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SYNONYMOUS CODON USAGE – A GUIDE FOR CO-TRANSLATIONAL PROTEIN FOLDING IN THE CELL

A. A. Komara,b,c,d,*

aCenter for Gene Regulation in Health and Disease and Department of Biological, Geological and Environmental Sciences, Cleveland State University, Cleveland, Ohio 44115, USA

bDepartment of Biochemistry and Center for RNA Science and Therapeutics, Case Western Reserve University, Cleveland, Ohio 44106, USA

^cGenomic Medicine Institute, Lerner Research Institute, Cleveland Clinic, Cleveland, OH, 44195, USA

^dDAPCEL, Inc., Cleveland, OH, 44106

Abstract

In the cell, protein folding begins during protein synthesis/translation and thus is a co-translational process. Co-translational protein folding is tightly linked to translation elongation, which is not a uniform process. While there are many reasons for translation non-uniformity, it is generally believed that non-uniform synonymous codon usage is one of the key factors modulating translation elongation rates. Frequent/optimal codons as a rule are translated more rapidly than infrequently used ones and vice versa. Over 30 years ago, it was hypothesized that changes in synonymous codon usage affecting translation elongation rates could impinge on co-translation protein folding and that many synonymous codons are strategically placed within mRNA to ensure a particular translation kinetics facilitating productive step-by-step co-translational folding of proteins. It was suggested that this particular translation kinetics (and, specifically, translation pause sites) may define the window of opportunity for the protein parts to fold locally, particularly at the critical points where folding is far from equilibrium. It was thus hypothesized that synonymous codons may provide a secondary code for protein folding in the cell. Although, mostly accepted now, this hypothesis appeared to be difficult to prove and many convincing results were obtained only relatively recently. Here, I review the progress in the field and explain, why this simple idea appeared to be so challenging to prove.

Keywords

genetic code; codon usage; synonymous codons; translation kinetics; nascent peptides; cotranslational protein folding

* a.komar@csuohio.edu .

INTRODUCTION

Protein folding is one of the most fundamental mechanisms in the cell. For any protein to function properly, the polypeptide chain produced by the ribosome, the protein-synthesizing factory of the cell, has to fold into the correct three-dimensional structure. Misfolding and aggregation are implicated in a number of diseases, including many neurodegenerative proteinopathies, such as Alzheimer's disease, prion diseases and Parkinson's disease to name a few [1-3]. Therefore, knowledge of how proteins acquire their spatial structure (folds) is extremely important.

For decades the mechanism by which the polypeptide chain acquires its native structure has been investigated mainly by in vitro denaturation/renaturation experiments [4, 5]. Seminal research performed by Christian Anfinsen and his colleagues on the reversible denaturation of ribonuclease (in the 1950-60's) prompted their suggestion that a protein's amino acid sequence contains all the information necessary to specify its unique three-dimensional structure [6]. These original experiments were followed by many in depth *in vitro* unfolding/ refolding as well as in silico studies and provided a wealth of information suggesting that protein folding obeys a sequential model which postulates a unique pathway with defined intermediates [4, 5, 7, 8]. Subsequently, the concept of the folding funnel has been developed further suggesting that there could be multiple pathways which guide protein folding to a native conformation with the lowest free energy minimum [7, 9]. The majority of these studies employed relatively small proteins that can be successfully refolded in aqueous solutions, and these observations supported the Anfinsen's principle [4, 5, 7-9]. Yet, many attempts to achieve *in vitro* 100% refolding of isolated denatured proteins were only partially successful [4]. In addition, in most cases reconstitution in a test tube was found to be exceedingly slow and not comparable with times required/expected for a protein to acquire its native structure in the cell [10].

