Novel eukaryotic cells and methods for recombinantly expressing a product of interest.

Fig. 1

1. eukaryotic cell line containing protein A
2. recombinant cell
3. recombinant cell line
4. host cell
5. recombinant eukaryotic cell line
6. recombinant cell line
7. recombinant eukaryotic cell line
8. recombinant eukaryotic cell line
9. recombinant eukaryotic cell line
10. recombinant eukaryotic cell line
11. recombinant eukaryotic cell line
12. recombinant eukaryotic cell line
13. recombinant eukaryotic cell line
14. recombinant eukaryotic cell line
15. recombinant eukaryotic cell line
16. recombinant eukaryotic cell line
17. recombinant eukaryotic cell line
18. recombinant eukaryotic cell line
19. recombinant eukaryotic cell line
20. recombinant eukaryotic cell line
21. recombinant eukaryotic cell line

Abstract: The disclosure pertains to novel eukaryotic cells suitable for recombinant production of a product of interest, wherein the genome of the host cell is altered so that the effect of protein FAM60A is impaired in said cell, e.g. by reducing or eliminating functional expression of gene FAM60A thereby improving the stability characteristics. Furthermore, the present disclosure provides associated technologies wherein such host cells are used in recombinant production technologies.
"Novβ1 eukaryotic cells and methods for recombinantly expressing a product of interest"

FIELD OF THE DISCLOSURE

[1] The present disclosure concerns the field of recombinant expression technologies. It inter alia provides altered eukaryotic cells which are capable of expressing a product of interest with increased stability as well as their use in recombinant expression methods. Furthermore, tools are provided which allow early in the selection process the identification of eukaryotic cells that express a recombinant product with improved stability based on the expression profile of the eukaryotic cell. The eukaryotic cell preferably is a mammalian cell.

BACKGROUND OF THE DISCLOSURE

[2] The market for biopharmaceuticals continues to grow at a high rate as biopharmaceuticals become more and more important for today’s medicine. Currently, an increasing number of biopharmaceuticals is produced in eukaryotic cells such as in particular mammalian cells. Successful and high yield production of biopharmaceuticals in eukaryotic cells is thus crucial. The time to generate such a cell line producing a therapeutic protein of interest is an essential part of the time needed to bring any biopharmaceutical in the clinic. Furthermore, also considering the production costs for biopharmaceuticals and other recombinant products it is important to have high but in particular also stably expressing recombinant eukaryotic cell lines, in particular mammalian cell lines.

[3] For biopharmaceutical production efficiency’s sake in particular on industrial scale, a tremendous effort is put into the clone selection process, with the goal to identify high producing clones with good stability and growth characteristics in a short amount of time. However, even if high expressing clones are identified in the course of the screening process, these initially high expressing clones often lose their advantageous expression characteristics and the expression yield decreases over time. This gradual loss of recombinant protein expression in cell clones during prolonged subculture is a common issue with many cell lines such as CHO cell lines and is referred to as instability. This instability seriously affects the industrial production process of recombinantly produced polypeptides. Causes of production instability are contemplated to be related to loss of recombinant gene copies due to genetic instability of the host cells and epigenetic silencing of transgene sequences. Furthermore, it was found that the instability rate can vary depending on the individual project, i.e. the individual product of interest to be expressed. Instability rates between 25% and up to almost 90% were observed in eukaryotic cell lines. Therefore, care must be taking in order to identify within the population of successfully expressing cells and even among the cell clones which express the protein of interest initially with a good yield those cells, respectively cell clones, which also have a high production stability during prolonged cultivation and therefore, are not prone to a gradual loss of recombinant protein expression. Such clones are also called “stable” clones. During prolonged culturing periods,
stable clones should not lose more than 30%, preferably not more than 25% of their initial productivity within a period of 8-12 weeks, e.g. 10 weeks. Productivity is defined as volumetric productivity, which is the expressed amount of protein per volume (e.g. g/L) at a certain time point of cultivation, respectively as cell specific productivity, which is the specific amount of expressed protein per cell per day (e.g. pg/cell/day). In order to avoid that a cell clone is selected for subsequent large scale production which is prone to instability and therefore will lose titer during prolonged culturing, usually extensive stability analyses are performed over several weeks up to several months in order to eliminate those cell clones which become unstable during that time period and to identify the stable clones. Therefore, the generation of recombinant cell clones for production of therapeutic proteins and other recombinant polypeptides that are produced on large scale usually comprises excessive screening of individual clones by time consuming stability studies in order to identify cell clones that show the expression stability necessary for large scale production. This screening practice for eliminating unstable clones and identifying stable clones prolongs the development of biotechnological production processes. Even when using highly stringent selection systems that favour survival of high expressing cells under the used selection conditions, finding a suitable production clone within the surviving population which combines a high expression rate with good growth and stability characteristics is difficult.

[4] It is an object of the invention to improve recombinant production of a product of interest in eukaryotic cells such as in particular mammalian cells. In particular, it is an object of the present invention to provide a novel eukaryotic cell line which upon stable transfection with a polynucleotide encoding a product of interest expresses the product of interest with improved stability characteristics. In particular, it is the object to provide a recombinant eukaryotic cell wherein the risk of a significant productivity loss during prolonged culturing is reduced. Additionally, it is an object to provide an improved method for recombinantly producing a product of interest using stably transfected eukaryotic cells, in particular mammalian cells. Furthermore, it is one object to provide analysis tools that allow discriminating between stable and unstable cell clones at an early stage of the development process.

**SUMMARY OF THE DISCLOSURE**

[5] The present disclosure is *inter alia* based on the unexpected finding that altering the genome of a eukaryotic cell to impair the effect of protein FAM60A in said cell, e.g. by reducing or eliminating the functional expression of the FAM60A gene, significantly increases the expression stability of a recombinant product of interest in said cells. With the FAM60A gene, a key gene was identified that influences the stability of recombinant expression. Impairing the effect of FAM60A in the cells allows to significantly improve the recombinant production of a product of interest by increasing the expression stability. As is shown in the examples, when using the novel eukaryotic cells described herein as host cells, recombinant cell clones are obtained after selection which show significantly improved stability characteristics. Pronounced losses in expression stability during prolonged culturing are rarer with respective host cells and furthermore, if occurring, result in a less dramatic decrease in the productivity compared to cells wherein the genome is not altered to impair the effect of
protein FAM60A in the cell. The abundance of stable clones is increased upon stable transfection. Therefore, stability analyses for identifying host cells that are not or are less prone to instability may be shortened or even skipped. This is an important advantage as it shortens the time that is required for obtaining stably expressing cell clones that express the recombinant product of interest with good yield over a prolonged time period and which accordingly, are suitable for large scale production purposes. Therefore, the present invention significantly reduces the screening effort and makes an important contribution to the prior art.

[6] According to a first aspect, the present disclosure provides an isolated eukaryotic cell, wherein the genome of the eukaryotic cell is altered so that the effect of protein FAM60A is impaired in said cell and wherein said cell comprises integrated into its genome a heterologous polynucleotide encoding a product of interest. The effect of protein FAM60A can be impaired in said cell e.g. by reducing or eliminating functional expression of the endogenous gene FAM60A, e.g. by gene silencing, gene deletion or by mutating the gene so that a non- or less functional protein is expressed. Other options are also described herein.

[7] According to a second aspect, a method is provided for selecting a host cell which recombinantly expresses a product of interest, comprising

(a) providing eukaryotic cells according to the first aspect as host cells; and
(b) selecting one or more host cells expressing the product of interest.

[8] According to a third aspect, a method is provided for recombinantly producing a product of interest, comprising using a eukaryotic cell according to the first aspect as host cell for recombinant expression of the product of interest. The product of interest is encoded by the heterologous polynucleotide that is stably integrated into the genome of the eukaryotic cell according to the first aspect. As described above, due to their favourable expression stability characteristics, these novel eukaryotic cells are particularly suitable as host cells for recombinant production of a product of interest.

[9] According to a fourth aspect, a method is provided for producing a eukaryotic cell suitable for recombinant production of a product of interest, comprising impairing the effect of protein FAM60A in an eukaryotic cell by altering the genome of said cell and stably transfecting into said cell at least one expression vector comprising a polynucleotide encoding a product of interest. The effect of FAM60A can be impaired e.g. by reducing or eliminating functional expression of gene FAM60A in said cell.

[10] According to a fifth aspect, a method is provided for analyzing eukaryotic cells for their suitability as host cells for stably expressing a recombinant product of interest, comprising analyzing directly or indirectly whether the effect of protein FAM60A is impaired in said cells. This method can be advantageously used e.g. in combination with the method according to the fourth aspect in order to identify whether a eukaryotic cell was obtained, wherein the effect of protein FAM60A is impaired in the cell. Furthermore, this method can be used as
analytical tool in order to discriminate early in the selection process between stable and unstable cell clones that express the product of interest.

[11] According to a sixth aspect, the present disclosure pertains to the use of an isolated eukaryotic cell for recombinantly expressing a product of interest, wherein the genome of the eukaryotic cell is altered so that the effect of protein FAM60A is impaired in said cell.

[12] Other objects, features, advantages and aspects of the present application will become apparent to those skilled in the art from the following description and appended claims. It should be understood, however, that the following description, appended claims, and specific examples, while indicating preferred embodiments of the application, are given by way of illustration only. Various changes and modifications within the spirit and scope of the disclosed invention will become readily apparent to those skilled in the art from reading the following.

**BRIEF DESCRIPTION OF THE FIGURES**

[13] **Fig. 1** provides a schematic overview of the telomeric region of chromosome 8 of Chinese hamster ovary (CHO) cells and genes located in said telomeric region. The genomic region depicted in the figure is the result of the merged scaffolds number 6 and 25 on chromosome 8. An overview over genes and putative genes on chromosome 8 of CHO cells can be found using the gene bank annotation file associated with the assembly of Brinkrolf et al. (Nature Biotechnology Volume 31, 694-695 (2013); see gene bank: APMK00000000, version APMK01 000000 as is described in said publication). Furthermore, the Beijing Genomics Institute also provided an annotation of this region (Xu et al, Nature Biotechnology, Volume 29, number 8, 735-741 (2011); see gene bank: AFTD00000000, version AFTD01 000000). Annotations which are marked with an ‘*’ in Fig. 1 are from gene bank file AFTD01 000000.

[14] A corresponding overview over the telomeric region of chromosome 6 of mouse can be found e.g. in the Ensembl database. Chromosome 6 of mouse has a structure which corresponds to chromosome 8 of Chinese hamster. The subsequent link of the Ensemble database shows the telomeric region of chromosome 6 of mouse which contains the FAM60A gene:

http://www.ensembl.org/Mus_musculus/Location/View?db=core;g=ENSMUSG00000039985; r=6:14892 1035-148946467

[15] The subsequent Table 1 provides an overview over abbreviations and alternative names (aliases) of genes and encoded products shown in Fig. 1 and indicates the corresponding annotation in mouse and Chinese hamster (according to Brinkrolf et al, 2013 and/or Xu et al, 2011) where feasible. Table 1 also lists alternative names used e.g. in different species. Wherein the present disclosure refers to a specific protein or gene name, this also refers to
and encompasses any alternative names of said protein or gene e.g. used to characterize the corresponding gene or protein in a different species. In particular, homologs and orthologs having the same function are encompassed thereby.

**Table 1**: Abbreviations and alternative names (aliases) of products encoded by genes located in chromosome 8 of Chinese hamster or chromosome 6 of mouse.

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>In public annotation of Chinese hamster, the gene product is annotated as</th>
<th>In public annotation of mouse the gene product is annotated as</th>
<th>Aliases (see <a href="http://www.genecards.org">www.genecards.org</a>)</th>
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<td>Coiled-coil domain containing 91</td>
<td>Coiled-Coil Domain Containing 91 P56 GGA-Binding Partner P56 Accessory Protein DKFZp779L1 558 FLJ1 1088 Coiled-Coil Domain-Containing Protein 91 GGA Binding Partner GGABP</td>
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<td>Far2</td>
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<td>Fatty acyl CoA reductase 2</td>
<td>Fatty Acyl CoA Reductase 2 MLSTD1 SDR1 0E2 Male Sterility Domain-Containing Protein 1 EC 1.2.1.N2 FLJ1 10462 Male Sterility Domain Containing 1 Fatty Acyl-CoA Reductase 2 Short Chain Dehydrogenase/Reductase Family 10E, Member 2</td>
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<td>Endoplasmic reticulum-Golgi intermediate compartment protein 2</td>
<td>ERGIC and golgi 2</td>
<td>ERGIC And Golgi 2 PTX1 Erv41 Cd002 CD14 Protein Endoplasmic Reticulum-Golgi Intermediate Compartment Protein 2 ERV41 CDA14</td>
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<td>RPS4Y2</td>
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<td>Ribosomal protein S4, Y linked 2</td>
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[16] Fig. 2 shows the relative expression levels of genes located in the telomeric region of chromosome 8 in a CHO cell line, namely TMTC1 (1), RPS4Y2 (2), IP08 (3), CAPRIN2 (4), FAM60A (5), Dennd5b (6), METTL20 (7), AMN1 (8), C12orf35 (9), Bicdl (10).

[17] Fig. 3 shows the results of stability tests performed for 7/8 weeks with three different cell clones (CHO wildtype and two FAM60A knock-out clones s16 and s23 derived from said wildtype) following stable transfection with an expression vector encoding an antibody as product of interest. (1) shows the stability results obtained with the parental wildtype cells (derived from CHO-K1); (2) shows the stability results with the FAM60A knock-out clone s16; (3) shows the stability results with the FAM60A knock-out clone s23. As can be seen, the
expression stability was significantly increased in the cell clones that derived from the FAM60A knock-out cells (see (2) and (3)). The number of stable clones was significantly increased when using the FAM60A knock-out cells for recombinant expression. Thus, impairing the effect of FAM60A in the host cell, here by gene knock-out, significantly improved the expression stability.

[18] Fig. 4A to L show FACS profiles obtained after reducing the expression of different target genes located in the telomeric region of chromosome 8 of Chinese hamster (CHO) cells using siRNAs. Cells that were stably transfected with an expression vector and expressed the encoded antibody as product of interest were fluorescently stained to detect the amount of recombinantly expressed antibody. The higher the intensity in the FACS profile, the more the antibody is expressed by the stained cell. The left peak shown in the FACS profile corresponds to the parental cell line (not transfected and hence not expressing the antibody) which was included for comparative purposes. The two other curves represent results obtained for a cell clone that was stably transfected with the expression vector and which recombinantly expresses the antibody. This cell clone was transfected with either a siRNA negative control (dark curve; no effect on expression of any gene) or with a siRNA that reduces the expression of a target gene (light grey curve). If silencing of the target gene does not have an effect on recombination expression of the antibody, the fluorescent curve for the siRNA control and the target siRNA overlap and remain the same. If silencing of the target gene increases the expression rate of the recombinantly expressed antibody, the intensity of the corresponding FACS profile increases and shifts to the right. A: gene Mettl20_1, 125 pmol, 24.9%; B: gene C12orf35_1, 125 pmol, 30.6%; C: gene C12orf35_2, 150 pmol, 31.7%; D: gene Caprin2_6, 100 pmol, 53.3%; E: FAM60A_3, 150 pmol, 48%; F: lpo8_1, 125 pmol, 20.3%; G: lpo8_2, 150 pmol, 57.5%; H: lpo8_3, 150 pmol, 21.5%; I: Dned5b_2, 100 pmol, 36.9%; J: Amn1_4, 125 pmol, 30.8%; K: TMTC1J, 150 pmol, 60.6%; L: TMTC1_2, 150 pmol, 53.4% (percentage values correspond to mRNA expression of the target gene between reference siRNA versus Ctrl siRNA). Fig. 4B and C show that a downregulation of gene C12orf35 significantly increases expression of the recombinant antibody and thus results in a higher productivity as is indicated by the clear shift of the FACS profile to the right (see light grey curve on the right, also marked with an arrow). Therefore, according to one embodiment, the effect of the expression product of gene C12orf35 is additionally impaired in the host cell in order to improve the yield.

[19] Fig. 5 and 6 show the mRNA expression levels of the antibody light and heavy chains of two different model polypeptides of interest (antibody 1 and 2) in different clones and pools in each case after reducing expression of gene C12orf35 in CHO cells by RNAi. The mRNA levels of the antibody chains are upregulated, if expression of gene C12orf35 is reduced by gene silencing. Thus, reduction of C12orf35 expression surprisingly leads to higher mRNA levels of HC and LC.

[20] Fig. 7 shows that upon silencing of gene C12orf35 using siRNA, significantly higher cell specific expression titers are obtained (calculated from days 3, 4, 5 and 6 of cultivation).
Fig. 8 shows that the 46 highest producing clones (black) derived from a CHO cell line in which the telomeric region comprising gene FAM60A and gene C12orf35 on chromosome 8 (q arm) is deleted (C8DEL) have higher titers compared to the 45 highest producing clones obtained from the parental cell line which were tested to be IP08 positive (grey).

Fig. 9 shows FACS profiles of stably transfected C8DEL cell pools after selection using a folate receptor/DHFR system. The concentration of MTX was increased from A to E (A: no MTX; B: 1nM MTX; C: 5nM MTX; D: 10nM MTX; E: 50nM MTX). Recombinant antibody expression was detected based on fluorescence. At 50nM MTX, predominantly high producing cells were comprised in the obtained pool as is demonstrated by FACS analysis. The obtained pool profile remarkably resembled the profile of a cell clone. This supports the extraordinary improvement the technology described herein achieves on the expression characteristics of a recombinant host cell.

DETAILED DESCRIPTION OF THE DISCLOSURE

The present disclosure is inter alia based on the surprising finding that eukaryotic cells, in which the effect of protein FAM60A is impaired, e.g. by reducing or eliminating functional expression of the endogenous FAM60A gene in said cell, e.g. by deleting said gene or by introducing mutations into the coding sequence, are upon stable transfection capable of expressing a recombinant product of interest with significantly improved stability characteristics. As is shown by the examples for mammalian cells, respectively altered cells surprisingly show very stable expression characteristics during prolonged culturing times, thereby allowing to shorten or even skip time-consuming stability analyses for identifying stable clones. In a population of successfully transfected host cells, the number of host cells which lose their favorable expression characteristics during prolonged culturing periods is significantly reduced when using respectively altered eukaryotic cells. Thus, because of their favorable stability characteristics, these altered eukaryotic cells are particularly suitable as host cells for recombinant production technologies and can be used for recombinant production of a product of interest. Based on this surprising finding that gene FAM60A has a strong impact on recombinant expression stability in eukaryotic host cells, the present disclosure also provides novel selection and production methods and associated technologies which allow improving the recombinant production of a product of interest. Therefore, the present disclosure makes an important contribution to the prior art.

The individual aspects and suitable and preferred embodiments thereof will now be described in detail.

A. Modified eukaryotic cells

According to a first aspect, the present disclosure provides an isolated eukaryotic cell, wherein the genome of the eukaryotic cell is altered so that the effect of protein FAM60A is impaired in said cell and wherein said cell comprises integrated into its genome a heterologous polynucleotide encoding a product of interest. As is shown in the examples, these altered cells show a significant higher production stability of the product of interest.
compared to unmodified cells wherein the effect of FAM60A is not impaired. The abundance of cells which exhibit stable expression characteristics is increased in the population of transfected cells as was shown for the preferred embodiment wherein a mammalian cell is used as eukaryotic cell. This improved expression stability allows to shorten or even skip time consuming long-term stability studies of high expressing cell clones. Further advantages are also described in the following and are also apparent from the examples. Thus, using these advantageous novel eukaryotic cell lines for recombinant production of a product of interest reduces screening efforts for identifying high expressing cells or cell clones with stable expression characteristics and in particular reduces the time needed for obtaining stable cell clones suitable for producing the product of interest on large scale. Thus, these altered eukaryotic cell lines have important advantages when being used as host cells for recombinant production technologies.

[26] FAM60A is a sub-unit of the SIN3-histone deacetylase (HDAC) complex (SIN3/HDAC complex) that functions in transcriptional repression (Munoz et al., 2012, THE JOURNAL OF BIOLOGICAL CHEMISTRY VOL. 287, NO. 39, pp. 32346-32353; Smith et al., 2012, Mol Cell Proteomics 11 (12): 1815-1828). Histone deacetylases (HDAC) catalyze the removal of acetyl groups from histones. Acetylation of histones on lysines is a major mechanism for modulating chromatin conformation. Histone acetylation promotes a relaxed, transcriptionally active chromatin state whereas deacetylation catalyzed by histone deacetylases (HDACs) favor a silent, inactive state. Database analysis revealed the presence of at least one FAM60A ortholog in most metazoans, but not in nematodes. The FAM60A gene is conserved in metazoans and can be found in all vertebrate and most invertebrate genomes that have been completely sequenced. E.g. a 100% sequence identity of FAM60A protein can be found between human, rat, mouse and cow. Sequence similarity research of FAM60A homologs indicates that predominantly, there is only a single representative member of this family in the genome. There are only few exceptions. According to the Smith et al, 2012, the FAM60A protein has a unique sequence lacking any known protein domains. Moreover, it was described by Smith et al 2012, that it does not exhibit any sequence homology to other known proteins in the human proteome. Sequence comparison between FAM60A proteins from different species showed that the FAM60A protein generally comprises three regions: (1) an N-terminus comprising highly conserved segments in all metazoans (2) a middle region which is highly conserved across vertebrates whereas in invertebrates it consists of a non-conserved spacer of a variable length (3) a C-terminus comprising highly conserved segments in all metazoans. Thus, highest conservation was observed in the FAM60A N- and C-terminal regions.

[27] As described above, research indicates that FAM60A associates with SIN3/HDAC complexes in various eukaryotic cell types such as in particular mammalian cells. However, to date, functional information about FAM60A is quite restricted. Recent functional studies (see Smith et al, 2012) indicate that FAM60A may repress gene expression and regulates a specific subset of genes. Smith et al 2012 report a role of FAM60A in the regulation of the TGF-beta signaling pathway, which plays a pivotal role in processes like cancer progression, metastasis, cell migration and immune surveillance. There are findings indicating that
FAM60A acts as a transcriptional repressor of components of the TGF-beta signaling pathway whereas this FAM60A function seems to be permitted via its role in the SIN3-HDAC complex. Depletion of FAM60A in different cancer cell lines using siRNA against FAM60A resulted in a change of normal cancer cell morphology. Furthermore, it was found that FAM60A protein levels do periodically change within the course of the cell cycle in U20S cells (Munoz et al, 2012). FAM60A knock-down experiments using FAM60A siRNA in U20S human bone osteosarcoma cells revealed that FAM60A restraints cyclin D1 gene expression. Against this scientific background, it was highly surprising to find that impairing the effect of protein FAM60A in a eukaryotic cell such as preferably a mammalian cell, significantly increases the stability of heterologous gene expression in said cell without negatively affecting other characteristics of the cell that are important for recombinant expression. This correlation between the effects of protein FAM60A and the expression stability during prolonged culturing of the cells was unexpected.

