

MOL.911 Cell Engineering

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² Cell Engineering

General strategies:

- Knock out of specific genes
 - Gene disruption
 - Mutagenesis
- Down-regulation of specific genes
 - Antisense expression
 - Manipulation of regulatory elements
- Overexpression of homologous genes
- Integration and expression of heterologous genes
 - Co-expression of helper proteins
 - Introduction of metabolic pathway steps



Engineering of E.coli for Expression of Proteins by adding tRNAs for rare codons

Rosetta™ and Rosetta 2 host strains are BL21 derivatives designed to enhance the expression of eukaryotic proteins that contain codons rarely used in *E. coli*. (13–17).

The original Rosetta strains supply tRNAs for the codons AUA, AGG, AGA, CUA, CCC, and GGA on a compatible chloramphenicol-resistant plasmid, pRARE (18).

The Rosetta 2 strains supply a seventh rare codon (CGG) in addition to the six found in the original Rosetta strains (19).

By supplying rare codons, the Rosetta strains provide for "universal" translation, where translation would otherwise be limited by the codon usage of *E. coli*. (15, 16, 20, 21).

The tRNA genes are driven by their native promoters (18).







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).

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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).

http://www.takara-bio.co.jp

Chaperone Plasmid Set

Cat.# 3340

v.0401

No.	Plasmid	Chaperone	Promoter	Inducer	Resistant Marker	References
1	pG-KJE8	dnaK-dnaJ-grpE	araB	L-Arabinose	Cm	2,3
		groES-groEL	Pzt1	Tetracyclin		
2	pGro7	groES-groEL	araB	L-Arabinose	Cm	2
3	pKJE7	dnaK-dnaJ-grpE	araB	L-Arabinose	Cm	2
4	pG-Tf2	groES-groEL-tig	Pzt1	Tetracyclin	Cm	3
5	pTf16	tig	araB	L-Arabinose	Cm	3

http://www.takara-bio.co.jp

Mammalian Chaperones

http://www.abvector.com/MultiChaperoneSystems.htm

Engineering for Expression of Proteins with Disulfide bonds

Thioredoxin (TRX) superfamily

- Consists of proteins containing one or more "TRXlike" domains
- Redox-active members have a "CXXC" catalytic motif
- TRX-like redox-active proteins can be:
 - reductants of disulfide bonds (eg. TRX itself)
 - oxidants of SH-groups (eg. PDI, DsbA)
 - disulfide isomerases (eg. PDI, DsbC)

Engineering for Expression of Proteins with Disulfide bonds

Origami™ host strains are K-12 derivatives that have mutations in both the thioredoxin reductase (*trxB*) and glutathione reductase (*gor*) genes, which greatly enhance disulfide bond formation in the cytoplasm.

Studies have shown that expression in Origami (DE3) yielded 10-fold more active protein than in another host even though overall expression levels were similar.

Origami hosts are compatible with ampicillin resistant plasmids and are ideal for use with pET-32 vectors, since the thioredoxin fusion tag further enhances the formation of disulfide bonds in the cytoplasm.

The *trxB* and *gor* mutations are selectable on kanamycin and tetracycline, respectively; therefore these strains cannot be used with plasmids carrying kanamycin- or tetracycline-resistance genes.

To reduce the possibility of disulfide bond formation between molecules, hosts containing the *trxB/gor* mutations are only recommended for the expression of proteins that require disulfide bond formation for proper folding.

<u>J Biol Chem.</u> 1997 Jun 20;272(25):15661-7.

The role of the thioredoxin and glutaredoxin pathways in reducing protein disulfide bonds in the Escherichia coli cytoplasm

Known components of the thioredoxin system (top) and glutaredoxin system (bottom). The genes encoding the components of these systems are shown inparentheses.

