SURVEY AND SUMMARY

Predictive biophysical modeling and understanding of the dynamics of mRNA translation and its evolution

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Received June 13, 2016; Revised August 07, 2016; Accepted August 19, 2016

ABSTRACT

mRNA translation is the fundamental process of decoding the information encoded in mRNA molecules by the ribosome for the synthesis of proteins. The centrality of this process in various biomedical disciplines such as cell biology, evolution and biotechnology, encouraged the development of dozens of mathematical and computational models of translation in recent years. These models aimed at capturing various biophysical aspects of the process. The objective of this review is to survey these models, focusing on those based and/or validated on real large-scale genomic data. We consider aspects such as the complexity of the models, the biophysical aspects they regard and the predictions they may provide. Furthermore, we survey the central systems biology discoveries reported on their basis. This review demonstrates the fundamental advantages of employing computational biophysical translation models in general, and discusses the relative advantages of the different approaches and the challenges in the field.

INTRODUCTION

Proteins are principal actors in all intra-cellular activities. The protein coding potential inscribed in a species' DNA is converted into proteins through the fundamental cellular process of gene expression, the two major cellular biophysical stages of which are the transcription of the gene to mRNA molecules by the RNA polymerase, and the translation of the mRNA molecules to proteins by the ribosome.

Surprisingly, generally there is only a limited degree of correlation between the levels of mRNAs and their encoded proteins (1–4), emphasizing the significant effect of post-transcriptional regulation on protein levels.

Furthermore, it was shown that translation and its efficiency have substantial effect on the organismal fitness, as it is the intracellular process with the highest energy consumption (5–7). Notably mRNA translation is a central intracellular process which can affect, be related, or interleaved with all central biological phenomena including protein folding, mRNA degradation, ribosomal collisions, abortion and allocation, transcript evolution, tumorigenesis, amongst others (7–23).

In recent years, various large scale techniques for estimating variables related to mRNA translation have been developed (1,3–4,24–41). Thus, in the recent and following years computational models of translation on the basis of high-throughput, genome-wide datasets, are expected to be instrumental in deciphering this process (1–2,7,12,19,42–70). Specifically, a central challenge in the field is to infer computational/mathematical biophysical models that consider various aspects of translation based on large scale data.

Indeed in recent years, dozens of studies aimed at modelling various aspects of the translation process. These models differ (and sometimes overlap) in their resolution, complexity, running times, assumptions, and more. Additionally, some of these models have been used for uncovering various fundamental biological phenomena.

The objective of this review is to report the advances in the field of computational large scale biophysical modelling of translation which are based on biological data. We first briefly review large scale biological experiments that can be used for inferring such models, and define what a biophysical model is. The main aim is to cover aspects such as the complexity of the models, their running time, the biological phenomena they uncover, and open challenges in the field. We believe that this is the first comprehensive review on this topic; thus, it is expected to be helpful to researchers studying translation via a quantitative manner.

It is important to mention that while there are some thorough previous reviews related to the field or to per-
tinent topics (see, for example, (71,72)), here we focus on various aspects that haven’t been comprehensively covered/examined before:

First, we review only models that consider biophysical aspects of the process. Second, we focus on studies that connect the model(s) to real large scale biological measurements and/or provide predictions related to translation of real genes. Third, we present some specific examples of the models’ parameter estimation. Fourth, we consider/discuss the complexity of the models and their running times (when possible), and compare the different models based on these measures. Finally, we review systems biology results suggested based on these models.

We believe that the review will be helpful for a very diverse audience. For example, computational/synthetic biologists reading the review will mainly learn about the open questions in the field, the advantages related to each model, and the data that can be used with these models. Systems biologists/genomics researchers will mainly learn about the type of questions that can be answered via these models. Experimentalists (e.g. people studying RNA, gene regulation, and molecular biology) are expected to gain some basic knowledge regarding TASEP based models, and what they can afford them.

Large scale experimental data related to translation

Data related to translation is a crucial factor in accelerating the development of predictive modeling of this process, and for improving its understanding. Thus, we begin with a brief description of the experimental technologies in the field.

Specifically, these data are used for inferring various free parameters of the models and evaluating their performances. Today, it is rather easy to perform large scale measurements of mRNA levels (73) while, for technical reasons, the technologies for performing large scale measurements of protein abundance lag behind. For example, the GEO database includes hundreds of thousands of large scale measurements of mRNA levels, whilst there are only a few such large scale measurements of protein abundance (1,3,29–30) (there are at least two orders of magnitude less measurements of post-transcriptional gene expression stages/aspects, for example, protein levels, translation rates, degradation rates of proteins and mRNA molecules, etc). Thus, researchers are frequently ‘forced’ to use mRNA levels, the rather rough proxy of protein abundance, instead of the protein abundance itself. Concurrently, technologies to measure various aspects related to translation are now rapidly emerging, transforming our understanding of this process (1,3,24–40). Among others, these include proteomic techniques to measure absolute protein levels (1,3,29–30,40,74), tRNA levels (7,26,39,75–76), mRNA folding (33), elongation dynamics (27), and fractionation of mRNA-ribosome complexes (polysomes) to estimate not only the number of ribosomes on each mRNA (24,77–78), but even the positions of individual ribosomes on each mRNA (25,54,79–91) (using an experimental technique called Ribo-Seq), translation initiation positions (92,93), mRNA degradation rates (36,37), protein degradation rates (94,95), and co-translational protein folding (96,97).

Most of the large scale measurements related to mRNA translation are based on endogenous genes.

The main disadvantage of analyzing endogenous genes is the fact that these genes are shaped by evolution, thus, hindering the ability to understand the causality of the observed correlation between various variables related to translation efficiency (40,98–100). For example, ribosomal proteins have certain amino acid contents which relate to their function, and are known to be highly expressed. As each amino acid is encoded by specific codons they induce/affacted various properties of the transcript (e.g. codon usage bias, GC content, mRNA folding). Thus, by analyzing such endogenous genes we may wrongly deduce that these transcript features affect translation efficiency. Furthermore as the different determinants of translation efficiency are correlated (e.g. highly expressed genes have both more efficient codons and more efficient initiation signals) and are under selection, it is difficult to evaluate the contribution of each of them to the translation speed (100).

An additional experimental strategy for studying the variables affecting protein levels and their regulation is via synthetic biology and heterologous gene expression. This approach includes manipulating the nucleotides of a transcript or parts of it, and/or measuring them out of their original endogenous setting, to understand the way translation aspects are encoded in the transcript. Recently, new datasets of protein abundance measurements of heterologous genes have emerged (40,99,101–104). Each of these libraries includes many versions of proteins; each such version includes an amino acid sequence identical to the original gene but with a different set of codons (i.e. synonymous modifications). Thus, by analyzing such datasets it is possible to overcome the two aforementioned problems (40,46,102). The heterologous genes datasets, on the other hand, may have two major problems of their own. First, in order to derive general conclusions about translation, many such datasets of heterologous genes based on numerous amino acid sequences should be generated (significantly more than those existing today). Second, in many cases heterologous genes are expressed in extreme and unnatural regimens, complicating the generalization of their analyses results to endogenous genes (6). Thus, these datasets instigate cautious analysis as well.

Together, these advances provide an enhanced account of basic cellular processes, with extensive implications for our understanding of the regulatory signals encoded in the transcript (e.g. the nature of codon biases), biophysical properties of transcripts, dynamics of translation, and transcript evolution.

However, it is important to mention that today there are no direct large scale in vivo measurements related to many fundamental aspects of translation. For example, we cannot directly measure initiation rates, codon decoding rates, ribosome abortion rates, etc. Thus, one challenge in the field is the inference of such variables from the available data mentioned above. In addition, most of the relevant experimental datasets include (sometimes extreme) non trivial biases (see, for example, (105)); thus, another fundamental challenge, interleaved with the objective of developing biophysical translation models, is the accurate filtering/handling of these biases.
The biophysical vs. machine learning approaches

As aforementioned, we survey the computational biophysical approaches for modeling mRNA translation based on these data. At this stage we would like to briefly mention that there are two major approaches for modelling/predicting translation, the machine learning approach and the biophysical approach. The biophysical approach is usually based on predictive simulations that are inspired by biophysical understanding of the studied processes. These simulations include biophysical phenomena such as diffusion, interactions between particles, folding of macro-molecules, etc. The machine learning approach, on the other hand, is based on statistical predictive inference of relations between sequence features and gene expression aspects, and it does not necessarily require prior knowledge of the biophysical gene expression mechanisms.

Machine learning is the more ‘traditional’ approach for practically predicting large scale gene translation, protein levels, or related variables (1–3,45,102,106–107). These models weigh the value of all the different variables to generate a prediction of, for example, protein levels. The main advantage of this approach is that it is relatively simple, no prior knowledge related to the biophysics of translation is required, and no knowledge related to the direction/causality of the relations is needed. This approach can often yield very significant predictions and improve our understanding regarding the transcript features relevant to translation efficiency.

The biophysical approach on the other hand, which captures the dynamics and physical nature of the process, is frequently used in (‘simple’) simulations without analyzing and modeling large scale biological data (108–111). The major advantage of the biophysical approach is the fact that it enables a better understanding of the biophysics of translation and its different regulatory stages than the machine-learning approach. Though theoretical physical models and simulations related to translation have been suggested nearly five decades ago (109,112), only recently have such approaches been implemented on real large-scale genomic data. Translation or protein synthesis consists of a complex system of chemical reactions, which ultimately results in the decoding of the mRNA and the production of a protein. The complexity of this reaction system makes it difficult to quantitatively connect its input parameters (such as translation factors, ribosome concentrations, tRNA concentrations, mRNA codon composition or energy availability) to output parameters (such as protein synthesis rates or ribosome densities) (71).