[28] As described, the FAM60A gene is endogenously expressed in metazoan and hence in mammalian species such as human, mouse, rat and hamster and the amino acid sequence of FAM60A is highly conserved in mammalian species as well as in vertebrates. The altered eukaryotic cell according to the first aspect is derived from a eukaryotic cell which endogenously expresses FAM60A. For the ease of simplicity, the protein FAM60A as well as the FAM60A gene encoding protein FAM60A is spelled herein in capital letters even though in some species a different spelling is used for the gene and/or the protein. The sequence listing shows exemplary amino acid sequences of known and/or predicted FAM60A proteins of different vertebrate species, namely Homo sapiens (SEQ ID NO: 1), Rattus norvegicus (SEQ ID NO: 2), Mus musculus (SEQ ID NO: 3), Cricetulus griseus (SEQ ID NO: 4), Gallus gallus (SEQ ID NO: 5), Pan troglodytes (SEQ ID NO: 6), Pongo abelii (SEQ ID NO: 7) and Bos taurus (SEQ ID NO: 8). The predicted FAM60A cDNA of Cricetulus griseus is shown in SEQ ID NO: 9 (coding sequence from 14-679; see also NCBI Reference Sequence: XM_003505482.1). Different names can be assigned for protein FAM60A or the FAM60A gene in different species and non-limiting alternative names (aliases) are also listed above in Table 1. The term "FAM60A" as used herein also encompasses any homologs and orthologs of FAM60A which have the same function as FAM60A. According to one embodiment the term "FAM60A" as used herein in particular refers a protein that shares at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% homology to one or more of the amino acid sequences shown in SEQ ID NO: 1 to 8. According to one embodiment, the foregoing percentage values refer to polypeptide identity instead of homology. Homology, respectively identity, may be calculated over the entire length of the reference protein. A respective protein preferably has the same function as the protein having an amino acid sequence that is shown in SEQ ID NO: 1 or one or more of SEQ ID NO: 2 to 8, preferably SEQ ID NO: 4. The FAM60A protein has not been described in detail in the literature. Thus, it was highly surprising that the expression stability of a recombinant host cell can be improved, if the genome of the host cell is altered so that the effect of endogenous protein FAM60A is impaired in the cell, as can be e.g. achieved by reducing or eliminating the functional
expression of the FAM60A gene in said cell. It was unexpected that FAM60A influences the expression stability of a recombinant product of interest.

[29] The FAM60A gene encoding the FAM60A protein can be modified as described herein in order to impair the effect of FAM60A in the cell. This can be achieved e.g. by genetic engineering technologies such as gene knock-out technologies. The genomic gene sequence of different mammalian species is known, and is e.g. described in Homo sapiens (NCBI Gene-ID: 58516); Rattus norvegicus (NCBI Gene-ID: 68661); Mus musculus (NCBI Gene-ID: 56306); Bos Taurus (NCBI Gene-ID: 538649) and others. Transcript variants may exist in a species-dependent manner and in different numbers. E.g. the human FAM60A gene expresses 3 putative transcript isoforms which differ in the UTRs but encode the same protein.

[30] The present disclosure *inter alia* pertains to modified eukaryotic cells, such as preferably mammalian cells, wherein the genome of the eukaryotic cell is altered so that the effect of protein FAM60A, which is endogenously expressed by a corresponding unmodified eukaryotic cell, is impaired. This modification allows to increase the number of stably expressing cells in the cell population that was stably transfected with an expression vector comprising a polynucleotide encoding a product of interest.

[31] There are several possibilities to modify the genome of a cell to impair the effect of protein FAM60A in said cell. The effect of FAM60A may be impaired e.g. on the gene level or on the protein level. The effect of FAM60A can be impaired, for example, by modification of the structure/sequence, the transcription, translation and/or interaction with other components forming the SIN3/HDAC complex. Non-limiting options are described in the following.

[32] According to one embodiment, the effect of protein FAM60A is impaired because the functional expression of gene FAM60A is reduced or eliminated in said cell. As is shown by the examples, altering the expression of gene FAM60A, e.g. by gene knock-out or by reducing the expression level, is a very efficient measure to provide altered cells that express a recombinant product of interest with improved stability characteristics.

[33] Reduction or elimination of functional expression of gene FAM60A may be achieved by various means. Functional expression can be reduced for example by reducing the expression level of FAM60A or by disrupting the function of FAM60A or by a combination of such methodologies. According to one embodiment, the cell is altered so that functional expression of the FAM60A gene is reduced or eliminated by gene knock-out, gene mutation, gene deletion, gene silencing or a combination of any of the foregoing. According to one embodiment, functional expression of gene FAM60A is reduced or eliminated in the cell by gene knockout. A gene knockout is a genetic technique by which a gene is made inoperative by disrupting its function. E.g. a nucleic acid can be inserted into the coding sequence, thereby disrupting the gene function. Furthermore, the complete FAM60A gene or a portion thereof can be deleted, whereby no or no functional protein is expressed by a respectively
altered cell. Another option is to introduce one or more knock-out mutations into the coding sequence, which render a non- or a less functional expression product. E.g. one or more frameshift mutations can be introduced into the coding sequence that result in a non- or less-functional expression product. Alternatively or additionally, one or more stop codons can be introduced into the coding sequence so that a truncated, non- or less functional protein is obtained. Furthermore, splicing sites may be altered. Hence, according to one embodiment, the FAM60A gene comprises one or more mutations which provide a non- or less functional expression product. According to one embodiment, one or more mutations are introduced into exon 1 of the FAM60A gene. According to one embodiment, due to the introduced one or more mutations, all or a part of the N-terminal or the C-terminal region of FAM60A is not present in the expression product. Other options include but are not limited to one or more mutations in the promoter, in the 5’- and/or 3’ UTRs or other regulatory elements. According to one embodiment, the promoter function of the FAM60A gene is disrupted, e.g. by introducing a promoter deletion or by introducing a construct between the promoter and the transcription start. Methods for achieving a gene knockout to suppress or eliminate expression of the target gene are also well-known to the skilled person and thus, do not need any detailed description herein. Some non-limiting examples are nevertheless described below.

[34] According to one embodiment, the FAM60A gene is functionally knocked out by genetic engineering. Examples include but are not limited to genome editing, such as genome editing with engineered nucleases (GEEN). This is a type of genetic engineering in which DNA is inserted, replaced or removed from a genome using artificially engineered nucleases, or "molecular scissors." The nucleases create specific double-stranded breaks (DSBs) at desired locations in the genome, and harness the cell's endogenous mechanisms to repair the induced break by natural processes of homologous recombination (HR) and nonhomologous end-joining (NHEJ). There are at least four families of engineered nucleases that can be used: Zinc finger nucleases (ZFNs), Transcription Activator-Like Effector Nucleases (TALENs), CRISPR, and engineered meganuclease re-engineered homing endonucleases. TALEN technology was also used in the examples to provide altered mammalian cells wherein the FAM60A gene was knocked out thereby impairing the effect of protein FAM60A in said cells.

[35] According to one embodiment at least one copy and optionally, if more copies of gene FAM60A are present in the genome of the eukaryotic cell, all copies are altered, e.g. knocked-out, deleted or otherwise rendered inoperative, to reduce or eliminate and hence impair the effect of protein FAM60A in the eukaryotic cell. Thus, according to one embodiment, at least one copy of gene FAM60A is deleted or functionally inactivated in the genome of the eukaryotic cell. For example, one or more mutations may be inserted into the one or more copies of the FAM60A gene to provide a non- or less functional expression product or to eliminate or reduce expression in toto and, hence impair the effect of FAM60A in the eukaryotic cell. Thereby, the FAM60A gene is basically inactivated in the genome. According to one embodiment, all copies of gene FAM60A are respectively altered in the eukaryotic cell, which preferably is a mammalian cell.
[36] According to one embodiment, the eukaryotic cell is a metazoan cell, a vertebrate cell or preferably, a mammalian cell. According to one embodiment, a portion of a chromosome is deleted in the said cell, wherein the deleted portion comprises gene FAM60A. According to one embodiment, a chromosomal portion comprising gene FAM60A is deleted in all chromosomes which comprise a copy of gene FAM60A if more than one copy is present. Thereby, all copies of gene FAM60A are deleted from the genome.

[37] According to one embodiment, a portion of the telomeric region of a chromosome is deleted, wherein the deleted portion comprises gene FAM60A. According to a preferred embodiment, the altered cell is a rodent cell. According to one embodiment, the cell is a hamster cell such as e.g. a CHO cell and at least a portion of the telomeric region of chromosome 8 is deleted in the genome, wherein said deleted portion comprises gene FAM60A. The meaning of the term "FAM60A" is explained above and non-limiting alternative names of homologs and orthologs that are also encompassed by the scope of said term are also indicated in Table 1. According to one embodiment, such deletion occurs in the q arm of chromosome 8 of a hamster cell, in particular a Chinese hamster cell, which comprises the FAM60A gene. As is shown by the examples, a CHO cell comprising a respective deletion in the telomeric region of chromosome 8 is particularly suitable as host cell for recombinant expression. After stable transfection with an expression vector, these cells show significant higher expression stability and productivity compared to cells wherein said portion of the telomeric region of chromosome 8 is not lost. Furthermore, the abundance and thus the proportion of stably expressing cells in the transfected cell population is significantly increased. Significant losses in titer during prolonged culturing are rarely observed. Thus, the stability of recombinant expression is significantly improved in such hamster cells which have lost said portion of the telomeric region in chromosome 8. Further important advantages are described in detail in the examples wherein CHO cells in which a respective portion of the telomeric region of chromosome 8 is deleted due to chromosome breakage are further characterized. The advantageous properties render these hamster cells particularly suitable as industrial production cell lines. Alternatively, the altered rodent cell may be a mouse cell wherein at least a portion of the telomeric region of chromosome 6 is deleted in the genome, wherein said deleted portion comprises gene FAM60A. The telomeric region of chromosome 6 of mouse is highly similar to the telomeric region of chromosome 8 of hamster.

[38] According to one embodiment, at least a portion of the telomeric region is deleted or not present in both chromosomes of chromosome pair 8 of hamster (or chromosome pair 6 in case of mouse cells), wherein the deleted portions comprise the FAM60A gene.

[39] According to one embodiment, at least a portion of the telomeric region is deleted in one chromosome of chromosome pair 8 of hamster (or chromosome pair 6 in case of mouse), wherein said deleted portion comprises the FAM60A gene and the expression of gene FAM60A in the other chromosome, if a further copy is present, is reduced or eliminated. Suitable ways to reduce or eliminate the expression of a gene are known to the skilled person and non-limiting examples are also described herein. According to one embodiment,
such deletion occurs in the q arm of chromosome 8 of hamster, in particular Chinese hamster.

[40] According to one embodiment, the deleted chromosomal region comprises the FAM60A gene and additionally comprises one or more or all genes selected from the group consisting of Bicl, C12orf35, Amn1, methyltransferase-like protein 20, Dennd5b, Caprin2 and Ipo8. According to one embodiment, all of the aforementioned genes are deleted. According to one embodiment the deleted chromosomal region additionally comprises at least a portion of or the full gene TmtC1. In hamster cells such as CHO cells, these genes may be located in the telomeric region of chromosome 8. According to one embodiment, the deleted chromosomal region additionally comprises gene RPS4Y2, if present. An overview over the telomeric region of chromosome 8 of the Chinese hamster genome wherein the location of aforementioned genes is deleted is shown is provided as Fig. 1. As is shown by the examples, CHO cells comprising a respective deletion in the telomeric region of chromosome 8 (q arm) have particular advantageous properties with respect to expression yield and expression stability. In mouse cells the aforementioned genes are located in the telomeric region of chromosome 6. Non-limiting alternative names of the aforementioned individual genes and/or encoded proteins including orthologs and homologs are also indicated in Table 1 above and the respective genes are encompassed by the scope of the terms used above for the individual genes.

[41] According to one embodiment, the deletion of the FAM60A gene is due to a chromosome breakage. A chromosome breakage can be induced e.g. by treating the mammalian cells with a toxic agent that promotes chromosome breakage, such as e.g. MTX, aphidicolin or hygromycin. Other options for inducing chromosome breakages include but are not limited to radiation, irradiation, mutagens, cancerogenic substances and bleomycin. Chromosome breakages may also occur spontaneously during transfection e.g. electroporation. Methods for inducing chromosome breakage are also known to the skilled person and thus, do not need any detailed description here. After inducing chromosome breakage, cells having the desired breakpoint (which results in a deletion of gene FAM60A) can be identified e.g. be analyzing the DNA or by using the method according to the fifth aspect of the present disclosure. For example, the expression profile of the treated cells can be analyzed to determine whether gene FAM60A or genes located centromeric of gene FAM60A are expressed, whether the expression is reduced or whether the genes are not expressed. For example, in case of mouse or hamster cells it can be analysed whether gene FAM60A is expressed and alternatively or in addition thereto, it can be analyzed whether one or more genes selected from the group consisting of Bicl, C12orf35, methyltransferase-like protein 20, Dennd5b, Caprin2, Ipo8, TmtC1 or genes that are located telomeric of the aforementioned genes (wherein telomeric in this respect means into the direction of the telomeric end) are expressed by the cell and/or whether the expression is reduced or eliminated. If the induced breakpoint is located centromeric of the respective gene(s) (wherein centromeric in this respect means further into the chromosome and hence further away from the telomeric end), the telomeric end comprising said genes is deleted which eliminates or reduces (if other copies of the gene exist elsewhere that are expressed) their
expression. As is evident from Fig. 1, gene FAM60A is located telomeric of the aforementioned genes Caprin2, Lpo8 and Trnd, i.e. it is located further into the direction of the telomeric end. Thus, if the aforementioned genes are deleted by a chromosome break, the deleted region also includes gene FAM60A. Thus, the above genes can be validly used as markers in order to basically indirectly determine whether the induced chromosome breakage resulted in a deletion of a chromosome portion that includes gene FAM60A. Furthermore, it was found that even though located telomeric of gene FAM60A, also other genes such as Bicdl or C12orf35 can be used as marker to determine whether a chromosome breakage was induced which resulted in a deletion of gene FAM60A. It was found in CHO cells that if e.g. gene Bicdl or gene C12orf35 is deleted because of a chromosome breakage, the deletion usually also includes gene FAM60A. It was confirmed by analyzing the expression characteristics of several hundred clones that the aforementioned genes can be validly used as markers in order to discriminate cell clones with high and stable expression characteristics from cell clones having low and unstable expression characteristics. The relative expression of the aforementioned genes in CHO cells is shown in Fig. 2. As can be seen from Fig. 2, genes Lpo8 (3), FAM60A (5) and C12orf35 (9) are relatively highly expressed in comparison to other genes that are located in the telomeric region of chromosome 8 in normal CHO-K1 cells, which do not comprise a deletion in the telomeric region of chromosome 8. Thus, it is advantageous to include one or more of the aforementioned genes in the analysis, as this simplifies the detection that their expression is eliminated or reduced. Non-limiting alternative names of the aforementioned individual genes and encoded proteins, including homologs and orthologs, are also indicated in Table 1 above and the respective genes are encompassed by the scope of the terms used above for the individual genes.

[42] According to one embodiment, the breakpoint on chromosome 8 is located centromeric of the FAM60A gene, centromeric of the Caprin2 gene, centromeric of the Lpo8 gene or centromeric of gene RPS4Y2. It was found that the breakpoint on chromosome 8 of the hamster genome is often located centromeric of the Lpo8 gene. According to one embodiment, the breakpoint on chromosome 8 is located within the Tn tcl gene and said gene is not expressed or its expression is low. According to one embodiment, the Ergic2 gene, which is located centromeric of the Tn tcl gene, is not deleted on chromosome 8. Thus, according to this embodiment, the breakpoint is telomeric of the Ergic 2 gene (wherein telomeric in this respect means downstream into the direction of the telomeric end) and the Ergic 2 gene is present.

[43] According to one embodiment, the genome of the cell is altered so that functional expression of gene FAM60A is reduced or eliminated. Functional expression of FAM60A can be influenced by various means, for example by altering the promoter and/or an enhancer of the FAM60A gene so that less or no transcript is produced, or by gene silencing technologies such as transcriptional or post-transcriptional gene silencing. According to one embodiment, the isolated eukaryotic cell comprises one or more mutations in the promoter region of the FAM60A gene. For example, the promoter region may be altered to provide a less functional or non-functional promoter, the promoter may also be completely eliminated. Alternatively or
in addition, it is possible to add a polynucleotide sequence encoding a polypeptide including a stop codon between the promoter and the start codon of the FAM60A gene which leads to the expression of the other polypeptide instead of FAM60A. Respective methods are well-known to the skilled person and thus, do not need any detailed description here.

[44] Reduction of functional gene expression may achieve a level wherein expression is even eliminated. Post-transcriptional gene silencing can be achieved e.g. by antisense molecules or molecules that mediate RNA interference. Non-limiting examples will be briefly described in the following.

[45] Antisense polynucleotides may be designed to specifically bind to RNA, resulting in the formation of RNA-DNA or RNA-RNA hybrids, with an arrest of reverse transcription or messenger RNA translation. Many forms of antisense have been developed and can be broadly categorized into enzyme-dependent antisense or steric blocking antisense. Enzyme-dependent antisense includes forms dependent on RNase H activity to degrade target mRNA, including single-stranded DNA, RNA, and phosphorothioate antisense. Antisense polynucleotides are typically generated within the cell by expression from antisense constructs that contain the antisense strand as the transcribed strand. Trans-cleaving catalytic RNAs (ribozymes) are RNA molecules possessing endoribonuclease activity. Ribozymes may be specifically designed for a particular target and may be engineered to cleave any RNA species site-specifically in the background of cellular RNA. The cleavage event renders the mRNA unstable and prevents protein expression. The genome of the eukaryotic cell can be altered so that a respective antisense molecule is e.g. permanently expressed.

[46] Another suitable option for reducing functional expression of gene FAM60A on a post-transcriptional level is based on RNA interference (RNAi). Methods for silencing genes by RNAi are well known to the skilled person and thus, do not need any detailed description here. Several embodiments and variations of siRNA compounds are known in the prior art and can be used to reduce expression of gene FAM60A. Suitable siRNAs targeting the chosen/identified target sequences of the target genes on the RNA level can be identified by using proper computational methods, applying certain design-algorithms. According to one embodiment, the RNAi inducing compound is expressed by a vector that is stably transfected into the eukaryotic cell and thus is integrated into the genome of the eukaryotic cell. For siRNA, this can be done e.g. by the introduction of a loop between the two strands, thus producing a single transcript, which can be then processed into a functional siRNA in the eukaryotic cell. Such transcription cassettes typically use an RNA polymerase III promoter (for example U6 or H1) which usually direct the transcription of small nuclear RNAs (shRNAs). It is assumed that the resulting shRNA transcript from the vector is then processed by dicer, thereby producing the double-stranded siRNA molecules, preferably having the characteristic 3' overhangs. According to one embodiment, such shRNA providing vector is stably integrated into the genome of the eukaryotic cell. This embodiment is advantageous, as the downregulation of gene FAM60A is due to the constantly produced siRNA stable and not transient and therefore, is feasible for providing a mammalian host cell
with improved expression stability. Cells comprising a respective shRNA providing vector can then be transfected with an expression vector comprising a polynucleotide encoding the product of interest. Alternatively, co-transfection strategies can be used, wherein the vector generating the shRNA is co-transfected with the expression vector comprising the polynucleotide encoding the product of interest.

[47] Transcriptional gene silencing may e.g. include epigenetic modifications. According to one embodiment, functional expression of gene FAM60A is reduced by epigenetic silencing. Furthermore, the sequence of the gene FAM60A can be changed to reduce the half-life of the FAM60A mRNA. Thereby, less FAM60A protein is obtained which also achieves a reduction in the effect of the FAM60A protein in the cell.

[48] According to one embodiment, functional expression of gene FAM60A is reduced or eliminated by targeting a regulatory element involved in the regulation of expression of the FAM60A gene. E.g. a transcription factor, promoter (see also above), enhancer, UTRs, or other regulatory elements can be targeted e.g. by knock-out, deletion, down-regulation or any other alteration that inactivates or reduces the activity of said regulatory element, thereby preventing or reducing functional expression of gene FAM60A and thereby impairing the effect of endogenous FAM60A in said cell.

[49] According to one embodiment, the genome of the eukaryotic cell is altered to impair the effect of FAM60A by heterologous expression of a mutant FAM60A which is non- or less functional than the endogenously expressed FAM60A protein. In this embodiment, the isolated eukaryotic cell comprises in addition to the heterologous polynucleotide encoding the polypeptide of interest a further heterologous polynucleotide encoding the mutant FAM60A. By overexpressing a respective non- or less functional mutant FAM60A, a dominant negative phenotype can be created. A further option to impair and hence reduce the effect of FAM60A in the cell is the heterologous expression of a protein such as an antibody which neutralizes FAM60A and hence impairs the effect of FAM60A in the cell. According to one embodiment, the effect of FAM60A is impaired in the cell by reducing or eliminating functional expression of molecules that functionally interact with FAM60A, e.g. by reducing or eliminating functional expression of one or more members of the SIN3/HDAC complex. Such embodiments also impair the effect of FAM60A, because one or more interaction partners of FAM60A that are required so that FAM60A can exert its biological effect, are not present in a functional form because their functional expression was reduced or eliminated.