It thus became evident that comprehensive understanding of the mechanism of protein folding requires elucidation of the folding pathway under native conditions, such as those that exist in vivo in the cell $[11-14]$. These conditions are quite different from those in a test tube. Firstly, folding in vivo takes place in a crowded cellular environment and thus is thought to be affected by a number of factors, such as concentration and activity/affinity of surrounding macromolecules and the presence of folding catalysts and accessory proteins [15, 16]. Systematic investigations of these effects revealed an important role played by folding catalysts and accessory proteins in acceleration of the rate of *in vivo* protein folding and in the prevention of protein misfolding and aggregation [15, 16]. However, it has been also suggested that both chaperones and folding catalysts are mainly involved in kinetic partitioning between proper folding and aggregation; thus, they thought to affect the yield (of the correctly folded protein) rather than the folding mechanism. Secondly, and most importantly, it became recognized that in vivo, protein folding begins co-translationally as nascent peptide chains emerge from the ribosome [17-25]. In the early 1960s and 1970s, the first observations were made [26-32] suggesting that *in vivo* protein folding starts while the growing peptide chain is still bound to the ribosome and that it is a vectorial process; i.e. the polypeptide chain is synthesized and is being folded predominantly from the N-terminal to the C-terminal end. Co-translational folding of a nascent polypeptide was

thus suggested to result in a sequential structuring of distinct regions of the polypeptide emerging from the ribosome at different points in time and hierarchical condensation has therefore been considered to be the most likely mechanism that governs assembly of the nascent polypeptide into the native protein during its synthesis in vivo. Earlier studies on the co-translational protein folding were followed by many modern experiments and are continuing to date, revealing many interesting features of the process ([17-25] and ref. therein).

It became also clear, that co-translational folding starts almost immediately after the first amino acid residues begin to polymerize at the ribosomal peptidyl transferase center ([17-25] and ref. therein), with alpha-helices forming inside the ribosome tunnel and some elements of the supersecondary and tertiary structure possibly forming already in the so-called vestibule (lower/wider) region of the exit tunnel [33-36]. Once the polypeptide chain emerges from the exit tunnel into the cytosol, the folding continues governed by the thermodynamics and kinetics of polypeptide chain, leading to the formation of subsequent co-translational folding intermediates and, finally, the native structure, which is usually being completely formed after the chain's release [17-25].

It should be noted that the ribosome reads the mRNA codons one-by-one and translates them into the sequence of amino acids of the protein. However, it became clear that not all codons are read with the same speed: periods of rapid translation are separated by translation pauses [17, 37, 38]. Therefore, it can't be excluded that variations in local translation rates may affect/facilitate protein folding by allowing ordered, sequential structuring of the discrete nascent polypeptide chain portions synthesized by the ribosome and that kinetics of protein synthesis may thus influence/ fine-tune the co-translational protein folding.

In the late 1980's Alistair Brown's group in the Institute of Genetic at Glasgow University, UK [39] and our group in the Department of Molecular Biology at Moscow State University, Russia [40-42] suggested that sequential folding events, which can take place during cotranslational folding of proteins, might be separated by translational pauses and that such regions of slowed translation might serve as interpunctuations during co-translational protein folding. This hypothesis was put forward based on observations that revealed a certain correlation between the locations of rare (slowly translated) codons in mRNA with either the domain boundaries in the encoded proteins [39, 41], or with the boundaries of the smaller structural units such as secondary or supersecondary structure elements [40, 42].

This hypothesis relied on several assumptions. First, it was based on the presumed assumption that non-uniformity in synonymous codon usage along mRNA would lead to a particular translation kinetics, resulting in ribosome pausing (at rare codon clusters) or ribosome acceleration (at frequent codon clusters), respectively ([17] for a review, [39-42]). Second, it assumed that synonymous codons are placed in mRNA non-randomly and strategically (thus facilitating ordered co-translational protein folding) and that changes in synonymous codon usage would lead to a different translation kinetics that in turn may alter protein folding ([17, 39-42]). Third, it also assumed that altered kinetics of translation will affect the conformation of the ribosome-bound nascent chains on the first place, subsequently potentially also changing the final conformation of the released protein

and/or altering the equilibrium between different protein conformers (native and near-native and/or non-native), which in turn could lead to e.g. enhanced protein aggregation and/or degradation (co- or post-translational), or a change of the protein's specific activity [17].

It should be noted that while overall hypothesis (broadly stating that synonymous codon usage along mRNA may serve as a kinetic guide for co-translational protein folding in the cell) has now been generally accepted by the scientific community, many of its postulates remain subjects of intense debate. Nevertheless, advances in modern techniques such as Next Generation Sequencing (NGS), single-molecule and time-resolved fluorescent approaches are transforming our ability to study co-translational protein folding and allowing to obtain key evidence in support of this hypothesis.

