Title: BACTERIAL HOST STRAIN COMPRISING A MUTANT SPr GENE AND A WILD-TYPE TSP GENE

Abstract: The present invention provides a recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene.
BACTERIAL HOST STRAIN COMPRISING A MUTANT SPR GENE AND A WILD-TYPE TSP GENE

The invention relates to a recombinant bacterial host strain, particularly *E. coli*. The invention also relates to a method for producing a protein of interest in such a cell.

**Background of the invention**

Bacterial cells, such as *E. coli*, are commonly used for producing recombinant proteins. There are many advantages to using bacterial cells, such as *E. coli*, for producing recombinant proteins particularly due to the versatile nature of bacterial cells as host cells allowing the gene insertion via plasmids. *E. coli* have been used to produce many recombinant proteins including human insulin.

Despite the many advantages to using bacterial cells to produce recombinant proteins, there are still significant limitations including the tendency of bacterial cells to lyse during expression of a recombinant protein of interest. This lysis phenotype may be seen in wild-type bacterial cells and also genetically engineered cell, such as cells which are deficient in bacterial proteases. Proteases play an important role in turning over old, damaged or miss-folded proteins in the *E. coli* periplasm and cytoplasm. Bacterial proteases act to degrade the recombinant protein of interest, thereby often significantly reducing the yield of active protein. Therefore, the reduction of protease activity is desirable to reduce proteolysis of proteins of interest. However, bacterial strains lacking proteases, such as Tsp (also known as Pre), also exhibit cell lysis.

Tsp (also known as Pre) is a 60kDa periplasmic protease. The reduction of Tsp (pre) activity is desirable to reduce the proteolysis of proteins of interest. However, it was found that cells lacking the protease pre show thermosensitive growth at low osmolarity. Hara et al isolated Tsp deficient strains which were thermoresistant revertants containing extragenic suppressor (spr) mutations (Hara et al., Microbial Drug Resistance, 2: 63-72 (1996)). Spr is an 18kDa membrane bound periplasmic protease and the substrates of spr are Tsp and peptidoglycans in the outer membrane involved in cell wall hydrolysis during cell division. The spr gene is designated as UniProtKB/Swiss-Prot P0AFV4 (SPR_ECOLI). Protease deficient bacterial strains carrying a mutant spr gene have been described in Chen et al (Chen C, Snedecor B, Nishihara JC, Joly JC, McFarland N, Andersen DC, Battersby JE, Champion KM. Biotechnol Bioeng. 2004 Mar 5;85(5):463-74) which describes the construction of *E. coli* strains carrying different combinations of mutations in pre (Tsp) and another protease, DegP, created by amplifying the upstream
and downstream regions of the gene and ligating these together on a vector comprising selection markers and a sprW174R mutation.

It has been surprisingly found that a gram-negative bacterial cell carrying a mutant spr gene and a wild-type Tsp gene provides a cell having reduced lysis. Accordingly, the present inventors have provided a new strain having advantageous properties for producing a protein of interest.

It was surprising that cells according to the present invention show advantageous growth and protein yield phenotype because spr and Tsp are known to be mutual suppressors and, therefore, it would be predicted that if one is allowed to dominate the cell may exhibit a poor growth phenotype, such as becoming leaky or show increase propensity to cell lysis. However, the cells of the present invention exhibited a significant reduction in cell lysis phenotype compared to wild-type cells and cells comprising a knockout mutated Tsp gene.

**Summary of the Invention**

The present invention provides a recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene.

In one embodiment, the genome of the cell according to the present invention is isogenic to the genome of a wild-type bacterial cell except for the mutated spr gene.

The cells provided by the present invention show advantageous growth and protein production phenotypes.

The present invention also provides a method for producing a recombinant protein of interest comprising expressing the recombinant protein of interest in a recombinant gram-negative bacterial cell as defined above.

**Brief Description of the Drawings**

Figure 1 shows the growth of MXE012 and MXE017 compared to the wild-type W3110 and MXE001.

Figure 2 shows the expression of the anti-TNFα Fab' in MXE012 and MXE017 compared to the wild-type W3110 and MXE001.

Figure 3 shows the growth profile of W3110 and MXE012 during a anti-TNFα Fab' producing fermentation.
Figure 4 shows periplasmic anti-TNFa Fab' accumulation (filled lines and symbols) and media Fab' accumulation (dashed lines and open symbols) for W3110 and MXE012 (W3110 spr H19A) during a anti-TNFa Fab' producing fermentation. Figure 5 shows the growth profile of anti-TNFa Fab' expressing strains W3110 and MXE012 and of anti-TNFa Fab' and recombinant DsbC expressing strains W3110 and MXE012. Figure 6 shows anti-TNFa Fab' yield from the periplasm (shaded symbols) and supernatant (open unshaded symbols) from anti-TNFa Fab' expressing strains W3110 and MXE012 and of anti-TNFa Fab' and recombinant DsbC expressing strains W3110 and MXE012. Figure 7 shows the results of a dsDNA assay of strains W3110, MXEOOl, MXE008 and MXE012. Figure 8 shows the results of a protein assay of strains W3110, MXEOOl, MXE008 and MXE012. Figure 9a shows the 5' end of the wild type ptr (protease III) and knockout mutated ptr (protease III) protein and gene sequences. Figure 9b shows the 5' end of the wild type Tsp and knockout mutated Tsp protein and gene sequences. Figure 9c shows a region of the wild type DegP and mutated DegP protein and gene sequences. Figure 10 shows the construction of a vector for use in producing a cell according to an embodiment of the present invention. Figure 11 shows the growth profiles of 200L fermentations of anti-TNFa Fab' and recombinant DsbC expressing strain MXE012. Figure 12 shows the anti-TNFa Fab' titres of 200L fermentations of anti-TNFa Fab' and recombinant DsbC expressing strain MXE012. Figure 13 shows the viabilities of 200L fermentations of anti-TNFa Fab' and recombinant DsbC expressing strain MXE012. Figure 14 shows the growth profiles of 3000L fermentations of anti-TNFa Fab' and recombinant DsbC expressing strain MXE012. Figure 15 shows the anti-TNFa Fab' titres of 3000L fermentations of anti-TNFa Fab' and recombinant DsbC expressing strain MXE012.
Brief Description of the Sequences

SEQ ID NO:1 is the DNA sequence of the wild-type Tsp gene including the 6 nucleotides ATGAAC upstream of the start codon.

SEQ ID NO:2 is the amino acid sequence of the wild-type Tsp protein.

SEQ ID NO:3 is the DNA sequence of a mutated knockout Tsp gene including the 6 nucleotides ATGAAT upstream of the start codon.

SEQ ID NO:4 is the DNA sequence of the wild-type Protease III gene.

SEQ ID NO:5 is the amino acid sequence of the wild-type Protease III protein.

SEQ ID NO:6 is the DNA sequence of a mutated knockout Protease III gene.

SEQ ID NO:7 is the DNA sequence of the wild-type DegP gene.

SEQ ID NO:8 is the amino acid sequence of the wild-type DegP protein.

SEQ ID NO:9 is the DNA sequence of a mutated DegP gene.

SEQ ID NO:10 is the amino acid sequence of a mutated DegP protein.

SEQ ID NO:11 is the amino acid sequence of the light chain variable region of an anti-TNF antibody.

SEQ ID NO:12 is the amino acid sequence of the heavy chain variable region of an anti-TNF antibody.

SEQ ID NO:13 is the amino acid sequence of the light chain of an anti-TNF antibody.

SEQ ID NO:14 is the amino acid sequence of the heavy chain of an anti-TNF antibody.

SEQ ID NO:15 is the sequence of the 3' oligonucleotide primer for the region of the mutated Tsp gene comprising the Ase I restriction site.

SEQ ID NO:16 is the sequence of the 5' oligonucleotide primer for the region of the mutated Tsp gene comprising the Ase I restriction site.

SEQ ID NO:17 is the sequence of the 3' oligonucleotide primer for the region of the mutated Protease III gene comprising the Ase I restriction site.

SEQ ID NO:18 is the sequence of the 5' oligonucleotide primer for the region of the mutated Protease III gene comprising the Ase I restriction site.

SEQ ID NO:19 is the sequence of the 5' oligonucleotide primer for the region of the mutated DegP gene comprising the Ase I restriction site.

SEQ ID NO:20 is the sequence of the 3' oligonucleotide primer for the region of the mutated DegP gene comprising the Ase I restriction site.
SEQ ID NO: 21 is the sequence of the wild-type spr gene including the signal sequence which is the first 26 amino acid residues. SEQ ID NO: 22 is the sequence of the non-mutated spr gene without the signal sequence.

SEQ ID NO: 23 is the nucleotide sequence of a mutated OmpT sequence comprising D210A and H212A mutations.

SEQ ID NO: 24 is the amino acid sequence of a mutated OmpT sequence comprising D210A and H212A mutations.

SEQ ID NO: 25 is the nucleotide sequence of a mutated knockout OmpT sequence.

SEQ ID NO: 26 is the nucleotide sequence of his-tagged DsbC.

SEQ ID NO: 27 is the amino acid sequence of his-tagged DsbC.

SEQ ID NO: 28 shows the amino acid sequence of CDRH1 of hTNF40.

SEQ ID NO: 29 shows the amino acid sequence of CDRH2 of hTNF40 which is a hybrid CDR wherein the C-terminal six amino acids are from the H2 CDR sequence of a human subgroup 3 germline antibody and the amino acid changes to the sequence resulting from this hybridisation are underlined as follows: WINTYIGEP YADSVVKG.

SEQ ID NO: 30 shows the amino acid sequence of CDRH3 of hTNF40.

SEQ ID NO: 31 shows the amino acid sequence of CDRL1 of hTNF40.

SEQ ID NO: 32 shows the amino acid sequence of CDRL2 of hTNF40.

SEQ ID NO: 33 shows the amino acid sequence of CDRL3 of hTNF40.

SEQ ID NO: 34 shows the amino acid sequence of CDRH2 of hTNF40.

SEQ ID NO: 35 shows the sequence of the OmpA oligonucleotide adapter.

SEQ ID NO: 36 shows the oligonucleotide cassette encoding intergenic sequence 1 (IGS1) for E. coli Fab expression.

SEQ ID NO: 37 shows the oligonucleotide cassette encoding intergenic sequence 2 (IGS2) for E. coli Fab expression.

SEQ ID NO: 38 shows the oligonucleotide cassette encoding intergenic sequence 3 (IGS3) for E. coli Fab expression.

SEQ ID NO: 39 shows the oligonucleotide cassette encoding intergenic sequence 4 (IGS4) for E. coli Fab expression.

**Detailed Description of the Preferred Embodiments of the Invention**

The present invention provides a recombinant gram-negative bacterial cell suitable for expressing a protein of interest which comprises a mutated spr gene and a non-recombinant wild-type chromosomal Tsp gene.
It has been surprisingly found that cells carrying a mutated spr and a non-recombinant wild-type chromosomal Tsp exhibit improved cell growth and exhibit reduced cell lysis phenotype compared to a wild-type cell or a cell comprising a mutated Tsp gene.

Further, in one embodiment cells carrying a mutant spr and a non-recombinant wild-type chromosomal Tsp exhibit increased yield of a recombinant protein of interest compared to a wild-type bacterial cell or a cell comprising a mutated Tsp gene. The improved protein yield may be the periplasmic protein yield and/or the supernatant protein yield. In one embodiment the cells of the present invention show improved periplasmic protein yield compared to a wild-type cell due to reduced leakage from the cell. The recombinant bacterial cells are be capable of prolonged expression of a recombinant protein of interest due to reduced cell lysis.

The cells according to the present invention preferably express a maximum yield in the periplasm and/or media of approximately 1.0g/L, 1.5g/L, 1.8g/L, 2.0g/L, 2.4g/L, 2.5g/L, 3.0g/L, 3.5g/L or 4.0g/L of a protein of interest.

A drawback associated with known genetically engineered strains, such as the protease deficient bacterial strains, previously created and used to express recombinant proteins involves the use of mutations of genes involved in cell metabolism and DNA replication such as, for example phoA, fliaA, lac, rec, gal, ara, org, thi and pro in E. coli strains. These mutations may have many deleterious effects on the host cell including effects on cell growth, stability, recombinant protein expression yield and toxicity. Strains having one or more of these genomic mutations, particularly strains having a high number of these mutations, may exhibit a loss of fitness which reduces bacterial growth rate to a level which is not suitable for industrial protein production. Further, any of the above genomic mutations may affect other genes in cis and/or in trans in unpredictable harmful ways thereby altering the strain's phenotype, fitness and protein profile. Further, the use of heavily mutated cells is not generally suitable for producing recombinant proteins for commercial use, particularly therapeutics, because such strains generally have defective metabolic pathways and hence may grow poorly or not at all in minimal or chemically defined media.

In a preferred embodiment of the invention, the cells carry only the minimal mutations to the genome required to introduce the spr mutant. In this embodiment, the genome of the bacterial cell only differs from the genome of a wild-type bacterial cell by
one or more mutations to the spr gene and do not carry any other mutations which may have deleterious effects on the cell’s growth and/or ability to express a protein of interest. Accordingly, one or more of the recombinant host cells according to the present invention may exhibit improved protein expression and/or improved growth characteristics compared to cells comprising further genetically engineered mutations to the genomic sequence. The cells provided by the present invention are also more suitable for use to produce therapeutic proteins compared to cells comprising further disruptions to the cell genome.

The present invention also provides a recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a mutant spr protein, wherein the genome of the cell is isogenic to the genome of a wild-type bacterial cell except for the mutated spr gene. In this aspect of the present invention, the cell carries a wild-type Tsp gene. The wild-type chromosomal Tsp gene is preferably a non-recombinant chromosomal Tsp gene.

In a preferred embodiment of the present invention, the cell further comprises a recombinant polynucleotide encoding DsbC.

The present invention will now be described in more detail.

The terms "protein" and "polypeptide" are used interchangeably herein, unless the context indicates otherwise. "Peptide" is intended to refer to 10 or less amino acids.

The terms "polynucleotide" includes a gene, DNA, cDNA, RNA, mRNA etc unless the context indicates otherwise.

As used herein, the term "comprising" in context of the present specification should be interpreted as "including".

The non-mutated cell or control cell in the context of the present invention means a cell of the same type as the recombinant gram-negative cell of the invention wherein the cell has not been modified to carry the mutant spr gene. For example, a non-mutated cell may be a wild-type cell and may be derived from the same population of host cells as the cells of the invention before modification to introduce the any mutations.

The expressions "cell", "cell line", "cell culture" and "strain" are used interchangeably.

The expression "phenotype of a cell comprising a mutated Tsp gene" in the context of the present invention means the phenotype exhibited by a cell harbouring a mutant Tsp gene. Typically cells comprising a mutant Tsp gene may lyse, especially at high cell densities. The lysis of these cells causes any recombinant protein to leak into the supernatant. Cells carrying mutated Tsp gene may also show thermosensitive growth at low osmolarity. For example, the cells exhibit no or reduced growth rate or the cells die in hypotonic media at a high temperature, such as at 40°C or more.

The term "isogenic" in the context of the present invention means that the genome of the cell of the present invention has substantially the same or the same genomic sequence compared to the wild-type cell from which the cell is derived except for mutated spr gene. In this embodiment the genome of the cell according to the present invention comprises no further non-naturally occurring or genetically engineered mutations. In one embodiment the cell according to the present invention may have substantially the same genomic sequence compared to the wild-type cell except for the mutated spr gene, taking into account any naturally occurring mutations which may occur. In one embodiment, the cell according to the present invention may have exactly the same genomic sequence compared to the wild-type cell except for the mutated spr gene.
In the embodiment of the present invention wherein the cell comprises a recombinant polynucleotide encoding DsbC, the polynucleotide encoding DsbC may be present on a suitable expression vector transformed into the cell and/or integrated into the host cell's genome. In the embodiment where the polynucleotide encoding DsbC is inserted into the host's genome, the cell of the present invention will also differ from a wild-type cell due to the inserted polynucleotide sequence encoding the DsbC. Preferably the polynucleotide encoding DsbC is in an expression vector in the cell thereby causing minimal disruption to the host cell's genome.

The term "wild-type" in the context of the present invention means a strain of a gram-negative bacterial cell as it may occur in nature or may be isolated from the environment, which does not carry any genetically engineered mutations. An example of a wild-type strain of E. coli is W31 10, such as W31 10 K-12 strain.