Glutaredoxin System

Prinz W A et al. J. Biol. Chem. 1997;272:15661-15667

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Disulfide isomerisation

- 1) nucleophilic attack of a disulfide bond by an thiolate anion
- 2) transition state
- 3) formation of a mixed disulfide between PDI and the substrate protein a substrate thiol is now free to attack another protein disulfide bond
- 4) isomerization reaction is driven by energy minimization: the native disulfide bond is favored and forms more quickly than the potential re-oxidation of the same bond
- 5) PDI is released unchanged in it's reduced state

Engineering of E.coli for Expression of Proteins with Disulfide bonds

E.coliSaccharomyces cerevisiae,
Pichia pastorisDsbC
DsbGPDI

E.coli

DsbA: Disulfide oxidase

PDI - Protein disulfide isomerase

- PDI family comprises of several members more than a dozen members in humans alone
- PDI (aka PDI-1) is the most abundant member and constitutes ~0.8% of total cellular proteins in mammalian cells and yeast
- PDI is a monomer containing 4 TRX-like domains:
 a, b, b', a'; linker (x) and a c-terminal extension domain (c)
 organized in the order: abb'xa'c
- only a and a have the catalytic CXXC motif

PDI - Protein disulfide isomerase

Oxidative folding in eukaryotes

Dsb – <u>Disulfide</u> bond proteins

- Dsb-family proteins in prokaryotes regulates the forming of disulfide bonds in the periplasmic space like PDI does in the ER, but...
- There are two pathways:
 - oxidation pathway: disulfide bonds are introduced by DsbA
 - isomerization pathway: rearrangement of incorrect disulfide bonds by dsbC (or dsbG)
- DsbA is a monomer (like PDI)
- DsbC and DsbG are homodimers

Dsb – <u>Disulfide</u> bond proteins

Comparison of PDI and dsbC, dsbG

- All of them have a chaperone activity, which is independent of their redox-properties, as they do not require the catalytic cysteines
- PDI is a monomer with 4 TRX-like domains, 2 of them with catalytic active CXXC-motifs
- DsbC and DsbG are homodimers, thus having also 2 catalytic active CXXC-motifs
- PDI is a multifunctional enzyme: It is able to function as a a disulfide oxidase as well as a chaperone and a disulfide isomerase
- DsbC and DsbG work as chaperones and isomerases

22.1.16

Glycosylation of Proteins in P. pastoris

First steps common in lower and higher eukaryotes

Hyperglycosylation in yeasts

Missing reactions in yeast (sialylation)

Problem for production of human therapeutic proteins activity determined by glycosylation stability and fate in human body

Benefit for industrial enzymes

N- and O-linked Protein Glycosylation

All N-linked carbohydrates are linked through **N-Acetylglucosamine** and the amino acid asparagine

The N-linked amino acid consensus sequence is Asn-any AA- Ser or Thr. The middle amino acid can not be proline (Pro).

Most O-linked carbohydrate

covalent attachments to proteins involve a linkage between the monosaccharide N- Acetylgalactosamine and the amino acids serine or threonine.

No consensus sequence defined for Olinked.

N-Glycosylation

Enzymes for Analysis of Glycoproteins

These are just a few of the enzymes available for carbohydrate analysis. Abbreviations are as follows: Asn--Asparagine, Gal--Galactose, GlcNAc--N-acetylglucosamine, GalNAc--N-acetylgalactosamine, and NeuAc--N-acetylneuraminic acid.

Enzyme	Type of enzyme	Specificity	
Endoglycosidase D	Endo	Cleaves various high mannose glycans	
Endoglycosidase F	Endo	Cleaves various high mannose glycans	
Endoglycosidase H	Endo	Cleaves various high mannose glycans	
β-galactosidase	Exo	Removes terminal galactosides from Gal-β1,3-GlcNAc, Gal-β1,4-GlcNAc or Gal-β1,3 GalNAc.	
Peptide:N-Glycosidase F	Endo	Glycoproteins between Asn and GlcNAc (removes oligosaccharides)	
Sialidases (Neuraminidases)	Exo	NeuAc-α2,6-Gal, NeuAc-α2,6- GlcNAc or NeuAc-α2,3-Gal	
Vibrio cholerae Clostridium perfringens Arthobacter ureafaciens Newcastle disease virus			