Below, I'll review some of this key evidence in support of each subsequent postulate of the hypothesis.

TRANSLATION IS A NON-UNIFORM PROCESS GOVERNED BY SYNONYMOUS CODON USAGE

The genetic code is degenerate [43]. With the exception of two amino acids (Met and Trp), all other amino acid residues are each encoded by multiple, so-called synonymous codons [37]. Synonymous codons are however not present at equal frequencies in individual mRNAs as well as entire genomes [37, 44-46]. This pattern of non-uniform codon use is known as codon usage bias [37, 44-46] (Fig. 1a). Codon usage bias varies between organisms and represents a unique feature of an organism [47-50] (Fig. 1a). This suggests that codon choice might have functional implications beyond amino acid coding [37, 44-46]. Organism-specific codon choice is related to organism-specific differences in populations of cognate tRNAs [44]. It was generally found that in both unicellular and multicellular organisms there exists a strong positive correlation between codon usage and cellular tRNA content [47, 51, 52], meaning that codon bias would likely have a direct impact on translation elongation rates. Indeed, frequently used codons were, as a rule, found to be translated more rapidly than infrequently used ones due to the more ready availability (during translation) of corresponding frequent cognate tRNAs and vise versa (see [22, 37, 53-55] for reviews) (Fig. 1b). Several additional lines of evidence supported this notion (i) highly expressed genes were found to harbor more preferred/frequent codons compared to lowly expressed genes, which were found to be enriched in synonymous un-preferred codons [56-59]; (ii) substitution of synonymous frequently used codons by infrequently used codons (or vice versa) affected protein expression levels (see [22, 37, 53-55] for reviews) (iii) alterations in the level of expression/abundance of particular tRNAs also altered protein expression levels [60-64].

While it was generally accepted that frequent codons would accelerate translation and rare codons would cause a translation pause, what appeared to be extremely challenging is to prove that a particular rare codon (or a cluster of rare codons) would determine the appearance of the corresponding ribosome-mediated translational pause at a particular place in mRNA.

Earlier experiments attempted to verify this notion by analyzing the sizes of nascent chains attached to the ribosome [65-68]. As discontinuous elongation rates were presumed to increase the residence time of a ribosome at particular positions along mRNA (enriched in rare codons), it was expected that this would lead to an increase in the amount of nascent peptides of the corresponding sizes present on polyribosomes. At first, the accumulation of nascent peptides of discrete sizes has been monitored by the use of gel-filtration [65, 66] and/or gel electrophoresis [67, 68]. Although, it was generally observed that enrichment in rare codons leads to enhanced pausing [65-68], precise determination and localization of translation pause sites appeared to be quite challenging due to low resolution of these methods. The development of a micrococcal nuclease protection assay [69, 70] and, subsequently, the ribosome profiling (which combined ribosome footprinting with deep sequencing of mRNA fragments protected by ribosomes during translation) have allowed the direct analysis of ribosome distribution along mRNA at codon resolution [71-73]. However, ribosome profiling experiments, at first, yielded puzzling results. While local variations in translation rates have been clearly detected, no reliable correlation between the position of ribosomes and rare codons has been originally observed [71]. It appeared however that detecting pause sites and their relation to rare codons in the original ribosome profiling data have been challenging, because the methods used to arrest translation, involved antibiotics (like cycloheximide), which skewed the position of ribosomes on messages and obscured the enrichment of ribosome density at non-optimal codons [71,74]. A systematically-revised ribosome profiling method revealed pause sites at rare codons at single-codon resolution [75]. The new data also revealed a clear negative correlation between ribosome density and codon adaptation index, consistent with the expectation that rare codons will be decoded by lower-abundance tRNAs more slowly than more abundant codons [75].