Any suitable gram-negative bacterium may be used as the parental cell for producing the recombinant cell of the present invention. Suitable gram-negative bacterium include Salmonella typhimurium, Pseudomonas fluorescens, Erwinia carotovora, Shigella, Klebsiella pneumoniae, Legionella pneumophila, Pseudomonas aeruginosa, Acinetobacter baumannii and E. coli. Preferably the parental cell is E. coli. Any suitable strain of E. coli may be used in the present invention but preferably a wild-type W3110 strain, such as K-12 W3110, is used.

In a preferred embodiment, the cell is isogenic to a wild-type E. coli cell, such as W3110, except for the mutated spr gene.

In one embodiment the cell of the present invention comprises a polynucleotide encoding the protein of interest. In this embodiment, the polynucleotide encoding the protein of interest may be contained within a suitable expression vector transformed into the cell and/or integrated into the host cell's genome. In the embodiment where the polynucleotide encoding the protein of interest is inserted into the host's genome, the genome of the present invention will also differ from a wild-type cell due to the inserted polynucleotide sequence encoding the protein of interest. Preferably the polynucleotide is in an expression vector in the cell thereby causing minimal disruption to the host cell's genome.

The cells according to the present invention carry a wild-type Tsp gene. In one aspect of the present invention the cells carry a wild-type non-recombinant chromosomal Tsp gene. The wild-type non-recombinant chromosomal Tsp gene refers to a
chromosomal Tsp gene that is not constructed, produced or inserted into the chromosome using recombinant DNA technology.

As used herein, "Tsp gene" means a gene encoding protease Tsp (also known as Pre) which is a periplasmic protease capable of acting on Penicillin-binding protein-3 (PBP3) and phage tail proteins. The sequence of the wild-type Tsp gene is shown in SEQ ID NO: 1 and the sequence of the wild-type Tsp protein is shown in SEQ ID NO: 2.

The spr protein is a 18kDa membrane bound periplasmic protease and the substrates of spr are Tsp and peptidoglycans in the outer membrane involved in cell wall hydrolysis during cell division.

The wild-type amino acid sequence of the spr protein is shown in SEQ ID NO:21 with the signal sequence at the N-terminus and in SEQ ID NO:22 without the signal sequence of 26 amino acids (according to UniProt Accession Number P0AFV4). The amino acid numbering of the spr protein sequence in the present invention includes the signal sequence. Accordingly, the amino acid 1 of the spr protein is the first amino acid (Met) shown in SEQ ID NO: 21.

The mutated spr gene is preferably the cell's chromosomal spr gene.

The mutated spr gene encodes a spr protein capable of suppressing the phenotype of a cell comprising a mutated Tsp gene. Cells carrying mutated Tsp gene may have a good cell growth rate but one limitation of these cells is their tendency to lyse, especially at high cell densities. Accordingly the phenotype of a cell comprising a mutated Tsp gene is a tendency to lyse, especially at high cell densities. Cells carrying mutated Tsp gene also show thermosensitive growth at low osmolarity. However, the spr mutations carried by the cells of the present invention, when introduced into a cell carrying a mutated Tsp gene suppress the mutant Tsp phenotype and, therefore, the cell exhibits reduced lysis, particularly at a high cell density. The growth phenotype of a cell may be easily measured by a person skilled in the art during shake flask or high cell density fermentation technique. The suppression of the cell lysis phenotype may be been seen from the improved growth rate and/or recombinant protein production, particularly in the periplasm, exhibited by a cell carrying spr mutant and Tsp mutant compared to a cell carrying the Tsp mutant and a wild-type spr.

Any suitable mutation or mutations may be made to the spr gene which results in a spr protein capable of suppressing the phenotype of a cell comprising a mutated Tsp gene. This activity may be tested by a person skilled in the art by creating a cell carrying a
mutant spr gene and mutant Tsp gene and comparing the phenotype to a cell carrying the mutant Tsp gene only. Suitable mutations to the Tsp gene are described in detail below. Reference to a mutated Tsp gene or mutated Tsp gene encoding Tsp, refers to either a mutated Tsp gene encoding a Tsp protein having reduced protease activity or a knockout mutated Tsp gene, unless otherwise indicated.

The expression "mutated Tsp gene encoding a Tsp protein having reduced protease activity" means that the mutated Tsp gene does not have the full protease activity compared to the wild-type non-mutated Tsp gene. The mutated Tsp gene may encode a Tsp protein having 50% or less, 40% or less, 30% or less, 20% or less, 10% or less or 5% or less of the protease activity of a wild-type non-mutated Tsp protein. The mutated Tsp gene may encode a Tsp protein having no protease activity. The cell is not deficient in chromosomal Tsp i.e. the Tsp gene sequence has not been deleted or mutated to prevent expression of any form of Tsp protein.

Any suitable mutation may be introduced into the Tsp gene in order to produce a protein having reduced protease activity. The protease activity of a Tsp protein expressed from a gram-negative bacterium may be easily tested by a person skilled in the art by any suitable method in the art, such as the method described in Keiler et al (Identification of Active Site Residues of the Tsp Protease* THE JOURNAL OF BIOLOGICAL CHEMISTRY Vol. 270, No. 48, Issue of December 1, pp. 28864-28868, 1995 Kenneth C. Keiler and Robert T. Sauer) wherein the protease activity of Tsp was tested.

Tsp has been reported in Keiler et al (supra) as having an active site comprising residues S430, D441 and K455 and residues G375, G376, E433 and T452 are important for maintaining the structure of Tsp. Keiler et al (supra) reports findings that the mutated Tsp genes S430A, D441A, K455A, K455H, K455R, G375A, G376A, E433A and T452A had no detectable protease activity. It is further reported that the mutated Tsp gene S430C displayed about 5-10% wild-type activity. Accordingly, the Tsp mutation to produce a protein having reduced protease activity may comprise a mutation, such as a missense mutation to one or more of residues S430, D441, K455, G375, G376, E433 and T452. Preferably the Tsp mutation to produce a protein having reduced protease activity may comprise a mutation, such as a missense mutation to one, two or all three of the active site residues S430, D441 and K455.

Accordingly the mutated Tsp gene may comprise:

- a mutation to S430; or
• a mutation to D441; or
• a mutation to K455; or
• a mutation to S430 and D441; or
• a mutation to S430 and K455; or
• a mutation to S430 and K455; or
• a mutation to D441 and K455; or
• a mutation to S430, D441 and K455.

One or more of S430, D441, K455, G375, G376, E433 and T452 may be mutated to any suitable amino acid which results in a protein having reduced protease activity. Examples of suitable mutations are S430A, S430C, D441A, K455A, K455H, K455R, G375A, G376A, E433A and T452A. The mutated Tsp gene may comprise one, two or three mutations to the active site residues, for example the gene may comprise:
• S430A or S430C; and/or
• D441A; and/or
• K455A or K455H or K455R.

Preferably, the Tsp gene comprises the point mutation S430A or S430C.

The expression "knockout mutated Tsp gene" means that the gene comprises one or more mutations thereby causing no expression of the protein encoded by the gene to provide a cell deficient in the protein encoded by the knockout mutated gene. The knockout gene may be partially or completely transcribed but not translated into the encoded protein. The knockout mutated Tsp gene may be mutated in any suitable way, for example by one or more deletion, insertion, point, missense, nonsense and frameshift mutations, to cause no expression of the protein. For example, the gene may be knocked out by insertion of a foreign DNA sequence, such as an antibiotic resistance marker, into the gene coding sequence.

The mutated Tsp gene may comprises a mutation to the gene start codon and/or one or more stop codons positioned downstream of the gene start codon and upstream of the gene stop codon thereby preventing expression of the Tsp protein. The mutation to the start codon may be a missense mutation of one, two or all three of the nucleotides of the start codon. Alternatively or additionally the start codon may be mutated by an insertion or deletion frameshift mutation. The Tsp gene comprises two ATG codons at the 5' end of the coding sequence, one or both of the ATG codons may be mutated by a missense mutation. The Tsp gene may be mutated at the second ATG codon (codon 3) to TCG, as
shown in Figure 9b. The Tsp gene may alternatively or additionally comprise one or more stop codons positioned downstream of the gene start codon and upstream of the gene stop codon. Preferably the knockout mutated Tsp gene comprises both a missense mutation to the start codon and one or more inserted stop codons. The Tsp gene may be mutated to delete "T" from the fifth codon thereby causing a frameshift resulting in stop codons at codons 11 and 16, as shown in Figure 9b. The Tsp gene may be mutated to insert an Ase I restriction site to create a third in-frame stop codon at codon 21, as shown in Figure 9b.

The knockout mutated Tsp gene may have the DNA sequence of SEQ ID NO: 3, which includes the 6 nucleotides ATGAAT upstream of the start codon. The mutations which have been made in the knockout mutated Tsp sequence of SEQ ID NO: 3 are shown in Figure 9b. In one embodiment the mutated Tsp gene has the DNA sequence of nucleotides 7 to 2048 of SEQ ID NO:3.

Accordingly, once a cell carrying a suitable mutant Tsp gene has been identified, suitable spr gene mutations can be identified which results in a spr protein capable of suppressing the phenotype of a cell comprising a mutated Tsp gene.

The cells according to a preferred embodiment of the present invention comprise a mutant spr gene encoding a spr protein having a mutation at one or more amino acids selected from N31, R62, 170, Q73, C94, S95, V98, Q99, R100, L108, Y115, D133, V135, L136, G140, R144, H145, G147, H157 and W174, more preferably at one or more amino acids selected from C94, S95, V98, Y115, D133, V135, H145, G147, H157 and W174. In this embodiment, the spr protein preferably does not have any further mutations. Preferably the mutant spr gene encodes a spr protein having a mutation at one or more amino acids selected from N31, R62, 170, Q73, C94, S95, V98, Q99, R100, L108, Y115, D133, V135, L136, G140, R144, H145, G147 and H157, more preferably at one or more amino acids selected from C94, S95, V98, Y115, D133, V135, H145, G147 and H157. In this embodiment, the spr protein preferably does not have any further mutations. Preferably, the mutant spr gene encodes a spr protein having a mutation at one or more amino acids selected from N31, R62, 170, Q73, S95, V98, Q99, R100, L108, Y115, D133, V135, L136, G140, R144 and G147, more preferably at one or more amino acids selected from S95, V98, Y115, D133, V135 and G147. In this embodiment, the spr protein preferably does not have any further mutations.

In one aspect of the present invention there is provided a gram-negative bacterial cell comprising a mutant spr gene encoding a spr protein having a mutation at one or more
amino acids selected from C94, S95, V98, Y115, D133, V135, H145, G147 and H157, preferably at one or more amino acids selected from S95, V98, Y115, D133, V135 and G147, and wherein the cell comprises a wild-type Tsp gene. In this embodiment, the spr protein preferably does not have any further mutations.

The wild-type chromosomal Tsp gene is preferably a non-recombinant chromosomal Tsp gene. Preferably, the cell further comprises a recombinant polynucleotide encoding DsbC.

The mutation to one or more of the above amino acids may be any suitable missense mutation to one, two or three of the nucleotides encoding the amino acid. The mutation changes the amino acid residue to any suitable amino acid which results in a mutated spr protein capable of suppressing the phenotype of a cell comprising a mutated Tsp gene. The missense mutation may change the amino acid to one which is a different size and/or has different chemical properties compared to the wild-type amino acid.

In one embodiment the mutation is to one, two or three of the catalytic triad of amino acid residues of C94, H145, and H157 (Solution NMR Structure of the NlpC/P60 Domain of Lipoprotein Spr from *Escherichia coli* Structural Evidence for a Novel Cysteine Peptidase Catalytic Triad, Biochemistry, 2008, 47, 9715-9717).

Accordingly, the mutated spr gene may comprise:

- a mutation to C94; or
- a mutation to H145; or
- a mutation to H157; or
- a mutation to C94 and H145; or
- a mutation to C94 and H157; or
- a mutation to H145 and H157; or
- a mutation to C94, H145 and H157.

In this embodiment, the spr protein preferably does not have any further mutations.

One, two or three of C94, H145 and H157 may be mutated to any suitable amino acid which results in a spr protein capable of suppressing the phenotype of a cell comprising a mutated Tsp gene. For example, one, two or three of C94, H145, and H157 may be mutated to a small amino acid such as Gly or Ala. Accordingly, the spr protein may have one, two or three of the mutations C94A, H145A and H157A. Preferably, the spr gene comprises the missense mutation H145A, which has been found to produce a spr protein capable of suppressing the phenotype of a cell comprising a mutated Tsp gene.
The designation for a substitution mutant herein consists of a letter followed by a number followed by a letter. The first letter designates the amino acid in the wild-type protein. The number refers to the amino acid position where the amino acid substitution is being made, and the second letter designates the amino acid that is used to replace the wild-type amino acid.

In a preferred embodiment the mutant spr protein comprises a mutation at one or more amino acids selected from N31, R62, 170, Q73, S95, V98, Q99, R100, L108, Y115, D133, V135, L136, G140, R144 and G147, preferably a mutation at one or more amino acids selected from S95, V98, Y115, D133, V135 and G147. In this embodiment, the spr protein preferably does not have any further mutations. Accordingly, the mutated spr gene may comprise:

- a mutation to N31; or
- a mutation to R62; or
- a mutation to 170; or
- a mutation to Q73; or
- a mutation to S95; or
- a mutation to V98; or
- a mutation to Q99; or
- a mutation to R100; or
- a mutation to L108; or
- a mutation to Y115; or
- a mutation to D133; or
- a mutation to V135; or
- a mutation to L136; or
- a mutation to G140; or
- a mutation to R144; or
- a mutation to G147.

In one embodiment the mutant spr protein comprises multiple mutations to amino acids:

- S95 and Y115; or
- N31, Q73, R100 and G140; or
- Q73, R100 and G140; or
\[ \begin{array}{l}
\text{• R100 and G140; or} \\
\text{• Q73 and G140; or} \\
\text{• Q73 and R100; or} \\
\text{• R62, Q99 and R144; or} \\
\text{• Q99 and R144.}
\end{array} \]

One or more of the amino acids N31, R62, 170, Q73, S95, V98, Q99, R100, L108, Y115, D133, V135, L136, G140, R144 and G147 may be mutated to any suitable amino acid which results in a spr protein capable of suppressing the phenotype of a cell comprising a mutated Tsp gene. For example, one or more of N31, R62, 170, Q73, S95, V98, Q99, R100, L108, Y115, D133, V135, L136, G140 and R144 may be mutated to a small amino acid such as Gly or Ala.

In a preferred embodiment the spr protein comprises one or more of the following mutations: N31Y, R62C, I70T, Q73R, S95F, V98E, Q99P, R100G, L108S, Y115F, D133A, V135D or V135G, L136P, G140C, R144C and G147C. Preferably the spr protein comprises one or more of the following mutations: S95F, V98E, Y115F, D133A, V135D or V135G and G147C. In this embodiment, the spr protein preferably does not have any further mutations.

In one embodiment the spr protein has one mutation selected from N31Y, R62C, I70T, Q73R, S95F, V98E, Q99P, R100G, L108S, Y115F, D133A, V135D or V135G, L136P, G140C, R144C and G147C. In this embodiment, the spr protein preferably does not have any further mutations.

In a further embodiment the spr protein has multiple mutations selected from:

\[ \begin{array}{l}
\text{• S95F and Y115F} \\
\text{• N31Y, Q73R, R100G and G140C;} \\
\text{• Q73R, R100G and G140C;} \\
\text{• R100G and G140C;} \\
\text{• Q73R and G140C;} \\
\text{• Q73R and R100G;} \\
\text{• R62C, Q99P and R144C; or} \\
\text{• Q99P and R144C.}
\end{array} \]

In one embodiment the spr protein has the mutation W174R. In an alternative embodiment the spr protein does not have the mutation W174R.
In a preferred embodiment the cell according to the present invention comprises the mutated spr gene and a recombinant polynucleotide encoding DsbC.

As used herein, a "recombinant polypeptide" refers to a protein that is constructed or produced using recombinant DNA technology. The polynucleotide sequence encoding DsbC may be identical to the endogenous sequence encoding DsbC found in bacterial cells. Alternatively, the recombinant polynucleotide sequence encoding DsbC is a mutated version of the wild-type DsbC sequence, for example having a restriction site removed, such as an EcoRI site, and/or a sequence encoding a his-tag. An example modified DsbC nucleotide sequence for use in the present invention is shown in SEQ ID NO: 26, which encodes the his-tagged DsbC amino acid sequence shown in SEQ ID NO: 27.