The simplest comprehensive (whole-cell) biophysical model of translation should consider the following aspects:

1. A (possibly) different speed/time for each codon, which is related to the local biophysical properties of the mRNA (e.g. its folding and the encoded amino acids), and their interaction with trans factors and/or the availability of such trans factors (e.g. tRNA levels).
2. Initiation rates (which are affected by the properties of the mRNAs and initiation factors, and global factors such as the concentrations of ribosomes and translation factors).
3. The fact that more than one ribosome can translate an mRNA at a certain time point.
4. Excluded volume interactions between ribosomes and possible traffic-jams.
5. The stochastic nature of the process.
6. The movement has a directionality (in the case of translation it is totally asymmetric).
7. The fact that different mRNA molecules compete for the same pool of ribosomes (and other translation factors); i.e. limited resources.

All the points above are known to be fundamental aspects of the canonical translation mechanism in all domains of life (5). The list above was chosen based on the majority of the previous studies in the field (as reviewed in this manuscript).

Of course, there are various (some less canonical) aspects of translation that are not mentioned above (e.g. ribosome recycling) that appear in some of the reviewed computational models. Importantly there are fundamental translation aspects that have not been considered in biophysical translation models (e.g. re-initiation after translating a uORF, frame shifts, numerous alternative initiation start sites, to name a few).

Currently there is no computational predictive model that considers all the aspects above; however, as described below, there are models that consider many of these aspects.

In the following sections we review the models suggested in the field sorted according to their complexity (see also, Figures 1 and 2).

Models that are based on averaging estimations of codon decoding rates

The simplest biophysical models of translation assume a ‘speed’ (or decoding rate) for each codon and average these speeds over the codons of a gene. The estimated speed of each codon can be inferred based on various gene expression variables such as tRNA levels, ribosome densities, and/or mRNA levels. Thus, these models consider only aspect 1 (but in some cases, as described below, they indirectly also consider aspect 7).

As a typical example of these models we would like to describe in further detail the tRNA Adaptation Index (tAI, 48,113), see Figures 1A and 2). While there were previous models that aimed at estimating the translation rate of a gene based on averaging weights related to the optimality of each codon (114–116) and based on their frequencies; the tAI is the first that aimed at considering basic aspects related to the biophysics of translation.

The tAI assumes that the relative concentrations of the tRNA molecules that recognize a codon have a significant effect on the codon translation rate, and it gauges the availability of the different tRNA molecules for each codon along an mRNA. Though this is clearly a very rough approximation of the reality and there are many additional relevant factors, the tAI has been shown to have high performance levels, and is employed as a main feature in many of the more sophisticated predictors (2,46–47).

This measure is determined by combining the estimated thermodynamic properties of the codon-anticodon interac-
Figure 1. Illustration of the biophysical models reviewed and envisioned. (A) The simplest biophysical model, the tRNA adaptation index (tAI), which gauges the availability of the different tRNA molecules for each codon along an mRNA; the output of this model can be an average over all the codons (one number per gene) or a profile of the nominal codon adaptation along the mRNA (a vector per gene). (B) The simplest TASEP (totally asymmetric simple exclusion process) model, which considers a single mRNA and its length $H$, "infinite" ribosomal pool (i.e. no feedback between the initiation rate and the ribosomes on the mRNAs), ribosome length $\ell$, single initiation rate $\alpha$, single elongation rate $\gamma$, single termination rate $\beta$. (C) An intricate TASEP model including all the canonical aspects necessary towards a whole cell simulation: finite ribosomal pool, ribosome length $\ell$, genome-wide mRNAs, their levels $n_i$, lengths $\eta_i$, and secondary structures, transcript specific initiation rates $\alpha_i$, codons specific elongation rates $\gamma_{cj}$ (based on numerous factors such as tRNA concentrations), transcript specific termination rates $\beta_i$, transcript specific re-initiation rates $\tau_i$. Each of the three main stages of gene translation, namely initiation, elongation, and termination, can be broken down to the multifarious steps comprising them, see, for example, Zhang et al. (51).

The absolute adaptiveness, $W_i$, for each codon $i$ is defined as (48,113):

$$W_i = \sum_{j=1}^{n_i} (1 - S_{ij})tCGN_{ij}$$

From $W_i$ we obtain $w_i$, which is the relative adaptiveness value of codon $i$, by normalizing the $W_i$'s values (dividing
them by the maximal of all the 61 $W_i$). The tAI of a gene is the geometric mean of the $w_i$ of its codons.

Recently novel indexes/models aimed at improving the estimation of codon decoding rates using similar ideas but in a more accurate manner:

The normalized translation efficiency (nTE) index (68) considers both the ‘supply’ (the number of tRNA of each type) and ‘demand’ (i.e. the number of codons of each type in the transcriptome) of tRNA species. This is achieved via normalizing the $w_i$ values of the tAI by the frequency of the corresponding codons in the transcriptome.

Another recent index named the mean of the typical codon decoding rate (MTDR) (117), aimed at estimating the typical translation rate of each codon based on Riboseq (25) (after filtering experimental biases and considering aspects such as ribosomal traffic jams). By definition this index also aims at considering both the ‘supply’ of tRNA species and their demand, in addition to other aspects of the process, such as amino acid availability and Aminoacyl-tRNA synthesis levels and the possible diffusion rates of different tRNAs, not considered in the other indexes mentioned above.

Both the nTE and the MTDR partially consider aspect 7 in the previous section: competition on limited elongation resources.

It is important to mention that many predictors and models integrated the tAI (or similar measures) into more complicated multivariate models (1–2,47,118–119). Such models try to predict gene expression aspects based on many transcript features, where the tAI (or similar measures) is only one of the features, but often has been one of the most predictive variables (2,47,118–119).

Finally, one fundamental disadvantage of the models described above is the fact that they attribute a single speed to each decoding region (which is presumably the mean of all the codons’ decoding rates). However, in recent years a natural generalization to these models has been used for studying various biological phenomena (7,68): compute a vector (or a profile) of estimated nominal decoding rates (the decoding rates when there are no interactions between ribosomes and traffic jams) which includes a value for each codon instead of an average of all these values (see Figures 1C and 3A).

**Rudimentary totally asymmetric simple exclusion process (TASEP) models**

The idea that protein synthesis involves unidirectional ribosomal movement along an mRNA molecule, was proposed by Warner and Rich in 1962 (71,120). The statistical properties of ribosome movement were explored initially by Zimmerman and Simha (121,122), and then refined by MacDonald and Gibbs (108,123). In these works mRNAs were considered as lattices on which ribosomes move with specific hopping probabilities, the latter being functions of the intrinsic kinetics of elongation; the ratio of initiation, elongation and termination rates (which determine ribosome density on the messenger RNA); and the probability that progress of a ribosome is unimpeded by preceding ribosomes (which is itself a function of the ribosome density on the message) (71). The statistical approach as developed by MacDonald and Gibbs (108,123) continued to be developed theoretically and eventually became known as the ‘Totally Asymmetric Simple Exclusion Process’ or TASEP model for translation. The model is called TASEP since it assumes that the movement of the ribosomes is only in one direction, from the 5’ end to the 3’ end (‘Totally Asymmetric’), a particle (ribosome) can only hop to the next position (codon) in the lattice (‘Simple’), and only if it is empty; i.e. there are excluded volume interactions (‘Exclusion’).

Characteristics of early versions of the TASEP include assumptions of limitless ribosome-supply (which is related to a very high probability of ribosome attachment to the transcript, which affects the initiation rate), a single, uniform elongation rate-constant along the mRNA, and a coarse-grained description of the elongation process, which is simply regarded as a ‘hopping-probability’ (71,124–127) (Figures 1B and 2).

In the case of the rudimentary TASEP models, the assumption is that the initiation rate is constant and/or not coupled/related to the number of available ribosomes. In other models, which will be discussed in the next section, the initiation rate is affected by the number of available ribosomes via the fact that the number of ribosomes is finite (increasing the number of ribosomes on one mRNA should cause a decrease in the number of ribosomes on other mRNAs). Specifically, if the number of mRNAs and ribosomes on the mRNA) is very small/large (respectively) relatively to the total number of mRNAs/ribosomes in the cell the initi-
The rate of translation of a single codon in the cell is determined by following four subsequent processes: (i) attachment of the amino acid to its cognate tRNA by the corresponding aaRS; (ii) aa-tRNA forms a ternary complex (TC) with GTP-bound elongation factor (EF-Tu in prokaryotes and eEF1A in eukaryotes); (iii) TC diffuses to a ribosome requesting it and transfers the amino acid to the growing polypeptide chain, whereby the tRNA is released; and (iv) the GDP-bound elongation factor is regenerated into its GTP-bound state by EF-Ts (prokaryotes) or EF-1B/C (eukaryotes) (see Figure 1C). In addition they modelled spatial diffusion of the TC complexes around ribosomes translating repetitive sequences using a Brownian random-walker model.

Mitarai et al. (69) developed and analyzed a translation model that captures in addition to the basic TASEP features, also the fact that ribosomes recycle. Specifically they added a parameter corresponding to the fact that with a certain probability a ribosome finishing translating the mRNA can immediately start another round of translation (e.g. due to the cyclicization of the mRNA molecule). Among others, they used their model to investigate the ef-

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fect of sequences of synonymous codons on ribosome traffic, based on codon-dependent rates estimated from experiments. They quantify the ability of codon usage to regulate ribosome traffic by simulating the translation of several natural genes, and demonstrate that those with a high codon usage bias, tend to have the ability to reduce ribosome collisions, hence optimizing the cellular investment in the translation apparatus. Recently, it was also demonstrated how ribosomal recycling can be modelled and analyzed via an RFM based approach (147).