Engineering of Glycosylation

Synthesis Route for Core Glycan Structure

TRENDS in Biotechnology

genome after OCH1 inactivation

FIG. 2. OCH1 inactivation vector. Upon digestion of pGlycoSwitchM8 with BstBI and transformation in *P. pastoris*, the construct integrates at the OCH1 locus. This results in a short OCH1 fragment that does not translate to a functional OCH1 gene and a promotorless fragment that cannot be translated because of the absence of a promoter and the presence of two in-frame nonsense codons.

Engineering of Glycosylation

Deletion of OCH1

Och1

Protein:Mannosyltransferase of the cis-Golgi apparatus, initiates the polymannose outer chain elongation of Nlinked oligosaccharides of glycoproteins

Engineering of Glycosylation

FIG. 5. Evaluation of hyperglycosylation after inactivation of *P. pastoris OCH1*. (A) Coomassie brilliant blue-stained SDS-PAGE gel containing supernatants of *P. pastoris* strains secreting *T. reesei* mannosidase. For the nonengineered strain (WT) a clear smear is visible, whereas this smear is not present for the strain with *och1* inactivated [och1 (M8)]. (B) FACE analysis of N-glycans derived from mannosidase secreted by a nonengineered strain (WT) and a strain with *och1* inactivated [och1 (M8)]. The bands with greater electrophoretic mobility are the Man8 and Man9 bands and represent core N-glycan structures. The hyperglycosyl structures are slowly migrating sugars. They are not present in the strain with *och1* inactivated.

Deletion of OCH1

Figure 5. (A) Stable integration of the gene for the *K. lactis* UDP-GlcNAc-transporter into the OCH1 locus of *P. pastoris.* Plasmid pJN407 is linearized with Sfil and integrated into the *P. pastoris* genome by double cross-over (knock-in). After marker recycling by selection on 5FOA, the gene of interest and a 'lacZ scar' are stably retained. (B) Integration of a gene of interest into the genome by single cross-over (roll-in). Because the roll-in method leads to duplication of the marker locus, a potential recombination event can lead to loss of the gene of interest and repair of the auxotrophic marker

Engineering of Glycosylation

Marker-free knock-out

N-linked glycosylation pathways in humans and yeast. Representative Nlinked glycosylation pathways in humans and *P. pastoris*(a). An alternative humanized N-Paucimannose linked glycosylation pathway Glycans in P. pastoris(b). Mns; α 1,2- mannosidase, MnsII; mannosidase II, GnTI; β 1,2-*N*acetylglucosaminyltransferase I. GnTII; β 1,2-*N*acetylglucosaminyltransferase II, GalT; β 1,4-galactosyltransferase, SiaT; α 2,6-sialyltransferase, MnT; mannosyltransferase. For simplicity the two GlcNAc residues present at the reducing end of all glycans have been omitted.

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Sialic Acids

The two most common sialic acid derivatives are Neu5Ac and Kdn.

Glycoengineering steps required for sialic acid transfer in the yeast Golgi. Endogenous UDP-GlcNAc, present in the yeast cytoplasm, is converted to CMP-sialic acid by UDP-*N*-acetylglucosamine-2-epimerase/*N*-acetylmannosamine kinase (GNE), *N*-acetylneuraminate-9-phosphate synthase (SPS), sialylate-9-P phosphatase (SPP) and CMP-sialic acid synthase (CSS). Subsequently, the product is translocated into the Golgi by the CMP-sialic acid transporter (CST) and sialic acid is transferred onto the acceptor glycan by sialyltransferase (ST). Enzymes are indicated by blue text and metabolic intermediates by black text. For simplicity the two GlcNAc residues present at the reducing end of all glycans have been omitted.

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