

# FEATURE ARTICLE

# Overcoming the codon bias of *E. coli* for enhanced protein expression

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ost amino acids are encoded by more than one codon, and each organism carries its own bias in the usage of the 61 available amino acid codons. In each cell, the tRNA population closely reflects the codon bias of the mRNA population (1, 2). When the mRNA of heterologous target genes is overexpressed in E. coli, differences in codon usage can impede translation due to the demand for one or more tRNAs that may be rare or lacking in the population (3-5). Insufficient tRNA pools can lead to translational stalling, premature translation termination, translation frameshifting and amino acid misincorporation (4).

In practice with the pET System and other high-level *E. coli* expression systems, the presence of a small number of rare codons often does not severely depress target protein synthesis. However, heterologous protein expression can be very low when a gene encodes clusters of and/or numerous rare *E. coli* codons. The most severe effects on expression have been observed when multiple consecutive rare codons are near the N-terminus of a coding sequence (6).

#### Rare codons in *E. coli*

Examination of codon usage in all 4,290 *E. coli* genes reveals a number of codons that are underrepresented (Table 1). In particular, Arg codons AGA, AGG, and CGA, Ile codon AUA, and Leu codon CUA all represent less than 8% of their corresponding population of codons. The codon usage of abundantly expressed genes ("Class II",

Table 1) demonstrates a more extreme bias in which the aforementioned low-usage codons are avoided, and codons for Gly (GGA), Arg (CGG) and Pro (CCC) fall to *continued on page 2* 

| Table 1. Arg, Gly, Ile, Leu and Pro codon usage in <i>E. coli</i> |       |                       |                         |  |  |  |
|---|-------|-----------------------|-------------------------|--|--|--|
| amino<br>acid   | codon | fraction in all genes | fraction<br>in Class II |  |  |  |
| Arg   | AGG   | 0.022                 | 0.003                   |  |  |  |
| Arg   | AGA   | 0.039                 | 0.006                   |  |  |  |
| Arg   | CGG   | 0.098                 | 0.008                   |  |  |  |
| Arg   | CGA   | 0.065                 | 0.011                   |  |  |  |
| Arg   | CGU   | 0.378                 | 0.643                   |  |  |  |
| Arg   | CGC   | 0.398                 | 0.330                   |  |  |  |
| Gly   | GGG   | 0.151                 | 0.044                   |  |  |  |
| Gly   | GGA   | 0.109                 | 0.020                   |  |  |  |
| Gly   | GGU   | 0.337                 | 0.508                   |  |  |  |
| Gly   | GGC   | 0.403                 | 0.428                   |  |  |  |
| Ile   | AUA   | 0.073                 | 0.006                   |  |  |  |
| Ile   | AUU   | 0.507                 | 0.335                   |  |  |  |
| Ile   | AUC   | 0.420                 | 0.659                   |  |  |  |
| Leu   | UUG   | 0.129                 | 0.034                   |  |  |  |
| Leu   | UUA   | 0.131                 | 0.055                   |  |  |  |
| Leu   | CUG   | 0.496                 | 0.767                   |  |  |  |
| Leu   | CUA   | 0.037                 | 0.008                   |  |  |  |
| Leu   | CUU   | 0.104                 | 0.056                   |  |  |  |
| Leu   | CUC   | 0.104                 | 0.080                   |  |  |  |
| Pro   | CCG   | 0.525                 | 0.719                   |  |  |  |
| Pro   | CCA   | 0.191                 | 0.153                   |  |  |  |
| Pro   | CCU   | 0.159                 | 0.112                   |  |  |  |
| Pro   | CCC   | 0.124                 | 0.016                   |  |  |  |
|   |       |                       |                         |  |  |  |

Codon usage is expressed as the fraction of all possible codons for a given amino acid. "All genes" is the fraction represented in all 4,290 coding sequences in the *E. coli* genome (6). "Class II" is the fraction represented in 195 genes highly and continuously expressed during exponential growth (7).

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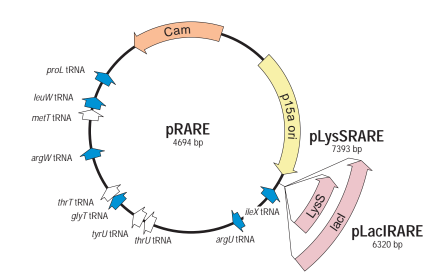
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less than 2% of their respective populations. Under typical growth conditions, target genes are expressed in *E. coli* at levels similar to (or exceeding) those for abundantly expressed endogenous genes. Thus, it is likely in many cases that the resident tRNA population available for target protein synthesis would more closely resemble that of the "Class II" genes in Table 1. Theoretically, modification of culture conditions (e.g. lowering the temperature, changing media composition, etc.) might shift the codon usage bias enough to alleviate some codon usage-based expression problems. However, it has been reported that the levels of most of the tRNA isoacceptors corresponding to rare codons remain unchanged at different growth rates (2). Translation problems similar to those caused by codon usage bias can also be created by high-level expression of proteins having an abundant amino acid. In these cases, expression may be improved by supplying the limiting amino acid in the culture medium (3).