In (61), the authors developed and analyzed a TASEP based model where the nominal codon decoding rate of each codon is affected by three different components: (i) the adaptation of the codon to the tRNA pool; (ii) the strength of the local mRNA folding of the 40 nucleotides downstream of the codon; (iii) the net charge of the 31 amino acids inside the exit tunnel—as the exit tunnel is negatively charged a positive charge is expected to slow down the ribosome.

In summary, the most important advantages of the TASEP (rudimentary and complex) models are related to the fact that they theoretically can capture and provide high resolution predictions of all the important local aspects of translation dynamics (aspects 1–6 in the third section). In addition, the fact that the different mRNAs are not ‘connected’ can enable accelerating and parallelizing the simulations. However, the models cannot capture aspects related to the coupling between the translation components (e.g. tRNAs, mRNAs, ribosomes) (aspect 7 in the third section) such as the ‘competition’ between mRNAs for ribosomes. It is clear that more complex translation models are preferred if the estimated parameters in the models are accurate and the running time is feasible, but this is obviously not always the case.

Whole cell simulations of translation

Currently there are no models that consider all the fundamental mRNA translation aspects at a cellular level. However, there are previous studies that contributed towards developing such a model. The aim of this section is to review some of these studies, specifically we review models which consider the dynamics of many interacting mRNA molecules. While there are quite a few previous studies that aimed at analyzing many mRNAs in the cell in a single model (51–52,129,148–157), many of them are based on mathematical analyses and simulations that are based on artificial genes and/or a small number of genes. As explained below, even the more comprehensive models suggested recently do not consider all the fundamental relevant translation aspects.

Many studies have demonstrated/suggested the importance of considering the ‘competition’ of mRNAs on a finite pool of resources (149–158). Specifically, unlike many other models of translation (46,53,65), which treat each mRNA molecule in isolation and assume an inexhaustible supply of free ribosomes that initiate the message at a constant rate, those models consider simultaneously every tRNA, mRNA, and ribosome molecule in the cell, and so capture the indirect effects of one gene’s translation on another’s. In particular, if many ribosomes are engaged in translating the mRNAs of one gene, this reduces the pool of free ribosomes and tRNAs available to translate other genes (Figures 1C and 3B).

For example, Siwiak et al. (67) developed a TASEP based comprehensive and quantitative model of translation, enabling the estimation of translation parameters which are difficult to measure experimentally. They model the number of transcripts, ribosome density, protein abundance, tRNAs’ decoding specificities, average cell volume, mean codon translation time, total transcript translation time, total time required for translation initiation and elongation, translation initiation rate, average number of active ribosomes in a cell, ribosome size/coverage, mRNA half-lives (optionally).

Shah et al. (62) developed a continuous-time, discrete-state Markov model of translation in a yeast cell. The model keeps track of every ribosome and tRNA molecule; according to the model, at any time point each ribosome/tRNA molecule is either freely diffusing in the cell or bound to a specific mRNA molecule at a specific codon position. Their model includes many parameters and describes translation dynamics at a single-nucleotide resolution for the entire transcriptome. In combination with ribosomal profiling data (Ribo-Seq), they aimed at using their model to infer the initiation rates of all abundant yeast transcripts, and systematically explore how the codon usage, transcript abundance, and initiation rates of genes jointly determine protein yield and cellular growth rate. The rates of initiation and elongation are based on physical parameters that have been experimentally determined in yeast, including the cell volume, the abundances of ribosomes and tRNAs, and their diffusion constants.

Another whole cell model which includes translation modeling was suggested by Karr et al. (156). This model was based on the human pathogen *Mycoplasma genitalium* and aimed at including all of its molecular components and their interactions (specifically translation). It is important to mention, however, that the resolution of translation modelling was not high and most aspects mentioned above (e.g. ribosome interactions and traffic jams during elongation, different codon decoding rates, initiation modelling, etc.) were not included. Similarly, Roberts et al. developed a whole cell simulation based on *E. coli* (159); also in this case they assume that a single rate-limiting event affects translation, ignoring the fundamental higher resolution details related to the translation process.

We conclude that currently there are no models that are based on real data that encapsulate all fundamental mRNA translation aspects. For example, the following canonical aspects (mentioned in the previous section) are not considered in (62,67,156,159): the different conformational steps of the ribosome during one iteration of elongation, the different sub-steps of the initiation stage (S′ end recognition, scanning, start codon recognition), the unfolding of the mRNA and interactions with mRNA folding, interactions between the nascent peptide and the ribosome, re-cycling of elongation factors, ribosome frame-shift, ribosome drop-off, to name a few.
Parameter estimation for the biophysical models

In this section, we provide some specific examples of parameter usage and estimation in some of the biophysical models described. There are variables such as protein levels (1,3,29–30,40,74), mRNA levels (160), number of ribosomes on the mRNA (24,77–78), that can be directly measured; however, as described here, other parameters can be estimated only via developing sophisticated approaches, which require a good understanding of the biophysics of translation, molecular evolution, machine learning, and more.

One fundamental parameter in biophysical translation models is the nominal decoding rate of codons. Various methods for estimating this parameter have been suggested in recent years. One approach is based on the tAI or normalized versions of it (2,46–48,61,68,113). The tAI was described above. One of the most challenging issues when computing the tAI is the selection of a meaningful set of $S_{\alpha}$’s, which are the selective constraints on the efficiency of the codon–anticodon coupling. Since translation efficiency, also in terms of adaptation to the tRNA pool, is expected to be higher for highly expressed genes, it was suggested to find the set of $S_{\alpha}$ values that maximize the correlation between expression levels and tAI values (48). This was first achieved based on microarray data from yeast (48); later, measurements of protein levels were used to optimize these values (7). Recently, it was suggested that these values can be estimated per organism, even when there are no gene expression measurements, based on optimizing the correlation with codon usage bias (113). Another component of the tAI is the tRNA concentration; today there are very few measurements of tRNA levels (7,26,39,75–76); however, it was suggested that tRNA copy numbers correlate with tRNA levels (48), and in many studies they are used as a proxy for tRNA levels. Codon decoding time estimation may be improved via considering the demand vs. supply of codons (68).

Recently, it was suggested that codon decoding rates can be estimated based on ribosome profiling data (161,162). To this end, ribosome profiling (Ribo-Seq) data should be normalized to control for the different initiation rates and mRNA levels of genes; then, based on the histogram of the normalized Ribo-Seq values of each codon the typical decoding rate can be estimated. In (161) a novel filter was suggested that enabled estimating the typical codon decoding rate via filtering various problems/phenomena such as pauses/biases/ribosomal traffic jams related to the Ribo-Seq protocol and the translation process. A newer study (163) has suggested inferring the elongation rates and initiation rates of mRNAs by finding parameters that yield the best fitting of the state variables (profiles of local ribosome densities) of the TASEP to the corresponding Ribo-Seq measurements.

An additional fundamental parameter which cannot be inferred in a trivial manner is the translation initiation rate. Translation initiation estimation is not trivial since it is based on many aspects such as the ribosome’s attempt to bind the mRNA, the concentration of initiation factors and the presence of secondary structures in the 5′ UTR region, the nucleotide composition surrounding the start codon, the number of START codons in the 5′ UTR and their nucleotide context, and more (5,54–55,92–93,98,137,164). An approach for estimating translation initiation was suggested in (54); they estimate mRNA-specific in vivo translation initiation rates on a genome-wide basis by integrating a computational model of mRNA translation with experimental datasets of ribosome occupancy. They first apply their TASEP based translation model which predicts the translation rate $J$ and the ribosome density $\rho$ on each mRNA as a function of the translation initiation rate $\alpha$. Then, by utilising genome-wide experimental data of ribosome density $\rho_{\psi}$ they identify the physiological translation initiation rate $\alpha_{\psi}$ as the one which, when used in their simulations, replicates the experimentally observed density:

$$\rho(\alpha_{\psi}) = \rho_{\psi}$$

This yields a translation initiation rate value $\alpha_{\psi}$ for each mRNA.

Recently, it was suggested that the ribosome drop off rate can be predicted based on the analysis of the decrease in ribosome density along coding regions via ribosome profiling data (165). These data can be used in the future when ribosome drop off rates are integrated into translation models.

Finally, a good source that includes many parameters related to translation (e.g. ribosomes’ size, initiation rates, codon decoding rates, diffusion coefficients, etc) based on measurements or estimations is the biological numbers database BioNumbers (166). Parameters from this database can be used in future development of translation models.

It is important to emphasize that some of the translation parameters (e.g. ribosome footprint size) are probably universal or very similar for various organisms; other parameters (e.g. initiation rates and ribosome drop off) can be estimated for various organisms when the related data (e.g. ribosome profiling) is available. However, there are parameters that currently are not available at a genome wide level for multiple organisms (e.g. ribosome recycling rates and non-programmed frame shift rates).

Comparison of different models in terms of running time and complexity

In the previous sections, we described various representative biophysical models of translation. In the Table 1 below we include a comparison of some of the recently suggested models. We focused on the following aspects: scope of the biophysical modelling, computational complexity (expected running time), number of parameters in the model, and properties of the model, etc.