[50] According to one embodiment, expression of gene FAM60A is reduced by at least 3 fold, at least 5 fold, at least 10 fold, at least 20 fold, at least 30 fold, at least 40 fold, at least 50 fold, at least 60 fold, at least 70 fold, at least 75 fold, at least 80 fold, at least 90 fold, at least 100 fold or at least 125 fold, at least 250 fold, at least 500 fold, at least 750 fold, at least 1000 fold, at least 1250 fold, at least 1500 fold, at least 1750 fold, at least 2000 fold, at least 2500 fold, at least 3000 fold or at least 3500 fold. Expression can be determined e.g. by using real-time RT-PCR or other sensitive RNA detection methods. Such reduction can be achieved e.g. in
comparison with the unmodified reference cell wherein the expression of endogenous gene FAM60A is not reduced. According to one embodiment, expression of gene FAM60A is 0.05% or less, 0.04% or less, 0.03% or less, 0.02% or less, 0.01% or less, 0.005% or less or 0.0025% or less compared to the expression of the 18S RNA (set as 100%) in the same cell. According to one embodiment, expression of gene FAM60A is even less, such as 0.001% or less, 0.0005% or less or even 0.0002 or less compared to the expression of the 18S RNA (set as 100%) in the same cell.

[51] According to one embodiment, the isolated eukaryotic cell which preferably is a mammalian cell, originates from a population of eukaryotic cells which are altered so that the effect of protein FAM60A is impaired in said cells and wherein said cells comprise stably integrated into their genome a heterologous polynucleotide encoding a product of interest, wherein on average at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 85% or at least 90% of the cells originating from said population do not lose more than 30%, preferably not more than 25%, of their product of interest expression titer over a time period of at least 8 weeks, preferably 10 weeks, more preferably over a time period of 12 weeks. As is shown in the examples, after transfection and identification of stably transfected cells, the amount of cells which do not show a gradual loss in productivity during prolonged culturing is increased when using the altered cells described herein, i.e. more stable cell clones are obtained from a selected cell population. The stability property can be tested by cultivating individual cells from said population as cell clones and determining the titer over the indicated time period. The stability can be tested using e.g. the assays described in the examples. As explained above, the stability rates can vary from project to project depending on the expressed protein and whether it is e.g. codon-optimized. However, with the altered eukaryotic cells according to the present disclosure, in all projects analyzed, a significant increase in stably expressing clones was observed compared to the unmodified wildtype cells. The percentage of cell clones that stably express the product of interest was in certain projects 80% or even higher in the analyzed clones during prolonged culturing for time spans in the range of 8 to 12 weeks. Therefore, the abundance of cells with stable expression characteristics was significantly increased in the population of successfully transfected host cells. Therefore, the risk that an unstable clone which gradually loses productivity during prolonged culturing is chosen for large scale production is significantly reduced with the teachings of the present disclosure. This important advantage allows to significantly reduce or even completely skip long term stability analyses in order to eliminate unstable clones.

[52] According to one embodiment, additionally the effect of the expression product of one or more genes selected from the group consisting of Bicdl, C12orf35, Amn1, methyltransferase-like protein 20, Dennd5b, Fam60a, Caprin2, lpo8, RPS4Y2 and Tmtc1 or one or more genes located telomeric of the aforementioned genes is impaired. Non-limiting alternative names of the aforementioned individual genes and/or the encoded proteins, including homologs and orthologs, are also indicated in Table 1 above and the respective genes encoding the respective proteins are encompassed by the scope of the terms used above for the individual genes or proteins encoded by said genes. Impairment of the effect
can likewise be achieved e.g. by reducing or eliminating the functional expression of the respective genes. Suitable technologies and embodiments are described above in conjunction with the FAM60A gene and likewise apply for any other target gene. As described above, said genes are located in the telomeric region of chromosome 8 of Chinese hamster and chromosome 6 of mouse. If a portion of said telomeric region is deleted, e.g. by inducing a chromosome breakage as described above, the deleted region usually comprises one or more of the aforementioned genes.

[53] According to a preferred embodiment, in said cell wherein the effect of protein FAM60A is impaired, additionally the effect of the expression product of gene C12orf35 is impaired in said cell, preferably by reducing or eliminating functional expression of gene C12orf35. A further unexpected finding was that impairing the effect of the expression product of gene C12orf35 in a eukaryotic cell, for example by reducing or eliminating functional expression of said endogenous gene, results in a significantly increase in the expression of the recombinant product of interest. Thus, a further key gene was identified that influences recombinant expression. Impairing the effect of the expression product of gene C12orf35 in a eukaryotic cell significantly increases the expression yield as is evidenced by the examples. Therefore, impairing the effect of FAM60A and C12orf35 is particularly advantageous, because host cells are provided which show improved characteristics with respect to expression stability and yield and hence, show particularly advantageous properties for the production of a recombinant product of interest. As described, the host cells are preferably mammalian cells.

[54] The C12orf35 gene is endogenously expressed in eukaryotic cells such as e.g. mammalian species such as human, mouse and hamster. The expression product of the C12orf35 gene is a rather large protein. The sequence listing shows exemplary amino acid sequences or putative amino acid sequences of the protein encoded by the endogenous C12orf35 gene of different mammalian species such as hamster (SEQ ID NO: 10 and 11), human (SEQ ID NO: 12 and 13), mouse (SEQ ID NO: 14), cattle (SEQ ID NO: 15) and wild boar (SEQ ID NO: 16). The CDS (Coding DNA Sequence) of C12orf35 from Chinese hamster is shown as SEQ ID NO: 17. Furthermore, a section of the 5'UTR (see SEQ ID NO: 18) and of the 3'UTR (see SEQ ID NO: 19) of the C12orf35 mRNA from Chinese hamster was sequenced. Gene C12orf35 is also referred to as C12orf35like or C12orf35 homolog in hamster or 281 047401 9Rik in mouse. Information about the gene, the coding sequence and the predicted C12orf35 protein is also disclosed for Cricetulus griseus in NCBI: XM_003512865, herein incorporated by reference. In human, it also referred to as KIAA1551. Different names can be assigned in different species for the protein or the gene and non-limiting alternative names (aliases) are also listed above in Table 1. Accordingly, the term "C12orf35 gene" as used herein in particular encompasses any endogenous gene which encodes a protein that shares at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% homology or identity to one or more of the amino acid sequences shown in SEQ ID NO: 10 to 16 or the protein encoded by SEQ ID NO 17. The protein encoded by such gene preferably has the same function as the protein having an amino acid sequence
as is shown in SEQ ID NO: 10 or one or more of SEQ ID NO: 11 to 16 or the protein encoded by SEQ ID NO: 17. Said gene can be modified as described herein in order to impair the function of the expression product that is expressed by the unmodified cell. The protein expressed by gene C12orf35 has not been described in detail in the literature. The terms “C12orf35 protein” or “expression product of endogenous C12orf35 gene” and similar expressions as used herein in particular encompass any protein that shares at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% homology or identity to one or more of the amino acid sequences shown in SEQ ID NO: 10 to 16 or the protein encoded by SEQ ID NO 17. Homology, respectively identity may be calculated over the entire length of the reference protein. According to one embodiment, expression of gene C12orf35 is reduced by at least 3 fold, at least 5 fold, at least 10fold, at least 20fold, at least 30fold, at least 40fold, at least 50fold, at least 60fold, at least 70fold, at least 75fold, at least 80fold, at least 90fold, at least 100fold or at least 125fold, at least 250fold, at least 500 fold, at least 750fold, at least 1000 fold, at least 1250fold, at least 1500fold, at least 1750fold or at least 2000fold. This can be determined e.g. by using real-time RT-PCR or other sensitive RNA detection methods. Such reduction can be achieved e.g. in comparison with the unmodified reference cell wherein the expression of endogenous gene C12orf35 is not reduced. According to one embodiment, expression of gene C12orf35 is 0.05% or less, 0.0475% or less, 0.045% or less, 0.0425% or less, 0.04% or less, 0.0375% or less, 0.035% or less, 0.0325% or less, 0.03% or less, 0.0275% or less, 0.025% or less, 0.0225% or less, 0.02% or less, 0.0175% or less, 0.015% or less compared to the expression of the 18S RNA (set as 100%) in the same cell. According to one embodiment, expression of gene C12orf35 is even less such as 0.001% or less, 0.0001% or less or even 0.00001% or less compared to the expression of the 18S RNA (set as 100%) in the same cell. The functional expression of gene C12orf35 is reduced such that it results in an increase in the expression of a recombinant product of interest if said modified eukaryotic cell is transfected with an expression vector encoding the product of interest compared to a corresponding cell wherein the functional expression of gene C12orf35 is not reduced or eliminated. According to one embodiment, expression of the recombinant product of interest is at least 1.5 times higher, at least 1.75 times higher, at least 2 times higher, at least 2.5 times higher, at least 3 times higher, at least 4 times higher or at least 5 times higher than the expression of a corresponding cell wherein the functional expression of gene C12orf35 is not reduced or eliminated. According embodiments, expression rates are obtained that are at least 8 times higher, at least 10 times higher or at least 15 times higher than the expression of a corresponding cell wherein the expression of gene C12orf35 is not reduced or eliminated. The expression rate of the recombinant product can be tested using e.g. the assays described in the examples.

[55] The eukaryotic cell is derived from a cell type which normally endogenously expresses FAM60A. Examples are described below. As explained above (see also Smith et al, 2012), FAM60A is endogenously expressed in eukaryotic cells such as all metazoans, in particular vertebrates but also invertebrates and in all mammalian cells. The term “isolated” is used to render clear that the eukaryotic cell is not contained in a living organism such as an animal or human. As described herein, the cell can be provided as cell culture, cell line, cell clone and
the like. Examples are also described below. As is described above, the eukaryotic cell is altered to impair the effect of FAM60A in said cell, e.g. by reducing or eliminating the effect of FAM60A compared to a corresponding, unmodified eukaryotic cell which endogenously expresses FAM60A. Impairment is preferably achieved by reducing or eliminating functional expression of gene FAM60A in the cell. Non-limiting embodiments are described above. In order to provide production cell lines with uniform and thus predictable stability characteristics, the genome of the eukaryotic cell is altered to achieve that result. Suitable embodiments are described above. The respectively altered eukaryotic cell may then be stably transfected with an expression vector comprising a polynucleotide encoding a product of interest in order to provide eukaryotic cells according to the first aspect which comprises integrated into the genome a heterologous polynucleotide encoding a product of interest. The eukaryotic cell preferably is a vertebrate cell, more preferred a mammalian cell. Thus, all embodiments described herein for eukaryotic cells in general apply to the preferred embodiment wherein mammalian cells are used. The eukaryotic cell may be e.g. selected from the group consisting of rodent cells, human cells and monkey cells. Preferred mammalian cells are rodent cells such as e.g. cells derived from hamster or mouse. They can be selected from the group consisting of a Chinese hamster cell such as a CHO cell, a BHK cell, a NSO cell, a C127 cell, a mouse 3T3 fibroblast cell, and a SP2/0 cell. Particularly preferred is a CHO cell, e.g. a CHO cell such as CHO-K1, CHO-S, CHO-K1SV, CHO-SSF3, CHO-DG44, CHO-DUXB11, or a cell line derived therefrom. As is shown in the examples, a knock-out of gene FAM60A in a CHO cell provides a population of CHO cells wherein the abundance of stably transfected cells with prolonged stability characteristics is increased. The FAM60A gene is also expressed in human cells. Thus, according to one embodiment, the mammalian cell is derived from a human cell, which may be e.g. selected from the group consisting of a HEK293 cell, a MCF-7 cell, a PerC6 cell, a CAP cell, hematopoietic cells and a HeLa cell. Another alternative are monkey cells, which, e.g. may be selected from the group consisting of a COS cells, COS-1, a COS-7 cell and a Vero cell. According to one embodiment, the eukaryotic cell, which preferably is a mammalian cell, is provided as cell clone or cell line.

[56] A eukaryotic cell wherein the genome is altered so that the effect of FAM60A is impaired in said cell and which does not comprise a heterologous polynucleotide encoding a product of interest, a heterologous polynucleotide encoding a selectable marker and/or a heterologous polynucleotide encoding a reporter polypeptide that is/are expressed, in particular secreted from said cell, can be used as starting material for preparing the eukaryotic cells according to the present disclosure. A respective "empty" altered eukaryotic cell can be used e.g. as cloning cell line for recombinant production technologies. A respective cell can be stably transfected with a heterologous polynucleotide encoding a product of interest, e.g. using an appropriate expression vector. Such "empty" eukaryotic cells in which the effect of FAM60A is impaired and which do not yet express and in particular do not secrete a recombinant product, can thus be transfected with different expression vectors, depending on the desired product of interest that is supposed to be recombinantly produced. Thus, such eukaryotic cell line can be used for different projects,
i.e. for the production of different products of interest, in particular different secreted polypeptides of interest.

[57] The eukaryotic cell according to the first aspect comprises a heterologous polynucleotide encoding a product of interest stably integrated into its genome. The product of interest is the recombinant product that is supposed to be expressed by the eukaryotic cell in large quantity. Preferably, the product of interest is a polypeptide. According to a preferred embodiment, the polypeptide of interest is secreted by the cell. The eukaryotic cell may additionally comprise a heterologous polynucleotide encoding a selectable marker and/or a heterologous polynucleotide encoding a reporter. This simplifies the selection of host cells which are successfully transfected and thus express the product of interest. Furthermore, the eukaryotic cell may comprise several polynucleotides encoding different selectable markers and/or reporter polypeptides.

[58] A "heterologous polynucleotide" or "heterologous nucleic acid" and likewise expressions used herein in particular refer to a polynucleotide sequence that has been introduced into the eukaryotic cell e.g. by the use of recombinant techniques such as transfection. A "polynucleotide" in particular refers to a polymer of nucleotides which are usually linked from one deoxyribose or ribose to another and refers to DNA as well as RNA, depending on the context. The term "polynucleotide" does not comprise any size restrictions.

[59] An expression vector can be used to introduce heterologous polynucleotides. The polynucleotides can be comprised in an expression cassette. The polynucleotide(s) encoding the product of interest and the polynucleotide(s) encoding a selectable marker or reporter polypeptide may be located on the same or different expression vectors. Introduction into the eukaryotic cell may be achieved e.g. by transflecting a suitable expression vector comprising the polynucleotide encoding the product of interest into the host cells. The expression vector integrates into the genome of the host cell (stable transfection). As is shown by the examples, the novel eukaryotic cells described herein are advantageous for stable transfection as the number of clones with prolonged stability is increased. Stable transfection is also the standard for generating high expressing cell clones for producing a product of interest such as a polypeptide of interest on industrial scale. This is particularly important for therapeutic or diagnostic polypeptides of interest. Several appropriate methods are known in the prior art for introducing a heterologous nucleic acid such as an expression vector into eukaryotic such as mammalian host cells and thus, do not need any detailed description herein. Respective methods include but are not limited to calcium phosphate transfection, electroporation, lipofection, biolistic- and polymer-mediated genes transfer and the like. Besides traditional random integration based methods also recombination mediated approaches can be used to transfer the heterologous polynucleotide into the host cell genome. As respective methods are well known in the prior art, they do not need any detailed description here. Non-limiting embodiments of suitable vector designs are also described subsequently and it is referred to the respective disclosure.
Expression vectors used to achieve expression of a recombinant product of interest usually contain transcriptional control elements suitable to drive transcription such as e.g. promoters, enhancers, polyadenylation signals, transcription pausing or termination signals usually as element of an expression cassette. If the desired product is a polypeptide, suitable translational control elements are preferably included in the vector, such as e.g. 5' untranslated regions leading to 5' cap structures suitable for recruiting ribosomes and stop codons to terminate the translation process. The resultant transcripts harbour functional translation elements that facilitate protein expression (i.e. translation) and proper translation termination. A functional expression unit, capable of properly driving the expression of an incorporated polynucleotide is also referred to as an "expression cassette". It is well-known to the skilled person how an expression cassette shall be designed in order to allow the expression in a eukaryotic cell, such as preferably in a mammalian cell.

The polynucleotide(s) encoding the product of interest and the polynucleotides encoding the selectable marker(s) and/or reporter polypeptide(s) as described herein are preferably comprised in expression cassettes. Several embodiments are suitable. For example, each of said polynucleotide(s) can be comprised in a separate expression cassette. This is also referred to as monocistronic setting. It is also within the scope of the present invention that at least two of the respective polynucleotides are comprised in one expression cassette. According to one embodiment, at least one internal ribosomal entry site (IRES) element is functionally located between the polynucleotides that are expressed from the same expression cassette. Thereby, it is ensured that separate translation products are obtained from said transcript. Respective IRES based expression technologies and other bi- and polycistronic systems are well known and thus need no further description here.

As described, the expression vector may comprise at least one promoter and/or promoter/enhancer element as element of an expression cassette. Promoters can be divided in two classes, those that function constitutively and those that are regulated by induction or derepression. Both are suitable. Strong constitutive promoters which drive expression in many cell types include but are not limited to the adenovirus major late promoter, the human cytomegalovirus immediate early promoter, the SV40 and Rous Sarcoma virus promoter, and the murine 3-phosphoglycerate kinase promoter, EF1a. According to one embodiment, the promoter and/or enhancer is either obtained from CMV and/or SV40. The transcription promoters can be selected from the group consisting of an SV40 promoter, a CMV promoter, an EF1 alpha promoter, a RSV promoter, a BROAD3 promoter, a murine rosa 26 promoter, a pCEFL promoter and a β-actin promoter. Also other promoters can be used if they result in expression of the product of interest in the eukaryotic cell which preferably is a mammalian cell.

Furthermore, an expression cassette may comprise at least one intron. Usually, introns are placed at the 5' end of the open reading frame but may also be placed at the 3' end. Said intron may be located between the promoter and or promoter/enhancer element(s) and the 5' end of the open reading frame of the polynucleotide encoding the product of interest to be
expressed. Several suitable introns are known in the state of the art that can be used in conjunction with the present disclosure.

[64] The product of interest can be any biological product capable of being produced by transcription, translation or any other event of expression of the genetic information encoded by the polynucleotide encoding the product of interest. The product of interest may be selected from the group consisting of polypeptides and nucleic acids, in particular RNA. The product can be a pharmaceutically or therapeutically active compound, or a research tool to be utilized in assays and the like. Preferably, the product of interest is a polypeptide. Any polypeptide of interest can be expressed with the method of the present invention. The term "polypeptide" refers to a molecule comprising a polymer of amino acids linked together by a peptide bond(s). Polypeptides include polypeptides of any length, including proteins (e.g. having more than 50 amino acids) and peptides (e.g. 2 - 49 amino acids). Polypeptides include proteins and/or peptides of any activity, function or size, and may include e.g. enzymes (e.g. proteases, kinases, phosphatases), receptors, transporters, bacterial and/or endotoxin-binding proteins, structural polypeptides, membrane-bound polypeptides, glycoproteins, globular proteins, immune polypeptides, toxins, antibiotics, hormones, growth factors, blood factors, vaccines or the like. The polypeptide may be selected from the group consisting of peptide hormones, interleukins, tissue plasminogen activators, cytokines, immunoglobulins, in particular antibodies or functional antibody fragments or variants thereof and Fc-fusion proteins. The polypeptide of interest that is expressed according to the teachings described herein may also be a subunit or domain of a polypeptide, such as e.g. a heavy chain or a light chain of an antibody or a functional fragment or derivative thereof. The terms "product of interest" or "polypeptide of interest" may refer to such individual subunit or domain or the final protein that is composed of the respective subunits or domains, depending on the context. In a preferred embodiment the polypeptide of interest is an immunoglobulin molecule, more preferably an antibody, or a subunit or domain thereof such as e.g. the heavy or light chain of an antibody. The term "antibody" as used herein particularly refers to a protein comprising at least two heavy chains and two light chains connected by disulfide bonds. The term "antibody" includes naturally occurring antibodies as well as all recombinant forms of antibodies, e.g., humanized antibodies, fully human antibodies and chimeric antibodies. Each heavy chain is usually comprised of a heavy chain variable region (VH) and a heavy chain constant region (CH). Each light chain is usually comprised of a light chain variable region (VL) and a light chain constant region (CL). The term "antibody", however, also includes other types of antibodies such as single domain antibodies, heavy chain antibodies, i.e. antibodies only composed of one or more, in particular two heavy chains, and nanobodies, i.e. antibodies only composed of a single monomeric variable domain. As discussed above, the polynucleotide encoding the polypeptide of interest may also encode one or more subunits or domains of an antibody, e.g. a heavy or a light chain or a functional fragment or derivative thereof, as polypeptide of interest. Said subunits or domains can be expressed either from the same or different expression cassettes. A "functional fragment or derivative" of an antibody in particular refers to a polypeptide which is derived from an antibody and is capable of binding to the same antigen, in particular to the same epitope as the antibody. It has been shown that the
antigen-binding function of an antibody can be executed by fragments of a full-length antibody or derivatives thereof. Examples of fragments or derivatives of an antibody include (i) Fab fragments, monovalent fragments consisting of the variable region and the first constant domain of each the heavy and the light chain; (ii) F(ab)₂ fragments, bivalent fragments comprising two Fab fragments linked by a disulfide bridge at the hinge region; (iii) Fd fragments consisting of the variable region and the first constant domain CH1 of the heavy chain; (iv) Fv fragments consisting of the heavy chain and light chain variable region of a single arm of an antibody; (v) scFv fragments, Fv fragments consisting of a single polypeptide chain; (vi) (Fv)₂ fragments consisting of two Fv fragments covalently linked together; (vii) a heavy chain variable domain; and (viii) multibodies consisting of a heavy chain variable region and a light chain variable region covalently linked together in such a manner that association of the heavy chain and light chain variable regions can only occur intermolecular but not intramolecular. According to one embodiment, the eukaryotic cell secretes the polypeptide of interest into the cell culture medium. According to one embodiment, the polypeptide of interest is not or does not comprise SIN3A. According to one embodiment, the polypeptide of interest is not or does not comprise FAM60A.

[65] The eukaryotic cell may or may not comprise an endogenous polynucleotide corresponding to, respectively being identical to the polynucleotide encoding the product of interest. According to one embodiment, the eukaryotic cell does not comprise an endogenous gene corresponding to the product of interest.

[66] As described, in embodiments, the eukaryotic cell comprises at least one heterologous polynucleotide encoding a selectable marker and/or a heterologous polynucleotide encoding a reporter polypeptide in addition to the heterologous polynucleotide encoding the product of interest.