#### Supplying the demand

As shown in Table 1, a subset of the codons for Arg, Ile, Gly, Leu, and Pro are very rarely used in highly expressed E. coli genes. Several laboratories have shown that expression yields of proteins whose genes contain rare codons can be dramatically improved when the cognate tRNA is increased within the host (8-10). tRNA levels can be elevated by increasing the copy number of the respective tRNA gene. This is typically accomplished by inserting the wild type tRNA gene on a multiple copy plasmid. The tRNA gene is either inserted into the expression vector itself or placed on a compatible plasmid. Early studies focused on the effect of increasing the gene dosage of argU (also known as dnaY, which encodes a tRNA that recognizes the AGG/AGA codons) on expression yield, plasmid stability and cell viability (9). The yield of human tissue plasminogen activator was increased approximately 10-fold (from 3% of total cell protein to 30%) in a strain that carried the *dnaY* gene on a compatible plasmid. Numerous subsequent studies also reported a substantial increase in protein yield when employing E. coli hosts with enhanced argU expression. Increasing the amount of other rare tRNAs has also been



#### Figure 1. Map of pRARE plasmid family

The basic structure of pRARE is indicated. pLysSRARE and pLacIRARE contain the genes encoding T7 lysozyme (LysS) and *lac* repressor (lacl), respectively. Also indicated are chloramphenicol resistance gene (Cam), origin of replication (p15a ori) and tRNA genes. tRNA genes corresponding to rare codons in *E. coli* are indicated in blue. pRARE is derived from pRIG (11).

shown to augment the yield and fidelity of heterologous proteins. tRNA genes for *ileX* (AUA), *leuW* (CUA), *proL* (CCC) or *glyT* (GGA) have all been used in this manner (reviewed in 3).

#### pRIG plasmid

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More recently, various combinations of rare tRNA genes have been assembled to optimize the expression of genes isolated from organisms with AT or GC rich genomes that have corresponding codon usage bias. One such assembly, the pRIG plasmid (11), encodes tRNA genes *argU*, *ileX* and *glyT* under their native promoters on a pACYC backbone, which carries the p15a origin of replication. The presence of pRIG in the host strain was shown to significantly enhance the expression of several genes derived from an AT-rich *Plasmodium* genome.

#### pRARE and the Rosetta<sup>™</sup> strains

To further extend the utility of pRIG for the expression of genes having rare E. coli codons, Novagen has added the *leuW* and proL tRNA genes to create pRARE (see Fig. 1). pRARE encodes tRNA genes for all of the "problematic" rarely used codons encoding Arg, Ile, Gly, Leu and Pro, except for Arg CGA/CGG. The entire RARE tRNA cassette was also added to the pLysS and pLacI plasmids (derived from pACYC184) to create pLysSRARE and pLacIRARE, respectively. The plasmids were transformed into various strains to create the Rosetta<sup>TM</sup> series of expression hosts (Table 2). These plasmids are compatible with Novagen's pET, pETBlue<sup>™</sup> and pTriEx<sup>TM</sup> expression vectors, and with expression vectors driven by other E. coli promoters (Table 3). These host strains are wellsuited to enhance protein expression from target genes containing rare E. coli codons that would otherwise impede translation.

#### Table 2. Rosetta host strains Antibiotic Derivation Strain Key Feature(s) Resistance Tuner™ BL21 lacYZ deletion, Rosetta Cam Rosetta(DE3) (B) Lacks lon and omp] Cam Rosetta(DE3)pLysS proteases Cam Rosetta(DE3)placI Cam RosettaBlue™ NovaBlue recA, endA, laclq Tet + Cam RosettaBlue(DE3) (K-12) High transformation Tet + Cam

| ettaBlue(DE3)pLysS<br>ettaBlue(DE3)pLacl                                   | ()                 | efficiency   | Tet + Cam<br>Tet + Cam   | yes<br>yes               |
|--|--------------------|--|--|--------------------------|
| etta-gami™<br>etta-gami(DE3)<br>etta-gami(DE3)pLysS<br>etta-gami(DE3)pLacI | Origami™<br>(K-12) | <i>trxBlgor</i> mutant, greatly facilitates cytoplasmic disulfide bond formation | Kan + Tet + Cam<br>Kan + Tet + Cam<br>Kan + Tet + Cam<br>Kan + Tet + Cam | yes<br>yes<br>yes<br>yes |