Systems biology studies based on computational biophysical translation models

In this section we briefly review some central biological results and discoveries derived from the translation biophysical models. A central objective is to demonstrate how biophysical models of translation can contribute to biomedical science; specifically, as we explain, they enable understanding fundamental topics related to genome evolution, translation biophysical dynamics, and regulatory processes related to translation (see also Table 2 and Figure 3).
Table 1. The table below summarizes and compares the biophysical models discussed, classifying them according to the features they encapsulate.

| Short description of the model | Reference | Relative running time | Parameters fit to real data | Additional info and data | Deterministic/ stochastic |
|--------------------------------|-----------|-----------------------|----------------------------|--------------------------|---------------------------|
| The tRNA adaptation index (tAI) | Dos Reis et al. (48) | 1 | 64 codon-anticodon interaction efficiencies; given the tRNA levels only 5 parameters should be estimated. | The parameters can be inferred by optimizing the correlation between the tAI and gene expression measurements. | Deterministic |
| A measure based on slow codon clusters | Zhang et al. (66) | 1 | tRNA concentration | The model was applied to E. coli and B. Subtilis. tRNA codon pairing ratios were inferred from experiments. | Deterministic |
| The Ribosome Flow Model (RFM), a mean field approximation of the TASEPC | Reuveni et al. (46) | 2 | 61 nominal codon decoding rates, initiation rate(s), codon chunk size. | For example, the parameters can be inferred based on Ribo-Seq or the tAI values. A global initiation rate can be used. | Deterministic |
| Deterministic TASEP | Zhang et al. (128) | 2 | Ribosome size, initiation rate, termination rate, codon decoding rates. Simple TASEP parameters only the termination rate equals the elongation rate; two degradation rates: one proportional to the fraction of the mRNA covered by the ribosomes and the other to the unshielded. | The model was applied to E. coli focusing on the lacZ mRNA. It simulates an in-vivo experiment. | Deterministic |
| Simple TASEP with mRNA decay | Deneke et al. (70) | 3 | Simple TASEP parameters only the termination rate equals the elongation rate; two degradation rates: one proportional to the fraction of the mRNA covered by the ribosomes and the other to the unshielded. Simple TASEP parameters. 2 sets of codon rates: uniform, and one of three values for each codon specific translation rate, average mRNA translation number. | The model was applied to E. coli and presents improved fitting of published decay assay data. | Stochastic |
| Simple TASEP with codon specific rates (TASEPC) | Mitarai et al. (53) | 3 | TASEP parameters, the codon dependent rates are composed of: tRNA recognition rate and codon translocation rate (codon independent), the initiation rates are predicted using experimental ribosomal densities. | This model was applied to S. cerevisiae, integrating genome-wide experimental data of ribosomal density. tRNA-capture rates can be estimated from data on tRNA abundances, which are assumed to be proportional to their gene copy numbers. | Stochastic |
| Simulating translation as a set of chemical equations via a version of the Gillespie SSA algorithm (208) | Chu et al. (140,208) | 3 | 61 codon frequencies, 61 tRNA levels, 20 tRNA re-charging parameters, forward/backwards rate of cognate tRNA binding to the ribosome and of near-cognate tRNA binding to the ribosome. | The model was parameterized for S. cerevisiae based on experiments. | Deterministic |
| A multi step model of translational elongation consisting of deterministic and stochastic parts based on the concentrations of ternary complexes and competing mRNAs on a finite ribosomal pool | Zhang et al. (51) | 3 | Experimentally quantified tRNA concentrations, fraction of charged tRNA, enzymatic parameters of tRNA acylation catalyzed by the aaRS, rate constant of the complex formation between the charged tRNA with the EF, overall rate constant of EF-GTP regeneration, number of ribosomes. | The model was applied to E. coli. The authors experimentally quantified tRNA concentrations, and polysomal profiles. | Deterministic and Stochastic |
| RFM network with a pool (RFMNP) | Raveh et al. (129) | 3 | RFM parameters, number of ribosomes, mRNAs levels. | Can be easily parallelized. This model was applied to S. cerevisiae here. | Deterministic |
| TASEPC with finite resources | Brackley et al. (149) | 4 | TASEP parameters, number of available tRNAs, tRNA recharging rate constants, mRNA levels. | This model was applied to some of the mRNAs in S. cerevisiae. | Stochastic |
| TASEPC with finite resources | Chi et al. (150,158) | 4 | TASEPC parameters, mRNAs levels, number of ribosomes, ribosome recycling rounds, rate of recharging tRNA, number of tRNAs. | This model was applied to S. cerevisiae. | Stochastic |
| TASEPC with finite resources | Siwiak et al. (67) | 4 | mRNAs decoding specificities, average cell volume, average number of active ribosomes in a cell, ribosome size/coverage. Each gene is attributed with a set of translational parameters, namely the absolute number of transcripts, ribosome density, mean codon translation time, total transcript translation time, total time required for translation initiation and elongation, translation initiation rate, mean mRNA lifetime (optionally), and absolute number of proteins produced by gene transcripts. | This model was applied to S. cerevisiae. | Stochastic |
| TASEPC with finite resources | Shah et al. (62) | 4 | Many parameters including, number of ribosomes, number of mRNAs, number of tRNAs, tRNA competition coefficient, diffusion coefficient of ribosome, gene specific initiation probability. | This model was applied to S. cerevisiae. | Stochastic |
Among others based on the tAI (48) it was suggested that genome size and tRNA gene redundancy determine the action of natural selection on codon usage in all living organisms (Figure 3H). Their findings suggest that an optimal combination of these factors exists, for which the action of translational selection is maximal. Moreover, they stress how the lack of duplicated tRNA genes might explain the absence of translationally selected codons in bacteria with small genomes.

A different study employed the tAI to suggest that cell cycle regulated genes tend to include codons that are not translationally efficient to generate cell cycle-dependent oscillations in protein levels (167). They suggest that due to oscillations of tRNA species during the cell cycle, these codons enable oscillations of the protein levels of these genes.

More recent studies based on genomic profiles (or profiles of single genes or gene groups) of measures of nominal translation elongation speed, such as the tAI, enabled discovering various translation signals that are selected for and are related to translation efficiency, ribosomal and other factors allocations, and co-translational protein folding. For example, it was suggested that the first 30–50 codons are slower to improve ribosomal allocation and protein folding (6–7,12,98,168) (Figure 3A), and that evolution tends to cluster codons that can be decoded by the same tRNA to improve tRNA recycling via improving their effective diffusion to the ribosome (6,12,42). Based on various methods for generating profiles of the normalized translation efficiency index, it was recently proposed that the folding of the protein is partially effected by the local translation elongation speed (66,68) (Figure 3D). For example, it was suggested that translation elongation is lower downstream of the protein domain boundaries probably to improve the fidelity of co-translational domain-wise protein folding (57,66). Similar ideas related to elongation and co-translational folding have been suggested based on additional techniques including the tAI and TASEP based models (98,169).

Numerous studies have employed various TASEP models to understand how evolution shapes the distribution of codons to improve ribosomal allocation and prevent ribosomal collisions. For example, it was suggested that clusters of rare codons near the 5′ of the coding region decreases the effective initiation rate, the number of ribosomes on the mRNA, and the probability of ribosomal collisions and abortions (7,128). Similarly, it was proposed that higher codon usage bias (or adaptation to the tRNA pool) helps prevent traffic jams and improves ribosome allocation (69,170), and that codon arrangement, rather than simply codon bias, has a key role in determining translational efficiency (54). Furthermore it was shown that codon bias may act as a gene expression regulator by favouring codons with high tRNA gene copy numbers in highly expressed genes, and with low tRNA gene copy numbers in lowly expressed genes (46,171).

Another line of research connects biophysical models of translation and coding sequence evolution (44,65,170). Among others, based on such models it was suggested that a combination of nonsense errors and codon usage bias can have a large effect on the probability that a ribosome will complete transcript translation. In addition, they showed position dependent selection on codons, and suggested that nonsense errors can play an important role in shaping codon usage bias and can be used for predicting protein levels.

Moreover, Ciandrini et al. (54) used TASEP to identify the initiation rate, ribosome traffic dynamics and response to ribosome availability, of different classes of mRNAs, demonstrating that this classification based on translational dynamic maps onto known classes of gene functionality (Figure 3B and G).

An important question studied with TASEP models is the relative rate limiting step of translation, these studies have demonstrated that depending on the condition or organism both initiation and elongation may be rate limiting (7,54,60,62). Specifically, nowadays there is much emphasis on initiation being the rate limiting step and the principle determinant of translation ‘efficiency’. However, the behavior of TASEP is a function of its three main parameters/stages (initiation, elongation, termination); two TASEP parameters/stages (initiation, elongation) are relevant when the termination is not rate limiting or is negligible (as usually assumed). A related topic is the phase diagram of the TASEP as a function of the initiation and typical/mean elongation rate (see, for example, (172)). It is common to divide the dynamics of the TASEP to three phases (see, Figure 3F): (i) Low density phase—which
is related to the case where the initiation is rate limiting (i.e., much lower than the minimal elongation rate); thus, the number of ribosomes on the mRNA and traffic jams is very low/negligible. (ii) High density phase—which is related to the case where the initiation is high (higher than the typical/mean elongation); thus, elongation is rate limiting and the number of ribosomes on the mRNA and traffic jams is high. (3) The third phase is maximal current—in this case, given the elongation rate, the initiation rate has similar levels which maximizes the translation rate. It is important to remember that ribosomes are extended objects that cover around 10 codons each; this contributes to the shrinking (relatively to the case where the TASEP particles occupy only one codon/cell) of the parameter space of the ‘low density’ phase, so that they render the cases where the initiation is ‘rate limiting’ less likely (111).

Furthermore, when considering only the mean measured initiation rate and elongation rates and a model of a single mRNA the initiation is clearly more rate limiting (123) (Figure 3G). However, more complicated models demonstrate that this is not the case (52, 54, 60, 71, 109). For example, recent experimental results show that eukaryotic ribosomes may translate mRNAs in multiple cycles before entering the free ribosome pool (71, 173). This would greatly affect the translation control, and compared with single-cycle models, it could transfer significant levels of control to the elongation stage. Another example is related to the fact that mRNAs actually ‘compete’ over a finite ribosome pool; thus, initiation and elongation are coupled: higher elongation rate is related to faster release of ribosomes and higher effective initiation rate (129, 149–150, 153–157).

