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# A Neural Network Approach for the Analysis of Reproducible Ribo–Seq Profiles

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**Abstract:** In recent years, the Ribosome profiling technique (Ribo–seq) has emerged as a powerful method for globally monitoring the translation process in vivo at single nucleotide resolution. Based on deep sequencing of mRNA fragments, Ribo–seq allows to obtain profiles that reflect the time spent by ribosomes in translating each part of an open reading frame. Unfortunately, the profiles produced by this method can vary significantly in different experimental setups, being characterized by a poor reproducibility. To address this problem, we have employed a statistical method for the identification of highly reproducible Ribo–seq profiles, which was tested on a set of *E. coli* genes. State-of-the-art artificial neural network models have been used to validate the quality of the produced sequences. Moreover, new insights into the dynamics of ribosome translation have been provided through a statistical analysis on the obtained sequences.

**Keywords:** Ribo–seq profiling; neural networks; prediction of translation speed; ribosome dynamics; CNN



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## 1. Introduction

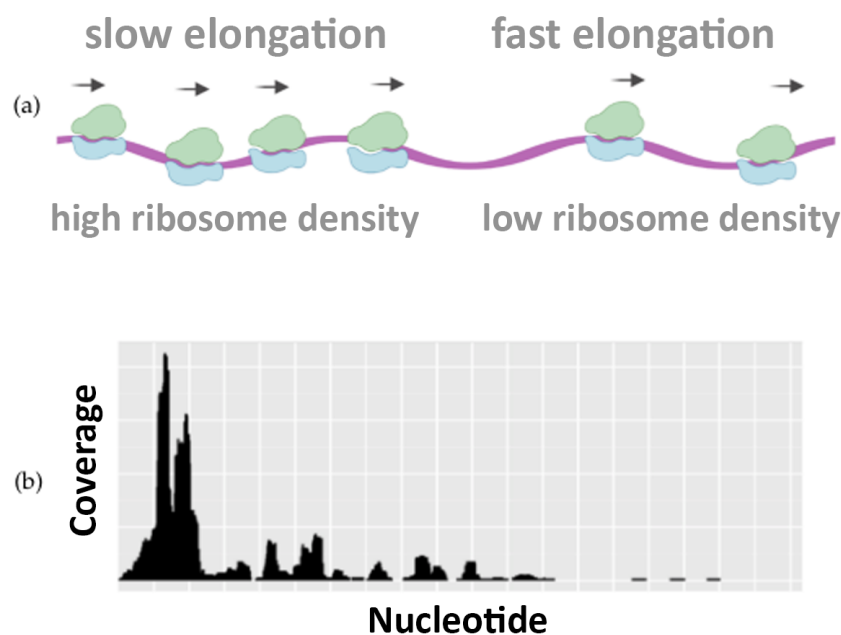
Ribosomes perform protein synthesis from mRNA templates by a highly regulated process called translation. Translation control plays a key role in the regulation of gene expression, both in physiological and pathological conditions [1].

The advent of high–throughput methods to measure the levels of gene expression has revealed the implications of multiple factors that might impact the rate at which an mRNA is translated. In recent years, the Ribosome profiling technique (Ribo–seq) has emerged as a powerful method for globally monitoring the translation process in vivo at single nucleotide resolution [2]. The application of this method to a different number of organisms subjected to different conditions, from the deprivation of nutrients in bacterial cells to the development of cancer in human cells, has allowed to investigate fundamental aspect of cell biology [3].

Interestingly, the nucleotide–level resolution of Ribo–seq experiments reveals the density of the ribosomes at each position along the mRNA template. Local differences in the density of Ribosome Protected Fragments (RPFs) along the Open Reading Frame (ORF) reflect differences in the speed of translation and elongation, determining regions where the translation is slower or faster.

Figure 1 illustrates how the translation speed is not uniform, highlighting the differences in ribosome occupancy. This piece of information is well visible in Ribo-seq profiling data and can be used to infer how the codon usage, the protein sequences, and other features can regulate the speed of translation [4].

Unfortunately, the reproducibility of Ribo-seq experiments can be affected by multiple variables due to the complexity of the experimental protocol and the lack of standardization in computational data analysis [5].



**Figure 1.** Ribosome footprint density along the mRNA. The schematic distribution of translating ribosomes along the mRNA (a) and their ribosome profiles (b). Ribo-seq data show differences in the density of ribosomes: regions of fast elongation accumulate fewer ribosomes (low density) with respect to regions of slow elongation (high density).

Therefore, our work aims to overcome the aforementioned limitations by introducing a new statistical approach, designed to extract a set of highly reproducible profiles.

In particular, inspired by the seminal work proposed in [6], we perform a novel analysis procedure for Ribo-seq data that allows to identify the reproducible Ribo-seq profiles emerging from the comparison of independent Ribo-seq experiments performed in different laboratories under the same conditions. These significantly reproducible profiles are then collected into a library of consensus sequences, in which sub-regions characterized by different translation speeds can be isolated. The aforementioned procedure has been applied to *E. coli* sequences (*Escherichia Coli* (*E. coli*) is a bacterium that lives in the lower intestine of warm-blooded animals and has a genome composed by approximately 4,600,000 base pairs. *E. coli* contains a total of 4288 genes, with coding sequences which are long, on average, 950 base pairs and separated, on average, from 118 bases. Considering the protein counterpart, in *E. coli*, the average length of a coding region is 316.8 codons, whereas less than 1.8% of the genes are shorter than 60 codons.), resulting in 40 highly reproducible profiles.

Differently from [6], based on the collected data, a statistical analysis has been carried out that gave new insights on the dynamics of the ribosome translation, showing a statistically significant difference in the nucleotide composition between sub-sequences characterized by different translation speeds. Moreover, to validate the procedure, the selected highly reproducible profiles have been analyzed through state-of-the-art Machine Learning (ML) models, accurately classifying subsequences according to their speed of translation (slow or fast). We have made our source code public available (<https://github.com>).