In one aspect of the present invention there is provided a gram-negative bacterial cell comprising a mutant spr gene encoding a mutant spr protein, a recombinant polynucleotide encoding DsbC and wherein the cell comprises a wild-type Tsp gene. The wild-type Tsp gene is preferably a non-recombinant chromosomal Tsp gene.

DsbC is a prokaryotic protein found in the periplasm of E.coli which catalyzes the formation of disulphide bonds in E.coli. DsbC has an amino acid sequence length of 236 (including signal peptide) and a molecular weight of 25.6 KDa (UniProt No. P0AEG6). DsbC was first identified in 1994 (Missiakas et al. The Escherichia coli dsbC (xprA) gene encodes a periplasmic protein involved in disulfide bond formation, The EMBO Journal vol 13, no 8, p2013-2020, 1994 and Shevchik et al. Characterization of DsbC, a periplasmic protein of Erwinia chrysanthemi and Escherichia coli with disulfide isomerase activity, The EMBO Journal vol 13, no 8, p2007-2012, 1994).

It is known to co-express proteins which catalyze the formation of disulphide bonds to improve protein expression in a host cell. WO98/56930 discloses a method for producing heterologous disulfide bond-containing polypeptides in bacterial cells wherein a prokaryotic disulfide isomerase, such as DsbC or DsbG is co-expressed with a eukaryotic polypeptide. US6673569 discloses an artificial operon comprising polynucleotides encoding each of DsbA, DsbB, DsbC and DsbD for use in producing a foreign protein. EP0786009 discloses a process for producing a heterologous polypeptide in bacteria wherein the expression of nucleic acid encoding DsbA or DsbC is induced prior to the induction of expression of nucleic acid encoding the heterologous polypeptide.
We have found that the specific combination of the expression of recombinant polynucleotide encoding DsbC in a bacterial cell which comprises a mutated spr gene and a wild-type Tsp gene provides an improved host for expressing proteins of interest. It was surprisingly found that the new strains exhibit increased cell growth rate and increased cell survival duration compared to a wild-type cell or a cell comprising a mutated Tsp gene. Specifically, cells carrying a recombinant DsbC gene, a spr mutation and a wild-type Tsp exhibit reduced cell lysis phenotype compared to cells carrying a mutated Tsp gene.

In one embodiment the cell according to the present invention also expresses one or more further proteins as follows:

- one or more proteins capable of facilitating protein folding, such as FkpA, Skp, SurA, PPIA and PPID; and/or
  - one or more protein capable of facilitating protein secretion or translocation, such as SecY, SecE, SecG, SecYEG, SecA, SecB, FtsY and Lep; and/or
  - one or more proteins capable of facilitating disulphide bond formation, such as DsbA, DsbB, DsbD, DsbG.

One of more of the above proteins may be integrated into the cell’s genome and/or inserted in an expression vector.

In one embodiment the cell according to the present invention does not comprise recombinant polynucleotide encoding one or more of the following further proteins:

- one or more proteins capable of facilitating protein folding, such as FkpA, Skp, SurA, PPIA and PPID;
  - one or more protein capable of facilitating protein secretion or translocation, such as SecY, SecE, SecG, SecYEG, SecA, SecB, FtsY and Lep; and
  - one or more proteins capable of facilitating disulphide bond formation, such as DsbA, DsbB, DsbD, DsbG.

In a preferred embodiment of the present invention the recombinant gram-negative bacterial cell further comprises a mutated DegP gene encoding a DegP protein having chaperone activity and reduced protease activity and/or a mutated ptr gene, wherein the mutated ptr gene encodes a Protease III protein having reduced protease activity or is a knockout mutated ptr gene and/or a mutated OmpT gene, wherein the mutated OmpT gene encodes an OmpT protein having reduced protease activity or is a knockout mutated OmpT gene.
In one embodiment the present invention provides a recombinant gram-negative bacterial cell comprising

a. a mutated spr gene;

b. a wild-type non-recombinant chromosomal Tsp gene; and

c. a mutated DegP gene encoding a DegP protein having chaperone activity and reduced protease activity and/or a mutated OmpT wherein the mutated OmpT gene encodes an OmpT protein having reduced protease activity or is a knockout mutated OmpT gene.

Preferably in this embodiment the cell is isogenic to a wild-type bacterial cell except for the above mutations.

In one embodiment the present invention provides a recombinant gram-negative bacterial cell comprising:

a. a mutated spr gene;

b. a wild-type non-recombinant chromosomal Tsp gene; and

c. a mutated ptr gene, wherein the mutated ptr gene encodes a Protease III protein having reduced protease activity or is a knockout mutated ptr gene and/or a mutated OmpT wherein the mutated OmpT gene encodes an OmpT protein having reduced protease activity or is a knockout mutated OmpT gene.

Preferably in this embodiment the cell is isogenic to a wild-type bacterial cell except for the above mutations.

In one embodiment the present invention provides a cell comprising

a. a mutated spr gene;

b. a wild-type non-recombinant chromosomal Tsp gene;

c. a mutated DegP gene encoding a DegP protein having chaperone activity and reduced protease activity;

d. a mutated ptr gene, wherein the mutated ptr gene encodes a Protease III protein having reduced protease activity or is a knockout mutated ptr gene; and

e. optionally a mutated OmpT wherein the mutated OmpT gene encodes an OmpT protein having reduced protease activity or is a knockout mutated OmpT gene.

Preferably in this embodiment the cell is isogenic to a wild-type bacterial cell except for the above mutations.

In one embodiment of the present invention the cell carries a mutated DegP gene.

As used herein, "DegP" means a gene encoding DegP protein (also known as HtrA),
which has dual function as a chaperone and a protease (Families of serine peptidases; Rawlings ND, Barrett AJ. Methods Enzymol. 1994;244:19-61). The sequence of the non-mutated DegP gene is shown in SEQ ID NO: 7 and the sequence of the non-mutated DegP protein is shown in SEQ ID NO: 8.

At low temperatures DegP functions as a chaperone and at high temperatures DegP has a preference to function as a protease (A Temperature-Dependent Switch from Chaperone to Protease in a Widely Conserved Heat Shock Protein. Cell, Volume 97 , Issue 3 , Pages 339 - 347. Spiess C, Beil A, Ehrmann M) and The proteolytic activity of the HtrA (DegP) protein from *Escherichia coli* at low temperatures, Skorko-Glonek J et al Microbiology 2008, 154, 3649-3658).

In the embodiments where the cell comprises the DegP mutation the DegP mutation in the cell provides a mutated DegP gene encoding a DegP protein having chaperone activity but not full protease activity.

The expression "having chaperone activity" in the context of the present invention means that the mutated DegP protein has the same or substantially the same chaperone activity compared to the wild-type non-mutated DegP protein. Preferably, the mutated DegP gene encodes a DegP protein having 50% or more, 60% or more, 70% or more, 80% or more, 90% or more or 95% or more of the chaperone activity of a wild-type non-mutated DegP protein. More preferably, the mutated DegP gene encodes a DegP protein having the same chaperone activity compared to wild-type DegP.

The expression "having reduced protease activity" in the context of the present invention means that the mutated DegP protein does not have the full protease activity compared to the wild-type non-mutated DegP protein. Preferably, the mutated DegP gene encodes a DegP protein having 50% or less, 40% or less, 30% or less, 20% or less, 10% or less or 5% or less of the protease activity of a wild-type non-mutated DegP protein. More preferably, the mutated DegP gene encodes a DegP protein having no protease activity. The cell is not deficient in chromosomal DegP i.e. the DegP gene sequences has not been deleted or mutated to prevent expression of any form of DegP protein.

Any suitable mutation may be introduced into the DegP gene in order to produce a protein having chaperone activity and reduced protease activity. The protease and chaperone activity of a DegP protein expressed from a gram-negative bacterium may be easily tested by a person skilled in the art by any suitable method such as the method described in Spiess et al wherein the protease and chaperone activities of DegP were tested.

DegP is a serine protease and has an active center consisting of a catalytic triad of amino acid residues of Hisl05, Aspl35 and Ser210 (Families of serine peptidases, Methods Enzymol., 1994, 244:19-61 Rawlings N and Barrett A). The DegP mutation to produce a protein having chaperone activity and reduced protease activity may comprise a mutation, such as a missense mutation to one, two or three of Hisl05, Aspl35 and Ser210.

Accordingly, the mutated DegP gene may comprise:

• a mutation to His105; or
• a mutation to Asp135; or
• a mutation to Ser210; or
• a mutation to His105 and Asp135; or
• a mutation to His105 and Ser210; or
• a mutation to Asp135 and Ser210; or
• a mutation to Hisl05, Aspl35 and Ser210.

One, two or three of Hisl05, Aspl35 and Ser210 may be mutated to any suitable amino acid which results in a protein having chaperone activity and reduced protease activity. For example, one, two or three of Hisl05, Aspl35 and Ser210 may be mutated to a small amino acid such as Gly or Ala. A further suitable mutation is to change one, two or three of His105, Aspl35 and Ser210 to an amino acid having opposite properties such as Aspl35 being mutated to Lys or Arg, polar His105 being mutated to a non-polar amino acid such as Gly, Ala, Val or Leu and small hydrophilic Ser210 being mutated to a large or hydrophobic residue such as Val, Leu, Phe or Tyr. Preferably, the DegP gene comprises the point mutation S210A, as shown in Figure 9c, which has been found to produce a protein having chaperone activity but not protease activity (A Temperature-Dependent Switch from Chaperone to Protease in a Widely Conserved Heat Shock Protein. Cell, Volume 97, Issue 3, Pages 339 - 347. Spiess C, Beil A, Ehrmann M).

DegP has two PDZ domains, PDZ1 (residues 260-358) and PDZ2 (residues 359-448), which mediate protein-protein interaction (A Temperature-Dependent Switch from Chaperone to Protease in a Widely Conserved Heat Shock Protein. Cell, Volume 97,
Issue 3, Pages 339 - 347. Spiess C, Beil A, Ehrmann M). In one embodiment of the present invention the degP gene is mutated to delete PDZ1 domain and/or PDZ2 domain. The deletion of PDZ1 and PDZ2 results in complete loss of protease activity of the DegP protein and lowered chaperone activity compared to wild-type DegP protein whilst deletion of either PDZ1 or PDZ2 results in 5% protease activity and similar chaperone activity compared to wild-type DegP protein (A Temperature-Dependent Switch from Chaperone to Protease in a Widely Conserved Heat Shock Protein. Cell. Volume 97, Issue 3, Pages 339 - 347. Spiess C, Beil A, Ehrmann M).

The mutated DegP gene may also comprise a silent non-naturally occurring restriction site, such as Ase I in order to aid in identification and screening methods, for example as shown in Figure 9c.

The preferred sequence of the mutated DegP gene comprising the point mutation S210A and an Ase I restriction marker site is provided in SEQ ID NO: 9 and the encoded protein sequence is shown in SEQ ID NO: 10. The mutations which have been made in the mutated DegP sequence of SEQ ID NO: 9 are shown in Figure 9c.

In the embodiments of the present invention wherein the cell comprises a mutated DegP gene encoding a DegP protein having chaperone activity and reduced protease activity, one or more of the cells provided by the present invention may provide improved yield of correctly folded proteins from the cell relative to mutated cells wherein the DegP gene has been mutated to knockout DegP preventing DegP expression, such as chromosomal deficient DegP. In a cell comprising a knockout mutated DegP gene preventing DegP expression, the chaperone activity of DegP is lost completely whereas in the cell according to the present invention the chaperone activity of DegP is retained whilst the full protease activity is lost. In these embodiments, one or more cells according to the present invention have a lower protease activity to prevent proteolysis of the protein whilst maintaining the chaperone activity to allow correct folding and transportation of the protein in the host cell.

The skilled person would easily be able to test secreted protein to see if the protein is correctly folded using methods well known in the art, such as protein G HPLC, circular dichroism, NMR, X-Ray crystallography and epitope affinity measurement methods.

In these embodiments, one or more cells according to the present invention may have improved cell growth compared to cells carrying a mutated knockout DegP gene preventing DegP expression. Without wishing to be bound by theory improved cell
growth maybe exhibited due to the DegP protease retaining chaperone activity which may increase capacity of the cell to process all proteins which require chaperone activity. Accordingly, the production of correctly folded proteins necessary for the cell's growth and reproduction may be increased in one or more of the cells of the present invention compared to cells carrying a DegP knockout mutation thereby improving the cellular pathways regulating growth. Further, known DegP protease deficient strains are generally temperature-sensitive and do not typically grow at temperatures higher than about 28°C. However, the cells according to the present invention are not temperature-sensitive and may be grown at temperatures of 28°C or higher, including temperatures of approximately 30°C to approximately 37°C, which are typically used for industrial scale production of proteins from bacteria.

In one embodiment of the present invention the cell carries a mutated ptr gene. As used herein, "ptr gene" means a gene encoding Protease III, a protease which degrades high molecular weight proteins. The sequence of the non-mutated ptr gene is shown in SEQ ID NO: 4 and the sequence of the non-mutated Protease III protein is shown in SEQ ID NO: 5.

Reference to the mutated ptr gene or mutated ptr gene encoding Protease III, refers to either a mutated ptr gene encoding a Protease III protein having reduced protease activity or a knockout mutated ptr gene, unless otherwise indicated.

The expression "mutated ptr gene encoding a Protease III protein having reduced protease activity" in the context of the present invention means that the mutated ptr gene does not have the full protease activity compared to the wild-type non-mutated ptr gene.

Preferably, the mutated ptr gene encodes a Protease III having 50% or less, 40% or less, 30% or less, 20% or less, 10% or less or 5% or less of the protease activity of a wild-type non-mutated Protease III protein. More preferably, the mutated ptr gene encodes a Protease III protein having no protease activity. In this embodiment the cell is not deficient in chromosomal ptr i.e. the ptr gene sequence has not been deleted or mutated to prevent expression of any form of Protease III protein.

Any suitable mutation may be introduced into the ptr gene in order to produce a Protease III protein having reduced protease activity. The protease activity of a Protease III protein expressed from a gram-negative bacterium may be easily tested by a person skilled in the art by any suitable method in the art.
The expression "knockout mutated ptr gene" in the context of the present invention means that the gene comprises one or more mutations thereby causing no expression of the protein encoded by the gene to provide a cell deficient in the protein encoded by the knockout mutated gene. The knockout gene may be partially or completely transcribed but not translated into the encoded protein. The knockout mutated ptr gene may be mutated in any suitable way, for example by one or more deletion, insertion, point, missense, nonsense and frameshift mutations, to cause no expression of the protein. For example, the gene may be knocked out by insertion of a foreign DNA sequence, such as an antibiotic resistance marker, into the gene coding sequence.

In a preferred embodiment the gene is not mutated by insertion of a foreign DNA sequence, such as an antibiotic resistance marker, into the gene coding sequence. Preferably the Protease III gene comprise a mutation to the gene start codon and/or one or more stop codons positioned downstream of the gene start codon and upstream of the gene stop codon thereby preventing expression of the Protease III protein.

A mutation to the target knockout gene start codon causes loss of function of the start codon and thereby ensures that the target gene does not comprise a suitable start codon at the start of the coding sequence. The mutation to the start codon may be a missense mutation of one, two or all three of the nucleotides of the start codon. Alternatively or additionally the start codon may be mutated by an insertion or deletion frameshift mutation.

In a preferred embodiment the ptr gene is mutated to change the ATG start codon to ATT, as shown in Figure 9a.

The knockout mutated ptr gene may alternatively or additionally comprise one or more stop codons positioned downstream of the gene start codon and upstream of the gene stop codon. Preferably the knockout mutated ptr gene comprises both a missense mutation to the start codon and one or more inserted stop codons.

The one or more inserted stop codons are preferably in-frame stop codons. However the one or more inserted stop codons may alternatively or additionally be out-of-frame stop codons. One or more out-of-frame stop codons may be required to stop translation where an out-of-frame start codon is changed to an in-frame start codon by an insertion or deletion frameshift mutation. The one or more stop codons may be introduced by any suitable mutation including a nonsense point mutation and a frameshift mutation. The one or more stop codons are preferably introduced by a frameshift mutation and/or an
insertion mutation, preferably by replacement of a segment of the gene sequence with a
sequence comprising a stop codon. For example an Ase I restriction site may be inserted,
which comprises the stop codon TAA.

