



Recoding the Genetic Code with Selenocysteine**

Markus J. Bröcker, Joanne M. L. Ho, George M. Church, Dieter Söll,* and Patrick O'Donoghue*

Abstract: Selenocysteine (Sec) is naturally incorporated into proteins by recoding the stop codon UGA. Sec is not hardwired to UGA, as the Sec insertion machinery was found to be able to site-specifically incorporate Sec directed by 58 of the 64 codons. For 15 sense codons, complete conversion of the codon meaning from canonical amino acid (AA) to Sec was observed along with a tenfold increase in selenoprotein yield compared to Sec insertion at the three stop codons. This high-fidelity sense-codon recoding mechanism was demonstrated for *Escherichia coli* formate dehydrogenase and recombinant human thioredoxin reductase and confirmed by independent biochemical and biophysical methods. Although Sec insertion at UGA is known to compete against protein termination, it is surprising that the Sec machinery has the ability to outcompete abundant aminoacyl-tRNAs in decoding sense codons. The findings have implications for the process of translation and the information storage capacity of the biological cell.

Sense-codon recoding is supposed to be impossible. Indeed, the fact that a codon can have more than one meaning was a dogma-breaking finding^[1] and is one reason why the mechanism of selenocysteine (Sec) insertion into proteins provoked intense biochemical investigation over the last three decades. In addition, selenium is an essential micronutrient for humans.^[2] Selenium in proteins is found in the form of Sec in enzymes that maintain the cell's redox balance, defending the cell against reactive oxygen species. Diseases involving Sec biosynthesis or selenoprotein malfunction have only recently surfaced because defects in these pathways are devastatingly detrimental to proper neuronal function and development.^[3]

The promise of sense-codon recoding is being actively pursued as a means to further expand the genetic code and synthesize proteins with multiple noncanonical amino acids (ncAAs).^[4] Nature expanded the genetic code with two amino

acids (Sec and pyrrolysine) by recoding or reassignment of stop codons.^[5] Thus, an efficient, naturally evolved machinery already exists that directs recoding of particular UGA stop codons to Sec (Scheme 1 A). Although Sec is not found in all organisms (notably lacking in plants, fungi, and most Archaea), the 21st amino acid is genetically encoded in all three domains of life. Sec is biosynthesized on its tRNA,^[6] and translational recoding of UGA requires the resulting Sec-tRNA^{Sec} product, a specialized elongation factor (SelB in *E. coli*), and a downstream mRNA hairpin motif known as the Sec insertion sequence (SECIS).

We investigated the possibility to site-specifically reassign multiple sense codons using the Sec machinery. Previous attempts to encode Sec with the Leu UUA codon^[7] and the Trp UGG codon^[8] produced lower selenoprotein yields compared to UGA-encoded Sec and suggested significant canonical AA contamination. To systematically investigate the recoding capacity of the Sec machinery, we created a library of *E. coli* formate dehydrogenase (FDH_H) mutants. Each gene variant had one of the 64 codons encoding a critical Sec residue (Sec140). In nature, FDH_H is part of the membrane-associated formate hydrogen lyase (FHL) complex that decomposes formate to H₂ and CO₂ under fermentative conditions. The FHL complex shuttles electrons from formate to hydrogenase 3 which reduces protons to hydrogen molecules during anaerobic respiration.^[9] Because the enzymatic activity of FDH_H is dependent on the active-site residue Sec140, which coordinates an active-site molybdopterin cofactor,^[10] sense-codon recoding is easily monitored in vivo and in vitro using the artificial electron acceptor benzyl viologen (BV). Active FDH_H reduces BV to give a purple color^[11] that is clearly visualized in living cells or monitored spectroscopically with purified FDH_H.