[com/pandrein/Ribo-Seq-analysis](https://www.mdpi.com/pandrein/Ribo-Seq-analysis), accessed on 29 June 2022). Furthermore, these experiments allowed to discover that the translation speed is modulated both by the nucleotide composition of the sequences and by the order in which they appear within each sequence.

The rest of the paper is organized as follows: Section 2 collects works from the literature dealing with related topics; Section 3 describes the ribosome profiling data extraction and preprocessing, together with their analyses based on both statistical and ML methods; Section 4 summarizes the results, discussing their meaning and their biological interpretation. Finally, Section 5 draws some conclusions and derives future perspectives.

## 2. Related works

The Ribosome profiling approach offers a promising method for developing unbiased translation models from data, but the quantitative analysis of ribosome profiling data is challenging, because of high measurement variance and the inability to distinguish the ribosome rate of translation. The Ribosome profiling strategy based on deep sequencing of ribosome-protected mRNA fragments enables genome-wide investigation of translation at the codon and sub-codon resolution [7].

In recent years, techniques based on machine learning have been employed with increasing frequency and intensity in many different fields, ranging from computer vision [8–11] to natural language processing [12–14] and bioinformatics [15,16]. The popularity of these approaches stems from their success in the automatic inference of complex functions directly from the data. In particular, both statistical and ML approaches have been successfully applied to non-biological *sequential data classification* tasks [17–19], in which each sequence is associated with a class label and the classification is performed on the whole sequence. Within bioinformatics, examples of ML applications include the prediction of splicing patterns and protein secondary structures, protein-protein interface prediction, protein subcellular localization, drug side-effect prediction, and DNA/RNA motif mining, to name just a few [20–23]. Moreover, ML and deep learning approaches have been used to process Ribo-seq data for gene annotation in prokaryotes [24], to predict ribosome stalling [25] and for micropeptide identification [26]. In particular, in [27], a deep learning based approach, called RiboMIMO, was proposed, based on a multi-input and multi-output framework, for modeling the ribosome density distributions of full-length mRNA Coding Sequence (CDS) regions. Through considering the underlying correlations in translation efficiency among neighboring and remote codons, and extracting hidden features from the input full-length coding sequence, RiboMIMO can accurately predict the ribosome density distributions along with the whole mRNA CDS regions, a problem strictly correlated with the one we intend to face in this paper. Indeed, we propose a machine learning-based approach to validate the extraction procedure of highly reproducible profiles, by classifying the translation speed of the extracted regions, which deeply depends on the ribosome density along mRNA.

## 3. Materials and Methods

In this work, a new software, written in Python, has been developed, which reproduces the procedure described in [6]. In this section, the method used to obtain a set of reproducible Ribo-seq profiles and their analysis through statistical and ML methods are presented. In particular, in Section 3.1, the procedure employed to extract the profiles is described, while in Section 3.3, a statistic analysis of the nucleotide composition is presented. Finally, in Section 3.4, two different ML approaches are proposed to analyze the data and assess their quality.

### 3.1. Ribosome Profiling Data Extraction

The Ribosomal profiling technique (Ribo-seq) is currently the most effective tool to study the protein synthesis process in vivo. The advantage of this method, over other approaches, lies in its ability to monitor translation by precisely mapping the position and number of ribosomes on an mRNA transcript. Ribo-seq involves the extraction of mRNA

molecules associated with ribosomes undergoing active translation and a digestion phase, during which the RNase enzyme processes all the RNA molecules, with the exception of the “protected” parts, to which ribosomes are attached. This step is followed by rRNA depletion and preparation of the sequencing library as in an RNA-seq approach. However, since reads are obtained relating only to actively transcribed mRNA molecules, Ribo-seq better reflects the translation rate than mRNA abundance alone, although it requires particularly laborious preparation and is only applicable to species that have a reference genome available. The strategy employed in our work consists in the identification of high resolution Ribo-seq profiles through the systematic comparison of Ribo-seq datasets referring to experiments performed independently in different laboratories and in different time periods.

In particular, the approach is composed by the following phases:

- **Preprocessing**—the ORF-specific ribosome profiling data from multiple datasets are collected and then processed by a bioinformatic pipeline;
- **Signal digitalization**—Ribo-seq profiles are digitalized by associating to each nucleotide a slow or fast label;
- **Comparison of digital profiles**—Digital profiles are used to quantify similarities and differences between Ribo-seq profiles of different datasets referring to the same ORF.
- **Identification of significantly reproducible Ribo-seq profiles**—A set of highly reproducible profiles is obtained and, among them, reproducible sub-sequences are identified.

### 3.1.1. Preprocessing of Ribosome Profiling Data

To illustrate the statistical procedure employed in this work, we analyse a set of *E. coli* Ribo-seq profiles. For this purpose, the data stored in the Gene Expression Omnibus [28] repository were used. Specifically, our analysis concerns a systematic comparison of Ribo-seq profiles, each belonging to a different GEO series (A series is a collection of datasets that include at least one group of data—sample—from Ribo-seq experiments performed on *E. coli* in various conditions according to the most used experimental protocol.), referring to experiments performed culturing wild-type *E. coli* strains under control conditions. In particular, our analysis regarded a subset of eight samples (labelled from Dataset 1 to Dataset 8) obtained through experiments characterised by K-12 MG1655 genotype and cultured in a MOPS-based medium. Table 1 reports the GEO Series ID and GEO sample ID of raw Ribo-seq dataset used in this experiment.