In a preferred embodiment the ptr gene is mutated to insert an in-frame stop codon
by insertion of an Ase I restriction site, as shown in Figure 9a. In a preferred embodiment
the knockout mutated ptr gene has the DNA sequence of SEQ ID NO: 6. The mutations
which have been made in the knockout mutated ptr gene sequence of SEQ ID NO: 6 are
shown in Figure 9a.

The above described knockout mutations are advantageous because they cause
minimal or no disruption to the chromosomal DNA upstream or downstream of the target
knockout gene site and do not require the insertion and retention of foreign DNA, such as
antibiotic resistance markers, which may affect the cell's suitability for expressing a
protein of interest, particularly therapeutic proteins. Accordingly, one or more of the cells
according to the present invention may exhibit improved growth characteristics and/or
protein expression compared to cells wherein the protease gene has been knocked out by
insertion of foreign DNA into the gene coding sequence.

In one embodiment the cells according to the present invention carry a mutated
OmpT gene. As used herein, "OmpT gene" means a gene encoding protease OmpT (outer
membrane protease T) which is an outer membrane protease. The sequence of the wild-
type non-mutated OmpT gene is SWISS-PROT P09169.

Reference to a mutated OmpT gene or mutated OmpT gene encoding OmpT, refers
to either a mutated OmpT gene encoding a OmpT protein having reduced protease activity
or a knockout mutated OmpT gene, unless otherwise indicated.

The expression "mutated OmpT gene encoding a OmpT protein having reduced
protease activity" in the context of the present invention means that the mutated OmpT
gene does not have the full protease activity compared to the wild-type non-mutated
OmpT gene. The mutated OmpT gene may encode a OmpT protein having 50% or less,
40% or less, 30% or less, 20% or less, 10% or less or 5% or less of the protease activity
of a wild-type non-mutated OmpT protein. The mutated OmpT gene may encode a OmpT
protein having no protease activity. In this embodiment the cell is not deficient in
chromosomal OmpT i.e. the OmpT gene sequence has not been deleted or mutated to
prevent expression of any form of OmpT protein.
Any suitable mutation may be introduced into the OmpT gene in order to produce a protein having reduced protease activity. The protease activity of a OmpT protein expressed from a gram-negative bacterium may be easily tested by a person skilled in the art by any suitable method in the art, such as the method described in Kramer et al (Identification of essential acidic residues of outer membrane protease OmpT supports a novel active site, FEBS Letters 505 (2001) 426-430) and Dekker et al (Substrate Specificity of the Integral Membrane Protease OmpT Determined by Spatially Addressed Peptide Libraries, Biochemistry 2001, 40, 1694-1701).


Accordingly, the OmpT mutation to produce a protein having reduced protease activity may comprise a mutation, such as a missense mutation to one or more of residues E27, D43, D83, D85, D97, S99, H101 E111, E136, E193, D206, D208, D210, H212 G216, K217, R218 & E250.

suitable mutated OmpT sequence comprising D210A and H212A mutations is shown in
SEQ ID NO: 23.

The expression "knockout mutated OmpT gene" in the context of the present
invention means that the gene comprises one or more mutations thereby causing no
expression of the protein encoded by the gene to provide a cell deficient in the protein
encoded by the knockout mutated gene. The knockout gene may be partially or
completely transcribed but not translated into the encoded protein. The knockout mutated
OmpT gene may be mutated in any suitable way, for example by one or more deletion,
insertion, point, missense, nonsense and frameshift mutations, to cause no expression of
the protein. For example, the gene may be knocked out by insertion of a foreign DNA
sequence, such as an antibiotic resistance marker, into the gene coding sequence.

In one embodiment the OmpT gene comprises a mutation to the gene start codon
and/or one or more stop codons positioned downstream of the gene start codon and
upstream of the gene stop codon thereby preventing expression of the OmpT protein. The
mutation to the start codon may be a missense mutation of one, two or all three of the
nucleotides of the start codon. Alternatively or additionally the start codon may be
mutated by an insertion or deletion frameshift mutation.

A suitable mutated knockout OmpT sequence is shown in SEQ ID NO: 24.

In one embodiment the gram-negative bacterial cell according to the present
invention does not carry a knockout mutated ompT gene, such as being deficient in
chromosomal ompT.

In one embodiment the gram-negative bacterial cell according to the present
invention does not carry a knockout mutated degP gene, such as being deficient in
chromosomal degP. In one embodiment the gram-negative bacterial cell according to the
present invention does not carry a mutated degP gene.

In one embodiment the gram-negative bacterial cell according to the present
invention does not carry a knockout mutated ptr gene, such as being deficient in
chromosomal ptr.

Many genetically engineered mutations including knockout mutations involve the
use of antibiotic resistance markers which allow the selection and identification of
successfully mutated cells. However, there are a number of disadvantages to using
antibiotic resistance markers.
In one embodiment of the present invention, the mutated genes may comprise one or more restriction marker site. Therefore, the spr gene and/or a mutated DegP gene encoding a DegP protein having chaperone activity but not protease activity and/or a mutated ptr gene and/or a mutated OmpT gene may be mutated to comprise one or more restriction marker sites. The restriction sites are genetically engineered into the gene and are non-naturally occurring. The restriction marker sites are advantageous because they allow screening and identification of correctly modified cells which comprise the required chromosomal mutations. Cells which have been modified to carry one or more of the mutated genes may be analyzed by PCR of genomic DNA from cell lysates using oligonucleotide pairs designed to amplify a region of the genomic DNA comprising a non-naturally occurring restriction marker site. The amplified DNA may then be analyzed by agarose gel electrophoresis before and after incubation with a suitable restriction enzyme capable of digesting the DNA at the non-naturally occurring restriction marker site. The presence of DNA fragments after incubation with the restriction enzyme confirms that the cells have been successfully modified to carry the one or more mutated genes.

In the embodiment wherein the cell comprises a knockout mutated ptr gene having the DNA sequence of SEQ ID NO: 6, the oligonucleotide primer sequences shown in SEQ ID NO: 17 and SEQ ID NO: 18 may be used to amplify the region of the DNA comprising the non-naturally occurring Ase I restriction site from the genomic DNA of transformed cells. The amplified genomic DNA may then be incubated with Ase I restriction enzyme and analyzed by gel electrophoresis to confirm the presence of the mutated ptr gene in the genomic DNA.

In the embodiment wherein the cell comprises a mutated DegP gene having the DNA sequence of SEQ ID NO: 9, the oligonucleotide primer sequences shown in SEQ ID NO: 19 and SEQ ID NO:20 may be used to amplify the region of the DNA comprising the non-naturally occurring Ase I restriction site from the genomic DNA of transformed cells. The amplified genomic DNA may then be incubated with Ase I restriction enzyme and analyzed by gel electrophoresis to confirm the presence of the mutated DegP gene in the genomic DNA.

The one or more restriction sites may be introduced by any suitable mutation including by one or more deletion, insertion, point, missense, nonsense and frameshift mutations. A restriction site may be introduced by the mutation of the start codon and/or mutation to introduce the one or more stop codons, as described above. This embodiment
is advantageous because the restriction marker site is a direct and unique marker of the knockout mutations introduced.

A restriction maker site may be inserted which comprises an in-frame stop codon, such as an *Ase I* restriction site. This is particularly advantageous because the inserted restriction site serves as both a restriction marker site and a stop codon to prevent full transcription of the gene coding sequence. For example, in the embodiment wherein a stop codon is introduced to the ptr gene by introduction of an *Ase I* site, this also creates a restriction site, as shown in Figure 9a.

A restriction marker site may be inserted by the mutation to the start codon and optionally one or more further point mutations. In this embodiment the restriction marker site is preferably an *EcoR I* restriction site. This is particularly advantageous because the mutation to the start codon also creates a restriction marker site. For example, in the embodiment wherein the start codon of the ptr gene is changed to ATT, this creates an *EcoR I* marker site, as shown in Figure 9a.

In the embodiment of the present invention wherein the cell carries a mutated OmpT gene, the one or more restriction sites may be introduced by any suitable mutation including by one or more deletion, insertion, point, missense, nonsense and frameshift mutations. For example, in the embodiment wherein the OmpT gene comprises the mutations D210A and H212A, these mutations introduce silent HindIII restriction site which may be used as a selection marker.

In the mutated spr gene and the mutated DegP gene, a marker restriction site may be introduced using silent codon changes. For example, an *Ase I* site may be used as a silent restriction marker site, wherein the TAA stop codon is out-of-frame, as shown in Figure 9c for DegP.

In the embodiments of the present invention, wherein the ptr gene is mutated to encode a Protease III having reduced protease activity, one or more marker restriction site may be introduced using silent codon changes.

The recombinant gram-negative bacterial cell according to the present invention may be produced by any suitable means.

The skilled person knows of suitable techniques which may be used to replace a chromosomal gene sequence with a mutated gene sequence in order to introduce the spr mutant gene. Suitable vectors may be employed which allow integration into the host chromosome by homologous recombination.

In the embodiment wherein the cell comprises a recombinant polynucleotide encoding DsbC, the skilled person knows suitable techniques which may be used to insert the recombinant polynucleotide encoding DsbC. The recombinant polynucleotide encoding DsbC may be integrated into the cell's genome using a suitable vector such as the pK03 plasmid.

In the embodiment wherein the cell comprises a recombinant polynucleotide encoding a protein of interest, the skilled person also knows suitable techniques which may be used to insert the recombinant polynucleotide encoding the protein of interest. The recombinant polynucleotide encoding the protein of interest may be integrated into the cell's genome using a suitable vector such as the pK03 plasmid.

Alternatively or additionally, the recombinant polynucleotide encoding DsbC and/or the recombinant polynucleotide encoding a protein of interest may be non-integrated in a recombinant expression cassette. In one embodiment an expression cassette is employed in the present invention to carry the polynucleotide encoding the DsbC and/or the protein of interest and one or more regulatory expression sequences. The one or more regulatory expression sequences may include a promoter. The one or more regulatory expression sequences may also include a 3' untranslated region such as a termination sequence. Suitable promoters are discussed in more detail below.
In one embodiment an expression cassette is employed in the present invention to carry the polynucleotide encoding the protein of interest and/or the recombinant polynucleotide encoding DsbC. An expression cassette typically comprises one or more regulatory expression sequences, one or more coding sequences encoding one or more proteins of interest and/or a coding sequence encoding DsbC. The one or more regulatory expression sequences may include a promoter. The one or more regulatory expression sequences may also include a 3' untranslated region such as a termination sequence. Suitable promoters are discussed in more detail below.

In one embodiment, the cell according to the present invention comprises one or more vectors, such as plasmid. The vector preferably comprises one or more of the expression cassettes as defined above. In one embodiment the polynucleotide sequence encoding a protein of interest and the polynucleotide encoding DsbC are inserted into one vector. Alternatively the polynucleotide sequence encoding a protein of interest and the polynucleotide encoding DsbC are inserted into separate vectors.

In the embodiment where the protein of interest is an antibody comprising both heavy and light chains, the cell line may be transfected with two vectors, a first vector encoding a light chain polypeptide and a second vector encoding a heavy chain polypeptide. Alternatively, a single vector may be used, the vector including sequences encoding light chain and heavy chain polypeptides. Alternatively, the polynucleotide sequence encoding the antibody and the polynucleotide encoding DsbC are inserted into one vector. Preferably the vector comprises the sequences encoding the light and heavy chain polypeptides of the antibody.

In the embodiment wherein the cell also expresses one or more further proteins as follows:

- one or more proteins capable of facilitating protein folding, such as FkpA, Skp, SurA, PPiA and PPiD; and/or
  - one or more protein capable of facilitating protein secretion or translocation, such as SecY, SecE, SecG, SecYEG, SecA, SecB, FtsY and Lep; and/or
  - one or more proteins capable of facilitating disulphide bond formation, such as DsbA, DsbB, DsbD, DsbG;

the one or more further protein may be expressed from one or more polynucleotides inserted into the same vector as the polynucleotide encoding DsbC and/or
the polynucleotide sequence encoding a protein of interest. Alternatively the one or more polynucleotides may be inserted into separate vectors.

The vector for use in the present invention may be produced by inserting one or more expression cassettes as defined above into a suitable vector. Alternatively, the regulatory expression sequences for directing expression of the polynucleotide sequence may be contained in the vector and thus only the encoding region of the polynucleotide may be required to complete the vector.

The polynucleotide encoding DsbC and/or the polynucleotide encoding the protein of interest is suitably inserted into a replicable vector, typically an autonomously replicating vector, for expression in the cell under the control of a suitable promoter for the cell. Many vectors are known in the art for this purpose and the selection of the appropriate vector may depend on the size of the nucleic acid and the particular cell type.

Examples of vectors which may be employed to transform the host cell with a polynucleotide according to the invention include:

- a plasmid, such as pBR322 or pACYC 184, and/or
- a viral vector such as bacterial phage
- a transposable genetic element such as a transposon

Such vectors usually comprise a plasmid origin of DNA replication, an antibiotic selectable marker, a promoter and transcriptional terminator separated by a multi-cloning site (expression cassette) and a DNA sequence encoding a ribosome binding site.

The promoters employed in the present invention can be linked to the relevant polynucleotide directly or alternatively be located in an appropriate position, for example in a vector such that when the relevant polypeptide is inserted the relevant promoter can act on the same. In one embodiment the promoter is located before the encoding portion of the polynucleotide on which it acts, for example a relevant promoter before each encoding portion of polynucleotide. "Before" as used herein is intended to imply that the promoter is located at the 5 prime end in relation to the encoding polynucleotide portion.

The promoters may be endogenous or exogenous to the host cells. Suitable promoters include Lac, tac, trp, PhoA, Ipp, Arab, Tet and T7.

One or more promoters employed may be inducible promoters.

In the embodiment wherein the polynucleotide encoding DsbC and the polynucleotide encoding the protein of interest are inserted into one vector, the nucleotide sequences encoding DsbC and the protein of interest may be under the control of a single
promoter or separate promoters. In the embodiment wherein the nucleotide sequences encoding DsbC and the protein of interest are under the control of separate promoters, the promoter may be independently inducible promoters.

Expression units for use in bacterial systems also generally contain a Shine-Dalgamo (S.D.) sequence operably linked to the DNA encoding the polypeptide of interest. The promoter can be removed from the bacterial source DNA by restriction enzyme digestion and inserted into the vector containing the desired DNA.

In the embodiments of the present invention wherein a polynucleotide sequence comprises two or more encoding sequences for two or more proteins of interest, for example an antibody light chain and antibody heavy chain, the polynucleotide sequence may comprise one or more internal ribosome entry site (IRES) sequences which allows translation initiation in the middle of an mRNA. An IRES sequence may be positioned between encoding polynucleotide sequences to enhance separate translation of the mRNA to produce the encoded polypeptide sequences.

The expression vector preferably also comprises a dicistronic message for producing the antibody or antigen binding fragment thereof as described in WO 03/048208 or WO2007/039714 (the contents of which are incorporated herein by reference). Preferably the upstream cistron contains DNA coding for the light chain of the antibody and the downstream cistron contains DNA coding for the corresponding heavy chain, and the dicistronic intergenic sequence (IGS) preferably comprises a sequence selected from IGS1 (SEQ ID NO: 36), IGS2 (SEQ ID NO: 37), IGS3 (SEQ ID NO: 38) and IGS4 (SEQ ID NO: 39).

The terminators may be endogenous or exogenous to the host cells. A suitable terminator is rrnB.

Further suitable transcriptional regulators including promoters and terminators and protein targeting methods may be found in "Strategies for Achieving High-Level Expression of Genes in Escherichia coli" Savvas C. Makrides, Microbiological Reviews, Sept 1996, p 512-538.

The DsbC polynucleotide inserted into the expression vector preferably comprises the nucleic acid encoding the DsbC signal sequence and the DsbC coding sequence. The vector preferably contains a nucleic acid sequence that enables the vector to replicate in one or more selected host cells, preferably to replicate independently of the host chromosome. Such sequences are well known for a variety of bacteria.
In one embodiment the DsbC and/or the protein of interest comprises a histidine-tag at the N-terminus and/or C-terminus.

The antibody molecule may be secreted from the cell or targeted to the periplasm by suitable signal sequences. Alternatively, the antibody molecules may accumulate within the cell's cytoplasm. Preferably the antibody molecule is targeted to the periplasm.