At what ribosome depletion level translation control is transferred to elongation depends in complex ways on the codon composition of the genome. The average speed of translation is not only a function of the number of slow codons in a message, but also of their distribution (46, 50, 54, 61, 66, 69, 71, 131, 174–177).

Another topic (which overlaps with the previously discussed) studied by a variety of models ranging from the simpler to the more complex, is the demand vs. supply of translation regimes in various genes, and the explanations for these differences, mRNA levels, and mRNA half lives. These include the study of the distributions of variables such as initiation rates and ribosomal densities and collisions, that cannot be easily measured, but can be estimated based on biophysical models and gene expression measurements (53, 62, 67, 70).

Another interesting line of research that can only be studied via biophysical models of translation is the relation between local mRNA folding dynamics and translation dynamics, although to date very few papers tackled this issue or introduced this aspect into biophysical translation modeling. It was suggested, for example, that strong local folding slows down the elongation rate—the ribosome has helicase activity and more time is needed to unfold regions with stronger local mRNA folding (61). Another study has recently also focused on the effect of the ribosome on the mRNA structure via its unfolding of the mRNA during translation (139) (Figure 3C). In (61) a TASEP model was devised where the nominal elongation rate of a codon is proportional to e\(^{kFE}\) where FE is the (negative) local folding energy of the 40nt downstream of the translated codon, and \(k\) is a parameter that was inferred based on fitting the model (ribosome density profile) to ribosome profiling data. In (139) they assume that the dwell time of a codon is proportional to the predicted mean base pairing probability of the codon’s three nucleotides. In addition, various studies connected the mRNA folding at the beginning of the ORF to translation initiation efficiency—specifically weak folding surrounding the start codon enables efficient recognition of the start codon and the initiation signals surrounding it by the pre-initiation complex, while strong folding may contribute to the pre-initiation complex missing the start codon (Figure 3E) (40, 98–99, 178). A thermodynamic model which calculates the initiation rate in prokaryotes based on the difference in Gibbs free energy before and after the 30S complex assembles onto an mRNA transcript was suggested by (179). In this model, various free energy terms are calculated and integrated, including: (i) The energy released when the last nine nucleotides of the 16S rRNA co-folds and hybridizes with the mRNA subsequence at the 16S rRNA-binding site. (ii) The energy released when the tRNA\(^{Met}\)\(^{5}\) anticodon hybridizes to the start codon. (iii) The folding energy of the mRNA subsequence surrounding the start codon prior to binding with the 30S complex. Finally, the effect of folding on translation, mentioned above, may suggest that the local mRNA folding landscape has shaped transcript evolution in various ways (98, 180–182). Thus, local mRNA folding is an important ‘intersection’ that connects the sequence composition of the transcript with the dynamics of translation initiation/elongation and the molecular spatial/temporal evolutionary patterns of the transcript.

Moreover, many studies based on the biophysical translation models aimed at predicting various gene expression variables such as mRNA levels, protein levels, protein degradation rates, and ribosome densities based only on the genomic sequences (46–47, 51, 54, 60, 66, 171).

**DISCUSSION AND CONCLUSIONS**

We reviewed dozens of biophysical computational/mathematical models related to ribosomal movement. We offer six major points for comparison among these models/papers:

1. **Number of parameters needed to be estimated**: on the one hand, models based on larger numbers of variables describe fine-tuned aspects of translation and capture larger fractions of the biological knowledge related to translation; on the other hand often there is no knowledge regarding the values of many of the parameters of the model, and/or estimation of these parameters is impossible or extremely noisy. Thus, subsequently these models cannot be used for providing predictions based on or related to real biological data. Furthermore, the major disadvantage when developing a model with many
parameters is overfitting (183), or accumulation of error due to individual errors in the different parameters.

2. Computational running times of model simulations: the running time of some of the measures/models is very short (48), while in others (111) the running time can be several orders of magnitude longer (as compared in (46)). The running time becomes an important issue when the objective is to analyze the entire genomes of complex eukaryotes with various/multifarious sets of variables.

3. The biophysical aspects modeled: which aspects of translation are modeled and which are not. The simplest models include mainly only the effect of tRNA levels on ribosomal codon decoding time, while the newer more complex models include many additional aspects. For example, notable extensions that were introduced to the TASEP include multiple competing mRNAs species (62,67,71,129,156), up to complete, genome-wide transcriptomes (51–52,71); mRNA transcription and decay (70,184); ribosome-induced peptidyl-tRNA hydrolysis in response to translational errors (‘ribosome editing’, (185)); aminoacyl-tRNA synthesis (19,46,51,54,62,66,69,131,134,140–141), ribosomal slow-down at mRNA secondary structures (57,66,138); competition for a finite pool of ribosomes (151–152,186); and the use of rate constants and species concentrations rather than rates (71,109), to mention a few.

4. Exact mathematical solutions versus simulations: The possibility of mathematically analyzing a model is a great advantage. Currently, there is no exact solution for a simple TASEP with non-homogenous hopping rates; thus, there is no solution for more comprehensive models based on the TASEP. However, in some cases a mathematical analysis is possible via various approximations, and answering questions in an indirect manner such as mean field approximations (see, for example, (46,72,167,168–190)).

5. Discrete models vs. an approximation of the models as continuous: The original TASEP based models are discrete; the number of ribosomes on the mRNA is an integer. However, there are various approximations for describing translation which are continuous (e.g. the RFM (46,191)). In these cases, the ribosomal densities are defined as real numbers (for example, it can be a probability to see a ribosome in a certain region). Continuous models are often easier to analyze mathematically (187), and are thus more advantageous in that sense, but are a cruder approximation of the reality.

6. What kind of predictions do the models provide in comparison to real data: connecting the model to real measurements is important. Specifically, the possibility to show that the model prediction correlates/matches with biological measurements should help evaluating the model, and predictive models have various advantages in biomedical research. Some of the models/papers mentioned here were not based on a comprehensive analysis of large scale biological data and validation of the predictions they provide, while others are more strongly connected to real data.

The points above demonstrate that it is not trivial to rank the different models mentioned in this review. Almost all the models have relative advantages in some cases. Thus, the user of such models should carefully assess his specific research aims before choosing a model.

We would like to emphasize that this review was focused on translation (and specifically translation elongation), since this is by far the most studied gene expression aspect in this context. We believe that in the near future models similar to the ones mentioned here can be used to study and focus on additional gene expression steps such as transcription (192), intracellular transport (193), and translation initiation (63,194–195).

There are many challenges and open questions in the field:

First, in many cases some of the predictions (e.g. correlation with protein levels) provided by biophysically more comprehensive models of mRNA translation are similar (or even inferior) to the ones obtained for much simpler models (1,118). However, there are many aspects that can be provided only by comprehensive biophysical translation models; for example, these include ribosomal traffic jams, interactions between ribosomes, ribosomal drop-off rate, stochasticity in translation rate, and more. One major challenge in this context is related to the fact that there are currently no measurements related to these variables, thus, they cannot be accurately evaluated.
Figure 3. Illustration of some central and fundamental biological questions that are studied with predictive biophysical translation modeling. (A) Ramp and ribosome allocation: the ramp is a region characterized by slower elongation speed and codons less adapted to the tRNA pool, and may provide several physiological benefits, such as increasing the distances between ribosomes, promoting improved ribosomal allocation, and reducing ribosomal collisions and jamming, thus reducing the cost of wasted ribosomes and of spontaneous or collision-induced abortions (7,211). Red indicates a slow codon decoding rate (as illustrated in H). (B) Competition between ribosomes: since the amount of ribosomes in a cell is finite, the ribosome consumption of one mRNA will affect all others. For example, mRNAs consuming many ribosomes due to their properties (such as strong folding, or relatively slow codons), prevent them from re-entering the pool, thus reducing the initiation rates of all the other mRNAs (129,212). (C) mRNA folding and translation elongation: mRNA folding tends to slow down ribosome elongation, however ribosomes also tend to unfold the mRNA (61,139). (D) Co-translational folding: local discontinuous translation rates temporally separate the translation of segments of the peptide chain and actively coordinate their co-translational folding, to promote accurate folding of the peptide. Pink represents the codon rates enabling correct folding of the α-helix domain, while orange the β-sheet domain (15,68). (E) The mRNA folding near the start codon affects translation initiation efficiency. Specifically, weak folding surrounding the start codon enables efficient recognition of the start codon and the initiation signals surrounding it by the pre-initiation complex (left), while strong folding may contribute to the pre-initiation complex missing the start codon (right). (F) The three different phases of the TASEP as a function of the initiation and elongation rate. (G) Rate limiting steps: for instance, efficient/non-efficient ribosome binding sites and weak/strong mRNA structure at the end of the 5′ UTR can promote either up/down regulation of translation (7,46,54,60–62,66,71,209,168,173). (H) The effect of the tRNA levels on codon decoding rates: codons that are recognized by low-abundance tRNA isoacceptors are decoded more slowly than those recognized by high-abundance tRNAs (19,42,46,51,54,66,68,140,149). Codon decoding rates are illustrated by the following colouring scheme - dark blue represents fast, red represents slow, light blue represents neutral.

As mentioned, one of the future challenges in the field is developing a whole cell simulation related to all mRNA translation aspects. Today there are models that consider the competition among many mRNAs (51–52,129,148–150,156–157) but omit additional central phenomena (e.g. interaction with the mRNA folding, the nascent peptide, ribosome drop off, initiation mechanisms, ribosome recycling, ribosome frame shift); some of these aspects were analyzed in simpler, single mRNA based, models (e.g. see 46,53–55,57,59,61,66,69,131,135,137–138,142–144,147). A later challenge includes developing comprehensive detailed models that connect all the gene expression steps and not only translation.