**Table 1.** The samples chosen for our analysis belong to different GEO series. Column 1: Dataset ID chosen to refer to the eight samples in this work; Column 2: GEO Series ID; Column 3: GEO Sample ID; Column 4: references. The listed ID can be used as access keys to the GEO Database (<https://www.ncbi.nlm.nih.gov/geo/>, accessed on 29 June 2022) to find a detailed description of both specific series and samples.

| Dataset ID | GEO Series ID | GEO Sample ID | Ref  |
|------------|---------------|---------------|------|
| Dataset 1  | GSE64488      | GSM1572266    | [29] |
| Dataset 2  | GSE90056      | GSM2396722    | [30] |
| Dataset 3  | GSE72899      | GSM1874188    | [31] |
| Dataset 4  | GSE53767      | GSM1300279    | [32] |
| Dataset 5  | GSE51052      | GSM1399615    | [33] |
| Dataset 6  | GSE77617      | GSM2055244    | [34] |
| Dataset 7  | GSE35641      | GSM872393     | [35] |
| Dataset 8  | GSE88725      | GSM2344796    | [36] |

To reconstruct the Ribo-seq profiles starting from raw Ribo-seq data, represented by the FASTA format (The FASTA format is a text-based format for representing nucleotide sequences in which base pairs are indicated using single-letter codes [A,C,G,T] where A = Adenosine, C = Cytosine, G = Guanine, T = Thymidine), the reads are mapped against

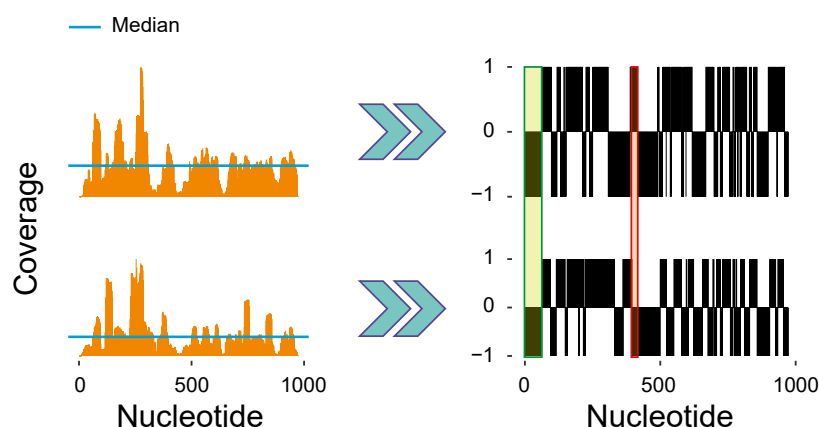
the whole set of coding sequences in *E. coli*, taken from the EnsemblBacteria database [37]. Then, we extracted and counted the number of reads mapping to each gene from the SAM alignment file (The Sequence Alignment Map (SAM) is a text-based format originally used for storing biological sequences aligned to a reference sequence.) using BEDTools [38]. The genomic coordinates are stored in a BED file (The BED file is a tab-delimited text file used to store genomic regions where each feature is described by chromosome, start, end, name, score, and strand.) to build the Ribo-seq profiles representing the input of the subsequent analysis. Each ORF can be associated to a specific Ribo-seq profile, a histogram that counts the number of reads that cover each nucleotide position. To realize the pairwise comparison of ORF-specific Ribo-seq profiles coming from independent datasets, we decided to proceed as described in the following.

### 3.1.2. Signal Digitalization Strategy

Firstly, we selected the ORFs in common between all the eight datasets highlighted in Table 1. For each ORF of each dataset, we generated a Ribo-seq profile. The Ribo-seq profiles (Figure 2, left side) are digitalized by comparing the profile heights at each nucleotide position (*coverage*) with its median value, computed along the entire ORF. We assign +1 to the positions having a coverage value higher than the median, -1 otherwise. The result is a *digital Ribo-seq profile* (Figure 2, right side) for each Ribo-seq profile, i.e., a vector having the length of the associated ORF and containing a sequence of -1 and +1.

### 3.1.3. Comparison of Digital Profiles

The digitalized profiles can be compared to detect matches, i.e., nucleotides characterized by an identical label (Figure 2). Calculating the relative number of matches (the ratio between the number of matches and the length of the ORF) yields the *matching score* ( $s_{i,k}$ ). Intuitively, a matching score close to one could indicate a high degree of similarity between a pair of digitalized profiles, whereas a score around one-half could mean a very poor overlap because the observed matches are likely to have occurred by chance. Given each score has a certain probability of being obtained by chance, we decided to implement a statistical test able to assess the significance of the matching score [6].



**Figure 2.** Pairwise comparison of two Ribo-seq profiles of the *ispB* gene. Two independent Ribo-seq profiles (left) are obtained by computing the coverage at each nucleotide position within the ORF. x-axis: position within the ORF (nucleotides); y-axis (top): relative coverage (number of mapping reads/total number of reads) mapping on the ORF. The Ribo-seq profiles are compared to the median coverage to produce the digitalized  $\pm 1$  profiles (right). The digitalized profiles can be easily compared to detect matches (e.g., green rectangle) and mismatches (e.g., red rectangle). The ratio between the number of matches and the total number of nucleotides in the ORF gives the matching score. A score equal to one means a perfect match between the two profiles, whereas a score equal to one-half means a poor matching [6].



For any Ribo-seq profile involved in a pairwise comparison, a large number of randomized profiles is generated by re-distributing the reads in random positions on the reference ORF. The randomized profiles are, in turn, compared pairwise yielding a large number of random matching scores that build each null distribution. Reiterating this process, we generated  $10^4$  pairs of random Ribo-seq profiles and an equal number of digitalized random profiles that, compared pairwise, yielded  $10^4$  random matching scores. These scores are used to build an ORF-specific null distribution which allows us to estimate the probability of obtaining each similarity score just by chance.