The polynucleotide encoding the protein of interest may be expressed as a fusion with another polypeptide, preferably a signal sequence or other polypeptide having a specific cleavage site at the N-terminus of the mature polypeptide. The heterologous signal sequence selected should be one that is recognized and processed by the host cell.

For prokaryotic host cells that do not recognize and process the native or a eukaryotic polypeptide signal sequence, the signal sequence is substituted by a prokaryotic signal sequence. Suitable signal sequences include OmpA, PhoA, LamB, PelB, DsbA and DsbC.

Construction of suitable vectors containing one or more of the above-listed components employs standard ligation techniques. Isolated plasmids or DNA fragments are cleaved, tailored, and re-ligated in the form desired to generate the plasmids required.

In a preferred embodiment of the present invention the present invention provides a multi-cistronic vector comprising the polynucleotide sequence encoding DsbC and the polynucleotide sequence encoding a protein of interest. The multicistronic vector may be produced by an advantageous cloning method which allows repeated sequential cloning of polynucleotide sequences into a vector. The method uses compatible cohesive ends of a pair of restrictions sites, such as the "AT" ends of AseI and NdeI restriction sites. A polynucleotide sequence comprising a coding sequence and having compatible cohesive ends, such as a AseI-Ndel fragment, may be cloned into a restrictions site in the vector, such as NdeI. The insertion of the polynucleotide sequence destroys the 5' restriction site but creates a new 3' restriction site, such as Ndel, which may then be used to insert a further polynucleotide sequence comprising compatible cohesive ends. The process may then be repeated to insert further sequences. Each polynucleotide sequence inserted into the vector comprises non-coding sequence 3' to the stop codon which may comprise an SspI site for screening, a Shine Dalgarno ribosome binding sequence, an A rich spacer and an Ndel site encoding a start codon.

A diagrammatic representation of the creation of a vector comprising a polynucleotide sequence encoding a light chain of an antibody (LC), a heavy chain of an
antibody (HC), a DsbC polynucleotide sequence and a further polynucleotide sequence is shown in Figure 10.

Successfully mutated strains may be identified using methods well known in the art including colony PCR DNA sequencing and colony PCR restriction enzyme mapping.

In the embodiment wherein the cell comprises two or more the mutated genes, the mutated protease may be introduced into the gram-negative bacterium on the same or different vectors.

In one embodiment the gram-negative bacterial cell according to the present invention does not carry a knockout mutated ompT gene, such as being deficient in chromosomal ompT.

The cell according to the present invention may further comprise a polynucleotide sequence encoding a protein of interest. The polynucleotide sequence encoding the protein of interest may be exogenous or endogenous. The polynucleotide sequence encoding the protein of interest may be integrated into the host's chromosome or may be non-integrated in a vector, typically a plasmid.

In one embodiment the cell according to the present invention expresses a protein of interest. "Protein of interest" in the context of the present specification is intended to refer to polypeptide for expression, usually a recombinant polypeptide. However, the protein of interest may be an endogenous protein expressed from an endogenous gene in the host cell.

As used herein, a "recombinant polypeptide" refers to a protein that is constructed or produced using recombinant DNA technology. The protein of interest may be an exogenous sequence identical to the endogenous protein or a mutated version thereof, for example with attenuated biological activity, or fragment thereof, expressed from an exogenous vector. Alternatively, the protein of interest may be a heterologous protein, not normally expressed by the host cell.

The protein of interest may be any suitable protein including a therapeutic, prophylactic or diagnostic protein.

In one embodiment the protein of interest is useful in the treatment of diseases or disorders including inflammatory diseases and disorders, immune disease and disorders, fibrotic disorders and cancers.

The term "inflammatory disease" or "disorder" and "immune disease or disorder" includes rheumatoid arthritis, psoriatic arthritis, still's disease, Muckle Wells disease,
psoriasis, Crohn's disease, ulcerative colitis, SLE (Systemic Lupus Erythematosus), asthma, allergic rhinitis, atopic dermatitis, multiple sclerosis, vasculitis, Type I diabetes mellitus, transplantation and graft-versus-host disease.

The term "fibrotic disorder" includes idiopathic pulmonary fibrosis (IPF), systemic sclerosis (or scleroderma), kidney fibrosis, diabetic nephropathy, IgA nephropathy, hypertension, end-stage renal disease, peritoneal fibrosis (continuous ambulatory peritoneal dialysis), liver cirrhosis, age-related macular degeneration (ARMD), retinopathy, cardiac reactive fibrosis, scarring, keloids, burns, skin ulcers, angioplasty, coronary bypass surgery, arthroplasty and cataract surgery.

The term "cancer" includes a malignant new growth that arises from epithelium, found in skin or, more commonly, the lining of body organs, for example: breast, ovary, prostate, lung, kidney, pancreas, stomach, bladder or bowel. Cancers tend to infiltrate into adjacent tissue and spread (metastasise) to distant organs, for example: to bone, liver, lung or the brain.

The protein may be a proteolytically-sensitive polypeptide, i.e. proteins that are prone to be cleaved, susceptible to cleavage, or cleaved by one or more gram-negative bacterial, such as E. coli, proteases, either in the native state or during secretion. In one embodiment the protein of interest is proteolytically-sensitive to a protease selected from DegP, Protease III and Tsp. In one embodiment the protein of interest is proteolytically-sensitive to the protease Tsp. In one embodiment the protein of interest is proteolytically-sensitive to the proteases DegP and Protease III. In one embodiment the protein of interest is proteolytically sensitive to the proteases DegP and Tsp. In one embodiment the protein of interest is proteolytically-sensitive to the proteases Tsp and Protease III. In one embodiment the protein of interest is proteolytically sensitive to the proteases DegP, Protease III and Tsp.

Preferably the protein is a eukaryotic polypeptide.

The protein of interest expressed by the cells according to the invention may, for example be an immunogen, a fusion protein comprising two heterologous proteins or an antibody. Antibodies for use as the protein of interest include monoclonal, multi-valent, multi-specific, humanized, fully human or chimeric antibodies. The antibody can be from any species but is preferably derived from a monoclonal antibody, a human antibody, or a humanized fragment. The antibody can be derived from any class (e.g. IgG, IgE, IgM, IgD or IgA) or subclass of immunoglobulin molecule and may be obtained from any
species including for example mouse, rat, shark, rabbit, pig, hamster, camel, llama, goat or human. Parts of the antibody fragment may be obtained from more than one species for example the antibody fragments may be chimeric. In one example the constant regions are from one species and the variable regions from another.

The antibody may be a complete antibody molecule having full length heavy and light chains or a fragment thereof, e.g. VH, VL, VHH, Fab, modified Fab, Fab', F(ab')2, Fv, scFv fragment, Fab-Fv, or a dual specificity antibody, such as a Fab-dAb, as described in PCT/GB2008/003331.

The antibody may be specific for any target antigen. The antigen may be a cell-associated protein, for example a cell surface protein on cells such as bacterial cells, yeast cells, T-cells, endothelial cells or tumour cells, or it may be a soluble protein. Antigens of interest may also be any medically relevant protein such as those proteins upregulated during disease or infection, for example receptors and/or their corresponding ligands. Particular examples of cell surface proteins include adhesion molecules, for example integrins such as β1 integrins e.g. VLA-4, E-selectin, P selectin or L-selectin, CD2, CD3, CD4, CD5, CD7, CD8, CDllα, CDllb, CD18, CD19, CD20, CD23, CD25, CD33, CD38, CD40, CD40L, CD45, CDW52, CD69, CD134 (OX40), ICOS, BCMP7, CD137, CD27L, CDCP1, CSF1 or CSF1- Receptor, DPCR1, DPCR1, dudulin2, FLJ20584, FLJ40787, HEK2, KIAA0634, KIAA0659, KIAA1246, KIAA1455, LTBP2, LTK, MAL2, MRP2, nectin-like2, NKCC1, PTK7, RAIG1, TCAM1, SC6, BCMP101, BCMP84, BCMP11, DTD, carcinoembryonic antigen (CEA), human milk fat globulin (HMFG1 and 2), MHC Class I and MHC Class II antigens, KDR and VEGF, and where appropriate, receptors thereof.

Soluble antigens include interleukins such as IL-1, IL-2, IL-3, IL-4, IL-5, IL-6, IL-8, IL-12, IL-13, IL-14, IL-16 or IL-17, such as IL17A and/or IL17F, viral antigens for example respiratory syncytial virus or cytomegalovirus antigens, immunoglobulins, such as IgE, interferons such as interferon α, interferon β or interferon γ, tumour necrosis factor TNF (formerly known as tumour necrosis factor-a), tumor necrosis factor-β, colony stimulating factors such as G-CSF or GM-CSF, and platelet derived growth factors such as PDGF-a, and PDGF-β and where appropriate receptors thereof. Other antigens include bacterial cell surface antigens, bacterial toxins, viruses such as influenza, EBV, HepA, B and C, bioterrorism agents, radionuclides and heavy metals, and snake and spider venoms and toxins.
In one embodiment, the antibody may be used to functionally alter the activity of the antigen of interest. For example, the antibody may neutralize, antagonize or agonise the activity of said antigen, directly or indirectly.

In one aspect of the present invention there is provided a recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a mutant spr protein, a wild-type Tsp gene and a polynucleotide sequence encoding an antibody or an antigen binding fragment thereof specific for TNF. The wild-type chromosomal Tsp gene is preferably a non-recombinant chromosomal Tsp gene. Preferably, the cell further comprises a recombinant polynucleotide encoding DsbC.

In a preferred embodiment the protein of interest expressed by the cells according to the present invention is an anti-TNF antibody, more preferably an anti-TNF Fab', as described in WO01/094585 (the contents of which are incorporated herein by reference).

In a one embodiment the antibody having specificity for human TNFα, comprises a heavy chain wherein the variable domain comprises a CDR having the sequence shown in SEQ ID NO:28 for CDRH1, the sequence shown in SEQ ID NO:29 or SEQ ID NO:34 for CDRH2 or the sequence shown in SEQ ID NO:30 for CDRH3.

In one embodiment the antibody comprises a light chain wherein the variable domain comprises a CDR having the sequence shown in SEQ ID NO:31 for CDRL1, the sequence shown in SEQ ID NO:32 for CDRL2 or the sequence shown in SEQ ID NO:33 for CDRL3.

The CDRs given in SEQ IDS NOS:28 and 30 to 34 referred to above are derived from a mouse monoclonal antibody hTNF40. However, SEQ ID NO:29 consists of a hybrid CDR. The hybrid CDR comprises part of heavy chain CDR2 from mouse monoclonal antibody hTNF40 (SEQ ID NO:34) and part of heavy chain CDR2 from a human group 3 germline V region sequence.

In one embodiment the antibody comprises a heavy chain wherein the variable domain comprises a CDR having the sequence shown in SEQ ID NO:28 for CDRH1, the sequence shown in SEQ ID NO:29 or SEQ ID NO:34 for CDRH2 or the sequence shown in SEQ ID NO:30 for CDRH3 and a light chain wherein the variable domain comprises a CDR having the sequence shown in SEQ ID NO:31 for CDRL1, the sequence shown in SEQ ID NO:32 for CDRL2 or the sequence shown in SEQ ID NO:33 for CDRL3.

In one embodiment the antibody comprises SEQ ID NO:28 for CDRH1, SEQ ID NO: 29 or SEQ ID NO:34 for CDRH2, SEQ ID NO:30 for CDRH3, SEQ ID NO:31 for
CDRL1, SEQ ID NO:32 for CDRL2 and SEQ ID NO:33 for CDRL3. Preferably the antibody comprises SEQ ID NO:29 for CDRH2.

The anti-TNF antibody is preferably a CDR-grafted antibody molecule. In a preferred embodiment the variable domain comprises human acceptor framework regions and non-human donor CDRs.

Preferably the antibody molecule has specificity for human TNF (formerly known as TNFa), wherein the light chain comprises the light chain variable region of SEQ ID NO: 11 and the heavy chain comprises the heavy chain variable region of SEQ ID NO: 12.

The anti-TNF antibody is preferably a Fab or Fab’ fragment.

Preferably the antibody molecule having specificity for human TNF is a Fab’ and has a light chain sequence comprising or consisting of SEQ ID NO: 13 and a heavy chain sequence comprising or consisting of SEQ ID NO: 14.

After expression, antibody fragments may be further processed, for example by conjugation to another entity such as an effector molecule.

The term effector molecule as used herein includes, for example, antineoplastic agents, drugs, toxins (such as enzymatically active toxins of bacterial or plant origin and fragments thereof e.g. ricin and fragments thereof) biologically active proteins, for example enzymes, other antibody or antibody fragments, synthetic or naturally occurring polymers, nucleic acids and fragments thereof e.g. DNA, RNA and fragments thereof, radionuclides, particularly radioiodide, radioisotopes, chelated metals, nanoparticles and reporter groups such as fluorescent compounds or compounds which may be detected by NMR or ESR spectroscopy. Effector molecular may be attached to the antibody or fragment thereof by any suitable method, for example an antibody fragment may be modified to attach at least one effector molecule as described in WO05/003171 or WO05/003170 (the contents of which are incorporated herein by reference). WO05/003171 or WO05/003170 also describe suitable effector molecules.

In one embodiment the antibody or fragment thereof, such as a Fab, is PEGylated to generate a product with the required properties, for example similar to the whole antibodies, if required. For example, the antibody may be a PEGylated anti-TNF- a Fab’, as described in WO05/094585, preferably having attached to one of the cysteine residues at the C-terminal end of the heavy chain a lysyl-maleimide-derived group wherein each of the two amino groups of the lysyl residue has covalently linked to it a methoxypoly(ethylene glycol) residue having a molecular weight of about 20,000 Da, such
that the total average molecular weight of the methoxypoly(ethyleneglycol) residues is about 40,000Da, more preferably the lysyl-maleimide-derived group is \[1-[[2-[[3-(2,5-dioxo-1-pyrrolidinyl)-1-oxopropyl]amino]ethyl]amino]-carbonyl]-1,5-pentanediyl]bis(iminocarbonyl).

The cell may also comprise further polynucleotide sequences encoding one or more further proteins of interest.

In one embodiment one or more *E.coli* host proteins that in the wild type are known to co-purify with the recombinant protein of interest during purification are selected for genetic modification, as described in Humphreys et al. "Engineering of *Escherichia coli* to improve the purification of periplasmic *Fab'* fragments: changing the pi of the chromosomally encoded PhoS/PstS protein", Protein Expression and Purification 37 (2004) 109-118 and WO04/035792 (the contents of which are incorporated herein by reference). The use of such modified host proteins improves the purification process for proteins of interest, especially antibodies, produced in *E.coli* by altering the physical properties of selected *E.coli* proteins so they no longer co-purify with the recombinant antibody. Preferably the *E.coli* protein that is altered is selected from one or more of Phosphate binding protein (PhoS/PstS), Dipeptide binding protein (DppA), Maltose binding protein (MBP) and Thioredoxin.

In one embodiment a physical property of a contaminating host protein is altered by the addition of an amino acid tag to the C-terminus or N-terminus. In a preferred embodiment the physical property that is altered is the isoelectric point and the amino acid tag is a poly-aspartic acid tag attached to the C-terminus. In one embodiment the *E.coli* proteins altered by the addition of said tag are Dipeptide binding protein (DppA), Maltose binding protein (MBP), Thioredoxin and Phosphate binding protein (PhoS/PstS). In one specific embodiment the pi of the *E.coli* Phosphate binding protein (PhoS/PstS) is reduced from 7.2 to 5.1 by the addition of a poly-aspartic acid tag (polyD), containing 6 aspartic acid residues to the C-terminus.