Another challenge is related to developing organism and tissue specific models of translation. As mentioned, a useful biophysical model of translation should include parameters that reflect the intracellular regime (e.g. number of ribosomes, concentration of translation factors, de-
coding rates, etc.), since these parameters vary among different organisms and cellular conditions an important step in the field will be to infer them for various organisms/tissues/conditions. Specifically, since most of the studies in the field are based on popular model organisms such as *E. coli* and *S. cerevisiae*, it will be important to develop such models for organisms from the three domains of life.

The organism specific models should provide organism/tissue specific answers to the biological questions and debates mentioned above (see Figure 3).

Moreover, there are central aspects related to translation that are today not considered in any of the models. The limited resources mentioned in the context of translation models in this review include mainly the ribosome pool and other translation factors. However, the cellular energy, i.e. ATP, is also a very important rate limiting factor for translation initiation and elongation which is not studied explicitly in our context.

Translation is known to be the most energy consuming intracellular process, consuming most (up to 75%) of the cellular energy (196).

Thus, it makes sense to develop models that explicitly consider the metabolic cost of translation together with the dynamics of the process. These models can, for example integrate metabolic modelling (197) with translation dynamics models to answer and consider aspects such as: the metabolic cost of traffic jams (e.g. due to ribosome drop off, and the energetic cost of jammed ribosomes); the energetic cost of synthesizing ribosomes (i.e. translating the ribosomal proteins); the energetic cost of generating other translation factors (e.g. tRNAs, ribosomal RNA); the relation/connection between competition of mRNAs for translation factors and their competition for cellular energy.

Initial attempts in this direction have been recently suggested (198); however, these studies haven’t considered the fundamental questions mentioned above.

The energetic aspects of translation, specifically its very high energetic cost, are clearly the central relation between the intracellular dynamics of translation and its effect on the way evolution shapes transcripts. Thus, considering explicitly energy consumption as a rate limiting resource is particularly important for molecular evolution studies that study translation via its biophysical modelling.

Another central challenge in the field is related to studying/modelling translation dynamics not under the assumption of steady state. While there are some theoretical papers that aimed at studying the dynamics of the translation models (199–201), currently there are no translation modelling studies that are based on biological data which focus on questions related to the dynamics of translation. The studies overviewed in this review focused on reporting steady state estimations related to the translation process. The assumptions in all these cases are that the number of translation factors, mRNAs, ribosomes, etc, are close to constant (small fluctuations around steady state), the mRNA life times are much longer than their translation times, and translation is studied not during the cell cycle or major changes in the system (e.g. G0—cell cycle arrest), etc. Thus, typical values of variables such as translation rates, initiation rates, and the number of ribosomes can be estimated and studied.

Little attention has been dedicated in the literature to study the processes beyond steady-state. For example, during transition from one state to the other of cell-cycle/cell division/cell growth we expect to observe significant absolute/relative changes in the levels of translation factors such as tRNAs, elongation factors, initiation factors, number of ribosomes, number of mRNAs and more (5,167,202–204). Thus, if we study the translation dynamics during the cell cycle it does not make sense to assume steady state and compute single typical values for each of the state variables during the entire cell cycle. Currently the main bottleneck related to the study of translation dynamics via computational biophysical models not during steady state is the data—there are no high resolution measurements of variables (e.g. tRNA levels, number of ribosomes, Ribo-Seq, mRNA levels, etc.) during different states of the cell, such as at different cell cycle steps, that enable inferring the relevant model parameters. In addition, it is important to remember that the data available today related to translation measurements (e.g. Ribo-Seq) is an average over many cells/mRNAs; accurate measurements of translation in real time (205–207) or single cell/single transcript measurements of translation should contribute towards the accurate modelling of translation dynamics without the steady state assumption.

**ACKNOWLEDGEMENT**

We would like to thank Alon Diament for helpful comments.

**FUNDING**

Israeli Ministry of Science, Technology and Space. Funding for open access charge: Israeli Ministry of Science, Technology and Space.

Conflict of interest statement. None declared.

**REFERENCES**

1. Vogel,C., de Sousa Abreu,R., Ko,D., Le,S.Y., Shaprio,B.A., Burns,S.C., Sandhu,D., Boutz,D.R., Marquette,E.M. and Penalva,L.O. (2010) Sequence signatures and mRNA concentration can explain two-thirds of protein abundance variation in a human cell line. *Mol. Syst. Biol.*, 6, 400.
2. Tuller,T., Kupiec,M. and Ruppin,E. (2007) Determinants of protein abundance and translation efficiency in *S. cerevisiae*. *PLoS Comput. Biol.*, 3, e248.
3. Ghaemmaghami,S., Huh,W.K., Bower,K., Howson,R.W., Belle,A., Dephoure,N., O’Shea,E.K. and Weissman,J.S. (2003) Global analysis of protein expression in yeast. *Nature*, 425, 737–741.
4. Cheng,Z., Teo,G., Krueger,S., Rock,T.M., Koh,H.W., Choi,H. and Vogel,C. (2016) Differential dynamics of the mammalian mRNA and protein expression response to misfolding stress. *Mol. Syst. Biol.*, 12, 855.
5. Alberts,B., Johnson,A., Lewis,J., Morgan,D., Raff,M., Roberts,K. and Walter,P. (2014) *Molecular Biology of the Cell*. Garland Science, NY.
6. Plotkin,J.B. and Kudla,G. (2011) Synonymous but not the same: the causes and consequences of codon bias. *Nat. Rev. Genet.*, 12, 32–42.
7. Tuller,T., Carmi,A., Vestiogian,K., Navon,S., Dorfan,Y., Zaborske,J., Pan,T., Dahan,O., Furman,I. and Pilpel,Y. (2010) An evolutionarily conserved mechanism for controlling the efficiency of protein translation. *Cell*, 141, 344–354.
22. Vogel, C. (2013) Protein expression under pressure.

21. Sharp, P. and Li, W.H. (1987) The rate of synonymous substitution in the tRNA pool to improve translation efficiency.

20. Bulmer, M. (1991) The selection-mutation-drift theory of synonymous codon usage.

19. Herschlag, D. (2003) Genome-wide analysis of mRNA translation profiles in Saccharomyces cerevisiae.

18. Ingolia, N.T., Ghaemmaghami, S., Newman, J.R.S. and Weissman, J.S. (2014) Genome-scale proteomics reveals Arabidopsis thaliana gene models and proteome dynamics. Science, 342, 938.

17. Rivas, A., Iverson, J.J., Whelan, J. and Wilhelm, T. (2007) Posttranscriptional expression regulation: what determines the tRNA pool to improve translation efficiency. mRNA levels.

16. Nissley, D.A. and O’Brien, E.P. (2014) Timing is everything: Unifying differences in human transfer RNA expression. PLoS Genet., 10, e1004413.

15. O’Brien, E.P., Vendruscolo, M. and Dobson, C.M. (2014) Kinetic modelling indicates that fast-translating codons can coordinate translational protein folding by avoiding misfolded intermediates. Nat. Commun., 5, 4988.

14. Nissley, D.A. and O’Brien, E.P. (2014) Timing is everything: Unifying translation rates and nascent proteome behavior. J. Am. Chem. Soc., 136, 17892–17898.

13. Bahir, I., Fromer, M., Prat, Y. and Linial, M. (2009) Viral adaptation to host: a proteome-based analysis of codon usage and amino acid preferences. Mol. Syst. Biol., 5, 311.

12. Fredrick, K. and Ibba, M. (2010) How the sequence of a gene can tune its translation. Cell, 141, 227–229.

11. Iost, I. and Dreyfus, M. (1995) The stability of Escherichia coli lacZ mRNA depends upon the simultaneity of its synthesis and translation. EMBO J, 14, 3252.

10. Zhang, F., Saha, S., Shabalina, S.A. and Kashina, A. (2010) Differential arginylation of actin isoforms is regulated by coding sequence-dependent degradation. Science, 329, 1534.

9. Kimchi-Sarfaty, C., Oh, J.M., Kim, I.W., Sauna, Z.E., Calcagno, A.M., Ambudkar, S.V. and Gottesman, M.M. (2007) A ‘silent’ polymorphism in the MDR1 gene changes substrate specificity. Science, 315, 352.

8. Stergachis, A.B., Haugen, E., Shafer, A., Fu, W., Vernot, B., Reynolds, A., Raubitschek, A., Ziegler, S., LeProust, E.M. and Akey, J.M. (2013) Exonic transcription factor binding directs codon choice and affects protein evolution. Science, 342, 1367–1372.

7. Kimchi-Sarfaty, C., Oh, J.M., Kim, I.W., Sauna, Z.E., Calcagno, A.M., Ambudkar, S.V. and Gottesman, M.M. (2007) A ‘silent’ polymorphism in the MDR1 gene changes substrate specificity. Science, 315, 352.

6. Nissley, D.A. and O’Brien, E.P. (2014) Timing is everything: Unifying translation rates and nascent proteome behavior. J. Am. Chem. Soc., 136, 17892–17898.

5. O’Brien, E.P., Vendruscolo, M. and Dobson, C.M. (2014) Kinetic modelling indicates that fast-translating codons can coordinate translational protein folding by avoiding misfolded intermediates. Nat. Commun., 5, 4988.

4. Nissley, D.A. and O’Brien, E.P. (2014) Timing is everything: Unifying translation rates and nascent proteome behavior. J. Am. Chem. Soc., 136, 17892–17898.