Given a pair of Ribo-seq profiles, the similarity score resulting from their comparison is tested for significance by comparing it to the corresponding ORF-specific null distribution. For each  $s_{i,k}$  and the corresponding null distribution, we computed a z-score  $z_{i,k}$ , mapping each similarity score on a standard normal distribution.

Subsequently, we computed the  $p$ -values  $p_{i,k}$ , as the integral:

$$p_{i,k} = \int_{z_{i,k}}^{+\infty} N_S(z) dz$$

where  $N_S(z)$  is the standard normal distribution. Mapping the matching score on the null distribution will yield the  $p$ -value. The results of this process can be summarised into a matrix (called *p-value matrix*) containing all the computed  $p$ -values and composed by one column for each pairwise comparison and one row for each considered ORF (for a total of 3588 rows and 28 columns). For the sake of simplicity, Table 2 collects a small extract of such matrix. Each  $p_{i,k}$  quantifies the probability of obtaining a similarity score at least as extreme as the corresponding  $s_{i,k}$ , given that the null hypothesis is true. In our context, the lower the  $p$ -value, the lower the probability that the similarity between the compared pairs of (digitalized) Ribo-seq profiles occurs by chance. If the  $p$ -value will result below a given threshold, the compared Ribo-seq profiles will exhibit a significant degree of similarity.

**Table 2.** Excerpt of the  $p$ -value matrix. Each column corresponds to a pairwise comparison between two datasets while each row contains the gene ID. For the sake of readability, only three columns and five rows are reported here.

|             | Dataset 1 vs. Dataset 2 | Dataset 1 vs. Dataset 3 | Dataset 1 vs. Dataset 4 |
|-------------|-------------------------|-------------------------|-------------------------|
| <b>alr</b>  | 0.769298564             | 0.122368427             | 0.632263895             |
| <b>modB</b> | 0.165522551             | 0.056591384             | 0.601754757             |
| <b>cysZ</b> | 0.005770742             | 0.00011569              | 0.2021111               |
| <b>dfp</b>  | 0.002343099             | 0.000384015             | 0.093624025             |
| <b>fruB</b> | 0.566785395             | 0.85548442              | 0.381131384             |

### 3.1.4. Identification of Significantly Reproducible Ribo-seq Profiles

Our strategy consists in inspecting each row of the  $p$ -value matrix. We define reproducible the Ribo-seq profiles referring to those rows featuring all the  $p$ -values below a chosen significance threshold. To cast our strategy into a more rigorous statistical framework, we exploit the False Discovery Rate (FDR) concept and the Benjamini-Hockberg (BH) correction method for multiple tests. In this experiment, for any given row of the  $p$ -value matrix, we set an FDR threshold of 0.01. This means that we accept that 1% of profiles are reproducible by chance. Then, we counted how many  $p$ -values in each row resulted significant according to the BH method, and we defined reproducible those Ribo-seq profiles associated with the rows where 80% of the  $p$ -values are significant. Following this strategy, we found that, out of 3588 genes that are common to the eight datasets, 40 genes, listed in Table 3, have a significantly reproducible Ribo-seq profile.

**Table 3.** Genes with significantly reproducible Ribo-seq profiles across the eight datasets. Column 1: Gene ID. Column 2: Annotation.

| Genes ID | Annotation   |
|----------|--|
| rodZ     | Cytoskeleton protein RodZ  |
| arcB     | Aerobic respiration control sensor protein ArcB                      |
| dld      | Quinone-dependent D-lactate dehydrogenase                            |
| dnaX     | DNA polymerase III subunit tau                                       |
| fhuA     | Ferrichrome outer membrane transporter/phage receptor                |
| glnA     | Glutamine synthetase   |
| gltB     | Glutamate synthase NADPH large chain                                 |
| hisS     | Histidine-tRNA ligase  |
| infB     | Translation initiation factor IF-2                                   |
| katG     | Catalase-peroxidase  |
| malF     | Maltose transport system permease protein MalF                       |
| metG     | Methionine-tRNA ligase   |
| mukB     | Chromosome partition protein MukB                                    |
| ompC     | Outer membrane protein C   |
| parC     | DNA topoisomerase 4 subunit A  |
| secY     | Protein translocase subunit SecY                                     |
| purL     | Phosphoribosylformylglycinamide synthase                             |
| rne      | Ribonuclease E   |
| sucA     | 2-oxoglutarate dehydrogenase E1 component                            |
| tufA     | Elongation factor Tu 1   |
| tufB     | Elongation factor Tu 2   |
| leuA     | 2-isopropylmalate synthase   |
| hokB     | Toxin HokB; Toxic component of a type I toxin-antitoxin (TA) system. |
| acnA     | Aconitate hydratase A  |
| ubiJ     | Ubiquinone biosynthesis protein UbiJ                                 |
| lptD     | LPS-assembly protein LptD  |
| rpnC     | Recombination-promoting nuclease RpnC                                |
| rpnA     | Recombination-promoting nuclease RpnA                                |
| fdoG     | Formate dehydrogenase-O major subunit                                |
| wbbH     | O-antigen polymerase   |
| wbbI     | Beta-1,6-galactofuranosyltransferase WbbI                            |
| wbbK     | Putative glycosyltransferase WbbK                                    |
| rpnE     | Inactive recombination-promoting nuclease-like protein RpnE          |
| lpoA     | Penicillin-binding protein activator LpoA                            |
| gspD     | Putative type II secretion system protein D                          |
| yfjI     | Uncharacterized protein YfjI; Phage or Prophage Related              |
| rlmL     | Ribosomal RNA large subunit methyltransferase K/L                    |
| rsxC     | Electron transport complex subunit RsxC                              |
| yfcI     | Recombination-promoting nuclease RpnB                                |
| gtrS     | Uncharacterized protein YfdI; Putative ligase                        |

As an example, Figure 3 shows the profile across all datasets of the gene *ompC* (EG10670). *OmpC*, also known as outer membrane (OM) protein C, is a porin of gram-negative bacteria tightly associated with the peptidoglycan layer. It has been recognized to have a crucial role in the non-specific diffusion of small solutes such as sugars, ions and amino acids across the outer membrane of the cell [39].