SYNONYMOUS CODONS ARE PLACED IN mRNA NON-RANDOMLY AND STRATEGICALLY

The neutral theory of molecular evolution suggests that synonymous codons (encoding the same amino acid) will be largely unaffected by the selective pressure and thus they should be distributed in mRNA and substituted during evolution randomly [76]. It appeared however that less then half of all synonymous substitutions are under neutral expectation and that synonymous mutations are subjects to constraints [77, 78]. Close examination of mRNA sequences revealed biases in the distribution of codons within mRNA open reading frames (ORFs), a phenomenon originally recognized as codon context [79-81]. In addition to biases in synonymous codon usage relative to neighboring codons in an mRNA (codon context [79-81] and the so-called codon pair bias [82-86]), it has also been established that codon choice (particularly for rare codons) is biased according to a codon's specific conserved location in an mRNA [87-94]. Rare/infrequent codons, specifically, have been shown to occur in clusters, enriched at a number of specific locations in mRNAs. These include clusters of rare codons located at 5' and 3' ORF termini [88-92], the so-called "+70" rare codon cluster (located \sim 35–40 codons downstream of the signal sequences (or transmembrane segments) in secreted proteins [87, 93], and many other internal clusters located at specific positions (88). Although earlier studies (focused on investigation of codon choice relative to codon placement at specific positions in mRNA [39-42, 94-97]) have not

been comprehensive enough (partially owing to a lack of sufficient structure and sequence information), it was nevertheless concluded that the locations of rare codon clusters along mRNAs are highly conserved throughout evolution, as for example, evidenced by their similarity across homologous protein families from different organisms [17, 41, 42, 88, 94, 98-100]. This observation supported the assumption that such placement of these rare codon clusters may be linked to protein structure [17, 88]. Strategically placed rare codon clusters were specifically observed to occur, for example, at regions encoding (or close to) domain/subdomain linkers (Fig. 2) and such location of these clusters were suggested to allow temporal separation of domain and/or sub-domain folding on the ribosome [17, 88, 94].

In general, genome-wide analyses of ORFeomes from prokaryotic and eukaryotic organisms revealed that rare codon clustering (i) is not limited to a particular set of genes or genotype, (ii) does not depend on and is not related to the overall GC content of the organism's genome, and (iii) is significantly more abundant than would be expected based on random selection [88, 98]. While the precise functional roles of many conserved codon clusters has yet to be determined, it is becoming evident that codon choice has functional implications beyond amino acid coding and support of the predefined translation levels of a protein and that mRNA (and thus a genetic code) might indeed contain a secondary information linked to protein structure/folding.

CHANGES IN SYNONYMOUS CODON USAGE AFFECT LOCAL TRANSLATION ELONGATION RATES

It is widely believed that the major influence of codon usage is on global translation rate. Approaches involving substitution of the majority (or a subset) of infrequently used codons with synonymous frequently used ones, have been widely used for optimization of protein expression [54, 101-103] and references therein). Use of gene sequences optimized through this strategy often yielded large amounts of recombinant proteins, indicating that protein synthesis/translation elongation rates of the engineered proteins have been substantially accelerated [54,101-103].

It must be noted however that in addition to the effects of codon usage on translation, synonymous codon choice can also impact the turnover/stability of mRNA template itself [55,104-106]. mRNA turnover is a critical determinant of gene expression, and mRNAs with longer half-lives would typically produce more protein. It was recently found that stable/ long-lived mRNAs harbor mainly preferred/optimal codons while many unstable/short-lived mRNAs have a higher frequency of un-preferred/non-optimal/rare codons within their ORFs [104-106]. Substitution of preferred codons with synonymous, un-preferred codons resulted in dramatic destabilization of the mRNA and vice versa (104-106).

These and other experiments brought about awareness of the scientific community to the impact of synonymous codon usage and codon adaptation index (CAI) (as a measure of synonymous codon usage bias [107]) on the efficiency of translation and protein expression ([101-107] and ref. therein). However, despite this general acceptance of the idea that substitution of synonymous codons in a gene can dramatically affect the rate/efficiency of

synthesis of the encoded protein and increase or decrease its yield, only limited studies have attempted to investigate, how local substitutions of synonymous codons would affect the elongation rate(s) at the particular region(s) of mRNA, where these changes were introduced. Nonetheless, such evidence has been obtained.