[67] A "selectable marker" allows under appropriate selective culture conditions the selection of host cells expressing said selectable marker. A selectable marker provides the carrier of said marker under selective conditions with a survival and/or growth advantage. Thereby, host cells successfully transfected with the expression vector can be selected under appropriate selection conditions. Typically, a selectable marker gene will confer resistance to a selection agent such as a drug, e.g. an antibiotic or other toxic agent, or compensate for a metabolic or catabolic defect in the host cell. It may be a positive or negative selection marker. For selecting successfully transfected host cells a culture medium may be used for culturing the host cells comprises a selection agent that allows selection for the selectable marker used. In other embodiments, the selection marker enables the host cell to survive and proliferate in the absence or reduction of a compound which is essential for survival and/or proliferation of the host cells lacking the selection marker. By cultivating the host cells in a medium which does not comprise the essential compound in a concentration high enough for survival and/or proliferation of the host cell or comprises a reduced amount of said essential compound, only host cells expressing the selection marker can survive and/or proliferate. According to one embodiment, the selectable marker is a drug resistance marker encoding a protein that confers resistance to selection conditions involving said drug. A
A variety of selectable marker genes have been described (see, e.g., WO 92/08796, WO 94/28143, WO2004/081 167, WO2009/080759, WO2010/097240). E.g. at least one selectable marker may be used which confers resistance against one or more antibiotic agents. The selectable marker may according to one embodiment be an amplifiable selectable marker. An amplifiable selectable marker allows the selection of vector containing host cells and may promote gene amplification of said vector in the host cells. Selectable marker genes commonly used with eukaryotic cells such as in particular mammalian cells include the genes for aminoglycoside phosphotransferase (APH), hygromycin phosphotransferase (hyg), dihydrofolate reductase (DHFR), thymidine kinase (tk), glutamine synthetase, asparagine synthetase, and genes encoding resistance to neomycin (G418), puromycin, hygromycin, zeocin, ouabain, blasticidin, histidinol D, bleomycin, pheomycin and mycophenolic acid. According to one embodiment, a folate receptor is used as selectable marker in conjunction with the novel eukaryotic cells described herein (see e.g. WO2009/080759), which preferably are mammalian cells. According to one embodiment, the eukaryotic cell which is dependent on folate uptake comprises a heterologous polynucleotide encoding a folate receptor as selectable marker and/or comprises a heterologous polynucleotide encoding a dihydrofolate reductase (DHFR) as selectable marker. This embodiment will also be described in detail below in conjunction with the selection method according to the second aspect. The eukaryotic cell may express endogenously DHFR and a folate receptor.

[68] A "reporter polypeptide" allows the identification of a cell expressing said reporter polypeptide based on the reporting characteristics (e.g. fluorescence). Reporter genes usually do not provide the host cells with a survival advantage. However, the expression of the reporter polypeptide can be used to differentiate between cells expressing the reporter polypeptide and those cells which do not. Therefore, also a reporter gene enables the selection of successfully transfected host cells. Suitable reporter polypeptides include but are not limited to as e.g. green fluorescence protein (GFP), YFP, CFP and luciferase. According to one embodiment, the reporter polypeptide has characteristics that enable the selection by flow cytometry.

[69] As described, the expression vector comprising the polynucleotide encoding the product of interest may also comprise more than one selectable marker and/or reporter gene. Furthermore, the one or more polynucleotides encoding the selectable marker(s) and/or the one or more polynucleotides encoding the reporter polypeptide(s) may also be provided on one or more different expression vectors which are co-transfected with the expression vector which comprises the polynucleotide encoding the product of interest. Such co-transfection strategies likewise enable selection as is well-known in the prior art.

[70] The expression vector or the combination of at least two expression vectors comprised in the eukaryotic cell may additionally comprise further vector elements. E.g. at least one additional polynucleotide encoding a further product of interest can be comprised. As explained above and as becomes apparent from the above described examples of polypeptides that can be expressed according to the present teachings, the final polypeptide
that is to be produced and preferably secreted by the host cell can also be a protein that is composed of several individual subunits or domains. A preferred example of a respective protein is an immunoglobulin molecule, in particular an antibody that comprises e.g. heavy and light chains. There are several options for producing a respective protein that is composed of different individual subunits or domains and appropriate vector designs are known in the art. According to one embodiment, two or more subunits or domains of said protein are expressed from one expression cassette. In this embodiment, one long transcript is obtained from the respective expression cassette that comprises the coding regions of the individual subunits or domains of the protein. According to one embodiment, at least one IRES element (internal ribosomal entry site) is functionally located between the coding regions of the individual subunits or domains and each coding region is preceded by a secretory leader sequence. Thereby, it is ensured that separate translation products are obtained from said transcript and that the final protein can be correctly assembled and secreted. Respective technologies are known in the prior art and thus, do not need any detailed description herein.

[71] For some embodiments such as the expression of antibodies it is even preferred to express the individual subunits or domains from different expression cassettes. According to one embodiment, the expression cassette used for expressing the product of interest is a monocistronic expression cassette. All expression cassettes comprised in the expression vector or combination of expression vectors may be monocistronic. According to one embodiment, accordingly, each expression cassette designed for expressing a product of interest comprises a polynucleotide encoding one subunit or domain of the protein to be expressed as polypeptide of interest. E.g. in case of antibodies, one expression cassette may encode the light chain of an antibody and another expression cassette may encode the heavy chain of the antibody. After expression of the individual subunits or domains from the individual expression cassettes, the final protein such as an antibody is assembled from said subunits or domains and secreted by the host cell. This embodiment is particularly suitable for expressing immunoglobulin molecules such as antibodies. In this case, a first heterologous polynucleotide encoding a product of interest encodes e.g. the heavy or the light chain of an immunoglobulin molecule and a second heterologous polynucleotide encoding a product of interest encodes the other chain of the immunoglobulin molecule. According to one embodiment, the expression vector or combination of at least two expression vectors used for transfecting the mammalian host cell comprises at least one expression cassette comprising a polynucleotide encoding the heavy chain of an immunoglobulin molecule or a functional fragment thereof and at least one expression cassette comprising a polynucleotide encoding the light chain of an immunoglobulin molecule or a functional fragment thereof. Said polynucleotides may be located on the same or on different expression vectors in case a combination of at least two expression vectors is used. Upon expression of said polynucleotides in the transfected host cell, a functional immunoglobulin molecule is obtained and preferably is secreted from the host cell. As is shown by the examples, using the novel cells and cell lines described herein wherein the genome of the host cell is altered so that the effect of FAM60A is impaired in said cell, preferably by reducing or eliminating functional expression of said gene, are particularly
suitable for expressing proteins due to their improved stability characteristics. Furthermore, in embodiments, in particular wherein the effect of the expression product of gene c12orf35 is additionally impaired, the expression yield also increases significantly. Therefore, the novel eukaryotic cell lines described herein have particular advantages when being used for recombinant expression of polypeptides, including proteins that are composed of several subunits of domains such as e.g. antibodies. Further advantages are also described in conjunction with the examples.

**B. Selection method**

[72] According to a second aspect, a method for selecting a host cell which recombinantly expresses a product of interest is provided, comprising

(a) providing eukaryotic cells according to the first aspect as host cells; and
(b) selecting one or more host cells expressing the product of interest.

[73] The eukaryotic host cells according to the first aspect, including suitable and preferred embodiments, as well as their advantages are described in detail above and it is referred to the respective disclosure which also applies here. As described, vertebrate cells, in particular mammalian cells, are preferably used as host cells. The advantageous properties of said eukaryotic cells simplify e.g. the selection and thus identification of suitable production clones. Furthermore, as the proportion of cells with favourable stability characteristics in the transfected cell population is significantly increased when using the eukaryotic cells described herein, less clones need to be analysed and screened for their production characteristics in order to identify suitable stable production clones. This saves time and furthermore, allows handling more projects in parallel. Furthermore, as is demonstrated by the examples, in embodiments, the number respectively proportion of high expressing cells with favourable characteristics is increased after stable transfection and selection steps. Such expressing cells, also called cell pools, produce significant quantities of the product of interest. Thus, such cell pools comprising high expressing cells can be e.g. used in order to produce a polypeptide of interest within a short timeframe. Therefore, the product of interest can be produced rapidly in respective cells.

[74] According to one embodiment, stage (a) of the selection method according to the second aspect comprises transfecting eukaryotic cells wherein the genome of said cells is altered so that the effect of protein FAM60A is impaired in said cells with a heterologous polynucleotide encoding the product of interest, thereby providing respective eukaryotic cells which comprise a heterologous polynucleotide encoding the product of interest stably integrated into the genome. As described, mammalian cells are preferably used as eukaryotic host cells. The polynucleotide encoding the product of interest may be comprised in an expression vector that is then transfected into the eukaryotic cell.

[75] Selection stage (b) may be a multi-step selection process comprising several selection steps in order to select and thus identify host cells that express a product of interest with high yield. For example, stage (b) may include one or more selection steps to identify cells that
were successfully transfected as well as one or more subsequent selection steps to select high expressing cells from the pool of successfully transfected cells. The appropriate selection strategy depends on the design of the expression vector that is used for introducing the polynucleotide encoding the product of interest and in particular depends on the used selection marker(s) and/or reporter(s). Non-limiting embodiments will be described in the following.

[76] As described above, the eukaryotic host cells may comprise at least one heterologous polynucleotide encoding a selectable marker. The polynucleotide encoding the selectable marker can be introduced into the host cell together with the polynucleotide encoding the product of interest using either the same or a different co-transfected expression vector. Stage (b) then comprises culturing said plurality of host cells under conditions providing a selection pressure to the host cells to identify successfully transfected host cells, e.g. using an appropriate selection medium. As used herein, a "selection medium" in particular refers to a cell culture medium useful for the selection of host cells that express the selectable marker. It may include e.g. a selection agent such as a toxic drug which allows to identify successfully transfected host cells. Alternatively, an essential compound can be absent or its concentration can be reduced in the selection medium. According to one embodiment, host cells which were not successfully transfected and hence, do not express the selection marker(s) or wherein expression is low cannot proliferate or die under the selective cultivation conditions. In contrast, host cells which were successfully transfected with the expression vector(s) and which express the selection marker(s) (and accordingly the co-introduced product of interest) with sufficient yield are resistant to or are less affected by the selection pressure and therefore, can proliferate, thereby outgrowing the host cells which were not successfully transfected or wherein the integration site into the genome of the cell is not favourable. According to one embodiment, the selectable marker is selected from the group consisting of antibiotic resistance markers, drug resistance markers and metabolic markers. Suitable examples for selectable markers and selection principles are described above in conjunction with the first aspect and appropriate selection conditions for the individual selectable markers are also well-known to the skilled person. Non-limiting but advantageous embodiments will be described briefly subsequently.

[77] As is shown by the examples, one advantage of the novel eukaryotic cells described herein, such as in particular a CHO cell line comprising a deletion in the telomeric region of chromosome 8 which includes gene FAM60A and therefore also gene C12orf35, is that already after one standard selection step, such as e.g. a G418/neo selection, the percentage of high-producing cells with stable characteristics is significantly increased. For example, when transfecting normal CHO cell lines, which are not respectively altered, often 60% or even up to 80% of the cells surviving G418/neo selection are non-producers. The number of non- or low producers is significantly reduced when using the novel cell lines according to the present disclosure. This allows to efficiently perform the selection in stage (b) by using only one selectable marker and thus only one selection step, such as for example a G418/neo selection, if desired. This increases the speed of selection and accordingly, reduces the time required for obtaining stably transfected cells that express the product of interest.
Furthermore, because the number of high and stable producers is increased, it was found that even in the absence of a specific selection for high producing cells, cell populations can be generated with clone like performance and thus, within very short time. Therefore, the product of interest can even be manufactured from such pools. Thus, for applications wherein e.g. only certain amounts of protein are needed quickly, for example for research or testing purposes, such a rapid selection system which quickly renders good producing cells or producing cell pools is highly advantageous.

[78] However, if it is desired to further increase the productivity, a multi-step selection in stage (b) is preferred. This is particularly the case when intending to establish a clonal cell line for producing the product of interest on large scale, in particular industrial scale.

[79] As described above, the expression vector or expression vectors introduced into the cells described herein may comprise two or more selectable marker genes and selection for the different selectable markers may be done simultaneously or sequentially for selecting host cells which were successfully transfected with the expression vector(s) and express the product of interest with high yield. The selection medium used for cultivation may comprise selection agents for all selectable markers used. In another embodiment, cultivation can be performed first with a selection medium only comprising the selection agent(s) of one or a subset of the selectable marker genes used, followed by the addition of one or more of the selection agents for the remaining selectable marker genes. In another embodiment, the host cells are cultivated with a first selection medium only comprising the selection agent(s) of one or a subset of the selectable marker genes of the vector(s), followed by cultivation with a second selection medium comprising the selection agent(s) of one or more of the selection agents of the remaining selectable marker genes. According to one embodiment, the second selection medium does not comprise the selection agent(s) used in the first selection medium.

[80] According to one embodiment, the eukaryotic cells provided in stage (a) comprise a heterologous polynucleotide encoding a dihydrofolate reductase (DHFR) as selectable marker and stage (b) comprises performing a selection step for DHFR. A respective selection step usually involves a selection medium comprising a DHFR inhibitor. Several suitable DHFR enzymes and accordingly DHFR genes are known in the prior art that can be used as selectable marker. The term DHFR refers to wild type DHFR as well as to DHFR enzymes having one or more amino acid sequence exchanges (e.g. deletions, substitutions or additions) with respect to the amino acid sequence of the corresponding wildtype DHFR enzyme, fusion proteins comprising a DHFR enzyme and DHFR enzymes which have been modified to provide an additional structure and/or function, as well as functional fragments of the foregoing, which still have at least one function of a DHFR enzyme. Such embodiments are well-known in the prior art and thus, do not need to be described in detail. For example, a DHFR enzyme may be used as selectable marker that is more or less sensitive to antifolates such as MTX than the wild type DHFR enzyme and/or the DHFR enzyme endogenously expressed by the host cell if expressed. Respective DHFR enzymes are well-known in the prior art and e.g. are described in EP 0 246 049 and other documents. The DHFR enzyme
can be derived from any species as long as it will be functional within the present invention, i.e. compatible with the mammalian host cell utilised. E.g. a mutant mouse DHFR with a major resistance to MTX has been extensively used as a dominant selectable marker in eukaryotic cells. A DHFR enzyme may be used as selectable marker which is less susceptible to a DHFR inhibitor such as MTX than the DHFR enzyme endogenously expressed in a DHFR + (plus) host cell and thus a host cell which comprises a functional endogenous DHFR gene. According to one embodiment, an intron or a fragment thereof is placed at the 3’ end of the open reading frame of the DHFR gene. The intron used in the DHFR expression cassette is leading to a smaller, non-functional variant of the DHFR gene (Grillari et al., 2001, J. Biotechnol. 87, 59-65). Thereby, the expression level of the DHFR gene is lowered which further increases the stringency of selection. Alternative methods making use of an intron to reduce the expression level of the DHFR gene are described in EPO 724 639 and could also be used.

[81] A “DHFR inhibitor” in particular refers to a compound which inhibits the activity of dihydrofolate reductase (DHFR). A respective inhibitor may for example compete with the DHFR substrate for binding to DHFR. Suitable DHFR inhibitors are for example antifolates such as methotrexate (MTX). Thus, according to one embodiment, stage (b) comprises performing a selection step for DHFR by cultivating said plurality of host cells in a selective culture medium which comprises at least one DHFR inhibitor, such as preferably an antifolate. Respective selection conditions are known to the skilled person. According to one embodiment, a selective culture medium is used in stage (b) which comprises an antifolate in a concentration of 1500nM or less, 1250nM or less, 1000nM or less, 750nM or less, 500nM or less, 250nM or less, 200nM or less, 150nM or less, 125nM or less, 100nM or less or 75nM or less. The used concentration of said inhibitor in the selective culture medium (which may also be increased gradually), determines the stringency of the selection conditions. Preferred concentration ranges for the antifolate may be selected from 500nM - 0.1 nM, 350nM - 1nM, 200nM - 2.5nM, 150nM - 5nM, 100nM - 7.5nM and 75nM - 10 nM. According to one embodiment, the selective culture medium comprises MTX as antifolate. Low MTX concentrations can be used in conjunction with the novel eukaryotic cells described herein for performing a DHFR selection, in particular if selection is performed in combination with a limiting concentration of folate.

[82] According to one embodiment, the host cells according to the present disclosure provided in stage (a) comprise a heterologous polynucleotide encoding a folate transporter as selectable marker. A folate transporter based selection system has several advantages when being used in conjunction with eukaryotic cells that are dependent on folate uptake such as mammalian cells. A folate transporter allows to import at least one folate from the culture medium into the host cell which preferably is a mammalian cell. According to one embodiment, the folate transporter is or comprises the reduced folate carrier (RFC). RFC is a ubiquitously expressed membrane glycoprotein that serves as the major transporter for the uptake of reduced folates such as 5-methyl-THF and 5-formyl-THF into the cell. However, RFC displays a very poor affinity for the oxidized folate, folic acid. Hence, cells that lack the expression of RFC or have been deleted from the genomic RFC locus can serve as
recipients for the transfection of the selectable marker gene RFC under conditions in which reduced folates such as 5-formyl-THF are gradually deprived from the growth medium, thereby allowing to identify cells that express increased levels of the this folate transporter and accordingly the product of interest. Suitable selection conditions when using RFC as selectable marker are known to the skilled person and are e.g. described in WO2006/059323.

[83] According to one embodiment which is also described in detail below and in the example section, the folate transporter used as selectable marker is a folate receptor. A folate receptor based selection system has several advantages. For example for selection, no toxic substances are needed (even though they can be used) and furthermore, the endogenous folate receptor of the host cell does not need to be knocked out. Furthermore, this expression system works particularly well in conjunction with the novel cells described herein, which preferably are mammalian cells.

[84] A “folate receptor” as used herein refers to a folate receptor that is functional and thus capable of import or uptake of a folate or derivative thereof into the eukaryotic cell. Preferably, the receptor is capable of unidirectional import or uptake of folate or a derivative of folate into the eukaryotic cell. Furthermore, a folate receptor as used herein is membrane-bound. Thus, the folate receptors described herein are functional membrane-bound folate receptors. Membrane anchorage can be achieved e.g. by a transmembrane anchor or by a GPI anchor. A GPI anchor is preferred as it corresponds to the natural setting of membrane-bound folate receptor. Folate receptors (FRs) are high-affinity folate-binding glycoproteins. They are encoded by three distinct genes FR alpha, FR beta and FR gamma. FR alpha and FR beta are glycosylphosphatidylinositol (GPI)-anchored, cell surface glycoproteins, whereas FR gamma is devoid of a GPI anchor and is a secreted protein. However, it can be genetically altered to include a transmembrane anchor or a GPI anchor. Such an altered form of a FR gamma that includes a membrane anchor is also considered a membrane-bound folate receptor if it is capable of import or uptake of a folate or derivative thereof into a eukaryotic cell. The term “folate receptor” also includes membrane-bound mutants of wildtype folate receptors that are capable of folate uptake and fusion proteins comprising a respective folate receptor.

[85] The folate receptor utilized can be derived from a folate receptor of any species as long as it will be functional within the present invention, i.e. it is compatible with the host cell that is utilized and when being expressed in the transfected host cell, incorporates folate, in particular folic acid, from the culture medium into the host cell that is dependent on folate uptake. In general, the folate receptor that is introduced into the host cell as selectable marker can be homologous or heterologous to an endogenous folate receptor of the host cell (if an endogenous folate receptor is present what is preferred). If it is homologous, it will be derived from the same species as the host cell which preferably is a mammalian host cell. If it is heterologous, it will be derived from another species than the host cell. For example a human folate receptor may be used as selectable marker for a rodent host cell, e.g. a hamster cell such as a CHO cell. Preferably, a folate receptor derived from a mammalian
species is used, for example derived from a rodent, such as mouse, rat or hamster, or, more preferred, derived from a human. The membrane-bound folate receptor used as selectable marker can be selected from the group consisting of a folate receptor alpha, a folate receptor beta and a folate receptor gamma. According to one embodiment, a human folate receptor alpha is used as selectable marker.

[86] The use of a membrane-bound folate receptor as folate transporter is advantageous because cells can be used, which endogenously express their own folate receptors and folic acid can be processed by the folate receptor. Suitable membrane-bound folate receptors that can be used as selectable markers for eukaryotic cells that are dependent on folate uptake such as mammalian cells and suitable selection conditions are also described in WO 2009/080759, herein incorporated by reference. According to one embodiment, the mature wild type human folate receptor alpha is used which comprises the following amino acid sequence shown in SEQ ID NO: 20 (1-letter code, shown in direction from N-terminus to C-terminus).

[87] Folate receptor alpha is naturally anchored to the cell membrane by a GPI anchor. The signal sequence for a GPI anchor is not shown in SEQ ID NO: 20. According to one embodiment, the folate receptor alpha which is derived from or comprises SEQ ID NO: 20 comprises a GPI anchor signal at the C-terminus. Any suitable GPI anchor signal may be used. The natural GPI anchor signal sequence of human folate receptor alpha is shown in SEQ ID NO: 21 (1-letter code, shown in direction from N-terminus to C-terminus).

[88] Membrane anchorage may alternatively be achieved by using a membrane anchor, e.g. a transmembrane anchor. In this embodiment, the folate receptor comprises a membrane anchor at its C-terminus. Suitable anchors are known in the prior art. The folate receptor alpha which is derived from or comprises SEQ ID NO: 20 may comprise a leader sequence at the N-terminus. Any suitable leader sequence can be used which ensures functional expression of the folate receptor.

[89] The full amino acid sequence including the natural leader sequence (at the N-terminus) and the natural GPI anchor signal sequence (at the C-terminus) of the wild type human folate receptor alpha is shown in SEQ ID NO: 22 (1-letter code, shown in direction from N-terminus to C-terminus).

[90] According to one embodiment, the membrane-bound folate receptor has or comprises the amino acid sequence of SEQ ID NO: 20, or 22 or is a functional variant of the foregoing which has at least 80%, at least 85%, at least 90%, at least 95%, at least 98% or at least 99% homology or identity to SEQ ID NO: 20 or 22, is membrane-bound and is capable of folic acid uptake into the cell.