To highlight which specific regions within the Ribo-seq profiles are similar to each other, we built a *consensus sequence*. The consensus sequence is a character string representing the nucleotides of the reference ORF and, in Figure 4, it is colored red in those positions where a peak is present in at least 80% of profiles (i.e., the digitalized profiles values are +1 and the ribosome proceeds slower), and green where a valley is located. The black color, instead, will be used in all other cases and the label assigned to these regions will be 0.

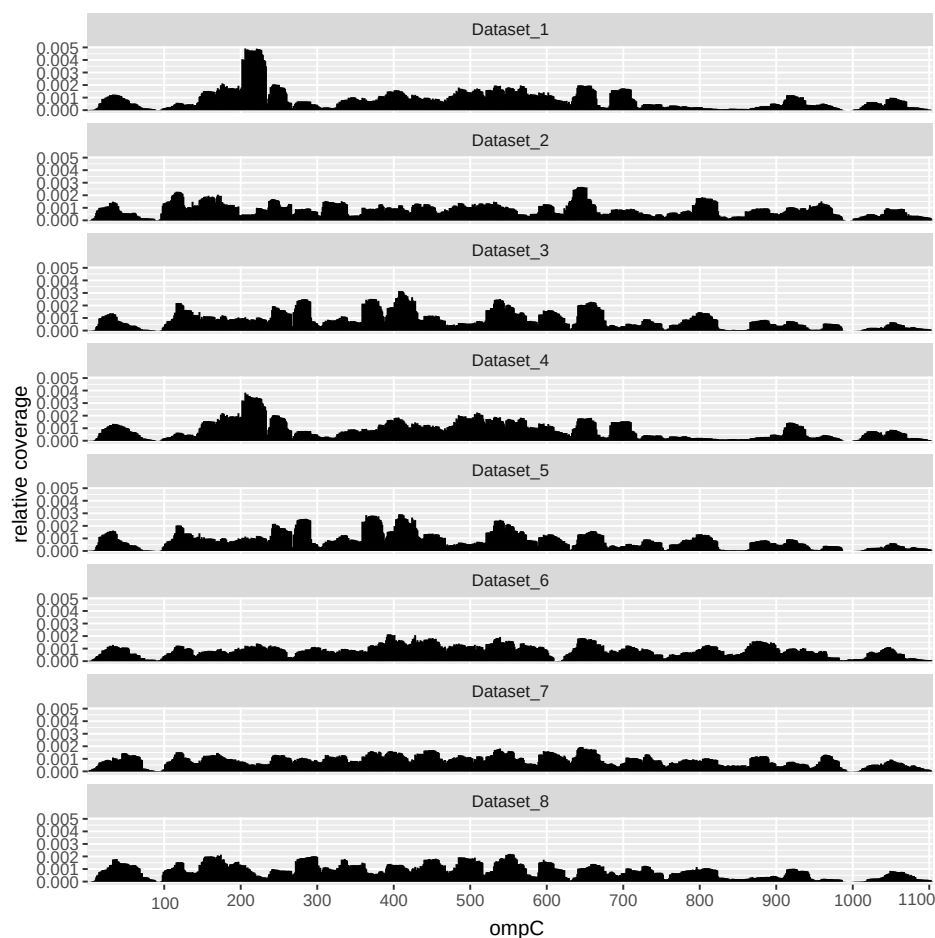


Figure 3. Example of a significantly reproducible Ribo-seq profile across the eight datasets (gene ompC, EG10670).

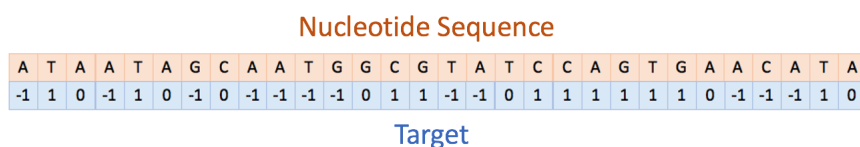


Figure 4. Part of a consensus sequence (computed for rsxC) indicating the nucleotides situated within fast (labelled with −1) and slow (labelled with +1) translation regions. On nucleotides labelled with 0, no reproducible results have been obtained [6].

### 3.2. Dataset

The presented analysis has been applied to the prokaryotic organism *E. coli* on 3588 Ribo-seq profiles across eight independent datasets, revealing that only 40 profiles are significantly reproducible. The digitalization process described above produces a target belonging to  $\{-1, +1, 0\}$  for each nucleotide of the sequence. Based on the obtained profiles, a dataset (“SubsequencesDataset”) has been constructed, consisting in mRNA reproducible sub-sequences with a uniform target (since they are reproducible, the target is not 0). In particular, 459 sub-sequences of variable length have been obtained, of which 264 are characterized by a slow translation (+1) and 195 by a fast translation (−1). In order to obtain a fixed-length vector of 36 elements, padding have been applied to the extracted regions, consisting of nucleotides of the original consensus sequence. Each nucleotide has been encoded with a 1-hot vector of 4 elements, representing one of the four possible nucleotides. The “SubsequencesDataset” is further splitted in a training set of 413 sub-sequences and a test set of 46 sub-sequences. Moreover, a validation set was built, randomly selecting 15% of the training set sub-sequences, to give an unbiased evaluation of the model performance



during training, and for keeping overfitting under control. The splitting was done ensuring that sub-sequences in the test set and in the training set are taken from different genes. Each sub-sequence is associated with a target which corresponds to its translation speed.