Earlier studies attempted to monitor increase and/or decrease in the residence time of a ribosome at particular positions along mRNA after changing synonymous codons via monitoring the increase/decrease of the amount of nascent peptides of the corresponding sizes [68, 108]. Renewed attempts have been made recently to estimate the ribosome residence time at different synonymous codons using ribosome profiling and time-resolved single-molecule fluorescence methods ([38, 75, 109, 110] and ref. therein). Ensemble realtime fluorescence approaches have been also used to determine how bulk substitutions of synonymous codons would affect the speed of ribosome movement [36, 111]. These experiments combined with the experiments that measured the influence of tRNA availability, clearly indicated that there is a direct link between synonymous codons usage and the local translation elongation rates and that changes in synonymous codon usage do affect translational kinetics.

FOLDING in vivo IS A CO-TRANSLATIONAL PROCESS

A direct demonstration of the influence of synonymous codon usage on co-translational protein folding required two pieces of evidence to be obtained: (i) indicating that codon usage may affect the final conformation of a protein and (ii) indicating that these conformational changes may originate within the nascent chains bound to the ribosome i.e. co-translationally. As has been mentioned above, the original hypothesis was put forward in the late 1980s [39-42]. However, at that time, even the basic idea that folding in vivo is a co-translational process was not widely accepted yet.

So, at first, it was necessary to obtain solid evidence in support of co-translational protein folding. The majority of the earlier experiments in support of co-translational folding involved isolation/fractionation of ribosome-bound nascent chain complexes through a sucrose density gradient, followed by assessment of the structural properties of the nascent chains through measurement of i) their specific enzymatic activities, ii) their recognition by specific/conformational antibodies, or iii) formation of correct disulfide cross-bridges within and/or between nascent chains ([23, 24] and ref. therein). Subsequently, other methods have been introduced ([23, 24] and ref. therein), such as those involving e.g., measurement of the resistance of ribosome attached nascent chains to proteolytic digestion and/or the ability of co-factors and ligands (such as heme) to bind the growing polypeptide chain (as an indication that a binding-competent conformation has been achieved).

Our group together with Alexander S. Spirin's laboratory at the institute of Protein Research in Pushchino used heme binding to probe co-translational folding of the α -globin chains [112,113]. Using in vitro translation reactions performed in the presence of $[3H]$ hemin and [³⁵S]methionine together with sucrose gradient centrifugation and puromycin treatment, we showed that ribosome-bound α -globin chains are capable of efficient heme binding [113]. In addition, we found that incomplete α–globin nascent chains attached to the ribosome are

capable of co-translational heme binding, indicating that a structure that allows for heme binding in the nascent chain is achieved prior to the completion of α–globin synthesis. These results provided strong support for co-translational folding of the α -globin molecule [113].

However, back at a time, one of the most solid supports of co-translational protein folding came from the work done by Kolb, Makeyev and Spirin, who pioneered the study of co-translational protein folding using real-time measurements [114, 115]. These authors developed a technique allowing to continuously monitor enzymatic activity of newly synthesized firefly luciferase in a cell-free system in a luminometer cuvette and showed that luciferase activity (indicative of folding of the protein) can be detected as soon as the full-length molecule was formed in the translation reaction [114]. Importantly, such rapid acquisition of the enzyme's activity was incompatible with a post-translational folding scenario [114]. Furthermore, subsequently, the same authors demonstrated that ribosomebound luciferase can be enzymatically active and concluded that folding of the firefly luciferase protein occurs during the course of translation [115].

More recently, a plethora of modern technologies, such as NMR spectroscopy, cryoelectron microscopy (cryo-EM) and fluorescent techniques (Fluorescence Resonance Energy Transfer (FRET) and fluorescence anisotropy/dynamic fluorescence depolarization) as well as some other approaches have been introduced to study co-translational folding ([19, 23, 24] and ref. therein). These and other experiments provided overwhelming evidence in support of co-translational protein folding and the idea of co-translational protein folding has now become widely accepted [17-25]. What remained unclear however, is to what extent the co-translational folding pathway is unique and whether it can be indeed influenced by the kinetics of translation?

Therefore, many researchers in the field turned their attention to the analysis of the influence of kinetics of translation on co-translational protein folding. These experiments appeared to be extremely challenging, because of the numerous quality control mechanisms existing in a cell, by which a cell monitors proteins to ensure that they are appropriately folded and if not - guides them to degradation [12-16]. It was thus possible that changes in co-translational folding caused by synonymous codon-driven alterations in elongation kinetics could not be substantial enough to overcome the effects of cellular quality control and chaperone network machineries and become detectable/visible.

