sequences that were not in training, indicating the package’s ability to recommend robust feature descriptors for the input problem.

Case study II—pipeline with annotated bacterial sequences

For this case, we extracted annotated bacterial sequences from databases, standard pipeline in several studies [37, 48, 49]. We used eight classes for this analysis: pre-miRNA, miRNA, snoRNA, snRNA, tmRNA, tRNA, rRNA and mRNA. Compared with case study I, we worked with specific types of sRNAs to study BioAutoML capacity for

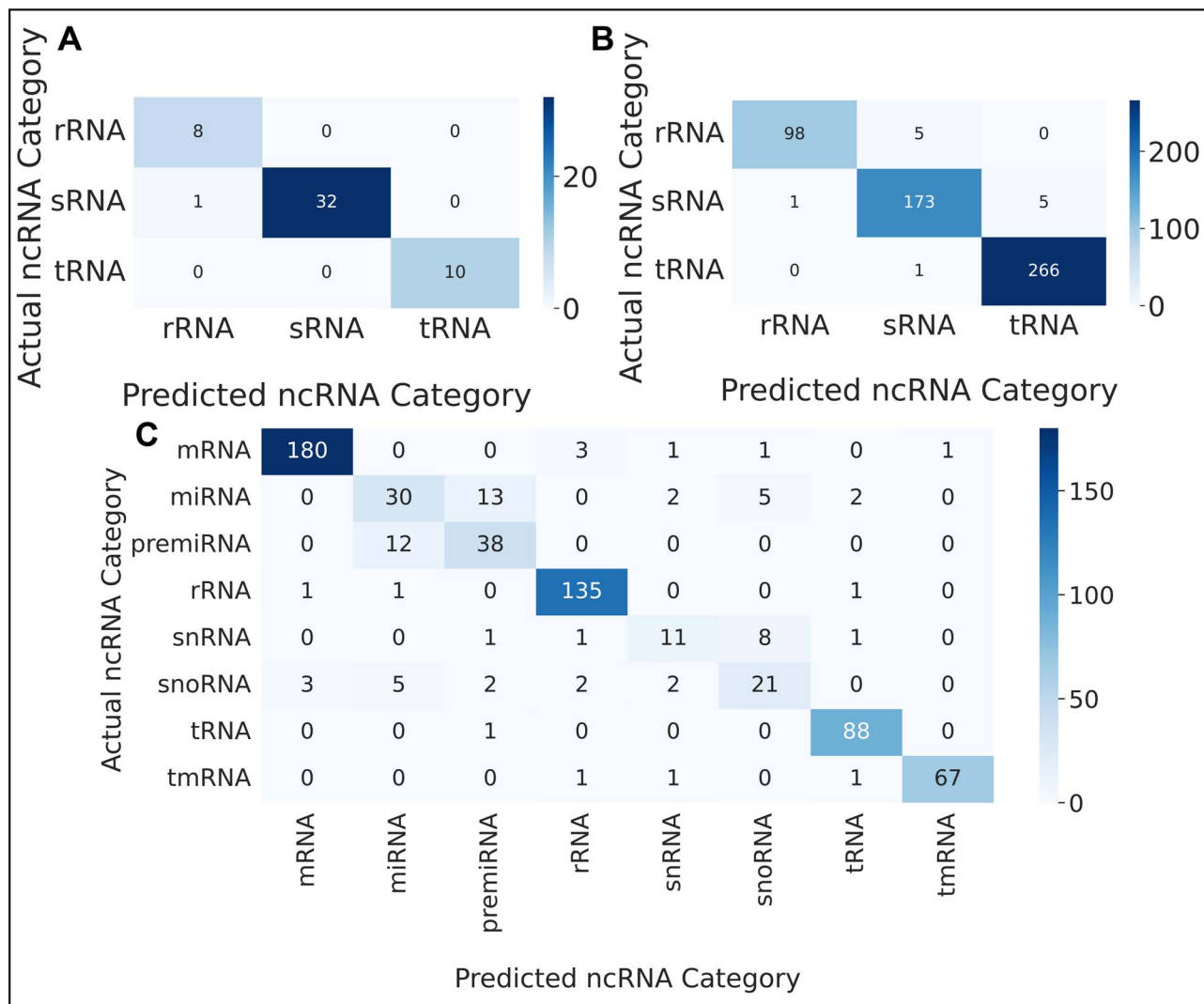


Figure 4. Confusion matrix of the experiments. (A) Case Study I—*E. Coli* K-12. (B) Case Study I—Bacterial phyla. (C) Case Study II—Annotated Bacterial Sequences.

dealing with more classes. In addition, we used mRNA as a counterpoint by containing coding regions compared with the ncRNAs. These classes can be separated into regulatory, and housekeeping ncRNAs [73]. We also demonstrate the performance metrics for the application considering recurrent problems such as the classification of pre-miRNA between miRNA [21, 74, 75], and the prediction of miRNA by itself [76]. There are few studies related to the prediction of miRNAs in bacteria [77] as the number of these annotated sequences is still small [78].