### 3.3. Statistical Analysis on the Nucleotide Composition of the Subsequences

Inferential statistics can provide insights on the specific patterns and characteristics of the data and highlight relationships between variables. The “SubsequencesDataset”, obtained with the procedure described in Section 3.2, is analyzed to explore the nucleotide composition of the mRNA sub-sequences, relating to the assigned labels (+1 and −1).

First, the relative frequency of each nucleotide (A, T, G, C) has been computed. To assess the significance of our results and to find out whether the obtained frequencies are typical of a certain speed or the correspondence is just obtained by chance, we built a statistical test. The aim of the test is to assign a probability value ( $p$ -value) to the following null hypothesis: the slow and fast sub-sequences are characterized by a random nucleotide composition. In order to test this hypothesis,  $10^4$  new profiles are generated by randomizing the nucleotide sequence, keeping fixed the +1 and −1 labels and therefore the length of the original sub-sequences. In each of the obtained sequence, the relative frequencies of the four nucleotides within the fast and slow sub-sequences can be calculated. This procedure leads to eight null distributions of relative frequencies, one for each nucleotide both for slow and fast translation. Comparing the null distributions against the original relative frequencies allows to calculate the  $p$ -values. In particular, if the original frequency is lower than the mean of the null distribution, we define the  $p$ -value as the probability of a random sub-sequence to show a relative frequency smaller than that observed in the original sub-sequence. On the contrary, if the original frequency is higher than the mean of the null distribution, the  $p$ -value is calculated as the probability of finding by chance a higher relative frequency value. If the  $p$ -value lies under the significance threshold of 0.05, the null hypothesis of completely random nucleotide frequencies is rejected. This indicates a statistically significant difference in the nucleotide composition between slow/fast and randomly generated sub-sequences.

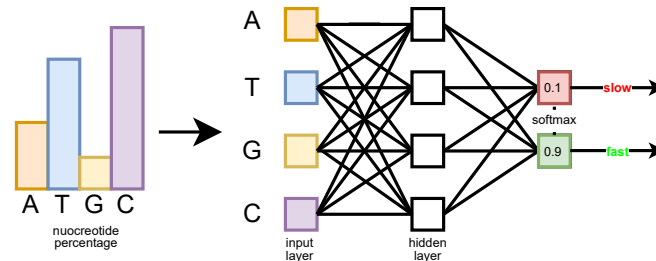
### 3.4. Data Validation with Neural Network Models

In this section, the informative content of the obtained data has been validated using a machine learning approach. This is achieved by employing common neural network architectures on the “SubsequencesDataset” (see Section 3.2) to predict the translation speed class: “slow” or “fast”. In our specific problem, exploiting a network architecture can reveal whether there is enough information in the data to classify the sub-sequences into slow and fast with high accuracy. To perform this task, we used two different types of data: vectors and sequences. In fact, we initially considered four-dimensional vectors that collect the relative frequencies of occurrence of the four nucleotides and then we considered the entire sequence, to evaluate whether the order in which the nucleotides are arranged helps to capture the translation rate signal. Consistently, the experiments were carried out by applying two different neural network architectures: *Multilayer Perceptrons* (MLPs) [40] and *Convolutional Neural Networks* (CNNs) [41]. While MLPs are good at processing vectors, indeed, CNNs are powerful machine learning models that can be used to directly process complex data, sequences in our case. Hence, we have first predicted the translation speed based only on the nucleotide composition of the sub-sequence while, in a second set of experiments, we employed a CNN to process each sub-sequence.

#### 3.4.1. MLP Analysis Based on the Nucleotide Frequencies

The MLP input is a four-dimensional vector whose elements correspond to the relative frequency value of each nucleotide in the sequence (in the order (A, T, G, C), see Figure 5). The goal is to determine how much information is carried by this simple statistic, regardless of the order in which the nucleotides appear in the sequence. The MLP has a single hidden layer with four neurons and an output layer with two neurons. A *Softmax* function is

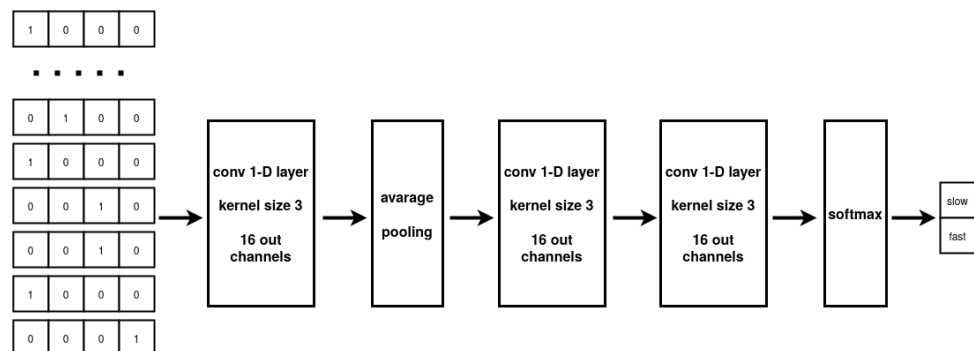
applied on the output layer, which produces the probability estimated by the network for each class (slow or fast) [42]. The network has been trained with the Adam optimizer, with an initial learning rate of 0.05.



**Figure 5.** The MLP architecture having the nucleotide frequencies as input, and predicting the sequence class probability distribution (slow or fast).

### 3.4.2. Convolutional Neural Network Analysis Based on Sub-Sequences

While the MLP architecture can only process vectorial data, CNNs can take full advantage of sequential information. The CNN multilayer design allows to extract a hierarchy of representations from the data while the implemented weight sharing guarantees to limit the number of parameters with respect to a fully connected architecture. In our experiments, the model input are the nucleotide sub-sequences collected in “SubsequencesDataset”. The CNN elaborates the input to produce a prediction about the translation speed associated with the sequence. Specifically, our CNN architecture comprises two successive 1-D convolutional layers, with average pooling employed between the layers, followed by two fully connected layers. The two-class probability distribution output of the network is produced by using a Softmax activation function on the output of the second fully-connected layer. The 1-D convolutional layers are based on 16 kernels with dimension three and a ReLU activation function, applied to each layer [43]. The optimization of the network parameters is performed by the Adam optimizer using early stopping [44]. The overall network architecture is described in Figure 6.