SYNONYMOUS CODON USAGE INFLUENCES CO-TRANSLATIONAL PROTEIN FOLDING

The choice of cell-free translation system(s), where the effects of the cellar quality control mechanisms could be less pronounced, or could be deliberately controlled by omitting members of the quality control machineries seemed to be therefore logical at first and is one of the preferred approaches for the analysis of the influence of synonymous codon usage on co-translational protein folding at present time. The *in vitro* translational systems and, especially, the fully reconstituted in vitro systems also allow easy control of many

In 1999, we have provided one the first observations showing that synonymous codon substitutions affect ribosome traffic and protein folding during *in vitro* translation of a model protein, chloramphenicol acetyltransferase (CAT) [108]. In this study, sixteen consecutive rare codons in the CAT gene have been replaced by frequent ones and this led to the acceleration of the ribosome traffic through the mutated region and at the same time affected the specific activity of the enzyme (in comparison with the wild-type protein) [108]. Since specific activity of a given protein could be considered as a measure of its proper folding, we have concluded that CAT folding was affected. We have further suggested that accelerated rates of translation of a selected CAT region potentially allowed the particular part of the polypeptide chain to appear earlier in time during translation and this might have led to an affected interaction of the extruded polypeptide region with the preceding one, which (we speculated) was not yet properly folded due to a lack of time [108]. This was one of the first experimental confirmations of the hypothesis stating that kinetics of protein translation can influence the in vivo protein folding pathway. However, the idea was yet not well accepted then.

The next breakthrough came 8 years later. In 2007, Chava Kimchi-Sarfaty, Michael Gottesman and their colleagues showed that substrate specificity of P-glycoprotein, the product of the multidrug resistance $1 \left(\text{MDR1} \right)$ gene, is altered by synonymous single nucleotide polymorphisms (SNPs) presumed to be silent [116]. The authors carefully investigated ex vivo (in transfected cells) the effect of naturally occurring polymorphic MDR1 variants on P-gp mRNA translation and protein expression, activity of the protein and its substrate specificity. They have concluded that a single synonymous mutation (C3435T; Ile-ATC>ATT, changing a relatively frequent ATC codon into a more rare ATT and, therefore, presumably affecting translation elongation rates in this region, although this was not demonstrated directly) can alter P-gp conformation and protein activity/substrate specificity [116]. This study was of immense importance as it for the first time demonstrated that naturally occurring synonymous/silent SNPs can lead to the synthesis of the protein product with the same amino acid sequence but different structural and functional properties. It also explained differences (observed previously in several clinical studies) in P-gp pharmacokinetics in individuals carrying this silent SNP [116].

The significance of synonymous codon usage for protein folding were highlighted by a number of subsequent studies showing that synonymous codon substitutions can affect proteins' sensitivity to limited proteolysis [117,118], phosphorylation profiles [118], spectroscopic properties [119], aggregation propensity [119-121] and specific activity [122], which ultimately can cause diseases [123-127]. Synonymous codon choice has been also suggested to affect efficient interaction of nascent polypeptides with the signal recognition particle [93], thus affecting protein secretion.

Of special interest is the study performed by Patricia Clark and colleagues [119], who took advantage of the so-called bimolecular fluorescence complementation (BiFC) assay [129, 130] to design a fluorescent protein consisting of three half-domains, where the N-

(yellow) and C-terminal (cyan) half-domains compete each other to interact with the central half-domain [119]. The outcome of this competition determined the fluorescence properties of the resulting folded structure. Using a cellular expression system and monitoring FRET between the YFP/CFP labels, they demonstrated that the rate at which a nascent protein emerges from the ribosome (which was affected by synonymous codon substitutions at the inter-half-domain linker) can specify the final folded conformation of a protein [119].