Each FDH_H gene variant (*fdhF*₁₄₀NNN) was co-expressed with *selA*, *selB*, and a *selC* mutant (expressing tRNA^{Sec}_{NNN})

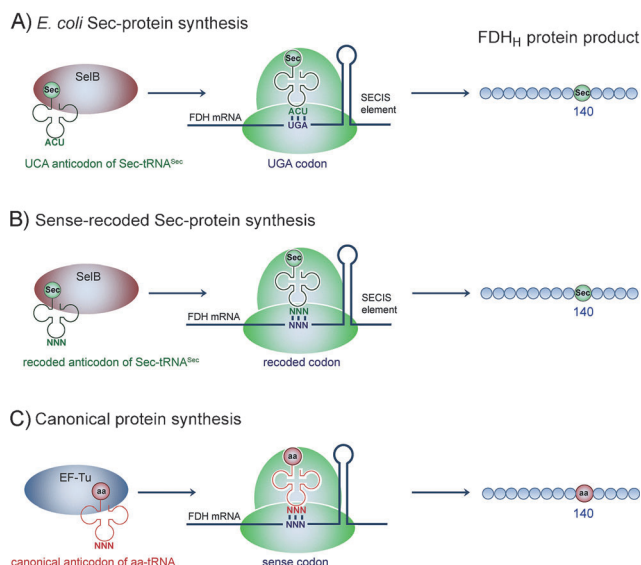
[*] Dr. M. J. Bröcker, Prof. D. Söll
Department of Molecular Biophysics & Biochemistry
Yale University, New Haven, CT 06520 (USA)
E-mail: dieter.soll@yale.edu
Prof. D. Söll
Department of Chemistry, Yale University
New Haven, CT 06520 (USA)
J. M. L. Ho, Prof. G. M. Church
Department of Genetics, Harvard Medical School
Boston, MA 02115 (USA)
Prof. P. O'Donoghue
Departments of Biochemistry and Chemistry
Western University, London, ON N6A 5C1 (Canada)
E-mail: patrick.odonoghue@uwo.ca

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Scheme 1. Codon recoding with Sec. A) Synthesis of the *E. coli* Sec-containing formate dehydrogenase (FDH_H) from the wild-type *fdhF* gene requires recoding the UGA stop codon at position 140 to Sec by Sec-tRNA^{Sec}, elongation factor SelB, and the SECIS mRNA hairpin. Replacing the UGA with any of the 64 NNN triplets at codon 140 in *fdhF* and co-expressing a cognate Sec-tRNA^{Sec}_{NNN} is expected to yield SelB and SECIS-dependent sense-codon-recoded Sec-containing FDH_H (B) that may compete with canonical AA insertion (C) directed by native aminoacyl-tRNAs and EF-Tu dependent protein synthesis. In this case (C), SECIS is still present in the *fdhF* mRNA, but SECIS is not expected to interact with EF-Tu directed decoding.

with the cognate anticodon (Scheme 1). We anticipated that these constructs could give rise to two different protein products. Sec-tRNA^{Sec}_{NNN} and SelB should compete with canonical aminoacyl-tRNA_{NNN} (AA-tRNA_{NNN}) and EF-Tu, which could give rise to Sec-containing FDH_H (Scheme 1 B), canonical AA containing FDH_H (Scheme 1 C), or a mixture of both protein species. The plasmid-borne FDH_H and *sel* genes complemented an *E. coli* $\Delta selA \Delta selB \Delta fdhF$ deletion strain (MH5)^[12] that is otherwise unable to produce selenoproteins. Quite unexpectedly, the in vivo assay shows that the Sec recoding machinery successfully alters the meaning of all three stop codons and nearly all sense codons to Sec (Figure 1). For 58 of the possible 64 codons, the coloration obtained by Sec-dependent BV reduction is as intense as that for the UGA (wild-type) Sec codon. Although this assay is qualitative, it demonstrates for the first time that most codons are recodable. At least one codon for each of the 20 canonical amino acids is recodable to Sec; so no particular AA-tRNA species is able to completely outcompete Sec insertion. For example, the meaning of all six Leu codons, five out of six Ser codons, and four out of six Arg codons was re-assigned to Sec.