[91] When using a eukaryotic cell that is dependent on folate uptake such as a mammalian host cell, which comprises a heterologous polynucleotide encoding a folate transporter, preferably a folate receptor, stage (b) comprises a selection step wherein said plurality of
host cells are cultured in a selective culture medium comprising folate in a limiting concentration. A "limiting concentration of folate" as used herein in particular refers to a concentration of folate(s) in the selective culture medium which provides a selective pressure on the host cell. Under such selection conditions, only host cells grow and proliferate that have incorporated the expression vector and thus, express the folate transporter as selectable marker. Accordingly, folates are not comprised in the selective culture medium in affluence, and this limitation of folate(s) in the culture medium provides a selection pressure on the host cells. The folate comprised in the selective culture medium in a limiting concentration is capable of being taken up into and being processed by the host cell, in particular by host cells that have incorporated the folate transporter that is used as selectable marker. Folates and in particular derivatives of folate which would not or cannot be processed by the host cell do not contribute to the selection pressure that is exerted to select host cells that have incorporated the folate transporter as selectable marker and accordingly do not contribute to the limiting concentration of folate. However, respective folates, such as e.g. antifolates, may be present and even preferably are present, if a combined selection with DHFR is performed as will be described subsequently. The folate present in the selective culture medium in a limiting concentration can e.g. be an oxidized folate or a reduced folate or a derivative thereof. Oxidized folates, such as folic acid, as well as reduced derivatives of folic acid, known as reduced folates or tetrahydrofolates (THF), are a group of B-9 vitamins that are essential cofactors and/or coenzymes for the biosynthesis of purines, thymidylate and certain amino acids in eukaryotic cells that are dependent on folate uptake such as mammalian cells. THF cofactors are particularly crucial for DNA replication and hence cellular proliferation. Preferably, the folate that is comprised in a limiting concentration in the selective culture medium is folic acid. Suitable concentration ranges for providing a limiting concentration of folate are described below.

[92] The folate transporter based selection system is based on the limited availability of folate, preferably folic acid, in the cell culture medium. Host cells that have not successfully incorporated the expression vector(s) and hence do not express sufficient quantities of the folate transporter, which preferably is a folate receptor, die or are impaired in growth under the selective culture conditions compared to host cells that have successfully incorporated the expression vector(s). As is shown by the examples, using a folate receptor based selection in combination with the novel cells described herein allows an accelerated selection, screening and establishment of cell clones that stably overexpress high levels of recombinant products of interest such as polypeptides.

[93] The selective culture medium that is used in stage (b), respectively used in a sub-step of stage (b), for selection against a folate receptor as selectable marker may comprise folate and in particular folic acid in a concentration selected from:

(a) about 5000nM - 0.1 nM;
(b) about 2500nM - 0.1 nM;
(c) about 1500nM - 0.1 nM;
(d) about 1000nM - 0.1 nM;
(e) about 750nM - 0.1 nM;
(f) about 500nM - 0.1 nM;
(g) about 250nM - 0.2nM; preferably about 250nM - 1 nM or about 250nM - 2.5nM;
(h) about 150nM - 0.3nM; preferably about 150nM - 1 nM or about 150nM - 2.5nM;
(i) about 100nM - 0.5nM; preferably about 100nM - 1 nM or about 100nM - 2.5nM;
(j) about 75nM - 0.6nM, preferably about 75nM - 1 nM or about 75nM - 2.5nM;
(k) about 50nM - 1nM; preferably about 50nM - 2.5nM or about 50nM - 5nM;
(l) about 35nM - 0.75nM; and
(m) about 25nM - 1nM or about 25nM - 2.5nM, about 20nM - 3nM about 15nM - 4nM or 10nM - 5nM.

[94] The concentrations and concentration ranges described above are particularly suitable for fast growing suspension mammalian cells, such as CHO cells, which is a preferred embodiment for commercial production cell lines. However, different cell lines may have different folic acid consumption properties. Suitable concentrations, however, can easily be determined experimentally by the skilled person. The folate comprised in the selective culture medium is preferably folic acid and according to one embodiment, folic acid is the only folate comprised in the selective culture medium that contributes to the limiting concentration of folate.

[95] According to one embodiment, the host cells provided in stage (a) comprise a heterologous polynucleotide encoding a folate transporter as first selectable marker and a heterologous polynucleotide encoding a second selectable marker which processes a substrate which is a folate, a derivative of a folate and/or a product that can be obtained by the processing of folate. E.g. the second selectable marker may be a dihydrofolate reductase (DHFR) or an enzyme operating downstream or in conjunction with DHFR such as thymidylate synthase (TS) and serine hydroxymethyltransferase (SHMT). Preferably, a membrane-bound folate receptor is used as first selectable marker and the second selectable marker is a dihydrofolate reductase (DHFR). According to this embodiment, stage (b) comprises culturing the host cells in a selective culture medium which comprises folate, preferably folic acid, in a limiting concentration and comprises at least one DHFR inhibitor such as e.g. MTX. As is shown by the examples, when using the novel cell lines described herein the number of high and stable expressing cells in the pool is significantly increased, obtaining suitable cell clones for large scale production from said cell pool is simplified. Less cell clones need to be made what reduces the screening effort. Suitable embodiments of the folate receptor and DHFR as well as suitable selection conditions and concentrations are described above and it is referred to the above disclosure which also applies here. The described concentrations and concentration ranges for folate and antifolate described above can be combined with each other. In one embodiment, a folate concentration of about 0.1 nM - 150nM, 1nM - 125M, 5nM - 100nM or 7.5nM to 75nM is used in combination with an antifolate concentration of 0.1 nM - 150nM, 1nM to 125nM, 2.5nM to 100nM, 5nM to 75nM or 7.5nM to 50nM in the selective culture medium. As described, folic acid is preferably used as folate and MTX as antifolate.
[96] Using a respective combination of a folate receptor and DHFR as selectable marker and applying in stage (b) the selection conditions for both markers simultaneously by using an appropriate selective culture medium, results in a very stringent selection system allowing the efficient enrichment of high producing cells from the transfected host cell population. The host cell's viability is considerably increased under selective conditions, if both selectable markers are strongly expressed. Thereby, eukaryotic cells dependent on folate uptake such as mammalian cells can be selected which show an increased expression rate of the product of interest. This concept of using a folate receptor as selectable marker in combination with a further selectable marker involved in the folate metabolism such as preferably a DHFR is disclosed in WO 2010/097240, herein incorporated by reference. As is shown by the examples, the high stringency of the selection system according to this embodiment can be combined advantageously with the novel cells described herein, in particular when mammalian cells are used. This combination further lowers the number of low producers and a more homogenous population of high producing cells can be obtained after selection. This *inter alia* simplifies single cell cloning of stable producing cells. Furthermore, this combination allows to significantly reduce the MTX concentration necessary for DHFR selection.

[97] According to one embodiment, two or more selection steps are performed in stage (b), wherein the two or more selection steps are based on the same or a different selection principle. For example, if an additional selectable marker is used in addition to the folate receptor and/or DHFR, the selective conditions for said additional selectable marker can be applied prior to (e.g. in a pre-selection step) or simultaneously with applying the selective conditions for the folate receptor and/or DHFR. For example, in case the neomycin phosphotransferase gene (neo) is used as additional selectable marker, stage (b) may comprise first growing cells in a medium e.g. containing G418 in order to select cells that have incorporated the expression vector or the combination of at least two expression vectors and then performing a selection step using a selective culture medium comprising a limiting concentration of folate and an inhibitor of the second selectable marker, such as e.g. MTX when using DHFR as second selectable marker.

[98] According to one embodiment, a flow cytometry based selection is performed in stage (b). A selection step employing flow cytometry, in particular fluorescence activated cell sorting (FACS), has the advantage that large numbers of cells can be screened rapidly for the desired characteristic expression yield.

[99] According to one embodiment, said flow cytometry based selection is performed in addition to, preferably after, one or more selection steps against one or more selectable marker gene(s) were performed. Thereby, high expressing cell clones can be identified in the population of successfully transfected cells and separated out.

[100] According to one embodiment, high expressing cells are identified by detecting the expression of a co-expressed reporter polypeptide such as e.g. green fluorescence protein (GFP), CFP, YFP, luciferase or other common reporter polypeptide that can be detected by flow cytometry. According to this embodiment, stage (a) comprises providing a plurality of
host cells comprising at least one heterologous polynucleotide encoding a reporter and stage (b) comprises identifying host cells expressing the reporter based on at least one characteristic of said reporter polypeptide using flow cytometry. In such reporter based selection, the expression of the reporter gene correlates with the expression of the product of interest. The reporter polypeptide may be intracellular^\textsuperscript{1} located, thereby marking the expressing cell. According to one embodiment, the expression of the reporter polypeptide is tightly linked to the expression of the product of interest which is a polypeptide. E.g. the reporter polypeptide and the polypeptide of interest may be expressed as separate polypeptides but from the same expression cassette, however, separated by an IRES element (internal ribosomal entry site). Furthermore, the reporter polypeptide and the polypeptide of interest may be expressed as fusion protein. According to one embodiment, in the expression cassette for expressing the polypeptide of interest, the polynucleotide encoding the polypeptide of interest is separated by at least one stop codon from the polynucleotide encoding the reporter. A fusion protein comprising the reporter polypeptide is only expressed if translation reads over the stop codon. As stop codon readthrough occurs only to a certain extent, which can be influenced by the number and design of the stop codon and the culture conditions, a certain proportion of the polypeptide of interest is produced as fusion protein comprising the reporter polypeptide which can be detected by flow cytometry. Using a respective strategy allows to tightly link the expression of the reporter polypeptide to the expression of the polypeptide of interest. The principle of obtaining fusion proteins by stop codon read through will also be explained subsequently in conjunction with an embodiment wherein a fusion protein (not necessarily comprising a reporter) is displayed on the cell surface and e.g. is stained by using a detection compound. For expressing a secreted polypeptide of interest one can additionally include a polynucleotide encoding a membrane anchor either between the stop codon and the polynucleotide encoding the reporter or downstream of the polynucleotide encoding the reporter. The membrane anchor ensures that the reporter polypeptide remains associated with the expressing cell. Thereby, the reporter polypeptide comprised in the fusion protein provides the expressing cells with a trait that is selectable by flow cytometry. The polypeptide of interest is expressed into the culture medium. The higher e.g. the fluorescence of the reporter polypeptide, the more fusion protein is produced and accordingly, the higher is the expression rate of the polypeptide of interest. A respective method is e.g. disclosed in WO 03/014361 to which it is referred.

[101] According to one embodiment, stage (b) comprises:

(i) performing at least one selection under conditions selective for one or more selectable markers expressed by transfected host cells; and
(ii) performing a flow cytometry based selection.

[102] Optionally, one or more further selection steps can be performed prior to or after step (i) and/or step (ii).

[103] According to one embodiment, the flow cytometry based selection comprises selecting a plurality of host cells expressing the polypeptide of interest with a desired yield
based upon the presence or amount of the polypeptide of interest. Preferably, the polypeptide of interest is a secreted polypeptide. According to one embodiment, the polypeptide of interest is detected on the cell surface using a detection compound that binds to the polypeptide of interest. According to one embodiment, the secreted polypeptide of interest is detected while it passes the cell membrane and accordingly is transiently associated with the plasma membrane during polypeptide secretion. A respective flow cytometry based selection system is e.g. disclosed in WO03/099996 to which it is referred.

[104] According to one embodiment, a portion of the polypeptide of interest is expressed fused to a membrane anchor and thus is expressed as membrane-bound fusion polypeptide. Thereby, a portion of the polypeptide is displayed as fusion polypeptide on the cell surface and can be stained using a detection compound. No reporter polypeptide is required for this type of selection even though it may be additionally used. Due to the presence of the membrane anchor, the polypeptide of interest is tightly anchored to the expressing cell. As the amount of produced fusion polypeptide correlates with the overall expression rate of the expressing cell, host cells can be selected via flow cytometry based upon the amount of fusion polypeptide displayed via the membrane anchor on the cell surface. This allows a rapid selection of high producing host cells. To allow efficient selection using flow cytometry, preferably by using FACS, it is advantageous to use special expression cassette designs for expressing the polypeptide of interest. Thus, according to one embodiment, the polynucleotide encoding the polypeptide of interest is comprised in an expression cassette that is designed such that a portion of the expressed polypeptide of interest comprises a membrane anchor. The polypeptide of interest which is fused to a membrane anchor is also referred to a fusion polypeptide or fusion protein. Several options exist to achieve that result.

[105] According to one embodiment, the host cells provided in stage (a) comprise
(i) a heterologous expression cassette comprising
   (i) a heterologous expression cassette comprising
       aa) the polynucleotide encoding a polypeptide of interest,
       bb) at least one stop codon downstream of the polynucleotide encoding the polypeptide
           of interest, and
       cc) a further polynucleotide downstream of the stop codon encoding a membrane anchor
           and/or a signal for a membrane anchor;

and
(ii) at least one heterologous expression cassette comprising a polynucleotide encoding a
     selectable marker;

and selection in stage (b) comprises
(i) culturing the host cells under conditions selective for the at least one selectable marker
     and allowing expression of the polypeptide of interest wherein at least a portion of the
     polypeptide of interest is expressed as fusion polypeptide comprising a membrane anchor,
     wherein said fusion polypeptide is displayed on the surface of said host cell;
(ii) performing a flow cytometry based selection, comprising selecting a plurality of host cells
     expressing the polypeptide of interest with a desired yield based upon the presence or
     amount of the fusion polypeptide displayed on the cell surface using flow cytometry.
[106] Transcription of the polynucleotide encoding the polypeptide of interest comprised in the above described expression cassette results in a transcript comprising in consecutive order at least
aa) a polynucleotide, wherein translation of said polynucleotide results in the polypeptide of interest;
bb) at least one stop codon downstream of said polynucleotide;
cc) a polynucleotide downstream of said stop codon, encoding a membrane anchor and/or a signal for a membrane anchor.

[107] At least a portion of the transcript is translated into a fusion polypeptide comprising the polypeptide of interest and the membrane anchor by translational read-through of the at least one stop codon. This design of the expression cassette that is used in this embodiment has the effect that through translational read-through processes (the stop codon is "leaky") a defined portion of the polypeptide of interest is produced as a fusion polypeptide comprising a membrane anchor. Thus, at least a portion of the transcript is translated into a fusion polypeptide comprising the polypeptide of interest and the membrane anchor by translational read-through of the at least one stop codon. Translational read-through may occur naturally due to the choice of the stop codon/design of the translation termination signal or can be induced by adapting the culturing conditions, e.g. by using a termination suppression agent. This read-through level results in a certain proportion of fusion polypeptides. These fusion polypeptides comprise a membrane anchor, which tightly anchors the fusion polypeptides to the cell surface. As a result, the fusion polypeptide is being displayed on the cell surface and cells displaying high levels of membrane-anchored fusion polypeptide can be selected by flow cytometry, preferably by FACS. Thereby, host cells are selected that express the polypeptide of interest with high yield. Details and preferred embodiments of this stop codon readthrough based technology are described in WO2005/073375 and WO 2010/022961 and it is referred to this disclosure for details.

[108] According to one embodiment, the expression cassette additionally comprises iv) a polynucleotide encoding a reporter polypeptide, such as e.g. GFP. Said polynucleotide encoding the reporter polypeptide is located downstream of the stop codon. Upon stop codon read-through a fusion polypeptide is obtained which comprises the reporter, thereby allowing selection by flow cytometry based on the characteristics of the reporter polypeptide such as e.g. its fluorescence. Details of said embodiment are already described above and it is referred to the above disclosure. Preferably, the polynucleotide encoding the reporter polypeptide is located downstream of the polynucleotide encoding a membrane anchor.

[109] According to an alternative embodiment, a portion of the polypeptide of interest is expressed as cell surface displayed fusion polypeptide using the technology described in WO2007/131774. Here, through transcription and transcript processing at least two different mature mRNAs (mRNA-polypeptide of interest) and (mRNA-polypeptide of interest-anchor) are obtained from the expression cassette encoding the polypeptide of interest. Translation of the mRNA-polypeptide of interest results in the polypeptide of interest. Translation of the
mRNA-polypeptide of interest-anchor results in a fusion polypeptide comprising the polypeptide of interest and a membrane anchor. As a result, this fusion polypeptide is again displayed on the cell surface and cells displaying high levels of membrane-anchored fusion polypeptide can be selected by flow cytometry, preferably FACS. Thereby, again host cells can be selected that have a high expression rate. According to one embodiment, the expression cassette additionally comprises a polynucleotide encoding a reporter polypeptide, such as e.g. GFP. Said polynucleotide encoding the reporter polypeptide is located downstream of the intron. Thereby, a fusion polypeptide is obtained which comprises the reporter polypeptide, thereby allowing selection by flow cytometry based on the characteristics of the reporter polypeptide such as e.g. its fluorescence. Preferably, the polynucleotide encoding the reporter polypeptide is located downstream of the polynucleotide encoding a membrane anchor. Thereby, the reporter is located inside the host cell.

[110] Both exemplary embodiments described above result in that a portion of the polypeptide of interest is expressed as fusion polypeptide that is displayed at the surface of the host cells, and cells displaying high levels of membrane-anchored fusion polypeptides (indicating a high level of secreted polypeptide) can be selected e.g. by flow cytometry, in particular by fluorescence activated cell sorting (FACS). Here, different embodiments are available. E.g. if a reporter polypeptide is comprised in the fusion polypeptide, high expressing host cells can be selected based upon a characteristic of the reporter polypeptide, e.g. its fluorescence. Furthermore, appropriately labelled detection compounds can be used as will be described briefly in the following.

[111] According to one embodiment, stage b) comprises selecting a plurality of eukaryotic host cells expressing the polypeptide of interest with a desired yield based upon the presence or amount of the fusion polypeptide displayed on the cell surface using flow cytometry by contacting the host cells with a detection compound binding the fusion polypeptide displayed on the cell surface and selecting a plurality of host cells expressing the polypeptide of interest with a desired yield based upon the presence or amount of the bound detection compound using flow cytometry. Thus, cells may be contacted with an appropriately labelled detection compound that binds the fusion protein, e.g. the portion corresponding to the polypeptide of interest. The detection compound used for binding to the fusion polypeptide may have at least one of the following characteristics: said compound may be labelled, in particular fluorescently labelled, it may be an antigen, it may be an immunoglobulin molecule or a binding fragment thereof or it may be protein-A, -G, or -L. The detection compound used for binding the fusion polypeptide at the cell surface can for example be an immunoglobulin molecule or a fragment thereof such as an antibody or antibody fragment, recognising the fusion polypeptide. Basically all accessible portions of the fusion polypeptide can be detected, thereunder also the portion corresponding to the polypeptide of interest which is secreted in parallel to the fusion polypeptide in soluble form. In order to allow detection and selection, said detection compound used for binding the fusion polypeptide may be labelled. The labelled detection compound that binds the fusion polypeptide displayed on the cell surface thereby labels, respectively stains the cell surface.
The higher the amount of fusion polypeptide that is expressed by the host cell, the more labelled detection compound is bound and the more intensive is the staining. This has the advantage that the flow cytometry based selection of the host cells can be easily performed as not only the presence but also the amount of the bound detection compound can be determined. To select high producing host cells, those host cells can be selected from the population of host cell which are most effectively, respectively intensively labelled by the detection compound. The label is suitable for flow cytometry based selection, in particular FACS selection. A fluorescent label is preferred as this allows easy detection by flow cytometry.

According to one embodiment, the expression cassette is constructed such that approximately ≤ 50%, ≤ 25%, ≤ 15%, ≤ 10%, ≤ 5%, ≤ 2.5%, ≤ 1.5%, ≤ 1% or less than ≤ 0.5% fusion polypeptide is obtained. The remaining portion is produced as the secreted polypeptide form not comprising the membrane anchor.

The membrane anchor may be of any kind as long as it enables anchorage of the polypeptide of interest to the cell membrane and thus allows the display of the fusion polypeptide on the cell surface. Suitable embodiments include but are not limited to a GPI anchor or a transmembrane anchor. A transmembrane anchor is preferred to ensure tight binding of the fusion polypeptide to the cell surface and to avoid shedding of the fusion protein. Particularly preferred, in particular when expressing antibodies as polypeptide of interest, is the use of an immunoglobulin transmembrane anchor. Other membrane anchors and preferred embodiments of an immunoglobulin transmembrane anchor are described in WO2007/131774, WO2005/073375 and WO 2010/022961.

According to one embodiment, the host cells express an immunoglobulin molecule such as an antibody as polypeptide of interest. The polynucleotide encoding the heavy chain of an immunoglobulin molecule and the polynucleotide encoding the light chain of an immunoglobulin molecule may be comprised in the same expression cassette or preferably, are comprised in separate expression cassettes as is described above in conjunction with the first aspect. When using an expression cassette design as described above, wherein a portion of the polypeptide of interest is produced as membrane-anchored fusion polypeptide by translational readthrough or alternative splicing, such design is used for expressing the antibody heavy chain.

According to one embodiment, two or more flow cytometry based selection cycles may be performed in stage (b) to select and enrich high expressing host cells.

In one embodiment, host cells expressing a high amount of polypeptide of interest which accordingly depict a high signal are sorted using fluorescence-activated cell sorting (FACS). FACS sorting is advantageous, since it allows rapid screening of large numbers of host cells to identify and enrich those cells which express the polypeptide of interest with a high yield. This embodiment is particularly suitable if the cells are selected based upon the expression of a fusion protein as described above. Those cells, showing the highest
fluorescence rate can be identified and isolated by FACS. A positive and statistically significant correlation between fluorescence, as determined by FACS, and the amount of produced polypeptide is found. Therefore, FACS sorting can be used not only for a qualitative analysis to identify cells expressing a polypeptide of interest in general, but can actually be used quantitatively to identify those host cells that express high levels of the polypeptide of interest. Thereby the best producing cells can be selected/enriched in stage (b).

[17] According to one embodiment, cells which express the polypeptide of interest with the desired yield, e.g. above a certain threshold and/or the top 15%, top 10%, top 5% or the top 2% of the host cells, are selected as pool. E.g. several high expressing cells, e.g. at least 10, at least 20, at least 30, at least 50, at least 100, at least 200, at least 300, at least 500, at least 1000 or at least 5000 high expressing cells can be selected in stage (b) and sorted into a cell pool. This cell pool comprising a plurality of different high expressing cells is also referred to as high expressing cell pool. Said cell pool comprising different individual high expressing cells can then be used in order to obtain individual cell clones for large scale production of the polypeptide of interest.