We collected ncRNA sequences from RNAcentral, a database of ncRNA sequences that provide a single access point to at least 44 RNA resources in its last version [23]. We accessed the RNAcentral Public Postgres database running SQL queries to filter active cross-reference sequences by type, limited to 1000 sequences, and restricting them to bacterial organisms. With the results from the queries, FASTA files for each class were created. Considering how collecting from diverse databases could bring some redundancy, we used CD-HIT Est [79] to cluster the sequences, removing redundancy

at 95% similarity. The same preprocessing pipeline was applied for mRNA, but we collected the sequences from GenBank [80], filtering for bacterial organisms. In Table 6, we show the numbers of examples collected from RNAcentral and GenBank, the numbers after applying CD-HIT Est with the preprocessing method used in BioAutoML, and how many of these sequences are used for training and testing (Hold-out 80% training and 20% test). The results generated by BioAutoML are presented in Table 7 and Figure 4-C.

Again, our package presented robust performance, even for eight classes, ranging from 0.79 to 0.89 (macro and weighted average). By analyzing each class individually, we observed a better performance for rRNA (F1-Score: 0.96), tRNA (F1-Score: 0.97), tmRNA (F1-Score: 0.97), pre-miRNA (F1-Score: 0.72) and mRNA (F1-Score: 0.97), but lower performance for miRNA, snoRNA and snRNA (F1-Score: both around 0.60). However, multi-class classification problems present more challenges than binary classification problems, e.g. an imbalanced dataset. Even so, BioAutoML recommended a good

Table 6. Number of sequences used for training and testing—case study II

RNA type	Samples	Preprocessing	Training	Testing
pre-miRNA	327	253	203	50
miRNA	464	263	211	52
snoRNA	331	178	143	35
snRNA	176	113	91	22
tmRNA	1000	350	280	70
tRNA	1000	445	356	89
rRNA	1000	687	549	138
mRNA	1000	702	514	188

Table 7. Results generated by BioAutoML in the case study II

	Precision	Recall	F1-Score
pre-miRNA	0.69	0.76	0.72
miRNA	0.63	0.58	0.60
snoRNA	0.60	0.60	0.60
snRNA	0.65	0.50	0.56
tmRNA	0.99	0.96	0.97
tRNA	0.95	0.99	0.97
rRNA	0.95	0.98	0.96
mRNA	0.98	0.97	0.97
Macro Average	0.80	0.79	0.80
Weighted Average	0.89	0.89	0.89

feature vector formed by the descriptors NAC, DNC, TNC, kGAP, ORF feature, Fourier and Tsallis entropy.

Case study III—comparing BioAutoML with other AutoML packages

In this last case study, we compared BioAutoML with well-known AutoML packages used in different classification tasks [81, 82]. In our literature review, we did not find any tool for biological sequence classification with automatic feature engineering, characterizing the innovative nature of BioAutoML. To allow the experimental comparison, we chose two packages using AutoML: RECIPE [24] and TPOT [50]. The major difference compared with BioAutoML is the lack of a feature extraction module for biological sequences.

Similar to BioAutoML, they include feature selection, algorithm recommendation and hyper-parameter tuning. These two packages are for any application domain, requiring a previously selected feature vector. For a fair comparison, we used the same output from the feature extraction module in the AutoML packages (all feature descriptors), feature descriptors recommended by BioAutoML and datasets from the previous case studies. These experiments assess whether BioAutoML can build predictive models with recommended feature vectors and ML algorithms as robust as RECIPE and TPOT, which are well known for the quality of their pipelines [81, 82]. The BioAutoML results shown in Table 8 are the average of 10 runs. All experiments, package configurations (default parameters) and datasets can be consulted in our repository (<https://github.com/Bonidia/BioAutoML-Case-Studies>). We performed the experiments using a machine with Intel Core i3-9100F CPU (3.60GHz), 16GB memory and running in Debian GNU/Linux 10.

As can be seen, we observed similar performance between BioAutoML and other tools (TPOT and RECIPE) in CS-I and CS-II, considering two different types of experiments: (i) with all feature descriptors, and (ii) with the vector recommended by BioAutoML. We also noted the improvement prediction of TPOT when the input was provided by vector recommended from BioAutoML (gain of 2% and 1% for CS-I and CS-II, respectively). Another interesting result is related to computational time to generate an ML model when both TPOT and RECIPE spent a huge computational effort (416.26 and 272.46 min, respectively in CS-II), while BioAutoML spent 85.02 min. BioAutoML also recommends the best vector be extracted automatically. It is important to highlight that both TPOT and RECIPE do not have any mechanism to recommend the best vector to be automatically extracted for biological sequences. Finally, the statistical significance was applied in this case study (difference in F1-Score (Weighted)), using Friedman’s test, indicating that there is no statistical significance in performance (P – value = 0.156, using α = 0.05), suggesting that our proposal is as robust as known methods in the literature.