Only six codons proved refractory to Sec insertion as indicated by the lack of FDH_H activity. Strikingly, four of these codons (CGA, AGU, AGG, and GGC) were found in the NGN boxes of the codon table. SDS gels and western blot analysis (Figures S1 and S2) show that full-length FDH_H of 80 kDa is produced for all codons except the arginine codons

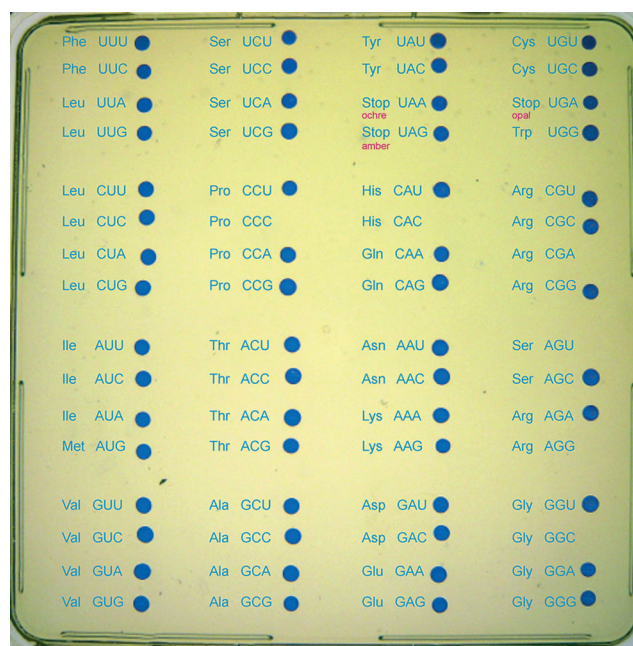


Figure 1. Recoding the genetic code with Sec. The canonical genetic code table is overlaid on a single agar plate spotted with 64 *E. coli* FDH_H variants. In each case, an *E. coli* $\Delta selA \Delta selB \Delta fdhF$ deletion parent strain was complemented with *E. coli* *selA*, *selB*, and each of the 64 *fdhF*₁₄₀NNN codon mutants and tRNA^{Sec}_{NNN} variants with the respective cognate anticodon. The capacity of each strain to recode the indicated codon to Sec is evidenced by an in vivo BV reduction assay in which purple colored cells express active Sec-containing FDH_H.

AGG and CGA. These codons show a truncated FDH_H product at 15.5 kDa, suggesting that protein synthesis terminates at position 140. These data indicate that canonical AA-tRNA completely outcompetes Sec insertion (Scheme 1 C) at AGU, GGC, CAC, and CCC, but the meaning of AGG and CGA is converted to stop by an unknown mechanism.

The Sec machinery has the inherent capability to recode nearly all codons (Figure 1), but this in vivo assay does not provide quantitative information regarding the degree of recoding. Under anaerobic conditions, 64 N-terminal His-tagged FDH_H proteins were over-expressed from the *fdhF*₁₄₀NNN gene variants and purified (Figures S1 and S2). The Sec insertion ratio was quantitated by comparison of the initial velocity of BV reduction in sense-codon-recoded versus wild-type FDH_H variants in vitro (Table S1). Although the same concentration of purified FDH_H protein is used in each assay, the specific activity values differ, which indicates the amount of active (i.e., Sec140) FDH_H versus the amount of inactive (i.e., canonical AA140) FDH_H protein in each preparation. The assays confirmed Sec incorporation in FDH_H for 58 codons with relative FDH_H-specific activities ranging from 12–100 % of the wild-type. Selenoprotein yield was greater for most of the recoded sense codons compared to Sec insertion at UGA or the other stop codons, UAA and UAG (Table S2).

Plotting FDH_H protein yield versus specific activity (Figure 2A) revealed six distinct types of recoding behavior.