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yes

yes

yes

yes

yes

ves

| Table 3. Vector compatibility of Rosetta host |
|---|
| strains                                       |

| Strain  | Compatible vectors*                             |
|---|---|
| Rosetta™  | <i>E. coli</i> promoter                         |
| RosettaBlue™  | based vectors, e.g.                             |
| Rosetta-gami™   | <i>tac, trc</i> , T5, λ                         |
| Rosetta(DE3)<br>Rosetta(DE3)pLysS<br>RosettaBlue(DE3)<br>RosettaBlue(DE3)pLysS<br>Rosetta-gami(DE3)<br>Rosetta-gami(DE3)pLysS | T7 and T7 <i>lac</i> promoter based pET vectors |
| Rosetta(DE3)pLacl   | T7 <i>lac</i> promoter based                    |
| RosettaBlue(DE3)pLacl   | pETBlue™, pTriEx™                               |
| Rosetta-gami(DE3)pLacl  | vectors   |

All vectors must carry a Cole i-based origin of replication (e.g. pBR, pUC) and lack chloramphenicol selection. RosettaBlue and Rosetta-gami hosts require the use of vectors that also lack tetracycline resistance, and tetracycline plus kanamycin resistance, respectively.

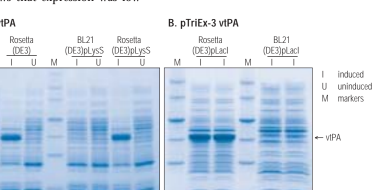
#### vtPA expression test

To evaluate the Rosetta host strains, a DNA fragment encoding vtPA ( $\Delta 6$ -175 deletion mutant of human tissue plasminogen activator) was cloned into the Nco I and Hind III sites of both pET-21d(+) and pTriEx<sup>TM</sup>-3 (12). Full length tPA was one of the original proteins whose expression was enhanced by additional copies of argUtRNA. The  $\Delta$ 6-175 vtPA construct encodes a 358 amino acid 40 kDa protein that contains 9 CCC, 8 AGA/AGG, 6 GGA, 6 CGG/CGA, 2 AUA, and 1 CUA for a total of 32 rare codons (8.9%). The pET construct was transformed into BL21(DE3), BL21(DE3)pLysS and the isogenic Rosetta strains, and the pTriEx plasmid was used with corresponding pLacI hosts. Fig. 2 shows an SDS-PAGE analysis of total cell extracts. In panel A, analysis of the pET constructs shows that expression was low

A. pET-21d(+) vtPA

Μ

(DE3)



#### Figure 2. Expression of △6-175 vtPA in different host strains

vtPA constructs in pET-21d(+) and pTriEx-3 were transformed into the indicated host strains. Cultures were grown at 37°C in LB + 0.5% glucose to an OD<sub>600</sub> of 0.6 to 1.0 and aliquots induced with 1 mM IPTG for 3 hours. Total cell protein samples were prepared and then analyzed by SDS-PAGE (4–20% gradient gels) and Coomassie blue staining. Panel A, pET-21d(+) vtPA; Panel B, pTriEx-3 vtPA. Duplicate induced cultures are shown in Panel B.

but detectable in BL21(DE3) and undetectable in BL21(DE3)pLysS. In contrast, vtPA was the major protein present in the induced samples from the Rosetta strains. A similar strong expression band was obtained when vtPA was expressed from pTriEx-3 in a Rosetta(DE3)pLacI host (panel B) but not in the host lacking the RARE cassette (BL21(DE3)pLacI).

In summary, Novagen's new Rosetta host strains allow researchers to alleviate low protein expression yields caused by codon usage bias. The strains circumvent the need to synthesize codon optimized genes, and enable rapid evaluation of codon usage as a possible strategy to enhance target protein expression. The strains are derivatives of popular *E. coli* expression hosts and versions are available for use with all of Novagen's *E. coli* and multisystem expression vectors. In addition, non- $\lambda$ DE3 lysogens are available for compatibility with other ColE1 origin, *E. coli* promoter based expression vectors.