The studies above, had however a number of drawbacks. As a rule, they investigated just only one aspect of the hypothesis and predominantly looked at the properties of the polypeptide chains released from the ribosome. Analyses of the released polypeptide chains in all the above experiments were done using indirect (e.g. specific activity, limited proteolysis), rather than direct (NMR, X-Ray) structure probing approaches. Also, few studies attempted to investigate the structure of ribosome bound nascent chains produced from the silently mutated mRNA in comparison with the wild-type. Changes in the kinetics of protein synthesis as a result of synonymous mutations were not usually simultaneously monitored in these studies and potential effects of miscoding (that could potentially arise from synonymous codon changes) were frequently neglected. Finally, none of the studies employed real-time measurements to demonstrate that synonymous mutations may not only affect kinetics of protein synthesis, but they, at the same time, may also affect the real-time kinetics of co-translational protein folding.

Recently, we have filled in the gaps above and in collaboration with Harald Schwalbe from the Johann Wolfgang Goethe-Universität in Frankfurt, Germany and Marina V. Rodnina from the Max Planck Institute for Biophysical Chemistry in Goettingen, Germany executed a study, which addressed all these outstanding questions [111].

To investigate how differential usage of synonymous codons affects translation kinetics, coand post-translational folding, and protein conformation and stability, we analyzed in vivo expression of the recombinant bovine eye lens protein gamma-B crystallin in *Escherichia* coli cells and in vitro in a completely reconstituted high-performance translation system from E. coli. We have chosen gamma-B-crystallin, because previously we have shown that translation of this two-domain protein is a non-uniform process [68]. We also suggested that the codon usage and translation rates in gamma-B-crystallin are optimized to tune the synthesis and folding of this protein in the cell [68]; however the direct experimental evidence in support of this suggestion was lacking. We therefore designed two variants of the mRNA coding for gamma-B crystallin, one with the codon usage that would be optimal for protein translation in E. coli (with an mRNA codon distribution (codon usage profile) similar to that found in B. taurus, which was expected to result in more natural translation kinetics) and the other with unaltered codon composition un-optimal for translation in E . coli [111].

Our analysis of the effects of synonymous codon choice on the translation of gamma-B crystallin mRNAs showed that codon choice alters local and global translation rates and results in the formation of alternative conformations of the protein [111)]. We showed using real time measurements, which employed fluorescence and FRET, that kinetics of synthesis and co-translational folding of gamma-B crystallin is indeed altered by

synonymous codon substitutions. Moreover, for the first time, we detected considerable structural heterogeneity of the purified, mature synonymous gamma-B crystallin protein variants by using direct structure elucidation approach (2D NMR). We concluded that the synonymous polymorphisms altered the distribution of populations within the landscape of accessible protein conformations both on the ribosome and after chain's release [111]. Importantly, we have carefully verified by using Mass Spectrometry and Microsequencing that synonymous gamma-B crystallin variants led to the synthesis of the polypeptide chains with identical amino acid sequence. Together our results provided a strong support to the hypothesis and showed that synonymous codons may indeed serve as a secondary code for protein folding in the cell [111]. We thus concluded that codon usage specifies a unique translation kinetics that affects the partitioning of the folding intermediates both on the ribosome and after chain's release and that non-natural codon usage and translation kinetics can result in a kinetically trapped folding intermediates. These intermediates can be converted, with (or without) the help of molecular chaperones (co- or post-translationally), to the native protein state through reshuffling reactions. However, such kinetically trapped intermediates could also remain stable and drive the overall folding into a non-native and/or aggregation-prone state. Non-productive, trapped species could be also degraded (co- or post-translationally) (Fig. 3).

Importantly, non-productive folding arising due to altered synonymous codon usage can also lead to a disease [123-127]. In collaboration with Chava Kimchi-Sarfaty, we have recently demonstrated that a synonymous mutation, c.459G>A (GTG>GTA, p.Val153Val or Val107Val (Val107 is the amino acid number after the prepro-petide cleavage on secretion)) that has been previously identified in $F9$ gene (encoding blood coagulation factor FIX) in patients with mild haemophilia B [128], alters FIX synthesis and affects its conformation resulting in decreased extracellular protein level [127]. As such, we were able to determine the pathogenic basis for a single synonymous mutation in the $F9$ gene associated with haemophilia B [127]. This case remained a mystery for about 10 years, since its discovery in 2008 [128], as back at a time it couldn't be explained by altered mRNA properties (mRNA levels, splicing/exon skipping or retention of introns, or stability), a common cause of many diseases associated with synonymous mutations [125, 126].