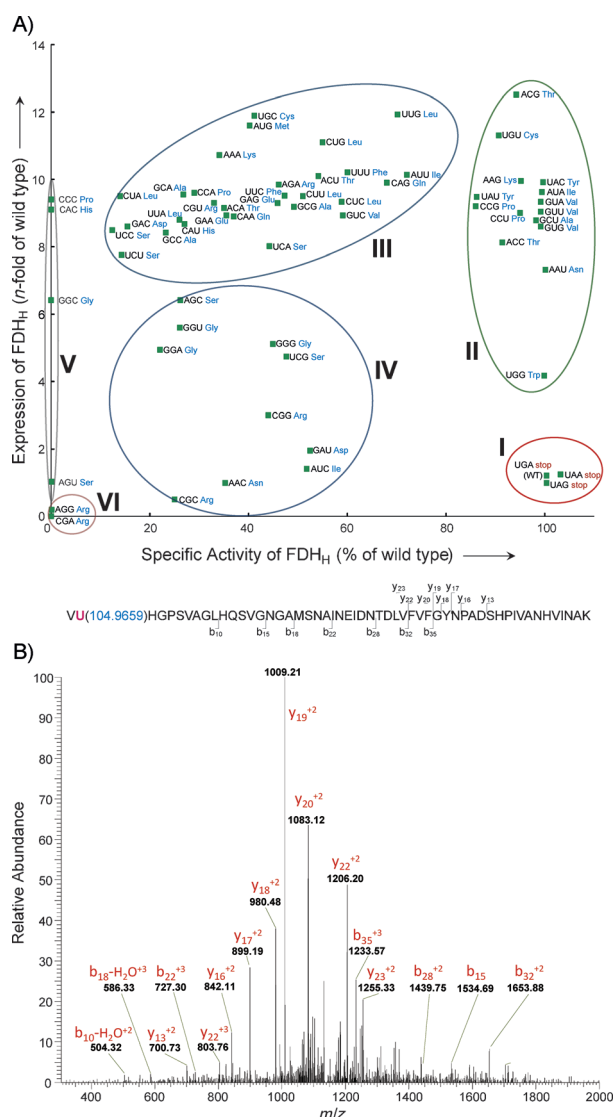


Figure 2. Quantitation of codon recoding with Sec. A) The relative yield of the 64 FDH_H $_{140}\text{NNN}$ variants (wild-type FDH_H $_{140}\text{UGA}=1$) is plotted versus specific activity (relative to wild-type FDH_H $_{140}\text{UGA}=100\%$). Activity, yield values, and standard deviations are shown in Table S1. Different levels of specific activity observed for the 64 FDH_H variants are the result of the partial incorporation of the respective canonical AAs (see Table S2). B) MS/MS identification of UAC-encoded Sec. The spectrum of the Sec-containing tryptic peptide, peptide sequence, and position of the observed b- and y-ions are indicated (see also Table S2). Sec (U) was identified by the MASCOT protein identification software as a Se-IAN-Cys modification of Cys; selenium alkylation ($\text{Se-CH}_2\text{CONH}_2$).

Including the three stop codons (Figure 2, group I), 18 codons led to fully active FDH_H (ca. 100% specific activity), indicating complete conversion of the codon meaning to Sec (groups I and II). For the two other stop codon variants ($\text{fdhF}_{140}\text{UAG}$ and $\text{fdhF}_{140}\text{UAA}$), specific activity and protein yield was nearly identical to UGA-encoded Sec. In contrast, 15 codons (group II) are also recoded at high fidelity, but they produce significantly more pure selenoprotein compared to UGA-encoded Sec. Relative FDH_H specific activities for codon group I were in the range of 86–100% with 7–13-fold

more Sec-containing FDH_H production. FDH_H variant $_{140}\text{ACG}$ (a Thr codon) showed the highest expression level, while the highest specific activity (i.e., highest-fidelity Sec recoding) was observed for the Tyr codon $_{140}\text{UAC}$. MS/MS analysis confirmed Sec incorporation at position 140 for the Tyr UAC codon (Figure 2B), and no Tyr-containing peptide was identified (Table S2 A,B).