Also preferred is the modification of specific residues of the contaminating *E.coli* protein to alter its physical properties, either alone or in combination with the addition of N or C terminal tags. Such changes can include insertions or deletions to alter the size of the protein or amino acid substitutions to alter pi or hydrophobicity. In one embodiment these residues are located on the surface of the protein. In a preferred embodiment surface residues of the PhoS protein are altered in order to reduce the pi of the protein. Preferably
residues that have been implicated to be important in phosphate binding (Bass, US5,304,472) are avoided in order to maintain a functional PhoS protein. Preferably lysine residues that project far out of the surface of the protein or are in or near large groups of basic residues are targeted. In one embodiment, the PhoS protein has a hexa poly-aspartic acid tag attached to the C-terminus whilst surface residues at the opposite end of the molecule are targeted for substitution. Preferably selected lysine residues are substituted for glutamic acid or aspartic acid to confer a greater potential pi change than when changing neutral residues to acidic ones. The designation for a substitution mutant herein consists of a letter followed by a number followed by a letter. The first letter designates the amino acid in the wild-type protein. The number refers to the amino acid position where the amino acid substitution is being made, and the second letter designates the amino acid that is used to replace the wild-type amino acid. In preferred mutations of PhoS in the present invention lysine residues (K) 275, 107, 109, 110, 262, 265, 266, 309, 313 are substituted for glutamic acid (E) or glutamine (Q), as single or combined mutations, in addition lysine(K)318 may be substituted for aspartic acid (D) as a single or combined mutation. Preferably the single mutations are K262E, K265E and K266E. Preferably the combined mutations are K265/266E and K110/265/266E. More preferably, all mutations are combined with the polyaspartic acid (polyD) tag attached at the C-terminus and optionally also with the K318D substitution. In a preferred embodiment the mutations result in a reduction in pi of at least 2 units. Preferably the mutations of the present invention reduce the pi of PhoS from 7.2 to between about 4 and about 5.5. In one embodiment of the present invention the pi of the PhoS protein of E.coli is reduced from 7.2 to about 4.9, about 4.8 and about 4.5 using the mutations polyD K318D, polyD K265/266E and polyD K110/265/266E respectively.

The polynucleotide encoding the protein of interest may be expressed as a fusion with another polypeptide, preferably a signal sequence or other polypeptide having a specific cleavage site at the N-terminus of the mature polypeptide. The heterologous signal sequence selected should be one that is recognized and processed by the host cell. For prokaryotic host cells that do not recognize and process the native or a eukaryotic polypeptide signal sequence, the signal sequence is substituted by a prokaryotic signal sequence. Suitable signal sequences include OmpA, PhoA, LamB, PelB, DsbA and DsbC.

Embodiments of the invention described herein with reference to the polynucleotide apply equally to alternative embodiments of the invention, for example
vectors, expression cassettes and/or host cells comprising the components employed therein, as far as the relevant aspect can be applied to same.

The present invention also provides a method for producing a recombinant protein of interest comprising culturing a recombinant gram-negative bacterial cell as described above in a culture medium under conditions effective to express the recombinant protein of interest and recovering the recombinant protein of interest from the periplasm of the recombinant gram-negative bacterial cell and/or the culture medium. In one embodiment wherein the cell comprises a recombinant polynucleotide encoding DsbC, the cell is cultured under conditions effective to express the recombinant polynucleotide encoding DsbC.

The gram negative bacterial cell and protein of interest preferably employed in the method of the present invention are described in detail above.

When the polynucleotide encoding the protein of interest is exogenous the polynucleotide may be incorporated into the host cell using any suitable means known in the art. The polynucleotide sequence encoding DsbC may also be incorporated into the host cell using any suitable means known in the art. Typically, the polynucleotide is incorporated as part of an expression vector which is transformed into the cell. Accordingly, in one aspect the cell according to the present invention comprises an expression cassette comprising the polynucleotide encoding the protein of interest and an expression cassette comprising the polynucleotide encoding DsbC.

The polynucleotide sequence encoding the protein of interest and the polynucleotide sequence encoding DsbC can be transformed into a cell using standard techniques, for example employing rubidium chloride, PEG or electroporation.

The method according to the present invention may also employ a selection system to facilitate selection of stable cells which have been successfully transformed with the polynucleotide encoding the protein of interest. The selection system typically employs co-transformation of a polynucleotide sequence encoding a selection marker. In one embodiment, each polynucleotide transformed into the cell further comprises a polynucleotide sequence encoding one or more selection markers. Accordingly, the transformation of the polynucleotide encoding the protein of interest and optionally the polynucleotide encoding DsbC and the one or more polynucleotides encoding the marker occurs together and the selection system can be employed to select those cells which produce the desired proteins.
Cells able to express the one or more markers are able to survive/grow/multiply under certain artificially imposed conditions, for example the addition of a toxin or antibiotic, because of the properties endowed by the polypeptide/gene or polypeptide component of the selection system incorporated therein (e.g. antibiotic resistance). Those cells that cannot express the one or more markers are not able to survive/grow/multiply in the artificially imposed conditions. The artificially imposed conditions can be chosen to be more or less vigorous, as required.

Any suitable selection system may be employed in the present invention. Typically the selection system may be based on including in the vector one or more genes that provides resistance to a known antibiotic, for example a tetracycline, chloramphenicol, kanamycin or ampicillin resistance gene. Cells that grow in the presence of a relevant antibiotic can be selected as they express both the gene that gives resistance to the antibiotic and the desired protein.

An inducible expression system or a constitutive promoter may be used in the present invention to express the protein of interest and/or the DsbC. In one embodiment, the expression of the polynucleotide sequence encoding a protein of interest and the recombinant polynucleotide encoding DsbC is induced by adding an inducer to the culture medium. Suitable inducible expression systems and constitutive promoters are well known in the art.

Any suitable medium may be used to culture the transformed cell. The medium may be adapted for a specific selection system, for example the medium may comprise an antibiotic, to allow only those cells which have been successfully transformed to grow in the medium.

The cells obtained from the medium may be subjected to further screening and/or purification as required. The method may further comprise one or more steps to extract and purify the protein of interest as required.

The polypeptide may be recovered from the strain, including from the cytoplasm, periplasm and/or supernatant.

The specific method (s) used to purify a protein depends on the type of protein. Suitable methods include fractionation on immuno-affinity or ion-exchange columns; ethanol precipitation; reversed-phase HPLC; hydrophobic-interaction chromatography; chromatography on silica; chromatography on an ion-exchange resin such as S-
SEPHEROSE and DEAE; chromatofocusing; ammonium-sulfate precipitation; and gel filtration.

In one embodiment the method further comprises separating the recombinant protein of interest from DsbC.

Antibodies may be suitably separated from the culture medium and/or cytoplasm extract and/or periplasm extract by conventional antibody purification procedures such as, for example, protein A-Sepharose, protein G chromatography, protein L chromatography, thiophilic, mixed mode resins, His-tag, FLAGTag, hydroxylapatite chromatography, gel electrophoresis, dialysis, affinity chromatography, Ammonium sulphate, ethanol or PEG fractionation/precipitation, ion exchange membranes, expanded bed adsorption chromatography (EBA) or simulated moving bed chromatography.

The method may also include a further step of measuring the quantity of expression of the protein of interest and selecting cells having high expression levels of the protein of interest.

The method may also including one or more further downstream processing steps such as PEGylation of the protein of interest, such as an antibody or antibody fragment.

One or more method steps described herein may be performed in combination in a suitable container such as a bioreactor.

Examples
Example 1 - Generation Cell Strain MXE001 (ΔTsp)
The MXE001 strain was generated as follows:

The Tsp cassette was moved as Sal I, Not I restriction fragments into similarly restricted pK03 plasmids. The pK03 plasmid uses the temperature sensitive mutant of the pSC101 origin of replication (RepA) along with a chloramphenicol marker to force and select for chromosomal integration events. The sacB gene which encodes for levansucrase is lethal to E. coli grown on sucrose and hence (along with the chloramphenicol marker and pSC101 origin) is used to force and select for de-integration and plasmid curing events.

This methodology had been described previously (Hamilton et al., 1989, Journal of Bacteriology, 111, 4617-4622 and Blomfield et al, 1991, Molecular Microbiology, 5, 1447-1457). The pK03 system removes all selective markers from the host genome except for the inserted gene.
The following plasmids were constructed.

pMXE191 comprising the knockout mutated Tsp gene as shown in the SEQ ID NO: 3 comprising EcoR I and Ase I restriction markers.


**Day 1** 40μl of E.coli cells were mixed with (1Opg) 1μl of pK03 DNA in a chilled BioRad 0.2cm electroporation cuvette before electroporation at 2500V, 25μF and 200Ω. 1000μl of 2xPY was added immediately, the cells recovered by shaking at 250rpm in an incubator at 30°C for 1 hour. Cells were serially 1/10 diluted in 2xPY before 100μl aliquots were plated out onto 2xPY agar plates containing chloramphenicol at 20μg/ml prewarmed at 30°C and 43°C. Plates were incubated overnight at 30°C and 43°C.

**Day 2** The number of colonies grown at 30°C gave an estimate of the efficiency of electroporation whilst colonies that survive growth at 43°C represent potential integration events. Single colonies from the 43°C plate were picked and resuspended in 10ml of 2xPY. 100μl of this was plated out onto 2xPY agar plates containing 5% (w/v) sucrose pre-warmed to 30°C to generate single colonies. Plates were incubated overnight at 30°C.

**Day 3** Colonies here represent potential simultaneous de-integration and plasmid curing events. If the de-integration and curing events happened early on in the growth, then the bulk of the colony mass will be clonal. Single colonies were picked and replica plated onto 2xPY agar that contained either chloramphenicol at 20μg/ml or 5% (w/v) sucrose. Plates were incubated overnight at 30°C.

**Day 4** Colonies that both grow on sucrose and die on chloramphenicol represent potential chromosomal replacement and plasmid curing events. These were picked and screened by PCR with a mutation specific oligonucleotide. Colonies that generated a positive PCR band of the correct size were struck out to produce single colonies on 2xPY agar containing 5% (w/v) sucrose and the plates were incubated overnight at 30°C.
Day 5 Single colonies of PCR positive, chloramphenicol sensitive and sucrose resistant *E. coli* were used to make glycerol stocks, chemically competent cells and act as PCR templates for a PCR reaction with 5' and 3' flanking oligos to generate PCR product for direct DNA sequencing using **Taq** polymerase.

Cell strain MXE001 was tested to confirm successful modification of genomic DNA carrying the mutated Tsp gene by PCR amplification of the region of the Tsp gene comprising a non-naturally occurring *Ase I* restriction site, as shown in Figures 1a, 1b and 1c, using oligonucleotides primers. The amplified regions of the DNA were then analyzed by gel electrophoresis before and after incubation with *Ase I* restriction enzyme to confirm the presence of the non-naturally occurring *Ase I* restriction site in the mutated genes. This method was carried out as follows:

The following oligos were used to amplify, using PCR, genomic DNA from prepared *E. coli* cell lysates from MXE001 and W3110:

- **6284 Tsp 3'** 5'-GCATCATATAATTTTCACCT-3' (SEQ ID NO: 15)
- **6283 Tsp 5'** 5'-GGGAATGAACCTGAGC-3' (SEQ ID NO: 16)

The lysates were prepared by heating a single colony of cells for 10 minutes at 95 °C in 20ul of 1x PCR buffer. The mixture was allowed to cool to room temperature then centrifugation at 13,200rpm for 10 minutes. The supernatant was removed and labeled as 'cell lysate'.

Each strain was amplified using the Tsp oligos pair.

The DNA was amplified using a standard PCR procedure.

- 5ul Buffer x1O (Roche)
- 1ul dNTP mix (Roche, 10mM mix)
- 1.5ul 5' oligo (5 pmol)
- 1.5ul 3' oligo (5 pmol)
- 2ul Cell lysate
- 0.5ul Taq DNA polymerase (Roche 5U/ul)
38.5ul  H20

PCR cycle.
94 °C  1 minute
5
94 °C  1 minute)  repeated for 30 cycles
55 °C  1 minute)
72 °C  1 minute)
72 °C  10 minutes

Once the reactions were complete 25ul was removed to a new microfuge tube for
digestion with Ase I. To the 25ul of PCR reaction 19ul of H20, 5ul of buffer 3 (NEB), 1ul
of Ase I (NEB) was added, mixed and incubated at 37 °C for 2 hours.

To the remaining PCR reaction 5ul of loading buffer (x6) was added and 20ul was loaded
onto a 0.8% TAE 200ml agarose gel (Invitrogen) plus Ethidium Bromide (5ul of 10mg/ml
stock) and run at 100 volts for 1 hour. 10ul of size marker (Perfect DNA marker 0.1-12Kb,
Novagen) was loaded in the final lane.

Once the Ase I digestions were complete 10ul of loading buffer (x6) was added and
20ul was loaded onto a 0.8% TAE agarose gel (Invitrogen) plus Ethidium Bromide (5ul of
10mg/ml stock) and run at 100 volts for 1 hour. 10ul of size marker (Perfect DNA marker
0.1-12Kb, Novagen) was loaded in the final lane. Both gels were visualized using UV
transluminator.

The genomic fragment amplified showed the correct sized band of 2.8Kb for Tsp.
Following digestion with Ase I this confirmed the presence of the introduced Ase I sites in
the Tsp deficient strain MXE001 but not in the W3110 control.

MXE001: genomic DNA amplified using the Tsp primer set and the resulting DNA was
digested with Ase I to produce 2.2 and 0.6 Kbps bands.

W3110 PCR amplified DNA was not digested by Ase I restriction enzyme.
Example 2 - Generation of spr mutants

The spr mutations were generated and selected for using a complementation assay.

The spr gene was mutated using the Clontech® random mutagenesis diversity PCR kit which introduced 1 to 2 mutations per 1000bp. The mutated spr PCR DNA was cloned into an inducible expression vector [pTTO CDP870] which expresses CDP870 Fab' along with the spr mutant. This ligation was then electro-transformed into an E.coli strain MXEOOl (ATsp) prepared using the method found in Miller, E.M. and Nickoloff, J.A., "Escherichia coli electrotransformation," in Methods in Molecular Biology, vol. 47, Nickoloff, J.A. (ed.), Humana Press, Totowa, NJ, 105 (1995). The following protocol was used, 40ul of electro competent MXEOOl, 2.5ul of the ligation (100pg of DNA) was added to a 0.2cm electroporation cuvette, electro-transformation was performed using as BioRad GenePulser Xcell with the following conditions, 2500V, 25μF and 200Ω. After the electro-transformation 1ml of SOC (Invitrogen) (pre-warmed to 37 °C) was added and the cells left to recover at 37 °C for 1 hour with gentle agitation.

The cells where plated onto Hypotonic agar (5g/L Yeast extract, 2.5g/L Tryptone, 15g/L Agar (all Difco)) and incubated at 40 °C. Cells which formed colonies were re-plated onto HLB at 43 °C to confirm restoration of the ability to grow under low osmotic conditions at high temperature to the MXEOOl strain. Plasmid DNA was prepared from the selected clones and sequenced to identify spr mutations.

Using this method eight single, one double mutation and two multiple mutations in the spr protein were isolated which complemented the ATsp phenotype as follows:

1. V98E
2. D133A
3. V135D
4. V135G
5. G147C
6. S95F and Y115F
7. I70T
8. N31T, Q73R, R100G, G140C
9. R62C, Q99P, R144C
Example 3 - Generation of Mutant *E. coli* cell strains carrying spr mutations

The individual mutations 1 to 5 identified in Example 2 and three catalytic triad mutations of spr (C94A, H145A, H157A) and W174R were used to generate new strains using either the wild-type W3110 *E. coli* strain (genotype: F- LAM- IN (rrnD-rrnE) rphl (ATCC no. 27325)) to create spr mutated strains carrying a wild-type non-recombinant chromosomal Tsp gene or MXEO01 (ATsp) strain from Example 1 to make combined ATsp/mutant spr strains.

The following mutant *E. coli* cell strains were generated using a gene replacement vector system using the pK03 homologous recombination/replacement plasmid (Link *et al.*, 1997, *Journal of Bacteriology*, 179, 6228-6237), as described in Example 1 for the generation of MXE001.

<table>
<thead>
<tr>
<th>Mutant Strain</th>
<th>E. coli Cell Strain</th>
<th>Genotype</th>
<th>Spr Vectors</th>
</tr>
</thead>
<tbody>
<tr>
<td>MXE001</td>
<td>ATsp</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>MXE008</td>
<td>ATsp, spr D133A</td>
<td>pMXE339, pK03 spr D133A (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE009</td>
<td>ATsp, spr H157A</td>
<td>pMXE345, pK03 spr H157A (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE010</td>
<td>spr G147C</td>
<td>pMXE338, pK03 spr G147C (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE011</td>
<td>spr C94A</td>
<td>pMXE343, pK03 spr C94A (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE012</td>
<td>spr H145A</td>
<td>pMXE344, pK03 spr H145A (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE013</td>
<td>spr W174R</td>
<td>pMXE346, pK03 spr W174R (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE014</td>
<td>ATsp, spr V135D</td>
<td>pMXE340, pK03 spr V135D (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE015</td>
<td>ATsp, spr V98E</td>
<td>pMXE342, pK03 spr V98E (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE016</td>
<td>ATsp, spr C94A</td>
<td>pMXE343, pK03 spr C94A (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE017</td>
<td>ATsp, spr H145A</td>
<td>pMXE344, pK03 spr H145A (-Sail)</td>
<td></td>
</tr>
<tr>
<td>MXE018</td>
<td>ATsp, spr V135G</td>
<td>pMXE341, pK03 spr V135G (-Sail)</td>
<td></td>
</tr>
</tbody>
</table>
The mutant spr integration cassettes were moved as Sal I, Not I restriction fragments into similarly restricted pK03 plasmids.