[18] According to one embodiment, the eukaryotic cells provided in stage (a) are mammalian cells, preferably rodent cells, more preferably hamster cells such as CHO cells. Suitable and preferred embodiments are described above in conjunction with the first aspect and it is referred to the above disclosure. As described above, according to one embodiment, said CHO cells have lost a portion of the telomeric region of chromosome 8 wherein said lost portion comprises gene FAM60A. Alternative embodiments are also described above. These cells have particularly favorable characteristics with respect to yield and stability. According to one embodiment, the method according to the second aspect comprises a step of analyzing whether expression of gene FAM60A is reduced or eliminated. Such analysis may be performed after selection, in particular DHFR selection. Details of such analytical method are already described above in conjunction with the first aspect of the present disclosure and are also described below in conjunction with the method according to the fifth aspect and it is referred to the respective disclosure.

[19] Further preferred embodiments in particular with respect to the eukaryotic cells according to the first aspect which preferably are mammalian cells, the expression vector, or combination of expression vectors are described in detail above. It is referred to the above disclosure.

[20] Cells obtained as a result of the selection method according to the second aspect can be isolated and cultured as individual cells. It is, however, also possible to use an enriched population of different host cells, i.e. a cell pool, in the downstream process. The obtained host cells can also be subjected to additional qualitative or quantitative analysis, or can be used e.g. in the development of a clonal cell line for protein production. A clonal cell line may be established from a selected host cell which stably expresses the product of interest with high yield.
Thus, according to one embodiment, selected cells are cultivated to provide cell clones, in particular in the form of clonal cell cultures. A clonal cell culture is a cell culture derived from one single ancestral cell. In a clonal cell culture, all cells are clones of each other. Preferably, all the cells in a cell culture contain the same or substantially the same genetic information. In certain embodiments, the amount or concentration of the polypeptide of interest in the cell culture is determined to determine the productivity. E.g. the titer can be measured by analysing the culture supernatant. According to one embodiment, after determining the productivity performance of each individual clone, a titer ranking is made to select the best producing clones as production clones. Furthermore, a stability study can be performed with the obtained cell clones. However, as is shown in the examples, using the novel cells described herein as host cells provide after selection recombinant cell clones showing a significantly improved stability. Thus, losses in expression stability are rarer with respective host cells and furthermore, if occurring, often only results in a less dramatic decrease in the productivity compared to when using cells wherein functional expression of gene FAM60A is not reduced or eliminated. Therefore, stability analyses may be shortened or even skipped which is an important advantage as it shortens the time that is required for obtaining stably expressing cell clones that can be used for large scale production of the product of interest.

C. Method for producing a product of interest

According to a third aspect, a method is provided for producing a recombinant product of interest, comprising using a eukaryotic cell according to the first aspect as host cell for recombinant expression of the product of interest.

As described above, the novel eukaryotic cells provided herein are particularly suitable as production host cells for recombinantly producing a product of interest such as a polypeptide of interest. Suitable and preferred examples of said eukaryotic cell, wherein the effect of protein FAM60A in said cell is impaired, preferably by reducing or eliminating the functional expression of gene FAM60A, as well as suitable and preferred examples of the product of interest are described in detail above and it is referred to the above disclosure which also applies here. As described above, the eukaryotic cell preferably is a vertebrate cell, more preferably a mammalian cell. Particularly advantageous is the embodiment wherein in addition to FAM60A, the effect of the expression product of gene C12orf35 is impaired, for example by reducing or eliminating functional expression of gene C12orf35 as it was found that thereby, the production yield can be significantly improved. By impairing the effect of protein FAM60A as well as of protein C12orf35, mammalian host cells can be provided which show improved characteristics with respect to both features, long-term stability as well as production yield, and hence key features important for the large-scale production of a product of interest, in particular a polypeptide of interest.

The eukaryotic host cell which preferably is a mammalian cell comprises stably integrated into the genome a polynucleotide encoding a product of interest. Introduction of
said polynucleotide can be achieved by stable transfection as described above. Selection of successfully transfected cells may occur using the method according to the second aspect.

[125] According to one embodiment, the method comprises
(a) culturing host cells according to the first aspect under conditions that allow for the expression of the product of interest;
(b) isolating the product of interest from said cell culture medium and/or from said host cells; and
(c) optionally processing the isolated product of interest.

[126] According to one embodiment, said host cells are cultured under serum-free conditions. The expressed product of interest may be obtained by disrupting the host cells. Preferably, the product of interest is a polypeptide. The polypeptide is preferably expressed, e.g. secreted, into the culture medium and can be obtained therefrom. For this purpose, an appropriate leader peptide is provided in the polypeptide of interest. Leader sequences and expression cassette designs to achieve secretion are well known in the prior art. Also a combination of the respective methods is possible. Thereby, polypeptides such as proteins can be produced and obtained/isolated efficiently with high yield.

[127] The product of interest which preferably is a polypeptide that is produced may be recovered, further purified, isolated, processed and/or modified by methods known in the art. For example, the polypeptide may be recovered from the nutrient medium by conventional procedures including, but not limited to, centrifugation, filtration, ultra-filtration, extraction or precipitation. Further processing steps such as purification steps may be performed by a variety of procedures known in the art including, but not limited to, chromatography (e.g. ion exchange, affinity, hydrophobic, chromatofocusing, and size exclusion), electrophoretic procedures (e.g., preparative isoelectric focusing), differential solubility (e.g. ammonium sulfate precipitation) or extraction. Furthermore, the isolated and purified product of interest may be further processed, such as e.g. formulated, into a composition, e.g. a pharmaceutical composition.

D. Method for producing a eukaryotic cell
[128] According to a fourth aspect, a method is provided for producing a eukaryotic cell suitable for recombinant production of a product of interest, comprising impairing the effect of protein FAM60A in an eukaryotic cell by altering the genome of said cell and stably transfecting into said cell at least one expression vector comprising a polynucleotide encoding a product of interest. Suitable and preferred embodiments to achieve that result are described above in conjunction with the eukaryotic cells according to the first aspect and it is referred to the above disclosure which also applies here. Non-limiting embodiments are again briefly described in the following.

[129] According to one embodiment, the method comprises altering the genome of the eukaryotic cell to reduce or eliminate the functional expression of gene FAM60A in said cell, thereby impairing the effect of the protein FAM60A in said cell. Suitable ways are described
above in conjunction with the eukaryotic cells according to the first aspect of the present disclosure and it is referred thereto.

[130] For example, a gene knock-out may be introduced into the FAM60A gene. According to one embodiment, such gene knock-out is introduced in all copies of the FAM60A gene if more than one copy is present. According to one embodiment, the FAM60A gene is deleted. All copies of the FAM60A gene may be deleted in the genome if more than one copy is present. According to one embodiment, the method comprises deleting a portion of a chromosome, wherein the deleted portion comprises gene FAM60A. The deleted portion may correspond to a telomeric region. However, due to genetic rearrangements, said region may also be located in a different region of the chromosome. Such deletion can be induced e.g. by using an agent that induces chromosome breakages. Here, the cells can be repeatedly treated with such agent in order to obtain cells in which the functional expression of gene FAM60A is reduced or eliminated, e.g. because all copies of said gene are deleted because of induced chromosome breaks.

[131] According to one embodiment, the eukaryotic cell is a metazoan cell, preferably a vertebrate cell, more preferred a mammalian cell. According to one embodiment, the mammalian cell is a rodent cell. Preferably, the rodent cell is a Chinese hamster cell such as a CHO cell, e.g. a CHO cell derived from CHO-K1. According to one embodiment, the method comprises deleting at least a portion of the telomeric region of chromosome 8 in a hamster cell, wherein said deleted portion comprises the FAM60A gene. As is shown by the examples, CHO cells comprising a respective deletion in the telomeric region of chromosome 8, here the q arm, have particularly favorable expression characteristics and thus are particularly suitable as host cells for recombinant production. According to one embodiment, the deleted telomeric region comprises the FAM60A gene and one or more or all genes selected from the group consisting of Caprin2, lpo8 and RPS4Y2. According to one embodiment, the deleted region additionally comprises at least a portion of or all of the Tmtc1 gene. According to one embodiment, the method comprises deleting at least a respective portion of the telomeric region in both chromosomes of chromosome pair 8. Non-limiting alternative names of the aforementioned individual genes and/or the encoded proteins are also indicated in Table 1 above and the respective genes are, as well as homologs and orthologs, encompassed by the scope of the terms used above for the individual genes.

[132] As described above, the telomeric region of chromosome 6 in mouse corresponds to the telomeric region of chromosome 8 in Chinese hamster. Thus, the above disclosure with respect to the telomeric region of chromosome 8 of hamster correspondingly applies to the telomeric region of chromosome 6 in mouse.

[133] According to one embodiment, the method comprises causing a chromosome break in the telomeric region of chromosome 8 of the hamster genome (or chromosome 6 of the mouse genome), wherein the breakpoint on chromosome 8 (or chromosome 6 of the mouse genome) is located centromeric of the Fam60a gene, centromeric of the Caprin2 gene,
centromeric of the lpo8 gene or centromeric of the RPS4Y2gene. Thereby, all genes that are present telomeric thereof, i.e. which lie further to the direction of the telomeric end, are deleted. Non-limiting alternative names of the aforementioned individual genes as well as homologs and orthologs are also indicated in Table 1 above and the respective genes and encoded proteins are encompassed by the scope of the terms used above for the individual genes and/or proteins. According to one embodiment, the obtained cells comprising a chromosome break in the telomeric region have one or more of the following characteristics:

a) the breakpoint is located centromeric of the lpo8 gene;
b) the breakpoint is located within the Tmtc1 gene.

As discussed above, all genes that are located telomeric of the breakpoint, i.e. further into the direction of the telomeric end, are comprised in the deleted region. This includes gene FAM60A. Methods for identifying respective mammalian cells having such breakpoint are described above are also described in the following in conjunction with the fifth aspect of the present disclosure. According to one embodiment, the Ergic2 gene is not deleted.

According to one embodiment, a CHO cell, preferably derived from the cell line K1, is used in order to produce an altered mammalian cell line wherein the effect of protein FAM60A is impaired, preferably by reducing or eliminating the functional expression of gene FAM60A. According to one embodiment, a telomeric region on the q arm of chromosome 8 comprising gene FAM60A is deleted. Details how to achieve that result are known to the skilled person and suitable embodiments are also described herein and it is referred to said disclosure. Particularly advantageous is the embodiment wherein additionally, the effect of the expression product of gene C12orf35 is impaired in said cell, for example by reducing or eliminating functional expression of gene C12orf35 as it was found that thereby, the production yield can be significantly improved. Therefore, according to one embodiment, the method according to the fourth aspect additionally comprises impairing the effect of the expression product of gene C12orf35 in said cell. Ways to achieve these results, preferably by reducing or eliminating functional expression of gene C12orf35 are described above and it is referred to the respective disclosure. Mammalian host cells that are altered so that the effect of both FAM60A and C12orf35 is impaired in said cells have particularly favorable expression characteristics. Thereby, mammalian host cells are provided which show improved characteristics with respect to both features, long-term stability as well as production yield, and hence key features important for the large-scale production of a product of interest, in particular a polypeptide of interest.

According to one embodiment, the method according to the fourth aspect comprises introducing into the eukaryotic cell in which the functional expression of gene FAM60A is reduced or eliminated at least one polynucleotide encoding a product of interest and preferably at least one polynucleotide encoding a selectable marker. According to one embodiment, the polynucleotide encoding a product of interest and the polynucleotide encoding a selectable marker are located on the same or on separate expression vectors. Suitable and preferred embodiments are described above and it is referred to the respective
disclosure which also applies here. The expression vector(s) integrate into the genome of the
host cells so that stably transfected cells are provided. Host cells that were successfully
transfected and express the product of interest can be selected using the method according
to the second aspect. It is referred to the above disclosure.

E. Method for analysing eukaryotic cells

According to a fifth aspect, a method is provided for analyzing eukaryotic cells for
their suitability as host cells for recombinant expression of a product of interest, comprising
analyzing directly or indirectly whether the function of the expression product of gene
FAM60A is impaired in said cells. As described above, the eukaryotic cell is preferably a
mammalian cell.

This analytical method can be advantageously used e.g. in combination with the
method according to the fourth aspect of the present disclosure in order to identify whether a
eukaryotic cell was produced, wherein the effect of protein FAM60A was impaired.
Furthermore, said method can be used in order to discriminate between stable or unstable
clones during the selection/screening process. Furthermore, in embodiments, this method
can additionally be used to discriminate between high and low producing clones. By using
this analytical method, clones can be identified early in the selection process, which have
favorable expression characteristics. This increases the probability to select stable and high
producing clones resulting in a higher proportion of high and stable producing clones.
Therefore, this analytical method has important applications and provides a further
improvement of recombinant expression technologies as it shortens the time required to
identify a suitable production clone.

According to a preferred embodiment, the method comprises analyzing whether the
functional expression of gene FAM60A is reduced or eliminated in said cells. The analysis
whether functional expression of gene FAM60A is reduced or eliminated can be performed
directly or indirectly. Non-limiting embodiments are described in the following. Which
analytical method is suitable also depends on how the cells are altered to achieve the
reduction or elimination of functional expression of endogenous gene FAM60A.

For example, when introducing a knock-out into the FAM60A gene in order to reduce
or eliminate expression of gene FAM60A, one can amplify the corresponding DNA section
and sequence the amplified DNA in order to confirm that the gene knock-out was introduced
in gene FAM60A. If functional expression of gene FAM60A is reduced or eliminated by
completely or partly deleting said gene, one can detect the deletion on the DNA level, e.g.
using suitable amplification based detection methods to detect the deletion (such methods
are known to the skilled person).

According to one embodiment, the expression profile of the eukaryotic cells is
analyzed to determine whether functional expression of gene FAM60A is reduced or
eliminated. For example, the analysis may comprise performing a qualitative or quantitative
RT (reverse transcription) PCR in order to detect the presence, absence, amount or length of
FAM60A mRNA. This would be an example of a direct analysis, as the analysis directly involves the transcript of gene FAM60A. Indirect analyses, wherein the expression status of gene FAM60A is indirectly determined by analyzing the expression profile of genes different from gene FAM60A and wherein accordingly, the analysis does not directly involve the analysis of gene FAM60A or its transcript are also suitable and thus encompassed by the term “analyzing whether the functional expression of gene FAM60A is reduced or eliminated". Such indirect analysis are e.g. suitable if a chromosomal portion comprising gene FAM60A (and other genes) is deleted by chromosome breakage and will be described subsequently. For a quantitative analysis, a comparison with a reference (e.g. unaltered corresponding cell) can be performed.

[142] According to one embodiment, it is additionally analyzed directly or indirectly whether the effect of the expression product of endogenous gene C12orf35 is impaired in said cells. This can be analyzed by determining whether the functional expression of gene C12orf35 is reduced or eliminated in said cells. This analysis can be performed *mutatis mutandis* as was described for gene FAM60A. It is referred to the above discussion.

[143] According to one embodiment, prior to analysis, cells are treated with an agent that induces chromosome breaks in order to delete a portion of the chromosome which comprises the FAM60A gene. As described above, all copies of gene FAM60A could be deleted thereby. The analysis may then comprise analyzing whether treatment with said agent resulted in a deletion of a portion of a chromosome which includes gene FAM60A. The cells are treated with an agent that induces chromosome breakage in an appropriate concentration so that chromosome breakage occurs. Here, also several treatment rounds can be performed. Chromosome breakage may be induced during the selection process if selection involves the use of an agent that induces chromosome breakages in a sufficiently high concentration. A non-limiting example of a suitable agent is e.g. MTX. In this case the cells may comprise a heterologous polynucleotide encoding DHFR as selectable marker in order to be able to survive MTX treatment. However, as discussed above also other agents can be used such as e.g. hygromycin and this was confirmed by examples.

[144] After treating the cells to induce chromosome breaks, the obtained cells can be analysed using the method according to the fifth aspect to determine whether treatment with said agent resulted in a deletion of a portion of a chromosome which includes gene FAM60A. Different embodiments are suitable for that purpose. According to one embodiment, the expression profile of the treated cells is analyzed. For example, it can be analysed whether gene FAM60A or one or more genes located centromeric of gene FAM60A (i.e. further away from the telomeric end into chromosome) are expressed. For example, in case of mouse or Chinese hamster cells, it can be analysed whether gene FAM60A is expressed and accordingly, if its mRNA can be detected (example of a direct analysis) and alternatively or in addition thereto, it can be analyzed whether one or more genes selected from the group consisting of Caprin2, Ipol, Tmtc1 or genes that are located telomeric of the aforementioned genes are expressed by the cell (example of indirect analysis). Non-limiting alternative names of the aforementioned individual genes are also indicated in Table 1 above and the
respective genes are encompassed by the scope of the terms used above for the individual genes and encoded products. If the induced breakpoint is located centromeric of the respective gene(s), said genes are deleted which eliminates or reduces (if other copies of the gene existed elsewhere that are expressed) their expression. As is evident from Fig. 1, gene FAM60A is located telomeric of the aforementioned genes. Thus, if the aforementioned genes are deleted, the deleted region also includes gene FAM60A. Thus, the above genes can be validly used as markers in order to basically indirectly determine whether the induced chromosome breakage resulted in a deletion of gene FAM60A and thus resulted in a reduction or elimination of expression of gene FAM60A. Thus, the analysis of whether expression of gene FAM60A was reduced or eliminated does not necessarily have to be based on a direct analysis of e.g. the FAM60A mRNA and such indirect methods are also encompassed by the method of the fifth aspect. Furthermore, it was found that even though located telomeric of gene FAM60A (i.e. further down into the direction of the telomeric end), also genes such as C12orf35 and Bicdl can be used as marker to determine whether a chromosome breakage was induced which resulted in a deletion of gene FAM60A. It was found in conjunction with the analysis of Chinese hamster cells such as CHO cells that if gene Bicdl or C12orf35 is deleted because of a chromosome breakage, the deleted telomeric region usually also includes FAM60A. It was confirmed by analyzing the expression characteristics of several hundred clones that the aforementioned genes can be validly used as markers in order to discriminate cell clones with high and stable expression characteristics from cell clones having low and unstable expression characteristics. The relative expression of the aforementioned genes in CHO cells is shown in Fig. 2.

[145] According to one embodiment, the method according to the fifth aspect is for analyzing hamster cells, preferably CHO cells, and the method comprises analyzing whether expression of gene FAM60A is reduced in said cells by analyzing whether the expression of one or more genes located in the telomeric region of chromosome 8 and being selected from the group consisting of the Trntd gene and genes located telomeric of the Trntd gene is reduced or eliminated in said cells. As described above, respective cells which after treatment with the agent that introduces chromosome breakage do no longer express the Tn1c1 gene and/or genes located telomeric of the Trntd gene usually comprise a deletion in the telomeric region of chromosome 8 which comprises said genes and in particular comprises the FAM60A gene. Genetic material telomeric of the breakpoint is lost.

[146] According to one embodiment, the selected host cells having the above described characteristics are transfected with an expression vector comprising at least one polynucleotide encoding a product of interest and preferably comprising at least one selectable marker. Suitable embodiments are described above in conjunction with the other aspects and it is referred to the above disclosure.

[147] According to one embodiment, prior to performing the analysis using the method according to the fifth aspect, the eukaryotic cells are transfected with at least one heterologous polynucleotide encoding a product of interest and at least one heterologous polynucleotide encoding a selectable marker, and prior to analysis at least one selection step
is performed to identify successfully transfected host cells. Suitable embodiments are described in detail above. According to one embodiment, selection involves the use of a selection agent that induces chromosome breakages. In this embodiment, the selectable marker may be DHFR and the selection agent may be MTX. Alternatively, the selection agent may be hygromycin and the selectable marker may be a gene that confers resistance against hygromycin such as e.g. the hph gene. In such applications, wherein the method is performed during or after the selection process, the method according to the fifth aspect can be used as analytical tool in order to identify within the selected cell population such cells, wherein the functional expression of gene FAM60A is reduced or eliminated. As described, such reduction or elimination may be caused or supported by the selection conditions, if e.g. a chromosome breakage is induced thereby which results in a deletion of gene FAM60A and the method according to the fifth aspect allows to identify such cells e.g. based on their expression profile. This allows the identification of such cells or cell clones which because of their expression profile, in particular due to the reduced or eliminated functional expression of gene FAM60A, are particularly suitable for establishing a recombinant production cell line as it can be expected that the high expression of the product of interest remains stable. This particularly, because if gene FAM60A is lost, gene C12orf35 is likewise lost and this significantly increases the expression yield as is described herein. As described, in case of hamster cells such as CHO cells wherein gene FAM60A is located in the telomeric region of chromosome 8 it is preferred that the cells have lost the telomeric region of chromosome 8, preferably the q arm, thereby reducing or eliminating expression of gene FAM60A. The analytical method can be performed after generating cell clones from cells comprised in the population of high expressing cells. According to one embodiment, a plurality of cell clones is analysed for discriminating between stable and unstable and/or between high and low producing cell clones.

[148] According to one embodiment, the method according to the fifth aspect comprises selecting at least one cell wherein the function of the expression product of gene FAM60A is impaired, preferably by reduction or elimination of functional expression of gene FAM60A, for recombinant expression of a product of interest, preferably a polypeptide of interest. Cells having the respective characteristics are particularly suitable for recombinant expression as is shown by the examples. Further embodiments of respective cells are also described in detail above. As described, preferably vertebrate cells such as most preferably mammalian cells are used as eukaryotic host cells.

F. Use of altered eukaryotic cells for recombinant production of a product of interest

[149] According to a sixth aspect, the present disclosure pertains to the use of an isolated eukaryotic cell for recombinantly expressing a product of interest, wherein the genome of the eukaryotic cell is altered so that the effect of protein FAM60A is impaired in said cell. Details with respect to the respectively altered eukaryotic host cells and embodiments suitable for achieving impairment of the effect of protein FAM60A in said cells, preferably by reducing or eliminating functional expression of gene FAM60A were described in detail above and it is referred to the above disclosure which also applies here. Non-limiting embodiments are described briefly below.
The eukaryotic cell may be selected from a metazoan, a vertebrate or mammalian cell. Preferably, the eukaryotic cell is a mammalian cell such as a rodent cell. Preferred are CHO cells. The genome of the eukaryotic cell may be altered as described in detail above. According to one embodiment, additionally the effect of the expression product of gene C12orf35 is impaired in said cell, preferably by reducing or eliminating functional expression of endogenous gene C12orf35. Details with respect to this embodiment and the associated advantages were described in detail above and it is referred to the above disclosure.