For a total of 11 codons, MS/MS analysis identified only the Sec-containing peptide (Table S2, Figure S3). To independently confirm this high-fidelity Sec insertion, purified FDH_H variants were analyzed by inductively coupled plasma (ICP)-MS, which provides an accurate estimation of the Se content of the protein sample. These data show 99.9% Sec insertion at UGA in FDH_H and confirm complete recoding for UAC (98.3% Sec) and slightly reduced Sec insertion at UAU (79.3%). The ICP-MS data correlate well with the FDH_H specific activities for all other samples tested (Tables S1 and S3). Sense-codon recoding with Sec requires expression of all the normal Sec insertion components including an intact SECIS, SelB, and Sec-tRNA^{Sec} formation on a tRNA with a “sense-cognate” anticodon (see the Supporting Information). Sec-tRNA^{Sec}_{UCA} (wild-type) is unable to decode $_{140}\text{UAC}$ or $_{140}\text{UAU}$ with Sec. In the absence of any component, approximately the same amount of full-length FDH_H is produced, but it is totally inactive and contains Tyr₁₄₀ (Figure S4).

Forty codons displayed ambiguous decoding at position 140 of FDH_H . Of these, 30 codons (group III) enabled partial recoding (12–72% Sec) as indicated by specific BV reduction activity (Figure 2, Table S1). These codons produce two protein variants, one containing Sec and one containing a canonical AA at position 140, which results in full-length but inactive protein (Figures S1 and S2). MS/MS confirmed ambiguous decoding for the Glu codon GAA, and for the three codons (Phe UUC, Leu CUC, Ala GCC) Sec insertion at position 140 was clearly identified (Figure S3). These variants displayed markedly increased FDH_H protein yields (8–12-fold) compared to UGA-encoded Sec. The ten group IV codons show 22–52% Sec incorporation and smaller increases in FDH_H yield (1–6-fold). Confirming the initial result (Figure 1), four codons were refractory to Sec-recoding (group V) and two Arg codons (CGA and AGG, group VI) produced little protein (Figure 2, Table S1), all of which is truncated at position 140 (Figure S1).

To test the generality of this novel sense-codon-recoding mechanism, we expressed a recombinant human selenoprotein, thioredoxin reductase (TrxR), with native UGA-encoded or UAC-encoded Sec (Figure 3A). Thioredoxin reductase is an essential selenoprotein involved in maintaining proper redox balance in human cells. The role of Se is also important in this enzyme as the Sec-containing TrxR is more resistant to oxidative inactivation than a Cys orthologue.^[13] Because of its ability to enhance cell survival by defending the cell against reactive oxygen species, TrxR1 over-expression is associated with numerous cancer cell lines, oncogenesis, and metastasis in lymph nodes.^[14] TrxR is an emerging target for novel cancer therapeutics.^[15] For these reasons, the ability to more efficiently produce TrxR is a significant advance for the field of redox biology, which is expected to impact future