The plasmid uses the temperature sensitive mutant of the pSClOl origin of replication (RepA) along with a chloramphenicol marker to force and select for chromosomal integration events. The sacB gene which encodes for levensucrase is lethal to E. coli grown on sucrose and hence (along with the chloramphenicol marker and pSClOl origin) is used to force and select for de-integration and plasmid curing events. This methodology had been described previously (Hamilton et al., 1989, Journal of Bacteriology, 171, 4617-4622 and Blomfield et al., 1991, Molecular Microbiology, 5, 1447-1457). The pK03 system removes all selective markers from the host genome except for the inserted gene.

The following pK03 vectors were constructed, comprising the mutated spr genes including a silent mutation within the spr sequence which removes a Sail restriction site for clone identification.

- pMXE336, pK03 spr S95F (−Sail)
- pMXE337, pK03 spr Y115F (−Sail)
- pMXE338, pK03 spr G147C (−Sail)
- pMXE339, pK03 spr D133A (−Sail)
- pMXE340, pK03 spr V135D (−Sail)
- pMXE341, pK03 spr V135G (−Sail)
- pMXE342, pK03 spr V98E (−Sail)
- pMXE343, pK03 spr C94A (−Sail)
- pMXE344, pK03 spr H145A (−Sail)
- pMXE345, pK03 spr H157A (−Sail)
- pMXE346, pK03 spr W174R (−Sail)

These plasmids were then transformed into chemically competent E. coli W3110 cells prepared using the method found in Miller, E.M. and Nickoloff, J.A., "Escherichia coli electrotransformation," in Methods in Molecular Biology, vol. 47, Nickoloff, J.A. (ed.), Humana Press, Totowa, NJ, 105 (1995) or into MXEOO1 strain from Example 1 to make combined ATsp/mutant spr strains, as shown in Table 1.
**Day 1** 40μl of electro-competent *E.coli* cells or MXE001 cells were mixed with (10pg) 1μl of pK03 DNA in a chilled BioRad 0.2cm electroporation cuvette before electroporation at 2500V, 25μF and 200Ω. 100μl of 2xPY was added immediately, the cells recovered by shaking at 250rpm in an incubator at 30°C for 1 hour. Cells were serially 1/10 diluted in 2xPY before 100μl aliquots were plated out onto 2xPY agar plates containing chloramphenicol at 20μg/ml prewarmed at 30°C and 43°C. Plates were incubated overnight at 30°C and 43°C.

**Day 2** The number of colonies grown at 30°C gave an estimate of the efficiency of electroporation whilst colonies that survive growth at 43°C represent potential integration events. Single colonies from the 43°C plate were picked and resuspended in 10ml of 2xPY. 100μl of this was plated out onto 2xPY agar plates containing 5% (w/v) sucrose pre-warmed to 30°C to generate single colonies. Plates were incubated overnight at 30°C.

**Day 3** Colonies here represent potential simultaneous de-integration and plasmid curing events. If the de-integration and curing events happened early on in the growth, then the bulk of the colony mass will be clonal. Single colonies were picked and replica plated onto 2xPY agar that contained either chloramphenicol at 20μg/ml or 5% (w/v) sucrose. Plates were incubated overnight at 30°C.

**Day 4** Colonies that both grow on sucrose and die on chloramphenicol represent potential chromosomal replacement and plasmid curing events. These were picked and screened by PCR plus restriction digest for the loss of a Sail site. Colonies that generated a positive PCR band of the correct size and resistance to digestion by Sail were struck out to produce single colonies on 2xPY agar containing 5% (w/v) sucrose and the plates were incubated overnight at 30°C.

**Day 5** Single colonies of PCR positive, chloramphenicol sensitive and sucrose resistant *E. coli* were used to make glycerol stocks, chemically competent cells and act as PCR templates for a PCR reaction with 5' and 3' flanking oligos to generate PCR product for direct DNA sequencing using Taq polymerase to confirm the correct mutation.
Example 4 - Generation of plasmid for Fab' and DsbC co-expression

A plasmid was constructed containing both the heavy and light chain sequences of an anti-TNF Fab' (an anti-TNF Fab' having a light chain sequence shown in SEQ ID NO: 13 and a heavy chain sequence shown in SEQ ID NO: 14) and the sequence encoding DsbC.

A dicistronic message was created of the anti-TNFa Fab' fragment (referred to as CDP870) described in WO 1/94585. The upstream cistron encoded the light chain of the antibody (SEQ ID NO: 13) whilst the downstream cistron encoded the heavy chain of the antibody (SEQ ID NO: 14). A DNA sequence encoding the OmpA signal peptide was fused to the 5' end of the DNA coding for each of the light chain and the heavy chain to allow efficient secretion to the periplasm. The intergenic sequence (IGS2) was used as shown in SEQ ID NO: 37.

Plasmid pDPH358 (pTTO 40.4 CDP870 IGS2), an expression vector for the CDP870 Fab' (an anti-TNF Fab') and DsbC (a periplasmic polypeptide), was constructed using conventional restriction cloning methodologies which can be found in Sambrook et al 1989, Molecular cloning: a laboratory manual. CSHL press, N.Y. The plasmid pDPH358 contained the following features; a strong tac promoter and lac operator sequence. As shown in Figure 10, the plasmid contained a unique EcoRI restriction site after the coding region of the Fab' heavy chain, followed by a non-coding sequence and then a unique Ndel restriction site. The DsbC gene was PCR cloned using W3I 10 crude chromosomal DNA as a template such that the PCR product encoded for a 5' EcoRI site followed by a strong ribosome binding, followed by the native start codon, signal sequence and mature sequence of DsbC, terminating in a C-terminal His tag and finally a non-coding Ndel site.

The EcoRI-Ndel PCR fragment was restricted and ligated into the expression vector such that all three polypeptides: Fab' light chain, Fab' heavy chain and DsbC were encoded on a single polycistronic mRNA.

The Fab light chain, heavy chain genes and DcbC gene were transcribed as a single polycistronic message. DNA encoding the signal peptide from the E. coli OmpA protein was fused to the 5' end of both light and heavy chain gene sequences, which directed the translocation of the polypeptides to the E. coli periplasm. Transcription was terminated using a dual transcription terminator rrnB tlt2. The lacIq gene encoded the constitutively
expressed Lac I repressor protein. This repressed transcription from the tac promoter until de-repression was induced by the presence of allolactose or IPTG. The origin of replication used was pi5A, which maintained a low copy number. The plasmid contained a tetracycline resistance gene for antibiotic selection.

Example 5 - Expression of anti-TNF Fab' or anti-TNF Fab' and DsbC in the E.coli strains

Expression of anti-TNF Fab' and DsbC

The wild-type W31 10 cell line, the MXE001 strain provided in Example 1 and the mutant strain MXE012 (HI45A spr mutant strain) provided in Example 3 were transformed with the plasmid generated in Example 4.

The transformation of the strains was carried out using the method found in Chung C.T et al Transformation and storage of bacterial cells in the same solution. PNAS 86:2172-2175 (1989).

Expression of anti-TNF Fab'

The wild-type W31 10 cell line, spr mutant strains MXE008, MXE012, MXE017 and MXE012 (HI45A spr mutant strain) provided in Example 3 and the MXE001 strain provided in Example 1 were transformed with plasmid pMXEl 17 (pTTO CDP870 or 40.4 IGS17), an expression vector for the CDP870 Fab' (an anti-TNF Fab' having a light chain sequence shown in SEQ ID NO: 13 and a heavy chain sequence shown in SEQ ID NO: 14), was constructed using conventional restriction cloning methodologies which can be found in Sambrook et al 1989, Molecular cloning: a laboratory manual. CSHL press, N.Y.

The plasmid pMXEl 17 (pTTO CDP870 or 40.4 IGS17) contained the following features; a strong tac promoter and lac operator sequence. The Fab light and heavy chain genes were transcribed as a single dicistronic message. DNA encoding the signal peptide from the E. coli OmpA protein was fused to the 5' end of both light and heavy chain gene sequences, which directed the translocation of the polypeptides to the E. coli periplasm.

Transcription was terminated using a dual transcription terminator rRNA tt2. The lacIq gene encoded the constitutively expressed Lac I repressor protein. This repressed transcription from the tac promoter until de-repression was induced by the presence of
aison lactose or IPTG. The origin of replication used was pI5A, which maintained a low copy number. The plasmid contained a tetracycline resistance gene for antibiotic selection.

The transformation of the strains was carried out using the method found in Chung C.T et al Transformation and storage of bacterial cells in the same solution. PNAS 86:2172-2175 (1989).

Example 6 - Expression of an anti-TNF Fab' in mutated E. coli strains using shake flask cultures

The following strains as produced by Example 5 expressing anti-TNF Fab': W3110, MXE001, MXE012 and MXE017 were tested in a shake flask experiment comparing growth and expression of the Fab'.

The shake flask experimental protocol used was performed as follows:

5ml Shake flask experiment
A single colony was picked into 5ml LB plus tetracycline at 1Oug/ml and grown overnight at 30°C with shaking at 250rpm.

The overnight culture was used to inoculate 100ml plus tetracycline to 0.1 OD600. (i.e. for OD of 4, 100/4x01 = 2.5mls in 100ml.)

3x5ml culture tubes were set up for every time point required using this master culture. A reference culture was set up to sample for OD measurement.

The cultures were shaken at 30°C 250rpm monitoring growth visually at first, then by sampling the reference culture to catch cultures at 0.5 OD600 (usually about 2hrs). IPTG was added to each culture tube to a concentration of 200uM (25ul of 0.04M) once the culture had achieved an OD greater than 0.5.

The culture tubes were removed at the required time points e.g. 1hr, 2hr, post induction and kept on ice.

After centrifugation at 13,200rpm for 5 minutes the cell pellet was re-suspended in 200ul of periplasmic extraction buffer (100mM Tris.Cl/10mM EDTA pH 7.4). Periplasmic extracts were agitated at 250rpm over night at 30°C. The next day, the extracts were centrifuged for 10 minutes at 13,200 rpm, the supernatant decanted off and stored at -20°C as 'periplasmic extract'. The spent cell pellet was discarded.
ELISA quantification.

96 well ELISA plates were coated overnight at 4°C with AB141 (rabbit anti-human CHI, UCB) at 2 µg/ml in PBS. After washing 3x with 300ul of sample/conjugate buffer (PBS, BSA 0.2% (w/v), Tween 20 0.1% (v/v)), serial ½ dilutions of samples and standards were performed on the plate in 100 µl of sample/conjugate buffer, and the plate agitated at 250 r.p.m at room temperature for 1 hour. After washing 3x with 300ul of wash buffer (PBS, Tween 20 0.1% (v/v)), 100 µl of the revealing antibody 6062 (rabbit anti-human kappa HRP conjugated, The Binding Site, Birmingham, U.K.) was added, after dilution at 1/1000 in sample/conjugate buffer. The plate was then agitated at 250 r.p.m at room temperature for 1 hour. After washing with 3x 300ul of wash buffer, 100 µl of TMB substrate was added (50:50 mix of TMB solution (Calbiochem): dH2O) and the A630 recorded using an automated plate reader. The concentration of Fab' in the periplasmic extracts were calculated by comparison with purified Fab' standards of the appropriate isotype.

Figure 1 shows the improved growth of MXE012 and MXE017 compared to the wild-type W3110 and MXE001.

Figure 2 shows improved expression of the Fab' in MXE012 and MXE017 compared to the wild-type W3110 and MXE001.

Example 7 - Growth of *E. coli* strains expressing; anti-TNF Fab' or anti-TNF Fab' and DsbC using high density fermentations

The following strains, as produced by example 5 were tested in fermentation experiments comparing growth and expression of an anti-TNFα Fab':

Strains expressing anti-TNF Fab' produced in Example 5:
W3100

MXE012 (H145A spr mutant strain)

Strains expressing anti-TNF Fab' and DsbC produced in Example 5:
W3110
Growth medium.

The fermentation growth medium was based on SM6E medium (described in Humphreys et al., 2002, Protein Expression and Purification, 26, 309-320) with 3.86 g/l NaH₂PO₄·H₂O and 112 g/l glycerol.

Inoculum. Inoculum cultures were grown in the same medium supplemented with 10 μg/ml tetracycline. Cultures were incubated at 30°C with agitation for approximately 22 hours.

Fermentation. Fermenters (2.5 litres total volume) were seeded with inoculum culture to 0.3-0.5 OD₆₀₀. Temperature was maintained at 30°C during the growth phase and was reduced to 25°C prior to induction. The dissolved oxygen concentration was maintained above 30% air saturation by variable agitation and airflow. Culture pH was controlled at 7.0 by automatic titration with 15% (v/v) NH₄OH and 10% (v/v) cone. H₂SO₄. Foaming was controlled by the addition of 10% (v/v) Struktol J673 solution (Schill and Seilacher).

A number of additions were made at different stages of the fermentation. When biomass concentration reached approximately 40 OD₆₀₀, magnesium salts and NaH₂PO₄·H₂O were added. Further additions of NaH₂PO₄·H₂O were made prior to and during the induction phase to ensure phosphate was maintained in excess. When the glycerol present at the beginning of fermentation had depleted (approximately 75 OD₆₀₀), a continuous feed of 80% (w/w) glycerol was applied. At the same point in the fermentation an IPTG feed at 170μM was applied. The start of IPTG feeding was taken as the start of induction. Fermentations were typically run for 64-120 hours at glycerol feed rates (ranging between 0.5 and 2.5 ml/h).

Measurement of biomass concentration and growth rate. Biomass concentration was determined by measuring the optical density of cultures at 600 nm.

Periplasmic Extraction. Cells were collected from culture samples by centrifugation. The supernatant fraction was retained (at -20°C) for further analysis. The cell pellet fraction was resuspended to the original culture volume in extraction buffer (100 mM Tris-Cl, 10 mM EDTA; pH 7.4). Following incubation at 60°C for approximately 16 hours the extract was clarified by centrifugation and the supernatant fraction retained (at -20°C) for analysis.
**Fab' quantification.** Fab' concentrations in periplasmic extracts and culture supernatants were determined by Fab' assembly ELISA as described in Humphreys et al., 2002, Protein Expression and Purification, 26, 309-320 and using Protein G hplc. A HiTrap Protein-G HP 1ml column (GE-Healthcare or equivalent) was loaded with analyte (approximately neutral pH, 30°C, 0.2µm filtered) at 2ml/min, the column was washed with 20mM phosphate, 50mM NaCl pH 7.4 and then Fab' eluted using an injection of 50mM Glycine/HCl pH 2.7. Eluted Fab' was measured by A280 on a Agilent 1100 or 1200 HPLC system and quantified by reference to a standard curve of a purified Fab' protein of known concentration.

Figure 3 shows the growth profile of W3110 and MXE012 expressing anti-TNF Fab' during fermentation with an extended run time. The data illustrates a small increase in initial growth rate of the spr strain relative to wild type during biomass accumulation and increased duration of survival of the spr mutant strain MXE012 relative to wild type strain W3110 in the last -20 hours of the fermentation.

Figure 4 shows periplasmic Fab' accumulation (filled lines and symbols) and media Fab' accumulation (dashed lines and open symbols) for W3110 and MXE012 (W3110 spr H145A) expressing anti-TNF Fab'during fermentation with an extended run time. The data show that the initial rates of periplasmic Fab' accumulation are very similar for the two strains, but that the wild type W3110 cells leak periplasmic Fab' later in the fermentation compared to MXE012.

Figure 5 shows the growth profile of anti-TNFα Fab' expressing strains W3110 and MXE012 and of anti-TNFα Fab' and recombinant DsbC expressing strains W3110 and MXE012. It can be seen that the strains expressing DsbC exhibit improved growth compared to the corresponding cell strains which do not express recombinant DsbC. It can also be seen that the presence of the spr mutation in the strains improves cell growth.