CONCLUSIONS AND FUTURE PERSPECTIVES

The work on the effects of synonymous codon usage undoubtedly deepens our understanding of protein folding in the cell. Importantly, it, indeed, helps to explain cases of genetic diseases linked to synonymous mutations; cases which couldn't be explained by previously known effects of synonymous mutations on mRNA splicing and/or mRNA stability [125, 126]. These studies further brought about increased awareness of the scientific community to the impact of the synonymous mutations on gene function and phenotype linked to protein folding and stability [123-127]. Finally, this work gave a novel tool to upscale the production of functionally active recombinant proteins and provided explanation for a long-standing paradigm, revealing that approaches involving substitution of the majority of infrequently used codons with synonymous frequently used ones (targeted to inflate CAI), which have been widely used for optimization of heterologous and homologous protein production, frequently yielded biologically inactive insoluble protein

aggregates [54]. Our work made it clear that maximizing the speed and output of translation may put conflicting demands on the protein synthesis machinery, resulting in improper protein folding.

However, much work has yet to be done. There is yet a limited understanding, of how exactly the structure of co-translational folding intermediates is affected by the synonymous mutations. Novel methods, such as single molecule FRET and/or time-resolved cryo-EM [131] may potentially allow improved analysis of the effects of synonymous mutations on the appearance and conformation of co-translational folding intermediates. However, application of time-resolved cryo-EM for analysis of nascent chain dynamics remains challenging at the present time and FRET per se will not allow direct visualization of nascent chain structures.

Nevertheless, our improved understanding of the impact of synonymous codon usage on protein folding strongly supports the view that synonymous codon usage serves as a guide for co-translational protein folding in the cell (Fig. 3).

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 (b)

MetThrGlyLysSerSerVal...

Figure 1. Genetic code redundancy and non-uniform/non-random codon utilization shape codon usage bias and govern non-uniform translation.

(a) Codon usage bias in Escherichia coli (EC), Saccharomyces cerevisiae (SC) and Homo sapiens (HS) ([https://www.kazusa.or.jp/codon/\)](https://www.kazusa.or.jp/codon/). These examples reveal substantial differences in usage of codons between the three species. Frequency per thousand codons is shown. (b) Preferentially used (frequent) codons are translated faster than infrequently used (rare) codons due to the more ready availability (during translation) of the corresponding frequent cognate tRNAs. Rare codons, as a rule, lead to a substantial increase in ribosome residence time due to an increased waiting period of the ribosome for a cognate tRNA. A simplified scheme is shown (omitting the elongation factor 1A, which promotes the GTP-dependent binding of aminoacyl-tRNA to the A-site of ribosomes during protein biosynthesis). Colors of tRNAs/codons correspond to the differential frequency of their usage.

Figure 2. Rare codon clusters are often non-randomly and strategically placed within mRNAs. Top – codon usage profile for Bovine (*Bos taurus*) β B2 crystallin revealing an extended cluster of rare codons partially encoding the domain linker and the adjacent downstream region of the protein structure. Bottom – backbone/cartoon structure of the βB2 crystallin (PDB 2BB2). The N-terminal domain is in blue, the C-terminal domain is in yellow and a portion of the linker connecting the two domains is shown in gray. Positions of Pro80 and Lys89 at the beginning and the end of the linker peptide connecting the domains are indicated; Asn95 marks the end of the first β-structure in the βB2 C-terminal domain.

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Figure 3. Synonymous codon usage alters kinetics of protein translation and can direct cotranslational folding towards different protein conformation(s).

A model illustrating the influence of codon usage/translation kinetics on the final conformation of the synthesized protein. (a) Natural (native) kinetics of translation leads to the efficient formation of the native structure through the number of productive co-translational intermediates. (b) Altered codon usage/translation kinetics might create kinetically trapped intermediates. These intermediates might then be converted, with (or without) the help of molecular chaperones (co- or post-translationally), to the native protein through reshuffling reactions. However, such kinetically trapped intermediates could also remain stable and drive the overall folding into a non-native and/or aggregation-prone state. Nonproductive, trapped species could be also degraded (co- or post-translationally). Various end-points of non-productive folding arising due to altered synonymous codon usage can lead to a disease.