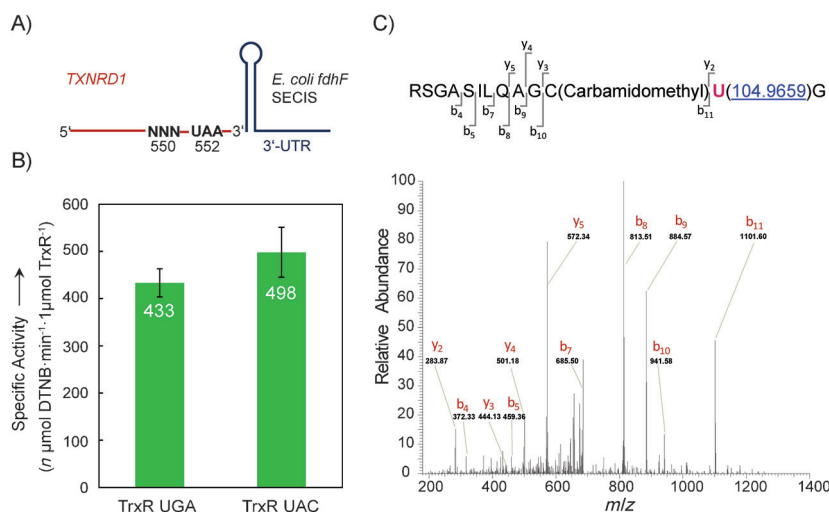


Figure 3. Recoding the Tyr codon UAC to Sec in human TrxR1. A) Schematic representation of the expression construct for human TXNRD1. An *E. coli fdhF* SECIS element is attached directly after the UAA codon of TXNRD1 in the 3'-untranslated region (UTR). B) In vitro activity assay of recombinant human TrxR₅₅₀UGA and TrxR₅₅₀UAC co-expressed with tRNA^{Sec}_{UCA} and tRNA^{Sec}_{GUA}, respectively. Error bars show standard deviations of quadruplicate experiments. C) MS/MS identification of Sec incorporation into TrxR₅₅₀UAC. The spectrum of the Sec-containing peptide, the peptide sequence and the positions of the observed b- and y-ions are indicated. Sec was identified by MASCOT as a Se-IAN-Cys modification of Cys; selenium alkylation (Se-CH₂-CONH₂).

therapeutic development. Similar to our observations of UAC-encoded Sec in FDH_H, TrxR₅₅₀UAC (4.1 mg L⁻¹ culture) leads to fourfold higher yield of pure selenoprotein compared to the UGA-encoded selenoenzyme (0.9 mg L⁻¹ culture). The specific activities are indistinguishable for TrxR produced from these two constructs (Figure 3B). No Tyr contamination was detected by MS/MS analysis (Figure 3C), and ICP-MS analysis independently confirmed 98.5% Sec incorporation in response to UGA and 95.1% Sec insertion for UAC in TrxR (Table S3).

It remains unclear why certain codons are more “recodable” than others. It is reasonable to assume that codon usage might correlate with recodability, because translation is believed to be less efficient with lower abundant codons. The intracellular concentration of AA-tRNA^[6] is another likely explanation. Perhaps codons read by less abundant AA-tRNA species are more easily recoded. Despite the logic of these ideas, there is no general correlation between recodability and codon usage or AA-tRNA concentration (Figure S5). There are potential complicating factors; perhaps the Sec-tRNA^{Sec}_{NNN} species are not decoded on the ribosome with equal efficiency due to compatibility of the tRNA body and anticodon loop. There is also the question of whether the nature of the mRNA codon (position 140) affects the structural integrity of the SECIS element. Furthermore, differences between AA-tRNA levels may not be sufficient enough to fundamentally alter competition with Sec-tRNA^{Sec}_{NNN}. Based on crystal structures, a model of the ribosome-bound Sec-decoding complex (i.e., ribosome, SelB, Sec-tRNA^{Sec}, mRNA with SECIS) indicates extensive interactions between SelB and the 30S subunit in regions that are known to alter the accuracy of mRNA decoding.^[17] Kinetic

and structural studies of sense-codon decoding with Sec will help define the mechanism discovered here.

In the normal Sec-decoding situation, the release factor (RF2 in *E. coli*) competes with Sec insertion,^[18] leading to truncated FDH_H (Figure S4). Encoding Sec with sense codons escapes competition with RFs, which leads to the significantly enhanced selenoprotein yield for all but ten codons. It is unclear why nature selected UGA as the “Sec” codon when most other codons lead to more selenoprotein production. Perhaps the scarcity of Se in the environment induced organisms to naturally limit the level of selenoprotein synthesis with RF competition. RF interaction may, nevertheless, affect the recodability of certain sense codons. Due to its similarity with UGA, UGG is a hotspot for premature termination by RF2,^[19] which may explain why this codon produces less Sec-containing FDH_H than all other group II codons (Figure 2).