3. Bahir, I., Fromer, M., Prat, Y. and Linial, M. (2009) Viral adaptation to host: a proteome-based analysis of codon usage and amino acid preferences. Mol. Syst. Biol., 5, 311.

2. van Weringh, A., Ragonnet-Cronin, M., Prancevičienė, E., Pavon-Eternod, M., Kleiman, L. and Xia, X. (2011) HIV-1 modulates the tRNA pool to improve translation efficiency. Mol. Biol. Evol., 28, 1827.

1. Shah, P. and Gilchrist, M.A. (2010) Effect of correlated tRNA abundances on translation errors and evolution of codon usage bias. PLoS Genet., 6, e1001128.

0. Bulmer, M. (1991) The selection-mutation-drift theory of synonymous codon usage. Genetics, 129, 897.

- Vogel, C. (2013) Protein expression under pressure. Science, 342, 1052–1053.

- Silveira, D., Formenti, S.C. and Schneider, R.J. (2010) Translational control in cancer. Nat. Rev. Cancer, 10, 254–266.

- Arava, Y., Wang, Y., Storey, J.D., Liu, C.L., Brown, P.O. and Herschlag, D. (2003) Genome-wide analysis of mRNA translation profiles in Saccharomyces cerevisiae. Proc. Natl. Acad. Sci. U.S.A., 100, 3889.

- Ingolia, N.T., Ghaemmaghami, S., Newman, J.R.S. and Weissman, J.S. (2009) Genome-wide analysis in vivo of translation with nucleotide resolution using ribosome profiling. Science, 324, 218.

- Dittmar, K.A., Sørensen, M.A., Elf, J., Ehrenberg, M. and Pan, T. (2005) Selective charging of tRNA isoacceptors induced by amino-acid starvation. EMBO Rep., 6, 151–157.

- Uemura, S., Aitken, C.E.A., Korlach, J., Flusberg, B.A., Turner, S.W. and Puglisi, J.D. (2010) Real-time tRNA transit on single translocating ribosomes at codon resolution. Nature, 464, 1012–1017.

- Taniguchi, Y., Choi, P.J., Li, G.W., Chen, H., Babu, M., Hearn, J., Emili, A. and Xie, X.S. (2010) Quantifying E. coli proteome and transcriptome with single-molecule sensitivity in single cells. Science, 329, 533.

- Newman, J.R.S., Ghaemmaghami, S., Ihmels, J., Breslow, D.K., Noble, M., DeRisi, J.L. and Weissman, J.S. (2006) Single-cell proteomic analysis of S. cerevisiae reveals the architecture of biological noise. Nature, 441, 840–846.

- Lu, P., Vogel, C., Wang, R., Yao, X. and Marcotte, E.M. (2006) Absolute protein expression profiling estimates the relative contributions of transcriptional and translational regulation. Nat. Biotechnol., 25, 117–124.
subcellular distribution of miRNAs and tRNA 3′ trailers. PLoS One, 5, e10563.
77. Kawaguchi,R., Girke,T., Bray,E.A. and Bailey-Serres,J. (2004) Differential mRNA translation contributes to gene regulation under non-stress and dehydration stress conditions in Arabidopsis thaliana. Plant J., 38, 823–839.
78. Mustroph,A., Zanetti,M.E., Jang,C.H., Holtan,H.E., Repetti,P.P., Galbraith,D.W., Girke,T. and Bailey-Serres,J. (2009) Profiling translatomes of discrete cell populations resolves altered cellular priorities during hypoxia in Arabidopsis. Proc. Natl. Acad. Sci. U.S.A., 106, 18843–18848.
79. Aspden,J.L., Eyre-Walker,Y.C., Phillips,R.J., Amin,U., Mumtaz,M.A.S., Brocard,M. and Cousin,J.-P. (2014) Extensive translation of small open reading frames revealed by Poly-Ribo-Seq. Elife, 3, e03528.
80. Michel,A.M. and Baranov,P.V. (2013) Ribosome profiling: a Hi-Def monitor for protein synthesis at the genome-wide scale. Wiley Interdiscip. Rev. RNA, 4, 473–490.
81. Bazzini,A.A., Johnston,T.G., Christiano,R., Mackowiak,S.D., Obermayer,B., Fleming,E.S., Veijar,C.E., Lee,M.T., Rajewsky,N. and Walther,T.C. (2014) Identification of small ORFs in vertebrates using ribosome footprinting and evolutionary conservation. EMBO J., e201488411.
82. Smith,J.E., Alvarez-Dominguez,J.R., Kline,N., Huynh,N.J., Geisler,S., Hu,W., Coller,J. and Baker,K.E. (2014) Translation of small open reading frames within unannotated RNA transcripts in Saccharomyces cerevisiae. Cell Rep., 7, 1588–1566.
83. Zupanic,A., Meplan,C., Grellscheid,S.N., Mathers,J.C., Kirkwood,T.B., Hesketh,J.E. and Shanley,D.P. (2014) Detecting translational regulation by change point analysis of ribosome profiling data sets. RNA, 20, 1507–1518.
84. Gerashchenko,M.V., Lobanov,A.V. and Gladyshev,V.N. (2012) Genome-wide ribosome profiling reveals complex translational regulation in response to oxidative stress. Proc. Natl. Acad. Sci. U.S.A., 109, 17394–17399.
85. Menschhaert,G., Van Criendinge,W., Noteelaerts,T., Koch,A., Crappe,J., Gevaert,K. and Van Damme,P. (2013) Deep proteome coverage based on ribosome profiling aids mass spectrometry-based protein and peptide discovery and provides evidence of alternative translation products and near-cognate translation initiation events. Mol. Cell. Proteomics, 12, 1780–1790.
86. Juntawong,P., Girke,T., Bazin,J. and Bailey-Serres,J. (2014) Translational dynamics revealed by genome-wide profiling of ribosome footprints in Arabidopsis. Proc. Natl. Acad. Sci. U.S.A., 111, E203–E212.
87. Han,Y., Gao,X., Liu,B., Wan,J., Zhang,X. and Qian,S.-B. (2014) Ribosome profiling reveals sequence-independent post-initiation pausing as a signature of translation. Cell Res., 24, 842–851.
88. Guttman,M., Russell,P., Ingolia,N.T., Weissman,J.S. and Lander,E.S. (2013) Ribosome profiling provides evidence that large noncoding RNAs do not encode proteins. Cell, 154, 246–251.
89. Brar,G.A., Yassour,M., Friedman,N., Regev,A., Ingolia,N.T. and Weissman,J.S. (2012) High-resolution view of the yeast mitotic program revealed by ribosome profiling. Science, 335, 552–557.
90. Ingolia,N.T., Brar,G.A., Stern-Ginossar,N., Harris,M.S., Talhouarme,G.J., Jackson,S.E., Wills,M.R. and Weissman,J.S. (2014) Ribosome profiling reveals pervasive translation outside of annotated protein-coding genes. Cell Rep., 8, 1365–1379.
91. Michel,A.M., Choudhury,K.R., Firth,A.E., Ingolia,N.T., Atkins,J.F. and Baranov,P.V. (2012) Observation of dually decoded regions of the human genome using ribosome profiling data. Genome Res., 22, 2219–2229.
92. Lee,S., Liu,B., Lee,S., Huang,S.-X., Shen,B. and Qian,S.-B. (2012) Global mapping of translation initiation sites in mammalian cells at single-nucleotide resolution. Proc. Natl. Acad. Sci. U.S.A., 109, E2424–E2432.
93. Ingolia,N.T., Lareau,L.F. and Weissman,J.S. (2011) Ribosome profiling of mouse embryonic stem cells reveals the complexity and dynamics of mammalian proteomes. Cell, 147, 789–802.
94. Yewdell,J.W., Raek,J.R., Reochster,M.C. and Nihitcha,C.Y. (2011) Out with the old, in with the new? Comparing methods for measuring protein degradation. Cell Biol. Int., 35, 457–462.

52. Mehra,A. and Hatzimanikatis,V. (2006) An algorithmic framework for genome-wide modeling and analysis of translation networks. Biophys. J., 90, 1136–1146.
53. Mitarai,N., Sneppe,K. and Pedersen,S. (2008) Ribosome collisions and translation efficiency: optimization by codon usage and mRNA destabilization. J. Mol. Biol., 382, 236–248.
54. Ciandrini,L., Stansfield,E. and Romano,M.C. (2013) Ribosome traffic on mRNAs maps to gene ontology: genome-wide quantification of translation initiation rates and polysome size regulation. PLoS Comput Biol, 9, e1002866.
55. Dimelou,R.J. and Wilkinson,S.J. (2009) Control of translation initiation: a model-based analysis from limited experimental data. J. Roy. Soc. Interf, 6, 51–61.
56. Fluitt,A., Pienaar,E. and Viljoen,H. (2007) Ribosome kinetics and aa-tRNA competition determine rate and fidelity of peptide synthesis. Comput. Biol. Chem., 31, 335–346.
57. Zhang,G., Hubalewska,M. and Ignatova,Z. (2009) Transient ribosomal attenuation coordinates protein synthesis and co-translational folding. Nat. Struct. Mol. Biol., 16, 274–280.
58. Shah,P., Ding,Y., Niemczyk,M., Kudla,G. and Plotkin,J.B. (2013) Rate-limiting steps in yeast protein translation. Cell, 153, 1599–1601.
59. Berthelot,K., Muldoon,M. and Walder,J. (2006) An algorithmic framework for genome-wide modeling and analysis of translation networks. PLoS Comput Biol, 2, 237–243.
60. Zhang,G. and Ignatova,Z. (2009) Generic algorithm to predict the speed of translational elongation: implications for protein biogenesis. PLoS One, 4, e5036.
61. Siwiak,M. and Zienlenkiewicz,P. (2010) A comprehensive, quantitative, and genome-wide model of translation. PLoS Comput. Biol., 6, e1000855.
62. Rotman,A., Markov,M., Eide,J. and Lander,E.S. (2013) Evolutionary conservation of codon optimization reveals hidden signatures of cotranslational folding. Nat. Struct. Mol. Biol., 20, 237–243.
63. Mitarai,N., Sneppen,K. and Pedersen,S. (2008) Ribosome collisions and translation efficiency: optimization by codon usage and mRNA destabilization. J. Mol. Biol., 382, 236–248.
64. Denecke,C., Lipowsky,R. and Valleriani,A. (2013) Effect of ribosome shielding on mRNA stability. Proc. Natl. Acad. Sci. U.S.A., 109, 1365–1379.
65. Liu,B., Gao,X., Liu,B., Wan,J., Zhang,X. and Qian,S.-B. (2014) Ribosome profiling reveals pervasive translation outside of annotated protein-coding genes. Cell Rep., 8, 1365–1379.
95. Belle, A., Tanay, A., Bitincka, L., Shamir, R. and O’Shea, E.K. (2006) Quantification of protein half-lives in the budding yeast proteome. *Proc. Natl. Acad. Sci. U.S.A.*, 103, 13004–13009.