Discussion

We assessed BioAutoML in three case studies with ncRNA sequences. We consider different ncRNA categories for multi-class classification tasks using ncRNA bacteria data. For case study I, we used Infernal, which builds statistical models of RNA secondary structure and sequence consensus called Covariance Models (CMs) [68]. Infernal is still widely used for genome annotation, especially for detecting ncRNA [83, 84]. However, one of its limitations for creating CM is the need for a secondary structure model for the RNA families. The experimental results from this case study shows the success of BioAutoML in using only primary structure features to predict what we found with Infernal.

For case study II, we considered eight classes, including miRNAs. Although some studies in the literature consider that prokaryotes do not have true miRNA as in eukaryotes [85, 86], recently, many similarities between the noncoding sequences were observed, indicating miRNA-like mechanisms in prokaryotes, which resulted in the annotated sequences used in our study [86, 87]. Prokaryotic miRNAs can also accumulate in the nucleolus as pre-miRNAs, and mature miRNAs [87], emphasizing the challenge for an accurate classification in these two classes. Other classes used, such as snoRNA [88] and snRNA [89], are also relatively rare in prokaryote organisms. Nevertheless, it is relevant to discover more of these noncoding sequences in bacteria with the advancements in RNA sequencing technology and ML-based algorithms.

Finally, in case study III, we observed the robust predictive performance of BioAutoML when compared with AutoML tools found in the literature, mainly due to the quality of their pipelines [81, 82]. The experimental

Table 8. Case study III—BioAutoML versus other AutoML packages

Dataset	Version	Precision (Weighted)	Recall (Weighted)	F1-Score (Weighted)	Time (min)
CS-I-phyla	BioAutoML	0.97	0.97	0.97	16.34
Recommended Feature Descriptors	RECIPE	0.97	0.97	0.97	32.48
Recommended Feature Descriptors	TPOT	0.99	0.99	0.99	72.41
All Feature Descriptors	RECIPE	0.96	0.96	0.96	30.46
All Feature Descriptors	TPOT	0.98	0.98	0.98	46.39
CS-II	BioAutoML	0.88	0.88	0.88	85.02
Recommended Feature Descriptors	RECIPE	0.87	0.61	0.68	272.46
Recommended Feature Descriptors	TPOT	0.89	0.89	0.89	416.28
All Feature Descriptors	RECIPE	0.77	0.36	0.38	151.12
All Feature Descriptors	TPOT	0.90	0.89	0.89	338.55

results indicated the efficiency of the feature extraction module, which can extract features based on different aspects, automated feature selection, algorithm(s) recommendation and tuning steps. Together, they predicted the categories of ncRNAs in bacteria with high predictive accuracy, even when the number of classes was increased.

Conclusion

In this article, we propose and experimentally evaluate a new package, BioAutoML, to classify biological sequences. BioAutoML uses AutoML to select the best feature vector from a set of descriptors extracted by the MathFeature package, to recommend the best ML algorithms, and tune the hyper-parameters of the recommended algorithm. For such, it initially performs automated feature engineering and metalearning for noncoding sequences in bacteria, which has the potential to accelerate new studies in bioinformatics. We develop a package that does not require specialized human assistance, supporting research on challenging problems in biological sequence analysis. Our findings support our hypothesis, showing the benefits of using automated feature engineering and metalearning. Although in this study, BioAutoML is applied only to ncRNA sequences in bacteria, it can be used in other DNA/RNA sequence scenarios. We focused exclusively on bacteria, due to the biotechnological potential existing in the investigated strains. Nevertheless, the first module of BioAutoML is an important task for providing feature descriptors for different types of sequences (nucleotides or proteins, i.e. prediction of structural features along the primary sequence of amino acids). We also used our previous framework, MathFeature, to extract features for BioAutoML. BioAutoML can be used for binary and multi-class classification problems, allowing its integration with many existing packages. Finally, in future work, we intend to expand the BioAutoML to proteins and add new feature extraction packages, e.g. iLearn, BioSeq-Analysis and BioSeq-BLM, testing other feature selection methods such as combining Bayesian Optimization and Lipschitz Optimization, Genetic Algorithm and Genetic programming.

Key Points

- The first study to propose an automated feature engineering and metalearning pipeline for ncRNA sequences in bacteria;
- BioAutoML can be used in multi-class and binary problems;
- BioAutoML can be employed in other DNA/RNA sequences scenarios;
- BioAutoML can accelerate new bioinformatics studies, reducing the feature engineering time-consuming stage and improving the design and performance of ML pipelines;
- BioAutoML reduces requirement of human expert assistance.

Availability of data and materials

BioAutoML, documentation and datasets are available in the Github repository: <https://github.com/Bonidia/BioAutoML>.

Acknowledgments

The authors would like to thank USP, CAPES, Google (LARA - 2021), CNPq, and FAPESP for the financial support for this research.

Funding

This project was supported by the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES) - Finance Code 001, Google (LARA - 2021), Universidade de São Paulo (USP) and São Paulo Research Foundation (FAPESP) - grants #2013/07375-0, #2021/08561-8.

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