**Figure 6.** The 1-D CNN exploited for sequence classification.

### 3.4.3. Ensemble Convolutional Neural Networks

In the last experiment, an ensemble [45] of seven CNNs has been employed to produce a more stable and accurate prediction. Each CNN receives as input the nucleotide sub-sequences contained in the “SubsequencesDataset”. The seven CNNs have been trained independently. In the test phase, their output has been averaged to produce the final result. More specifically, the Softmax CNN output for each class (slow and fast) is averaged among the seven networks that compose the ensemble.

#### 4. Results and Discussion

In the following, we report the results of the experimental analysis described above. In particular, in Section 4.1, the outcome of the statistical analysis on the frequency of nucleotides is presented while, in Section 4.2, the results of the machine learning analysis is outlined.

##### 4.1. Statistical Analysis on the Nucleotide Frequencies

The first step of the statistical analysis consists in computing the relative frequencies of each nucleotide in the fast and slow sub-sequences, respectively. As it can be observed in Table 4, showing the nucleotide frequencies in the fast sub-sequences, adenine has a concentration which is significantly larger than those of the other nucleotides. In particular, adenine shows a relative frequency of approximately 0.32, while cytosine has the lowest frequency (0.2). Instead, Table 5 reports the frequency values in slow sub-sequences. In the latter case, both guanine and cytosine have the highest relative nucleotide frequencies (0.27 and 0.26, respectively).

**Table 4.** Relative nucleotide frequency across fast sub-sequences.

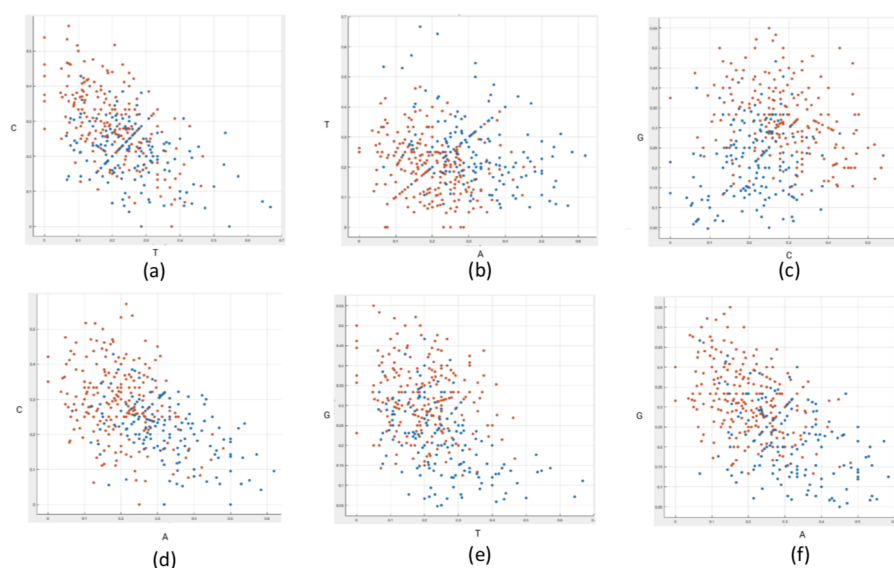
| Nucleotide | Frequency |
|------------|-----------|
| A          | 0.328     |
| T          | 0.257     |
| G          | 0.216     |
| C          | 0.209     |

**Table 5.** Relative nucleotide frequency across slow sub-sequences.

| Nucleotide | Frequency |
|------------|-----------|
| A          | 0.240     |
| T          | 0.228     |
| G          | 0.274     |
| C          | 0.258     |

To assess the significance of the results, the statistical test described in Section 3.3 has been performed. The purpose of the test is to determine whether the frequencies obtained are specific characteristics of fast and slow sub-sequences or they are simply a product of chance. Actually, the results show a  $p$ -value equal to zero for every nucleotide, for both fast and slow sub-sequences. In our context, the lower the  $p$ -value, the lower the probability that the nucleotide frequencies for the two classes occur by chance. The obtained results suggest that the relative frequencies of the four nucleotides are significantly different from those occurring randomly. In particular, we can observe that nucleotides A and T have a higher frequency in fast sequences than G and C. Instead, the frequency of nucleotides G and C is significantly higher in slow sequences than those of A and T. Based on this evidence, the proposed method for the identification of reproducible Ribo-Seq profiles was able to correctly detect sub-sequences characterized by a higher information content with respect to random sub-sequences, confirming the validity of our new approach.

Moreover, some insights are provided by Figure 7, which reports the distribution of the sub-sequences across the whole dataset, based on their nucleotide composition. In particular, each dot on the plane represents a sequence, characterized by the relative frequencies of pairs of nucleotides. The color represents the translation speed: orange and blue dots correspond to slow and fast sub-sequences, respectively. Interestingly, it can be observed that selecting the pairs of nucleotides A–T and G–C, the fast and slow sub-sequences identify two clusters located in distinct regions of the plane.



**Figure 7.** Representation of the sub-sequences on the plane based on the relative frequencies of two nucleotides. Each sub-sequence is represented by a dot on the plane. More specifically, the relative frequency of thymine and cytosine (a), adenine and thymine (b), cytosine and guanine (c), adenine and cytosine (d), thymine and guanine (e), and adenine and guanine (f), respectively, is shown. The colors represent the speed of translation: orange and blue dots indicate slow and fast sub-sequences, respectively.