According to one embodiment, the product of interest is a polypeptide. Preferably, the polypeptide of interest is upon expression in the eukaryotic cell secreted into the cell culture medium. Details with respect to the polypeptide of interest were described above and it is referred to the respective disclosure. To allow expression of the product of interest, the eukaryotic host cell may be stably transfected with an expression vector comprising a polynucleotide encoding the polypeptide of interest. Details are described above and it is referred to the above disclosure. Preferably, a eukaryotic host cell according to the first aspect is used as eukaryotic host cells. Said cells are described in detail above and it is referred to the above disclosure.

Numeric ranges described herein are inclusive of the numbers defining the range. The headings provided herein are not limitations of the various aspects or embodiments of this disclosure which can be read by reference to the specification as a whole. According to one embodiment, subject-matter described herein as comprising certain elements also refers to subject-matter consisting of the respective elements. In particular, the polynucleotides described herein as comprising certain sequences may also consist of the respective sequences. It is preferred to select and combine preferred embodiments described herein and the specific subject-matter arising from a respective combination of preferred embodiments also belongs to the present disclosure.

The present application claims priority of prior US provisional application no. US 61/919340 filed on 20 December 2013, the entire disclosure of which is incorporated herein by reference.

EXEMPLARY EXAMPLES

The following examples serve to illustrate the present invention without in any way limiting the scope thereof. In particular, the examples relate to preferred embodiments of the present invention.

**Example 1: Knock-out of FAM60A in CHO cells using TALEN technology**

Two cell clones on the basis of CHO cells derived from the cell line CHO-K1 were made that comprise a knock-out mutation in the FAM60A gene. For creating the FAM60A mutant cells, TALEN (Transcription Activator-Like Effector Nuclease) technology was used.
For the knock-out of FAM60A, a coding region (presumed exon 1) of the gene FAM60A was targeted. The CHO-K1 cells used as parental cell only contain one copy of FAM60A. Thus, a single knock-out per cell is sufficient to impair the effect of FAM60A in said cell.

1. Design/production and use of TALENs which are specific for FAM60A

The following genomic DNA exon sequence of gene FAM60A of the CHO parental cell line was targeted:

```
atgtttgttttcacaagccaaagtgattagaggggtgtatctgcagacgcaagtcctccagctctcggttc
acggacagtaacgttgtaaaagacttccagagctgttttgg  (SEQ ID NO: 23)
```

The nucleotides of the TALEN binding sites are marked in bold. Two TAL Fok I targeting the coding sequence of FAM60 shown above (SEQ ID NO: 23) were designed. TALEN TAL-L targets and binds the marked 25 nucleotides on the 5' (forward) DNA strand and TALEN TAL-R targets and binds to the marked 25 nucleotides on the 3' (reverse) DNA strand of gene FAM60A as is shown above for SEQ ID NO: 23 (see also Table 2, which additionally shows primer sequences that were subsequently used for identifying knock-outs). The two binding sites are separated by the sixteen nucleotides of the cutting site. Plasmids coding for the two TALENS TAL-L and TAL-R were obtained.

**Table 2**: TALEN target sequences for FAM60A gene knockout and primer sequences

<table>
<thead>
<tr>
<th>TAL-L</th>
<th>TGTACCGAAGTATAGAGGGCTGCTG (SEQ ID NO:24)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TAL-R</td>
<td>TGTCCGTGAACCGAGGCTGGAGGA (SEQ ID NO: 25)</td>
</tr>
<tr>
<td>Primer 1:</td>
<td>GTCCAGCCTCATGAGGAT (SEQ ID NO: 26)</td>
</tr>
<tr>
<td>Primer 2:</td>
<td>CCTCCTAGCTGCTGATTTT (SEQ ID NO:27)</td>
</tr>
<tr>
<td>Primer 3:</td>
<td>GAGGACTTGCTGCTGAG (SEQ ID NO: 28)</td>
</tr>
<tr>
<td>Primer 4:</td>
<td>TACCACGCACAGCAGCCCGAT (SEQ ID NO: 29)</td>
</tr>
</tbody>
</table>

A portion of the genomic DNA sequence of gene FAM60A which encompasses the targeted coding sequence shown in SEQ ID NO: 23 and which also extends over the primer binding sites for primers 1, 2 and 4 which are located in intron sections is shown as SEQ ID NO: 30.

2. Transfection of the TALEN plasmid

Transfection was carried out using a standard transfection protocol involving electroporation using the parental CHO cells in exponential growth phase with viability over 95% and 5 µg of the each of the TALEN plasmids.
3. Cel-I-Assay and cell sorting

The Cel-I-assay was performed according to the manual of SAFC Biosciences. The Cel-I-assay is a standard assay in order to determine the cutting efficiency. In brief, after several days of cultivation, genomic DNA was isolated from the cells and a PCR was performed using primers 1 and 2 (see Table 2). The amplification product was denatured and allowed to renature. Then, nuclease S and nuclease S enhancer were added and incubated. The digested product was analyzed. If TALEN activity occurred, two smaller bands are present indicating TALEN activity within that region of the genome and therefore, supporting that cells wherein the FAM60A gene is knocked-out are present in the analyzed cell pool. From the positive cell pools, single cells were sorted in 96 well plates by limiting dilution.

4. Screening strategy

Genomic DNA (gDNA) was extracted from each clone in 96 well plates. The gDNA was analyzed by standard procedure to identify knock-out clones by PCR analysis. For this purpose, primers 3 and 4 (see Table 2) were used. In case of a mutation in the cutting region, primer 3 will not bind so that no PCR product is generated. The PCR products resulting from PCR with primers 1 and 2 of gDNA (see above) of positive clones were sequenced in order to analyze the introduced mutation.

Two cell clones with a knock-out mutation were obtained: FAM60A_ko_s16 (s16), with a deletion of 14 nucleotides and FAM60A_ko_s23 (s23) with a deletion of 5 nucleotides. The mutated sequences of the cell clones are shown in Table 3. Each of the deletions results in a frame-shift. Due to the frame-shift within the targeted sequence of FAM60A, stop codons are provided within the reading frame (highlighted in Table 3 by nucleotides in italic letters and underlining). Thus, it is expected that an abnormally short and less-or non-functional FAM60A expression product is expressed by the obtained FAM60A knock-out clones.

**Table 3**: DNA sequence of presumed exon 1 of FAM60A in CHO wildtype (WT - derived from CHO-K1) and two knock-out cell clones (s16 and s23) derived from said WT. Nucleotides of TALEN binding sites are highlighted in bold and nucleotides of premature stop codons in italic letters and underlining.

<table>
<thead>
<tr>
<th></th>
<th>DNA sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>atgttttgtttcccagcaagccaaagatgaccaagtatatagggcgtctgtagtgcagagccaatcctcaca</td>
</tr>
<tr>
<td></td>
<td>gctgcgtgttcacggacaagtataacggtatgaaagaagctccagagcgttttgg</td>
</tr>
<tr>
<td></td>
<td>(SEQ ID NO: 23)</td>
</tr>
<tr>
<td>s16(del14)</td>
<td>atgttttgtttcccagcaagccaaagatgaccaagtatatagggcgtctgtagtgcagagccaatcctcaca</td>
</tr>
<tr>
<td></td>
<td>gctgcgtgttcacggacaagtataacggtatgaaagaagctccagagcgttttgg</td>
</tr>
<tr>
<td></td>
<td>(SEQ ID NO: 31)</td>
</tr>
<tr>
<td>s23: (del5)</td>
<td>atgttttgtttcccagcaagccaaagatgaccaagtatatagggcgtctgtagtgcagagccaatcctcaca</td>
</tr>
<tr>
<td></td>
<td>gctgcgtgttcacggacaagtataacggtatgaaagaagctccagagcgttttgg</td>
</tr>
<tr>
<td></td>
<td>(SEQ ID NO: 32)</td>
</tr>
</tbody>
</table>
5. Stability analysis

The parental WT cell line from which the FAM60A knock-out clones were obtained and the obtained FAM60A knock-out cells were stably transfected with an expression vector encoding an antibody as polypeptide of interest. The transfected expression vector comprised an expression cassette comprising a polynucleotide encoding a neomycin phosphotransferase as selectable marker, an expression cassette encoding DHFR as selectable marker, an expression cassette comprising a polynucleotide encoding the light chain of an antibody and an expression cassette comprising a polynucleotide encoding the heavy chain of an antibody so that a complete antibody was expressed from said expression vector. All expression cassettes in the expression vectors were oriented in the same direction. The expression cassette used for the heavy chain was designed such that a portion of the heavy chain was expressed due to stop codon read through as fusion polypeptide comprising a membrane anchor. The fusion protein was displayed on the cell surface, thereby simplifying FACS analysis (see description). The transfected cells were selected for recombinant expression using G418 and MTX (1 μM) selection. From the selected pools of each stably transfected cell line (CHO WT, s16 and s23), cell clones that expressed the product of interest with good yields were obtained and were cultured for several weeks (7 weeks for WT CHO parental cell line (45 clones) and 8 weeks for the FAM60A knock-out cells (13 clones for s16 and 18 clones for s23)) in order to analyse their expression stability during prolonged culturing. To ensure that production cell lines can be upscaled to high volume bioreactors, stability studies of 12 weeks, in particular additional analysis of the expression stability during 12 week culturing, were also performed. Clones were classified as unstable if they lost more than 25% of their initial volumetric expression titer over the stability period analyzed. Within the usual level of variation some clones are just above or under the border line of 25%. The higher proportion of unstable cones in week 7/8 compared to week 12 for parental cell line can be explained with the variation in the productivity assay for clones which are close to the 25% threshold.

Table 4 compares the stability results that were obtained with the cell lines. As can be seen, the percentage of clones with stable volumetric titer is considerably higher in the clones derived from either FAM60A knock-out cell line compared to the clones that are derived from the wildtype cell line. This demonstrates that impairing the effect of endogenous FAM60A in the cell, here by introducing a gene knock-out, significantly improves the stability results during prolonged culturing. The ratio of stable versus unstable clones is significantly increased when using the cells of the invention so that more stable clones are obtained that maintain their favorable high expression characteristics during prolonged culturing. The antibody that was recombinantly expressed in this example was not codon-optimized and showed in the parental cell line a very high degree of instability. Because of this significant instability, this project was chosen as example for comparison because it demonstrates the significant benefits that are achieved with the present invention even when being confronted with difficult projects wherein instability rates are with the wildtype unmodified cell line high. However, as discussed above, with other projects, the instability rates are with the parental CHO wildtype cell line less high. However, in all cases analysed, the host cells according to the present disclosure, wherein the effect of FAM60A is impaired in said cells, achieve in
comparison with the unmodified wildtype a significant increase in the amount of stable cells. The stability rates can reach up to 60% or more, 70% or more, 80% or more, 85% or more, or even 90% or more, depending on the project. With the cells according to the present disclosure, which e.g. comprise a gene knock-out in the FAM60A gene or wherein a portion of the telomeric region comprising gene FAM60A is lost due to chromosome breakage, instable clones appeared irrespective of the project analysed significantly less frequently and even if appearing, the loss in volumetric productivity was less pronounced compared to corresponding cells wherein the effect of FAM60A is not impaired in the cell. Therefore, due to the increased percentage of stable clones that are obtained after transfection and selection, the cells according to the present disclosure allow to significantly shorten or even skip long-term stability studies. The stability studies of the FAM60A knock-out clones confirm the beneficial results that are achieved with the technology of the present disclosure.

Table 4: Results of stability studies

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Stable clones</th>
<th>Unstable clones</th>
<th>Stable clones</th>
<th>Unstable clones</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent cell line (7 and 12 weeks)</td>
<td>13.3%</td>
<td>86.7%</td>
<td>24.3%</td>
<td>75.7%</td>
</tr>
<tr>
<td>FAM60A knock-out cell line s16 (8 and 12 weeks)</td>
<td>61.5%</td>
<td>38.5%</td>
<td>61.5%</td>
<td>38.5%</td>
</tr>
<tr>
<td>FAM60A knock-out cell line s23 (8 and 12 weeks)</td>
<td>44.4%</td>
<td>55.6%</td>
<td>47.1%</td>
<td>52.9%</td>
</tr>
</tbody>
</table>

The results are also shown in Fig. 3 and demonstrate the important advantages that are achieved when impairing the effect of FAM60A in the cells, here by gene knock-out.

Example 2: Reducing C12orf35 gene expression by RNA interference (RNAi) increases expression yield

As described above, it is particularly preferred to additionally impair the effect of the expression product of gene C12orf35 in the eukaryotic cell wherein the effect of the FAM60A protein is impaired. Suitable methods for achieving impairment are described above and include, but are not limited to, reducing or eliminating functional expression of the endogenous C12orf35 gene. It was found that impairing the effect of the expression product of gene C12orf35 in a eukaryotic cell, such as preferably in a mammalian cell, surprisingly significantly increases the expression yield of a recombinant polypeptide of interest that is expressed from the respective cells. This beneficial effect with respect to expression yield that is obtained when reducing functional expression of gene C12orf35 is demonstrated in this example 2.

In order to demonstrate that reducing expression of gene C12orf35 results in an increase in volumetric and specific productivity, siRNAs were designed against different target genes located in the telomeric region of chromosome 8 of the Chinese hamster.
genome analysed (CHO-K1). siRNAs were designed against the following target genes listed in Table 5:

Table 5: siRNAs against different target genes

<table>
<thead>
<tr>
<th>Target gene</th>
<th>Sense</th>
<th>Antisense</th>
</tr>
</thead>
<tbody>
<tr>
<td>METTL20_1</td>
<td>CCCUGAUUGUUGUUGAGGATT (SEQ ID NO: 33)</td>
<td>UCCUCUAACAACUUCAGGTT (SEQ ID NO: 34)</td>
</tr>
<tr>
<td>C12orf35_1</td>
<td>CAUCCAGACAAUCUCUCATT (SEQ ID NO: 35)</td>
<td>UGUAAGAUUGGUGAGGTT (SEQ ID NO: 36)</td>
</tr>
<tr>
<td>C12orf35_2</td>
<td>CCAGAAAGUAACUCUCATT (SEQ ID NO: 37)</td>
<td>UGUAAGAUUCUCUCUGGTA (SEQ ID NO: 38)</td>
</tr>
<tr>
<td>Caprin2_6</td>
<td>UGGCUUGGGCGGAGGATT (SEQ ID NO: 39)</td>
<td>UUCUUCAGGCGGACGT (SEQ ID NO: 40)</td>
</tr>
<tr>
<td>FAM60A</td>
<td>GCUCUGCCUCUACAAGATT (SEQ ID NO: 41)</td>
<td>UUCUGUAGCGAAGGCGCA (SEQ ID NO: 42)</td>
</tr>
<tr>
<td>IPO8_1</td>
<td>GACCGGAACUUGACCUATT (SEQ ID NO: 43)</td>
<td>UAGGGUCAAAGCUUGGCTG (SEQ ID NO: 44)</td>
</tr>
<tr>
<td>IPO8_2</td>
<td>CGGAGACUCUCAAUGATT (SEQ ID NO: 45)</td>
<td>UCAAAUGAGAGUCUCGGA (SEQ ID NO: 46)</td>
</tr>
<tr>
<td>IPO8_3</td>
<td>GCCUGAUUGACGAGGATT (SEQ ID NO: 47)</td>
<td>UCCUCGUCUCAUACGGGTT (SEQ ID NO: 48)</td>
</tr>
<tr>
<td>Dennd5b_2</td>
<td>GGGUCUCUCUAAUCAAGATT (SEQ ID NO: 49)</td>
<td>UCUAGAAUAGGGAGACCTG (SEQ ID NO: 50)</td>
</tr>
<tr>
<td>Amn1_4</td>
<td>GCUCGUAAGUAUCACGATT (SEQ ID NO: 51)</td>
<td>UCAGGAUAUACUGACGCCA (SEQ ID NO: 52)</td>
</tr>
<tr>
<td>TMTC1_1</td>
<td>GUAUACCGUGAGAUAACATT (SEQ ID NO: 53)</td>
<td>UGUAUAAUCACAGGUAUACAT (SEQ ID NO: 54)</td>
</tr>
<tr>
<td>TMTC1_2</td>
<td>CGUGAAGUCAUCUCUCATT (SEQ ID NO: 55)</td>
<td>UGUAAGGAUCACUUCACCCGA (SEQ ID NO: 56)</td>
</tr>
</tbody>
</table>

siRNA negative control (no effect on expression – referred to as siRNA control) | Silencer Negative Control siRNA #5 (50uM) (Ambion, Cat#AM4642)

[168] The used siRNAs were validated using real-time RT-PCR to confirm that they reduce the expression of the target genes by gene silencing. Gene expression was normalized to 18S RNA. The gene expression observed when transfecting the siRNA negative control was set as 100%. The relative reduction of expression of the target gene is shown in the subsequent Table 6 for the two different siRNAs against target gene C12orf35:

Table 6

<table>
<thead>
<tr>
<th>Concentration</th>
<th>Gene expression (siRNA 1)</th>
<th>Gene expression (siRNA 2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>100pmol</td>
<td>Approx. 28%</td>
<td>Approx. 37%</td>
</tr>
<tr>
<td>125pmol</td>
<td>Approx. 25%</td>
<td>Approx. 38%</td>
</tr>
<tr>
<td>150pmol</td>
<td>Approx. 35%</td>
<td>Approx. 30%</td>
</tr>
</tbody>
</table>

[169] Furthermore, it was confirmed that the target siRNAs used in this experiment does not inhibit growth of the transfected cells. Furthermore BLAST (Basic Local Alignment Search Tool) analysis based on the available Chinese hamster genome data does not indicate any off-target effects.
The following cell lines were transfected: A CHO cell line derived from CHO-K1 was used as parental cell line. Said cell line expresses the above genes as is shown in Fig. 2. This parental cell line was not transfected with an expression vector and served as control. CHO cells (clones and pools) derived from said parental cell line which comprised an expression vector encoding an antibody as protein of interest stably integrated into the genome was used to determine the effect of the siRNAs. The expression vector comprised in said cell clone comprised selectable marker genes and the antibody heavy chain and the antibody light chain were expressed from different expression cassettes. The expression cassette used for the heavy chain was designed such that a portion of the heavy chain was expressed due to stop codon read through as fusion polypeptide comprising a membrane anchor. The fusion protein was displayed on the cell surface, thereby simplifying FACS analysis (see description). Said CHO cells expressed the above-mentioned siRNA target genes similar to the parental cell line which was determined by microarrays for hundreds of clones and pools. A CHO clone which recombinantly expressed the antibody was used in order to determine whether a downregulation of one or more of the above target genes results in an increase of the expression of the polypeptide of interest. If this was the case, an increase of the volumetric antibody productivity of said cell clone would be seen which is detectable using FACS analysis.

The CHO clone comprising the expression vector stably integrated into the genome was transfected either with a siRNA control (not having an effect on gene expression) or with one of the above-mentioned siRNAs against the target genes. After transfection of the siRNAs it was analyzed whether the reduction of the expression of the target gene results in an increase of the expression of the antibody. Inter alia, the transfected cells were stained by using a fluorescent detection compound and analysed by FACS in order to determine the expression rate of the antibody. The more antibody is produced, the more displayed fusion protein can be stained using a labeled compound and accordingly, the higher is the fluorescence signal detected by FACS. Therefore, the higher the intensities in the FACS profiles, the more antibody is expressed.

The results are shown in Fig. 4A to L for the different siRNAs tested. The left peak in the profiles corresponds to the signal obtained for the parental cell line, which does not express the antibody. The two other curves represent the results obtained with the antibody expressing cell clone which was transfected with either the siRNA-negative control (no effect on expression) or with the tested siRNA which reduces the expression of its target gene. If the tested siRNA and accordingly, the downregulation of the target gene does not have an effect on the expression of the antibody (i.e. no upregulation of volumetric productivity), the obtained fluorescence curve for the siRNA-negative control and the fluorescent curve obtained for the target siRNA overlap and thus are basically identical. This was in essence the case for all tested target genes on clone level, except for gene C12orf35. For FAM60A, however, a slight shift was seen on pool level which is assumed to be attributable to the increased expression stability (data not shown).
As can be seen from the results obtained with the siRNAs against C12orf35 (see Fig. 4B and C), the fluorescence peaks obtained for the siRNA-negative control and the siRNA against gene C12orf35 clearly separate upon silencing gene C12orf35 using RNAi. The fluorescent peak obtained for the cell clone transfected with the siRNA against gene C12orf35 clearly shifts to the right (marked by arrows), which means that the fluorescence is significantly increased. This observed increase in fluorescence is attributable to a higher expression of the antibody as more fusion protein is present and thus is stained on the cell surface. Therefore, this experiment clearly shows that a downregulation of the functional expression of gene C12orf35 directly results in a significant upregulation of the recombinant expression of the antibody (yield of antibody). The same remarkable shift in the FACS profiles was observed when using said siRNAs against C12orf35 in all three concentrations. A prolonged reduction of the expression of gene C12orf35 by RNA interference can be achieved if for example stably integrated in an expression vector which expresses an RNAi inducing transcript, as is for example described in the description. Furthermore, a reduction or elimination of the expression of gene C12orf35 can be achieved by gene knockout or gene deletion/mutation, according to one embodiment by deleting a portion of the telomeric region of chromosome 8 in case of hamster cells such as CHO cells, as described in the description. Furthermore, as is described therein, it is also feasible to reduce or eliminate the effect of the expression product e.g. by introducing one or more mutations which result in a non-functional or less functional protein.

The achieved increase in the expression of the recombinant polypeptide of interest was also confirmed by analysis of the mRNA expression level of the heavy and the light chain (which were expressed from separate expression cassettes, see above). The results are shown for two different polypeptides of interest (antibody 1 and 2) in Figs. 5 and 6. The data shown is normalized to the siRNA negative control (125pmol). It was found that reduction of expression of gene C12orf35 leads to significantly higher mRNA levels of the heavy chain and the light chain of the expressed antibody. In comparison, reduction of expression of other tested target genes did not have an impact on the mRNA expression level of the heavy and the light chain. Thus, reduction of the expression of gene C12orf35 results in a significant increase in the mRNA level of the recombinant polypeptide of interest. Furthermore, it was observed that also other introduced genes such as selection markers are upregulated if gene C12orf35 is silenced. An experiment, in which the gene silencing effect was measured over time starting on day 3, showed that siRNA1 (see siRNA C12orf35_1 in Table 7) had a longer effect compared to siRNA2 (see siRNA C12orf35_2 in Table 7). In addition, cell numbers and titer were determined throughout the time course experiment and revealed that downregulation of C12orf35 leads to significantly higher specific productivities (see Fig. 7).