#### REFERENCES

- 1. Ikemura, T. (1981) *J. Mol. Biol.* **146**, 1–21.
- Dong, H., Nilsson, L. and Kurland, C.G. (1996) *J. Mol. Biol.* 260, 649–663.
- 3. Kane, J.F. (1995) *Curr. Opin. Biotechnol.* **6**, 494–500.
- 4. Kurland, C. and Gallant, J. (1996) *Curr. Opin. Biotechnol.* **7**, 489–493.
- Goldman, E., Rosenberg, A.H., Zubay, G. and Studier, F.W. (1995) *J. Mol. Biol.* 245, 467–473.

- Nakamura, Y., Gojobori, T. and Ikemura, T. (2000) *Nucl. Acids Res.* 29, 292.
- Henaut, A. and Danchin, A. (1996) in *Escherichia coli and Salmonella typhimurium cellular and molecular biology, vol. 2,* pp. 2047–2066. Neidhardt, F., Curtiss III, R., Ingraham, J., Lin, E., Low, B., Magasanik, B., Reznikoff, W., Riley, M., Schaechter, M. and Umbarger, H. (eds.) American Society for Microbiology, Washington, DC
- Rosenberg, A.H., Goldman, E., Dunn, J.J., Studier, F.W. and Zubay, G. (1993) *J. Bacteriol.* **175**, 716–722.
- 9. Brinkmann, U., Mattes, R.E. and Buckel, P. (1989) *Gene* **85**, 109–114.
- Seidel, H.M., Pompliano, D.L. and Knowes J.R. (1992) *Biochemistry* 31, 2598–2608.
- 11. Baca, A.M. and Hol, W.G.J. (2000) *Int. J. Parasitology* **30**, 113–118.
- Novy, R., Yaeger, K., Monsma, S. and Scott, M. (1999) *inNovations* 10, 1–5.

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| Product   | Size           | Cat. No.           |
|---|----------------|--------------------|
| Rosetta™<br>Competent Cells<br>(guaranteed efficiency > 2 × 10 <sup>6</sup> cfu/µg)               | 0.4 ml<br>1 ml | 70953-3<br>70953-4 |
| RosettaBlue™<br>Competent Cells<br>(guaranteed efficiency > 1 × 10 <sup>8</sup> cfu/µg)           | 0.4 ml<br>1 ml | 71058-3<br>71058-4 |
| Rosetta-gami™<br>Competent Cells<br>(guaranteed efficiency > 2 × 10 <sup>6</sup> cfu/µg)          | 0.4 ml<br>1 ml | 71054-3<br>71054-4 |
| Rosetta(DE3)<br>Competent Cells<br>(guaranteed efficiency > 2 × 10 <sup>6</sup> cfu/µg)           | 0.4 ml<br>1 ml | 70954-3<br>70954-4 |
| Rosetta(DE3)pLysS<br>Competent Cells<br>(guaranteed efficiency > 2 × 10 <sup>6</sup> cfu/µg)      | 0.4 ml<br>1 ml | 70956-3<br>70956-4 |
| RosettaBlue(DE3)<br>Competent Cells<br>(guaranteed efficiency > 1 × 10 <sup>8</sup> cfu/µg)       | 0.4 ml<br>1 ml | 71059-3<br>71059-4 |
| RosettaBlue(DE3)pLysS<br>Competent Cells<br>(guaranteed efficiency > 1 × 10 <sup>8</sup> cfu/µg)  | 0.4 ml<br>1 ml | 71034-3<br>71034-4 |
| Rosetta-gami(DE3)<br>Competent Cells<br>(guaranteed efficiency > 2 × 10 <sup>6</sup> cfu/µg)      | 0.4 ml<br>1 ml | 71055-3<br>71055-4 |
| Rosetta-gami(DE3)pLysS<br>Competent Cells<br>(guaranteed efficiency > 2 × 10 <sup>6</sup> cfu/µg) | 0.4 ml<br>1 ml | 71057-3<br>71057-4 |
| Rosetta(DE3)pLacl<br>Competent Cells<br>(guaranteed efficiency > 2 × 10 <sup>6</sup> cfu/µg)      | 0.4 ml<br>1 ml | 70920-3<br>70920-4 |
| RosettaBlue(DE3)pLacl<br>Competent Cells<br>(guaranteed efficiency > 1 × 10 <sup>s</sup> cfu/µg)  | 0.4 ml<br>1 ml | 71060-3<br>71060-4 |
| Rosetta-gami(DE3)pLacl<br>Competent Cells<br>(guaranteed efficiency > 2 × 10 <sup>6</sup> cfu/µg) | 0.4 ml<br>1 ml | 71056-3<br>71056-4 |

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