Encoding Sec with sense codons will become a useful method for efficiently producing selenoproteins; yet this discovery opens the door to recode many codons, not

only with Sec, but possibly also with a diverse array of ncAAs of biomedical and biotechnological interest. The human proteome contains many essential modified proteins that arise by posttranslational modification (PTM), and mismodified proteins may lead to defects in cellular signaling and protein aggregates that form the molecular basis for diseases. In order to biochemically define the role of PTMs and to produce therapeutic agents that specifically target mismodified proteins, there is an urgent need to synthesize proteins containing several different, distinct ncAAs. Progress in the last decade allowed site-specific incorporation of one or two ncAAs into recombinant proteins by reassignment of non-sense codons.^[4b] In vitro protein synthesis methods were recently enhanced by creating dual-meaning initiation and sense codons to synthesize potential chemotherapeutic peptides with two ncAAs,^[20] and amino acid starvation methods permitted production of proteins with three ncAAs for biological imaging applications.^[21] Furthermore, unlike the Sec machinery that only recodes codons associated with SECIS, current genetic code expansion methods are not site-specific because they lead to global reassignment of stop codons; this contaminates the natural proteome with ncAAs, resulting in growth defects.^[22]

In order to synthesize proteins with more than 23 amino acids, additional “recodable” codons are needed.^[23] We anticipate that mutagenesis of the Sec insertion machinery will enable protein synthesis with multiple ncAAs, which will require engineering SelB, SECIS, and also orthogonal tRNA synthetases to acylate tRNA^{Sec} and tRNA^{Sec}-like molecules with ncAAs. The fact that SepCysS forms Cys-tRNA^{Sec}, which actively decodes UGA with Cys instead of Sec, is an encouraging example.^[24]

The fact that the Sec insertion machinery has the intrinsic capability to alter the meaning of 58 of the 64 codons indicates that the biological cell will be able to genetically encode far more amino acids than previously recognized. Although the Sec insertion machinery recodes some codons poorly, it is possible that the recoding fidelity may be enhanced through selection. Given that certain codons are recodable at high fidelity, it is conceivable that in nature Sec is encoded by codons other than UGA, suggesting the existence of unknown selenoproteins. Indeed, SECIS-like structures that are not associated with in-frame UGA codons have been identified bioinformatically and dismissed as evolutionary remnants.^[25] If other RNA signals, like SECIS, exist in nature or if they can be engineered in the laboratory, there is potentially no limit to the number of amino acids that the cell can encode and no reason to stop at 20 or even 23.