Figure 6 shows total Fab yield from the periplasm (shaded symbols) and supernatant (open unshaded symbols) from anti-TNFα Fab' expressing *E. coli* strains W3110 and MXE012 and from anti-TNFα Fab' and recombinant DsbC expressing *E. coli* strains W3110 and MXE012. It can be seen from this graph that the strains expressing recombinant DsbC
produced a high yield of anti-TNFα Fab' with strain MXE012 producing over 3.0 g/L in approximately 92 hours. It can also be seen that the MXE012 strains carrying a mutant spr gene exhibited reduced lysis compared to the W3110 strains which can be seen as less supernatant anti-TNFα Fab' (open symbols).

Example 8 - Determination of DNA leakage and total protein quantity in strains

dsDNA assay:
The double-stranded DNA leakage into the supernatant of strains W3110, MXE001, MXE008 and MXE012 was determined using the Quant-IT Picogreen dsDNA assay kit (Invitrogen, Ref: PI 1496). A standard curve was prepared by diluting the DNA standard provided in the range of 1-1000 ng/mL. Samples were diluted in TE buffer, so that the fluorescence reading fell within the linear range of the method (500 to 1000 times). In a 96-well plate, 100 µL of diluted sample or standard were mixed with 100 µL of the Picogreen reagent, and the plate was incubated for 5 minutes at room temperature, protected from light. The fluorescence counts were measured for 0.1s using a 485nm excitation filter, and a 535nm emission filter. The results are shown in Figure 7.

Protein Assay:
The total proteins concentration of strains W3110, MXE001, MXE008 and MXE012 was determined using the Coomassie Plus Bradford assay kit (Pierce, Ref: 23236). A standard curve was made by diluting Bovine Serum Albumin standard over a range of 25-1000µg/mL. Samples were diluted in water so that the optical density fell within the linear range of the method (5 to 10 times), and 33 µL of sample or standard were mixed with 1 mL of coomassie reagent. After incubating for 10 minutes at room temperature, the \( \text{OD}_{535} \) was read on a spectrophotometer with coomassie reagent as a blank. The total proteins concentration was calculated based on the standard curve. The results are shown in Figure 8.

Example 9 Growth of \textit{E. coli} strains expressing anti-TNF Fab' and DsbC using large scale fermentations

The following strain, as produced by example 5 was tested in fermentation experiments comparing growth and viability of the strain and the expression of an anti-TNFα Fab'.
MXE012 (spr H145A mutant) expressing anti-TNF Fab' and DsbC produced in Example 5

The fermentations were carried out as follows:

The MXE012 expressing anti-TNF Fab' and DsbC cells were grown initially using a complex medium of yeast extract and peptone in shake flask culture. The cells were then transferred to a seed stage fermenter using a chemically defined medium. The cells were grown under non-nutrient limiting conditions until a defined transfer point. The cells were then transferred to a 250L production fermenter using a similar chemically defined medium with a final volume of approximately 230L. The culture was initially grown in batch mode until carbon source depletion. After carbon source depletion a feed limiting the carbon source was fed at an exponentially increasing rate. After the addition of a defined quantity of carbon source the rate of feed solution addition was decreased and IPTG was added to induce expression of the Fab'. The fermentation was then continued and the Fab' accumulated in the periplasm. At a defined period after induction the culture was harvested by centrifugation and the Fab' was extracted from the cells by resuspending the harvested cells in a Tris and EDTA buffer and heating to 59°C.

The growth profiles of the fermentations were determined by measuring the optical density of culture at 600 nm.

The Fab' titres were determined by Protein G HPLC as described in Example 7 above except that during the periplasmic extraction fresh cells were used and 1mL of extraction buffer was added to the cell culture. The supernatant and periplasmic Fab' were measured as described in Example 7. Figure 12 shows the periplasmic Fab' titre.

The cell culture viability was measured by flow cytometry using Fluorescence-Activated Cell Sorting.

Figure 11 shows the growth profiles of 200L fermentations of anti-TNFa Fab' and recombinant DsbC expressing strain MXE012.
Figure 12 shows the periplasmic anti-TNFα Fab' titres of 200L fermentations of anti-TNFα Fab' and recombinant DsbC expressing strain MXE012.

Figure 13 shows the viabilities of 200L fermentations of anti-TNFα Fab' and recombinant DsbC expressing strain MXE012.

Example 10 Growth of *E. coli* strains expressing anti-TNF Fab' and DsbC using large scale fermentations

The following strain, as produced by example 5 was tested in fermentation experiments comparing growth of the strain and the expression of an anti-TNFα Fab':

MXE012 expressing anti-TNF Fab' and DsbC produced in Example 5

The fermentations were carried out as described in Example 9 with a 3000L production fermenter containing a final volume of approximately 2650L.

The growth profiles of the fermentations were determined by measuring the optical density of culture at 600 nm.

The Fab' titres were determined by Protein G HPLC as described in Example 9 above.

Figure 14 shows the growth profiles of 3000L fermentations of anti-TNFα Fab' and recombinant DsbC expressing strain MXE012.

Figure 15 shows the periplasmic anti-TNFα Fab' titres of 3000L fermentations of anti-TNFα Fab' and recombinant DsbC expressing strain MXE012.

While this invention has been particularly shown and described with reference to preferred embodiments, it will be understood to those skilled in the art that various changes in form and detail may be made without departing from the scope of the invention as defined by the appendant claims.
Claims:

1. A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene.

2. A cell according to claim 1, wherein the mutant spr gene encodes a spr protein having a mutation at one or more amino acids selected from H145, N31, R62, 170, Q73, C94, S95, V98, Q99, R100, L108, Y115, D133, V135, L136, G140, R144, G147, H157 and W174.


4. A cell according to claim 3, wherein the one or more spr protein mutations are selected from S95F, V98E, Y115F, D133A, V135D, V135G and G147C.

5. A cell according to claim 4, wherein the mutant spr gene encodes a spr protein having the mutations S95F and Y115F.

6. A cell according to claim 3, wherein the spr protein mutation is H145A.

7. A cell according to any preceding claim, wherein the cell is isogenic to a wild-type bacterial cell except for the mutated spr gene.

8. A cell according to any preceding claim, wherein the cell further comprises a recombinant polynucleotide encoding DsbC.

9. A cell according to any of claims 1 to 8, wherein the cell further comprises one of more of the following mutated genes:

   a. a mutated DegP gene encoding a DegP protein having chaperone activity and reduced protease activity;
b. a mutated ptr gene, wherein the mutated ptr gene encodes a Protease III protein having reduced protease activity or is a knockout mutated ptr gene; and

c. a mutated OmpT gene, wherein the mutated OmpT gene encodes a OmpT protein having reduced protease activity or is a knockout mutated OmpT gene.

10. A cell according to any preceding claim, wherein the cell is E. coli.

11. A cell according to any preceding claim, wherein the cell comprises a polynucleotide sequence encoding a protein of interest.

12. A cell according to claim 11, wherein the cell comprises a vector comprising the recombinant polynucleotide encoding DsbC and the polynucleotide sequence encoding a protein of interest.

13. A cell according to claim 12, wherein the vector comprises a promoter which controls the expression of the recombinant polynucleotide encoding DsbC and the polynucleotide sequence encoding a protein of interest.

14. A cell according to any of claims 11 to 13, wherein the protein of interest is an antibody or an antigen binding fragment thereof.

15. A cell according to claim 14, wherein the antibody or antigen binding fragment thereof is specific for TNF.

16. A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a mutant spr protein, a wild-type Tsp gene and a polynucleotide sequence encoding an antibody or an antigen binding fragment thereof specific for TNF.

17. A cell according to claim 16, wherein the cell comprises a recombinant polynucleotide encoding DsbC.

18. A method for producing a recombinant protein of interest comprising culturing a recombinant gram-negative bacterial cell as defined in any of claims 1 to 17 in a culture medium under conditions effective to express the recombinant protein of
interest and recovering the recombinant protein of interest from the periplasm of
the recombinant gram-negative bacterial cell and/or the culture medium.

19. A method according to claim 18, wherein the recombinant protein of interest is
recovered from the periplasm and/or the supernatant.

20. A method according to claim 18 or 19, wherein the cell comprises a recombinant
polynucleotide encoding DsbC and the cell is cultured under conditions effective
to express the recombinant polynucleotide encoding DsbC.

21. A method according to claim 20, wherein the expression of the polynucleotide
sequence encoding a protein of interest and the recombinant polynucleotide
encoding DsbC is induced by adding an inducer to the culture medium.

22. A method according to claim 20 or claim 21, wherein the method further
comprises separating the recombinant protein of interest from DsbC.
Figure 7

Effect of spr on DNA leakage to the supernatant

![Graph showing DNA leakage over time post induction for different strains.]

- ◯ W3110 (wild type)
- ★ MXE012 (W3110 spr H145A)
- ■ MXE001 (W3110 Δtsp)
- ● MXE008 (W3110 Δtsp spr D133A)
Figure 8

Effect of spr on proteins leakage to the supernatant

- W3110 (wild type)
- MXE012 (W3110 spr H145A)
- MXE001 (W3110 Δtsp)
- MXE008 (W3110 Δtsp spr D133A)
Figure 9a

**Wild type ptr (protease III) 5’.**

* M P R S T W F K A L L L L V
    TGA ATG CCC CGC AGC ACC TGG TTC AAA GCA TTA TTG TTG TTA GTT
    A L W A P L S
    GCC CTT TGG GCA CCC TTA AGT

**Mutated Δ ptr (protease III) 5’.**

_EcoRI_

* I P R S T W F K A L L L L V
    TGA ATT CCC CGC AGC ACC TGG TTC AAA GCA TTA TTG TTG TTA GTT

_AseI_

    A L W A H * C
    GCC CTT TGG GCA CAT TAA TGT

Figure 9b

**Wild type Tsp 5’.**

M N M F R L T A L A G L L A
     ATG AAC ATG TTT TTT AGG CTT ACC GCG TTA GCT GCC CTG CTT GCA
     I A G Q T F A
     ATA GCA GGC CAG ACC TTC GCT

**Mutated Δ Tsp 5’.**

_EcoRI_

M N S F L G L P R * L A C L Q
     ATG AAT TCG TTT TTA GGC TTA CCG CGT TAG CTG GCC TGC TTG CAA

_AseI_

* Q A R H * L
     TAG CAG GCC AGA CAT TAA TTG
Figure 9c

Wild type DegP

202 D A A I N R G N S G G
949 GAT GCA GCG ATC AAC CGT GGT AAC TCC GGT GGT

Mutated DegP S210A

Ase I

202 D A A I N R G N A G G
Figure 10
**INTERNATIONAL SEARCH REPORT**

**Box No. I  Nucleotide and/or amino acid sequence(s) (Continuation of item 1.b of the first sheet)**

1. With regard to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the claimed invention, the international search was carried out on the basis of:
   a. (means)
      □ on paper
      □ in electronic form
   b. (time)
      □ in the international application as filed
      □ together with the international application in electronic form
      □ subsequently to this Authority for the purpose of search

2. □ In addition, in the case that more than one version or copy of a sequence listing and/or table relating thereto has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished.

3. Additional comments:

Form PCT/ISA/21 0 (continuation of first sheet (1)) (July 2009)
**INTERNATIONAL SEARCH REPORT**

**Box No. II**  Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☐ Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:

2. ☐ Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. ☐ Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

**Box No. III**  Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1. ☐ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. ☐ As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of additional fees.

3. □ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:

   6(completely) ; 1-3, 7-22 (partially)

4. □ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:  

**Remark on Protest**

☐ The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.

☐ The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.

☒ No protest accompanied the payment of additional search fees.
A. CLASSIFICATION OF SUBJECT MATTER
   INV. C07K16/24 C12N9/48 C12N15/70 C12P21/02

ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
C07K C12N C12P

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, BIOSIS, EMBASE, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<td>X</td>
<td>CHEN CHRISTINA ET AL: &quot;High-l evel accumulation of a recombinant anti body fragment in the periplasm of Escheri chi coli requires a triple-mutant (degP pre spr) host strain n.&quot;, BIOTECHNOLOGY AND BIOENGINEERING, vol. 85, no. 5, 5 March 2004 (2004-03-05), pages 463-474, XP002630315, ISSN: 0006-3592 cited in the application on the whole document</td>
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<td>W0 02/48376 A2 (GENENTECH INC [US]) 20 June 2002 (2002-06-20) page 33, line 21 - line 31 the whole document page 30, line 17 - line 23</td>
<td>1-3,6-22</td>
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</table>

Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents:
- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier document published on or after the international filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed

Date of the actual completion of the international search: 8 June 2011

Date of mailing of the international search report: 20/06/2011

Name and mailing address of the ISA:
European Patent Office, P.B. 5818 Patentlaan 2 NL-2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016

Authorized officer: Giebel er, Kathar ina
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<td>DATABASE Uni Prot [Online] 10 February 2009 (2009-02-10), SubName: Full name of predicted peptidase, outer membrane lipoprotein; XP002630316, retrieved from EBI accession no. UNI PROT: B7UFJ2 Database accession no. B7UFJ2 compound</td>
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Form PCT/ISA/210 (continuation of second sheet) (April 2008)
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<td>A</td>
<td>HARA H ET AL: &quot;OVERPRODUCTION OF PENICILLIN-BINDING PROTEIN 7 SUPPRESSES THERMOSENSITIVE GROWTH DEFECT AT LOW OSMOLARITY DUE TO AN SPR MUTATION OF ESCHERICHIA COLI &quot;. MICROBIAL DRUG RESISTANCE, LI EBERT, US, vol. 2, no. 1, 1 January 1996 (1996-01-01), pages 63-72, XP008015427, ISSN: 1076-6294 cited in the application on the whole document</td>
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<td>A</td>
<td>SIMMONS LAURA C [US]; KLIMOWSKI LAURA [US]; REI LLY 8 August 2002 (2002-08-08) the whole document</td>
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<td>A</td>
<td>ARAMINI JAMES M ET AL: &quot;Solution NMR structure of the NlpC/P60 domain of lipoproteins from Escherichia coli: Structural evidence for a novel cysteine protease catalytic triad&quot;, BIOCHEMISTRY, vol. 47, no. 37, September 2008 (2008-09), pages 9715-9717, XP002630320, ISSN: 0006-2960 cited in the application on the whole document page 28, line 13 - page 29, line 6</td>
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| US 7662587                             | 16-02-2010       | NONE                     |

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|                                        |                  | JP 2004530419 T          | 07-10-2004       |
|                                        |                  | MX PA03005273 A          | 25-09-2003       |
This International Search Authority found multiple (groups of) inventions in this international application, as follows:

1. claims: 6 (completely); 1-3, 7-22 (partially)

   A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a H145 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

   ---

2. claims: 1-3, 7-22 (all partially)

   A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a N31 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

   ---

3. claims: 1-3, 7-22 (all partially)

   A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a R62 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

   ---

4. claims: 1-3, 7-22 (all partially)

   A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a 170 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

   ---

5. claims: 1-3, 7-22 (all partially)

   A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a Y73 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

   ---

6. claims: 1-3, 7-22 (all partially)

   A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a C94 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

   ---

7. claims: 1-5, 7-22 (all partially)

   A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a S95 mutant spr protein and
wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

8. claims: 1-4, 7-22 (partially)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a V98 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

9. claims: 1-3, 7-22 (partially)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a Q99 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

10. claims: 1-3, 7-22 (partially)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a R100 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

11. claims: 1-3, 7-22 (partially)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a L108 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

12. claims: 1-5, 7-22 (partially)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a Y115 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

13. claims: 1-4, 7-22 (partially)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a D133 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

14. claims: 1-4, 7-22 (partially)

A recombinant gram-negative bacterial cell comprising a
FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

15. claims: 1-3, 7-22 (al 1 partly)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a V135 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

16. claims: 1-3, 7-22 (al 1 partly)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a L136 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

17. claims: 1-3, 7-22 (al 1 partly)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a G140 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

18. claims: 1-4, 7-22 (al 1 partly)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a R144 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

19. claims: 1-3, 7-22 (al 1 partly)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a G147 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---

20. claims: 1-3, 7-22 (al 1 partly)

A recombinant gram-negative bacterial cell comprising a mutant spr gene encoding a W174 mutant spr protein and wherein the cell comprises a non-recombinant wild-type chromosomal Tsp gene; methods relating thereto.

---