96. Johnson, A.E. (2005) The co-translational folding and interactions of nascent protein chains: a new approach using fluorescence resonance energy transfer. *FEBS Lett.*, 579, 916–920.

97. Clark, P.L. and Ugrinov, K.G. (2009) Measuring cotranslational folding of nascent polypeptide chains on ribosomes. *Methods Enzymol.*, 466, 576–590.

98. Tuller, T. and Zur, H. (2015) Challenges and obstacles related to solving the codon bias ridges. *Biochem. Soc. Trans.*, 42, 155–159.

99. Burgess-Brown, N.A., Sharma, S., Sobott, F., Loenarz, C., Oppermann, U. and Gileadi, O. (2008) Codon optimization can improve expression of human genes in Escherichia coli: A multi-gene study. *Protein Expr. Purif.*, 59, 94–102.

100. Weich, M., Govindarajan, S., Ness, J.E., Villalobos, A., Gurney, A., Minshull, J. and Gustafsson, C. (2009) Design parameters to control synthetic gene expression in Escherichia coli. *PLoS One*, 4, e7002.

101. Goodman, D.B., Church, G.M. and Kosuri, S. (2013) Causes and effects of N-terminal codon bias in bacterial genes. *Science*, 342, 475–479.

102. Ceroni, F., Algar, R., Stan, G.-B. and Ellis, T. (2015) Quantifying cellular capacity identifies gene expression designs with reduced burden. *Nat. Methods*, 12, 415–418.

103. Diamant, E. and Tuller, T. (2016) Estimation of ribosome profiling performance and reproducibility at various levels of resolution. *Biol. Direct*, 11, 1.

104. Margaliot, M. and Tuller, T. (2012) Quantification of codon–tRNA interactions based on codon usage bias. *Biomed. Res. Int.*, 2012, 974635.

105. Shaw, L.B., Sethna, J.P. and Lee, K.H. (2004) Mean-field approaches to the totally asymmetric exclusion process with quenched disorder and large particles. *Phys. Rev. E*, 70, 021901.

106. Che, T. and Lakatos, G. (2004) Clustered bottlenecks in mRNA translation and protein synthesis. *Phys. Rev. Lett.*, 93, 198101.

107. Zouridis, H. and Hatzimanikatis, V. (2008) Effects of codon distributions and tRNA competition on protein translation. *Biophys. J.*, 95, 1018–1033.

108. Zouridis, H. and Hatzimanikatis, V. (2010) The origins of time-delay in template biopolymerization processes. *PLoS Comput. Biol.*, 6, e1000726.

109. Sharma, A.K. and Chowdhury, D. (2011) Stochastic theory of protein synthesis and polysome: Ribosome profiling on a single mRNA transcript. *J. Theor. Biol.*, 289, 36–46.

110. Basu, A. and Chowdhury, D. (2007) Traffic of interacting ribosomes: effects of single-machine mechanochemistry on protein synthesis. *Phys. Rev. E*, 75, 021902.

111. Seyfried, A. and Anderson, D.A. (2008) Dynamic model of the ribosome exclusion process with extended objects: A model for protein synthesis. *Bull. Math. Biol.*, 70, 1771–1793.

112. Biro, N. and Tuller, T. (2015) Modelling the efficiency of ribosomoprophiling data indicates a major role for proline in stalling translation. *Genome Res.*, 25, 4190–4198.

113. Shaw, L.B., Zia, R. and Lee, K.H. (2003) Totally asymmetric exclusion process with quenched disorder. *J. Stat. Phys.*, 112, 21–34.

114. Skjøndal-Bara, N. and Morris, D.R. (2007) Dynamic model of the process of protein synthesis in eukaryotic cells. *Bull. Math. Biol.*, 69, 361–393.

115. Dong, J., Schmittmann, B. and Zia, R.K. (2007) Towards a model for protein production rates. *J. Stat. Phys.*, 128, 21–34.

116. Arteri, C.G. and Fraser, H.B. (2014) Accounting for biases in riboprofiling data indicates a major role for proline in stallling translation. *Genome Res.*, 24, 1011–2011.

117. Liu, J. and Deutsch, C. (2008) Electrostatics in the ribosomal tunnel modulation chain elongation rates. *J. Mol. Biol.*, 384, 73–86.

118. Marshall, E., Stansfeld, I. and Romano, M. (2014) Ribosome recycling induces optimal translation rate at low ribosomal availability. *J. Roy. Soc. Interface*, 11, 20140589.
computationally design mRNA sequences with desired expression levels in prokaryotes. BMC Syst. Biol., 4, 1.
196. Lane, N. and Martin, W. (2010) The energetics of genome complexity. Nature, 467, 929–934.
197. Bordbar, A., Monk, J.M., King, Z.A. and Palsson, B.O. (2014) Constraint-based models predict metabolic and associated cellular functions. Nat. Rev. Genet., 15, 107–120.
198. Thiele, I., Fleming, R.M., Que, R., Bordbar, A., Diep, D. and Palsson, B.O. (2012) Multiscale modeling of metabolism and macromolecular synthesis in E. coli and its application to the evolution of codon usage. PLoS One, 7, e45635.
199. Margaliot, M., Sontag, E.D. and Tuller, T. (2014) Entrainment to periodic initiation and transition rates in a computational model for gene translation. PLoS One, 9, e96039.
200. De Jong, H. (2002) Modeling and simulation of genetic regulatory systems: a literature review. J. Comput. Biol., 9, 67–103.
201. Nagar, A., Vallieriani, A. and Lipowsky, R. (2011) Translation by ribosomes with mRNA degradation: Exclusion processes on aging tracks. J. Stat. Phys., 145, 1385–1404.
202. Higareda-Mendoza, A.E. and Pardo-Galván, M.A. (2010) Expression of human eukaryotic initiation factor 3f oscillates with cell cycle in A549 cells and is essential for cell viability. Cell Division, 5, 1.
203. Patil, A., Dyavaiah, M., Joseph, F., Rooney, J.P., Chan, C.T., Dedon, P.C. and Begley, T.J. (2012) Increased tRNA modification and gene-specific codon usage regulate cell cycle progression during the DNA damage response. Cell Cycle, 11, 3656–3665.
204. Spellman, P.T., Sherlock, G., Zhang, M.Q., Iyer, V.R., Anders, K., Eisen, M.B., Brown, P.O., Botstein, D. and Futcher, B. (1998) Comprehensive identification of cell cycle–regulated genes of the yeast Saccharomyces cerevisiae by microarray hybridization. Mol. Biol. Cell, 9, 3273–3297.
205. Yan, X., Hoek, T.A., Vale, R.D. and Tanenbaum, M.E. (2016) Dynamics of translation of single mRNA molecules in vivo. Cell, 165, 976–989.
206. Wu, B., Eliscovich, C., Yoon, Y. and Singer, R.H. (2016) Translation dynamics of single mRNAs in live cells and neurons. Science, 352, 1430–1435.
207. Morisaki, T., Lyon, K., DeLuca, K.F., DeLuca, J.G., English, B.P., Zhang, Z., Lavis, L.D., Grimm, J.B., Viswanathan, S., Looger, L.L. et al. (2016) Real-time quantification of single RNA translation dynamics in living cells. Science, 352, 1425–1429.
208. Gillespie, D.T. (1977) Exact stochastic simulation of coupled chemical reactions. J. Phys. Chem., 81, 2340–2361.
209. Chu, D., Kazana, E., Bellanger, N., Singh, T., Tuite, M.F. and Haar, T. (2014) Translation elongation can control translation initiation on eukaryotic mRNAs. EMBO J., 33, 21–34.
210. Warner, J.R. (1999) The economics of ribosome biosynthesis in yeast. Trends Biochem. Sci., 24, 437–440.
211. Lauria, F., Tebaldi, T., Lunelli, L., Struffi, P., Gatto, A., Brigotti, M., Montanaro, L., Cribilli, Y. and Inga, A. (2015) RiboAbacus: a model trained on polyribosome images predicts ribosome density and translational efficiency from mammalian transcriptomes. Nucleic Acids Res., 43, e153.
212. Gorochowski, T.S., Avciar-Kucukgoze, I., Bovenberg, R.A., Roukos, I.A. and Ignatova, Z. (2016) A minimal model of ribosome allocation dynamics captures trade-offs in expression between endogenous and synthetic genes. ACS Synth. Biol., 5, 710–720.