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- [1] F. Zinoni, A. Birkmann, W. Leinfelder, A. Böck, *Proc. Natl. Acad. Sci. USA* **1987**, *84*, 3156–3160.
- [2] M. P. Rayman, *Lancet* **2000**, *356*, 233–241.
- [3] a) O. Agamy, B. B. Zeev, D. Lev, B. Marcus, D. Fine, D. Su, G. Narkis, R. Ofir, C. Hoffmann, E. Leshinsky-Silver, H. Flusser, S. Sivan, D. Söll, T. Lerman-Sagie, O. S. Birk, *Am. J. Hum. Genet.* **2010**, *87*, 538–544; b) S. Seeher, Y. Mahdi, U. Schweizer, *Curr. Protein Pept. Sci.* **2012**, *13*, 337–346.
- [4] a) R. Krishnakumar, L. Prat, H. R. Aerni, J. Ling, C. Merryman, J. I. Glass, J. Rinehart, D. Söll, *ChemBioChem* **2013**, *14*, 1967–1972; b) P. O'Donoghue, J. Ling, Y. S. Wang, D. Söll, *Nat. Chem. Biol.* **2013**, *9*, 594–598.
- [5] A. Ambrogelly, S. Palioura, D. Söll, *Nat. Chem. Biol.* **2007**, *3*, 29–35.
- [6] J. Yuan, P. O'Donoghue, A. Ambrogelly, S. Gundllapalli, R. L. Sherrer, S. Palioura, M. Simonovic, D. Söll, *FEBS Lett.* **2010**, *584*, 342–349.
- [7] M. J. Berry, J. W. Harney, T. Ohama, D. L. Hatfield, *Nucleic Acids Res.* **1994**, *22*, 3753–3759.
- [8] J. Xu, V. Croitoru, D. Rutishauser, Q. Cheng, E. S. Arnér, *Nucleic Acids Res.* **2013**, DOI: 10.1093/nar/gkt764.
- [9] G. Sawers, *Antonie van Leeuwenhoek* **1994**, *66*, 57–88.
- [10] J. C. Boyington, V. N. Gladyshev, S. V. Khangulov, T. C. Stadtman, P. D. Sun, *Science* **1997**, *275*, 1305–1308.
- [11] M. J. Axley, D. A. Grahame, T. C. Stadtman, *J. Biol. Chem.* **1990**, *265*, 18213–18218.
- [12] C. Aldag, M. J. Bröcker, M. J. Hohn, L. Prat, G. Hammond, A. Plummer, D. Söll, *Angew. Chem.* **2013**, *125*, 1481–1485; *Angew. Chem. Int. Ed.* **2013**, *52*, 1441–1445.
- [13] G. W. Snider, E. Ruggles, N. Khan, R. J. Hondal, *Biochemistry* **2013**, *52*, 5472–5481.
- [14] Y. C. Chen, K. S. Prabhu, A. M. Mastro, *Nutrients* **2013**, *5*, 1149–1168.
- [15] J. Lu, A. Holmgren, *Antioxid. Redox Signaling* **2012**, *17*, 1738–1747.
- [16] a) H. Dong, L. Nilsson, C. G. Kurland, *J. Mol. Biol.* **1996**, *260*, 649–663; b) K. A. Dittmar, M. A. Sorensen, J. Elf, M. Ehrenberg, T. Pan, *EMBO Rep.* **2005**, *6*, 151–157.
- [17] S. Yoshizawa, L. Rasubala, T. Ose, D. Kohda, D. Fourmy, K. Maenaka, *Nat. Struct. Mol. Biol.* **2005**, *12*, 198–203.
- [18] J. B. Mansell, D. Guevremont, E. S. Poole, W. P. Tate, *EMBO J.* **2001**, *20*, 7284–7293.
- [19] a) D. V. Freistoffer, M. Kwiatkowski, R. H. Buckingham, M. Ehrenberg, *Proc. Natl. Acad. Sci. USA* **2000**, *97*, 2046–2051; b) J. Sund, M. Ander, J. Aqvist, *Nature* **2010**, *465*, 947–950.
- [20] Y. Goto, M. Iseki, A. Hitomi, H. Murakami, H. Suga, *ACS Chem. Biol.* **2013**, DOI: 10.1021/cb400549p.
- [21] S. Lepthien, L. Merkel, N. Budisa, *Angew. Chem.* **2010**, *122*, 5576–5581; *Angew. Chem. Int. Ed.* **2010**, *49*, 5446–5450.
- [22] I. U. Heinemann, A. J. Rovner, H. R. Aerni, S. Rogulina, L. Cheng, W. Olds, J. T. Fischer, D. Söll, F. J. Isaacs, J. Rinehart, *FEBS Lett.* **2012**, *586*, 3716–3722.
- [23] M. J. Lajoie, A. J. Rovner, D. B. Goodman, H. R. Aerni, A. D. Haimovich, G. Kuznetsov, J. A. Mercer, H. H. Wang, P. A. Carr, J. A. Mosberg, N. Rohland, P. G. Schultz, J. M. Jacobson, J. Rinehart, G. M. Church, F. J. Isaacs, *Science* **2013**, *342*, 357–360.
- [24] J. Yuan, M. J. Hohn, R. L. Sherrer, S. Palioura, D. Su, D. Söll, *FEBS Lett.* **2010**, *584*, 2857–2861.
- [25] Y. Zhang, H. Romero, G. Salinas, V. N. Gladyshev, *Genome Biol.* **2006**, *7*, R94.