## 4.2. Performance of the Neural Network Models

### 4.2.1. MLP Classification Based on Nucleotide Frequencies

In this experiment, a Multi-Layer Perceptron is used to predict the translation speed of sequences based only on their nucleotide composition. The model performance is summarized across five runs and reported in Table 6. It can be noted that the network reaches on average 82.24% accuracy over the test set, while the standard deviation between different runs is 2.20%. In fact, these results are surprising, given that the only information used by the model is the nucleotide composition of the sub-sequences. Moreover, the results are compared with those obtained by training the MLP on random sub-sequences, in order to demonstrate that the reproducible sub-sequences are significantly more informative. Indeed, as reported in Table 6, the performance obtained on reproducible sub-sequences is significantly higher than that obtained on random sub-sequences, validating the significance of our approach.

**Table 6.** Summary of the results obtained with the MLP model. We collect the obtained test set metrics computed over five different runs. The last two rows report the average over the runs and the corresponding standard deviation.

| MLP           |           |        |          |          |
|---------------|-----------|--------|----------|----------|
| Run           | Precision | Recall | F1-Score | Accuracy |
| 1             | 70.83     | 89.47  | 79.07    | 81.63    |
| 2             | 70.80     | 89.51  | 79.06    | 80.62    |
| 3             | 72.00     | 94.74  | 81.82    | 83.67    |
| 4             | 66.67     | 94.74  | 78.26    | 79.59    |
| 5             | 75.00     | 94.74  | 83.72    | 85.71    |
| Average       | 71.06     | 92.64  | 80.39    | 82.24    |
| Standard Dev. | 2.68      | 2.57   | 2.06     | 2.20     |

#### 4.2.2. CNN Classification Based on the Entire Sequence

The model performance is summarized across five runs in Table 7.

**Table 7.** Summary of the results obtained with the 1-D CNN model. We collect the test set metrics computed over five different runs. The last two rows report the average over the runs and the corresponding standard deviation.

| CNN           |           |        |          |          |
|---------------|-----------|--------|----------|----------|
| Run           | Precision | Recall | F1-Score | Accuracy |
| 1             | 96.00     | 90.00  | 93.00    | 91.84    |
| 2             | 96.00     | 87.00  | 91.00    | 89.80    |
| 3             | 93.00     | 90.00  | 92.00    | 89.80    |
| 4             | 100.00    | 77.00  | 87.00    | 85.71    |
| 5             | 93.00     | 90.00  | 92.00    | 89.80    |
| Average       | 95.60     | 90.00  | 91.00    | 89.39    |
| Standard Dev. | 2.88      | 5.20   | 2.35     | 2.24     |

The achieved classification accuracy is 89.39%. It is worth noting that, by training a significantly more complex network than the MLP, and by providing a sequential data input, the accuracy increases by approximately eight percentage points. The obtained results clearly show that this model can extract useful information from a limited amount of data in a better way than MLPs, achieving a very high accuracy. Nonetheless, the standard deviation between the runs is 2.24%, which proves that the results are rather steady but still influenced by the parameter initialization. Therefore, to further improve performance, we have set up a more complex architecture consisting of seven CNNs: each of them provides a different prediction, i.e., a pair of probabilities describing the class membership of a sequence (slow or fast). The results of the experiments performed by the CNN ensemble are summarised in Table 8.

**Table 8.** Summary of the results obtained with the CNN-ensemble model. We collect the obtained test set metrics computed over five different runs. The last two rows report the average over the runs and the corresponding standard deviation.

| ENSEMBLE: 7 CNN |           |        |          |          |
|-----------------|-----------|--------|----------|----------|
| Run             | Precision | Recall | F1-Score | Accuracy |
| 1               | 96.00     | 90.00  | 93.00    | 91.84    |
| 2               | 96.00     | 87.00  | 91.00    | 89.80    |
| 3               | 96.00     | 90.00  | 93.00    | 91.84    |
| 4               | 96.00     | 90.00  | 93.00    | 91.84    |
| 5               | 93.00     | 90.00  | 92.00    | 89.80    |
| Average         | 95.40     | 90.00  | 92.40    | 91.02    |
| Standard Dev.   | 1.34      | 1.22   | 0.89     | 1.12     |

Our model reaches an accuracy of 91.02%, with a standard deviation of 1.12%. As expected, the accuracy of the ensemble CNN is improved while the variance is significantly decreased. Indeed, the ensemble model is very effective in the sub-sequence classification and also provides very stable results across different training runs.

## 5. Conclusions

In this paper, we have proposed an innovative method to characterize highly reproducible Ribo-seq profiles. Ribo-seq data have been analysed through statistical analyses and state-of-the-art machine learning models, extremely effective in predicting the ribosome translation speed. In fact, using neural network architectures capable of processing both plain and sequential data, we have been able to obtain high accuracy, also proving that fundamental information is contained both in the nucleotide composition of the sequences and in the order in which nucleotides appear within each sequence. In this way, our work opens new exciting frontiers in the analysis of the ribosome translation dynamics in different organisms. Indeed, we have conducted a preliminary analysis of Ribo-seq profiles referring to liver tumours and their adjacent noncancerous normal liver tissues from ten patients with hepatocellular carcinoma (HCC) [46], achieving promising results. For what concerns the machine learning analysis, once we have obtained the consensus sequences, we carried on a preliminary study exploiting the same neural architectures employed on the *E. coli* datasets. Nonetheless, given the increased complexity of the human data, we believe that a further analysis is necessary, in particular defining ad-hoc neural architectures and with a specialized hyperparameter search, to improve performance. Finally, it is worth mentioning that our method represents an effective approach for any kind of Ribo-seq data, to investigate an extremely relevant open questions in biology, i.e., which features can influence the speed of the ribosome during translation.

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