Furthermore, expression of the C12orf35 gene relative to 18S RNA was analysed in an antibody expressing clone upon repression with siRNA 1 and 2 and compared to different controls. The results are shown in the subsequent Table 7:
Table 7

<table>
<thead>
<tr>
<th>siRNA C12orf35_1</th>
<th>18S</th>
<th>97%</th>
<th>0.0129%</th>
<th>0.8517%</th>
<th>3.4639%</th>
</tr>
</thead>
<tbody>
<tr>
<td>siRNA C12orf35_2</td>
<td>91%</td>
<td>0.0186%</td>
<td>1.0643%</td>
<td></td>
<td>2.9623%</td>
</tr>
<tr>
<td>siRNA control</td>
<td>93%</td>
<td>0.0617%</td>
<td>0.3098%</td>
<td></td>
<td>0.9195%</td>
</tr>
<tr>
<td>cells untreated</td>
<td>100%</td>
<td>0.0782%</td>
<td>0.2713%</td>
<td></td>
<td>0.7840%</td>
</tr>
<tr>
<td>cells lipofectamin</td>
<td>114%</td>
<td>0.0648%</td>
<td>0.2633%</td>
<td></td>
<td>0.8019%</td>
</tr>
</tbody>
</table>

Example 3: Generation of a CHO cell line which comprises a deletion in the telomeric region of chromosome 8 which deletes gene FAM60A and gene C12orf35

[176] A novel CHO cell line (C8DEL) was generated, which comprises a deletion in the telomeric region of the q arm of chromosome 8. The deletion was induced by chromosome breakage. The deleted portion comprised gene FAM60A as well as among others gene C12orf35 which is located telomeric from gene FAM60A (see Fig. 1). Said novel cell line was obtained from a parental cell line derived from CHO-K1. Said cell line with a chromosome break in chromosome 8 was prepared as follows. The parental CHO cells were split at 2E5 cells/ml in culture medium comprising 0.5 μM, 1 μM or 2 μM MTX. After six days the cell viabilities were around 30-40%. Cells were centrifuged at 180xg for 5 min and cultivated in culture medium without MTX to allow the cells to recover until viabilities were above 95% (after ca. 21 days). This procedure was repeated two more times. Single cell clones were obtained from cell pools. Overall 561 cell clones were grown and DNA was isolated using the "Extract-N-Amp Blood PCR Kit". PCR screening using primers detecting the gene lpo8 was performed. Three out of the 561 clones were lpo8 negative indicating the loss of telomeric region of chromosome 8 which includes the lpo8 gene. The lpo8 gene is located centromeric of the FAM60A gene (see Fig. 1). Thus, if the lpo8 gene is deleted due to chromosome breakage, all genes located telomeric of the lpo8 gene (and accordingly also the FAM60A gene and gene C12orf35) are deleted as well. These three clones were expanded and further evaluated. One of these clones is referred to as "C8DEL" cell line. Using PCR technique the breakpoint of the telomeric region of chromosome 8 from the C8DEL cell line could be determined. The breakpoint was determined between two PCRs, called PCR20 and 28:

PCR20: fwd 5'-ACC AGT GAA TAA TCG TGT TT-3' (SEQ ID NO: 57), rev 5'-CTA TGA GTC AAT GTC CCA AG-3' (SEQ ID NO: 58);

PCR28: fwd 5'-CAC ACA CAA CCT CCT AAC AAC CC-3' (SEQ ID NO: 59), rev 5'-TTC CGC ACC GAC TCA GTT CT-3' (SEQ ID NO: 60)

[177] The breakpoint lies within the Tmtc1 gene. Additionally it could be demonstrated that the identified breakpoint of C8DEL cell line is stable over several weeks of cultivation (determined via PCR). Transfecting this novel cell line with an expression vector encoding a product of interest increases the chances of selecting stable producing clones with high expression capacity as is shown in the following. Transfection and MTX treatment of C8DEL
cell line apparently has no effect on the breakpoint (no further genetic material is lost) as was determined based on the analysis of transfected clones.

**Example 4: Analysis of the characteristics of the cell line C8DEL**

[178] The cell line C8DEL was analyzed for its performance when recombinantly expressing a polypeptide of interest and compared to the parental cell line from which the cell line C8DEL was derived. As described above, said parental cell line does not comprise a corresponding deletion in the telomeric region of chromosome 8.

**4.1. Analysis of productivity**

[179] The volumetric productivity of C8DEL was evaluated in comparison to the parental cell line from which it was derived. Stable as well as transient transfections were performed.

**Stable transfection**

[180] Cell cultivation, transfection and screening were carried out in shake flasks using suspension growing CHO cells in a chemically defined culture medium. Cells were transfected by electroporation with different expression vectors encoding a variety of antibodies and therapeutic proteins. Expression vectors used comprised an expression cassette comprising a polynucleotide encoding a neomycin phosphotransferase as selectable marker and an expression cassette encoding a DHFR as selectable marker. The expression vectors used for expressing antibodies additionally comprised an expression cassette comprising a polynucleotide encoding the light chain of an antibody and an expression cassette comprising a polynucleotide encoding the heavy chain of an antibody so that a complete antibody was expressed from said expression vector. The expression vectors used for expressing a polypeptide which was not an antibody, comprised an expression cassette comprising a polynucleotide encoding the polypeptide in addition to the selection markers. All expression cassettes in the expression vectors were oriented in the same direction. The vector was suitable for FACS selection and details of such vector are described above.

[181] Depending on the cell viability, the first selection step was started 24-48 h after transfection by adding G418 selective medium to the cells. As soon as cells recovered to a viability of above 80%, a second selection step was applied by passaging the cells to 500nM MTX or 1μM MTX.

[182] Volumetric productivity of the selected cell populations was analyzed after G418 and MTX selection steps via overgrown shake flask batch cultures in medium with G418 or MTX. The G418 batch was done in 30ml (125ml flask) and MTX fed-batch in 100ml (500flask). G418 batch cultures were seeded at 1E5 vc/ml in shake flask and cultivated in a shaker cabinet (not humidified) at 150 rpm and 10% C02. Fed-batches were seeded at 4E5 vc/ml. Viability of cells had to be >90% when starting the assay. Titer determination took place at day 14. Antibody titers in the cell culture supernatant were determined by protein-A HPLC 14 days after starting the culture.
After the first selection step (G418 selection) a massive volumetric titer increase (12-35 fold) could be detected for the stably transfected C8DEL pools in comparison to the stably transfected parental pools. After the second selection step (MTX selection) the C8DEL pools were expressing 4-7 fold more polypeptide of interest in comparison to the transfected parental cells, as exemplified by 2 antibodies (antibody 1 and antibody 2). Volumetric G418 and MTX (fed-batch) titers of C8DEL in comparison to the parental CHO cell line not comprising the deletion in the telomeric region of chromosome 8 are exemplarily displayed for two antibody projects in the following Tables 8.a and 8.b. Tables 8.a and 8.b show the volumetric G418 and MTX fed-batch pool titer (antibody) produced in C8DEL in comparison to the parental pools (average of 4 pools/condition are shown).

<table>
<thead>
<tr>
<th>Table 8a: Pool titers of example antibody 1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cell line</strong></td>
</tr>
<tr>
<td>Parental cell line derived from CHO-K1</td>
</tr>
<tr>
<td>C8DEL</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 8.b: Pool titers of example antibody 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cell line</strong></td>
</tr>
<tr>
<td>Parental cell line derived from CHO-K1</td>
</tr>
<tr>
<td>C8DEL</td>
</tr>
</tbody>
</table>

The results support the conclusion that a deletion in the telomeric region of chromosome 8 which includes the FAM60A gene and the C12orf35 gene is correlated with higher volumetric productivity. Already with respect to volumetric pool titer, the C8DEL cell line is outperforming the parental cell line that does not comprise a respective deletion in the telomeric region of chromosome 8. Considering the siRNA results from example 2, it is believed that the volumetric titer increase is due to the loss of gene C12orf35 located on the telomeric part that was lost.

**Transient transfection**

C8DEL cells and parental cell line cells grown in culture medium were transiently transfected in triplicates with expression plasmids encoding either eGFP or an Fc fusion protein as model protein of interest. Polyethyleneimine (PEI) was used as transfection reagent. The titer of the model protein in the medium supernatant was measured by Protein A HPLC on day 3 and day 6 after transfection. The expression of the model protein was approx. 3 fold higher in C8DEL. The percentage of eGFP-expressing cells was measured 48 h after transfection by flow cytometry with non-transfected cells acting as negative control. Cells exhibiting a fluorescence level greater than 99% of the negative control cells were regarded as “transfected”. Cells exhibiting fluorescence level of more than 1000-fold the
intensity of the negative control cells were regarded as "highly fluorescent". The number of high fluorescent cells was 2-3 fold higher using the C8DEL cell line compared to the parental cell line from which C8DEL was derived.

This example shows that the advantages of increased volumetric productivity are also achieved when performing a transient transfection with the C8DEL cell line wherein a portion of the telomeric region of chromosome 8 is lost due to chromosome breakage.

4.2. Stability analysis

The stability characteristics of 46 C8DEL derived clones and 37 clones derived from the parental cell line (which were tested to be IP08 positive and hence did not lose the telomeric region of chromosome 8) was analysed after stable transfection. All clones recombinantly expressed the same antibody as product of interest and were classified as stable if they were not losing more than 25% antibody titer (volumetric) in 12 weeks. 76% of the analyzed clones from the parental cell line lost more than 25% of titer (volumetric) within 12 weeks of cultivation. Only 24% of the analysed clones were classified as stable. Thus, instability rates were high. In comparison, 67% of the C8DEL clones could be classified as stable and only 33% went unstable as is shown in Table 9:

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Stable clones</th>
<th>Unstable clones</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent cell line (12 weeks)</td>
<td>24%</td>
<td>76%</td>
</tr>
<tr>
<td>C8DEL (deletion of gene FAM60A due to chromosome breakage) (12 weeks)</td>
<td>67%</td>
<td>33%</td>
</tr>
</tbody>
</table>

Using a x2-test with a Yates-correction a p-value of 0.0002 could be calculated which supports that C8DEL derived clones have a significant higher tendency to be stable producers. Thus, a significant higher number of stable producing clones for the C8DEL cell line were found. This further supports in particular in conjunction with the knock-out experiments of example 1 that hamster cells such as CHO cells, wherein a portion of the telomeric region of chromosome 8 comprising gene FAM60A is deleted, show superior stability characteristics. Thus, using such cell line for recombinant expression increases the chance that high and stable producing recombinant cells are identified. Furthermore, the volumetric productivity of said clones was analysed (see 4.3.).

4.3. Further analyses of the characteristics of C8DEL

The characteristics of the CHO cell line comprising a deletion in the telomeric region of chromosome 8 wherein said deletion comprises gene FAM60A as well as gene C12orf35 were analysed in additional experiments which demonstrate further advantages of said cell line.

Less single cell cloning required for selecting high producers

An advantageous characteristic of the C8DEL cell line is the greater proportion of high producing clones after single cell cloning. It was found that the C8DEL pools contain an
enlarged proportion of high producing cells (resulting in an increased volumetric pool titer) compared to the parental cell line derived from CHO-K1. After single cell cloning of C8DEL pools using FACS technology, a significantly greater proportion of clones expressing high quantities of antibody were selected compared to pools derived from the parental WT cell line. Table 10 shows that most clones derived from the parental cell line had a "volumetric 96 well titer" of 0-20 mg/L. In contrast, the majority of clones derived from C8DEL cell line had an average volumetric titer of 80-100 mg/L what is a significant improvement. One advantage of using C8DEL pools is the reduced number of clones which have to be generated to obtain a comparable amount of high and stable producing clones. This significantly reduces the screening effort.

<table>
<thead>
<tr>
<th>96 well titer (mg/L)</th>
<th>Parental cell line</th>
<th>C8DEL</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-20</td>
<td>80.3%</td>
<td>0.8%</td>
</tr>
<tr>
<td>20-40</td>
<td>6.1%</td>
<td>3.1%</td>
</tr>
<tr>
<td>40-60</td>
<td>5.4%</td>
<td>5.3%</td>
</tr>
<tr>
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Using the C8DEL cell line as production cell line results not only in an enlarged proportion of high producing clones, also the volumetric titer of the individual C8DEL clones is higher. Considering the results of example 2, it is believed that this increase in yield is attributable to the deletion of gene C12orf35. Fig. 8 shows the volumetric titer of the 45 highest producing clones from the parental cell line derived from CHO-K1 (all Lpo8 positive) and C8DEL from an antibody project (results of the stability analysis of said clones is shown in 4.2). As can be seen, the average volumetric titer for the clones derived from C8DEL is higher compared to the parental cell line, and additionally, also the highest antibody producer clones are originated from C8DEL cell line.

**Bioreactor suitability**

Additional tests to evaluate C8DEL cell line in comparison to the parental cell line derived from CHO-K1 were performed *inter alia* to determine their suitability for upscaling. Bioreactor runs have shown that C8DEL cell line is suitable for upscaling. C8DEL cell line cultivated in bioreactors had a viable cell density that is suitable for large scale production. Furthermore, it was found that the viability was better than that for the parental cell line. Overall, the C8DEL cell line is suitable for upscaling and is outperforming the parental cell line from which it is derived regarding viability. The viability of the C8DEL cell line stays longer at a higher level.
**Improved time lines from transfection to stable pool production**

[193] Another advantage of the C8DEL cell line is the faster recovery from MTX selection. The recovery of pools after MTX incubation was accomplished 7-8 days faster compared to the parental cell line wherein no portion of the telomeric region of chromosome 8 comprising gene FAM60A is deleted. Overall, it was found that the cell crisis is significant lower with the cells according to the present disclosure.

**Example 5: Selection using a folate receptor as selectable marker**

[194] The C8DEL cell line was used in different settings in conjunction with the folic acid receptor as selectable marker and shows particular advantages in conjunction with said selection system. In particular, a combined selection against the folate receptor and DHFR as selectable markers is beneficial. Here, the transfected cells comprised a human folate receptor alpha and DHFR as selectable markers and expressed an antibody. Whereas selection of the parental cell line with very low amounts of folic acid (50 nM folic acid (FA)/50nM MTX) encountered difficulties due to the selection stringency (cells did not always recover), the combination of C8DEL and the folate receptor as selectable marker is very powerful under such stringent conditions and resulted in a significant volumetric titer increase. Table 11 highlights the volumetric titer differences between the parental cell line and C8DEL as well as the additional volumetric titer increase that is achieved when using the folate receptor as a selection marker in combination with low amounts of folic acid instead of 500nM MTX selection step. Thus, the use of the mammalian cells described herein wherein the expression of gene FAM60A and gene C12orf35 is reduced or eliminated allows in combination with the folate receptor/DHFR selection system to use very stringent selection conditions that do not require the use of high amounts of toxic agents.

### Table 11

<table>
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<th>Cell line</th>
<th>Selection conditions</th>
<th>Pool titer (mAb g/L) – shake flask batch culture</th>
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<td>Parental cell line derived from CHO-K1</td>
<td>0.8 g/L G418/500nM MTX</td>
<td>Approx. 0.07</td>
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<tr>
<td>C8DEL</td>
<td>0.8 g/L G418/500nM MTX</td>
<td>Approx. 0.83</td>
</tr>
<tr>
<td>C8DEL</td>
<td>50 nM FA/50nM MTX</td>
<td>Approx. 1.61</td>
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[195] Furthermore, C8DEL cells were transfected (nucleofection) with an expression vector which comprised an expression cassette comprising a polynucleotide encoding a human folate receptor alpha and an expression cassette comprising a polynucleotide encoding DHFR. Thus, both selectable markers FRalpha and DHFR were on the same expression vector. Furthermore, the expression vector comprised an expression cassette comprising a polynucleotide encoding the light chain of an antibody and an expression cassette comprising a polynucleotide encoding the heavy chain of an antibody. The expression cassette for the antibody heavy chain was designed such that a portion of the heavy chain was due to stop codon readthrough produced as membrane-anchored fusion, thereby
facilitating FACS selection (see above). Five different selection conditions using 100nM folic acid (FA) and different concentrations of MTX were tested. The selection media are summarized in subsequent Table 12. After selection, the selected cell pools were transferred to complete medium and grown in shake flask batch cultures. At day 13 of the culture, samples of the culture medium were taken and analyzed for antibody content by Protein-A HPLC. The results are also shown in Table 12.

Table 12

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<tr>
<th>Selection condition</th>
<th>Approx. antibody concentration [g/L]</th>
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<tr>
<td>100nM FA/no MTX</td>
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<tr>
<td>100nM FA/1nM MTX</td>
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<tr>
<td>100nM FA/5nM MTX</td>
<td>0.46</td>
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<td>100nM FA/10nM MTX</td>
<td>1.44</td>
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<td>100nM FA/50nM MTX</td>
<td>1.57</td>
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</table>

As can be seen, a MTX concentration already as low as 5nM provided a selection advantage. Increasing the selection stringency also increases the volumetric pool titer. Thus, the volumetric antibody productivities are significantly increased. Furthermore, compared to standard MTX selections, significantly lower concentrations of MTX can be used during selection. This is an important advantage considering that MTX is a toxic agent. Furthermore, it was analyzed how the selection stringency influences the volumetric pool titers and the time for selection. It was found that increasing the selection stringency by increasing the concentration of MTX prolongs the recovery time. Thus, the selection stringency can be adjusted according to the needs of different applications (time versus titer).

Furthermore, analysis of the pools obtained after folic acid/MTX selection for surface expression of the antibody by FACS show that using this selection system in combination with the novel cell line significantly increases the abundance of high producers in the cell pool as is apparent from the obtained fluorescent profiles shown as Fig. 9A to E. The concentration of MTX was increased from A to E (A: no MTX; B: 1nM MTX; C: 5nM MTX; D: 10nM MTX; E: 50nM MTX). When increasing the MTX concentration, the number of high expressing cell clones in the cell pool was increased as can be derived from the increase of the peak size on the right hand side (higher fluorescence correlating with a higher antibody expression rate). Using 50nM folic acid in combination with 10nM MTX (see Fig. 9D) already resulted in a cell pool predominantly comprising high producing cell clones (one dominant peak on the right hand side). Furthermore, when increasing the MTX concentration to 50nM (see Fig. 9E), basically exclusively high-producing cells were comprised in the obtained pool. These results are remarkable, because when using the C8DEL cell line in combination with
the folate receptor/DHFR selection system, one obtains a pool profile after FACS analysis, which closely resembles more that of a cell clone (comprising genetically identical cells) than that of a cell pool (comprising genetically different cells). It appears that the deletion of the C12orf35 gene comprised in the lost telomeric region of cell line C8DEL results in a significant increase of the volumetric productivity, so that basically the majority of the cells in the cell pools obtained after folic acid/MTX selection under appropriate conditions were high producers according to the FACS profiles.

Furthermore, it was found that when cultivating stably transfected clones obtained from the C8DEL cell line (gene FAM60A is lost, see above) in selective medium (50nM folic acid, 10nM MTX) stability rates of up to 80% and up to almost 100% could be obtained in projects. Significant high stability results were also achieved in a semi-selective medium, which only comprised a limited concentration of folic acid (50nM), however, no MTX. Here, stability rates up to 87% were achieved with this cell line. In certain projects, stability rates of up to almost 100% were obtained.

Example 6: Validation tool to identify high and stable producers based on the expression profile

A real-time RT-PCR analytical tool was developed to predict clone productivity and stability at an early stage of development pipeline process. Real-time RT-PCR was implemented for four genes: C12orf35, Dennd5b, Fam60a and lpo8 (all localized at telomeric region on the q arm of chromosome 8). After selection and clone generation, several hundred clones stably expressing an antibody as polypeptide of interest were analyzed with respect to the presence and expression level of these four genes at telomeric region of chromosome 8 and the expression yield. A clear correlation was found between stability, volumetric productivity and a loss in the telomeric region of chromosome 8. A study was conducted to determine if there is a correlation between stability and presence of telomeric region of chromosome 8. Clones were classified as stable if they were not losing more than 25% titer (volumetric) in 12 weeks. A significant correlation between loss of telomeric region of chromosome 8 and clone stability is existent (p-value: 4.67E-06 based on x2-test). Consequently, the loss of telomeric region on chromosome 8 which includes gene FAM60A can be used as a prediction tool for stability. Analysing the presence or absence of telomeric region of chromosome 8 via real-time RT-PCR increases the probability to select a higher proportion of stable clones in pipeline projects.

Furthermore, it was found by analyzing several hundred clones that have lost a portion of the telomeric region of chromosome 8, that there appear to be several breakpoints in the telomeric region of chromosome 8 existent that can be induced. In most analysed cases, the breakpoint was located centromeric of the lpo8 gene. Breakpoints were also detected between FAM60A and lpo8. The deleted region comprised in all cases gene C12orf35 (which is located telomeric of gene encoding methyltransferase-like protein 20) what is associated with an increase in volumetric productivity.
CLAIMS

1. An isolated eukaryotic cell, wherein the genome of the eukaryotic cell is altered so that the effect of protein FAM60A is impaired in said cell and wherein said cell comprises integrated into its genome at least one heterologous polynucleotide encoding a product of interest.

2. The isolated eukaryotic cell according to claim 1, wherein the effect of protein FAM60A is impaired because functional expression of gene FAM60A is reduced or eliminated in said cell.

3. The isolated eukaryotic cell according to claim 2, wherein the eukaryotic cell is altered to reduce or eliminate functional expression of gene FAM60A by gene knock-out, gene mutation, gene deletion, gene silencing or a combination of any the foregoing.

4. The isolated eukaryotic cell according to claim 3, wherein the FAM60A gene comprises as gene mutation one or more mutations in the FAM60A gene which provide a non- or less functional expression product.

5. The isolated eukaryotic cell according to claim 4, wherein at least one copy of gene FAM60A is deleted or functionally inactivated in the genome of the eukaryotic cell.

6. The isolated eukaryotic cell according to one or more of claims 1 to 5, wherein the eukaryotic cell is a metazoan cell, a vertebrate cell or a mammalian cell.

7. The isolated eukaryotic cell according to claim 6, wherein the eukaryotic cell is a mammalian cell.

8. The isolated eukaryotic cell according to one or more of claims 1 to 7, wherein a portion of a chromosome is deleted, wherein the deleted portion comprises gene FAM60A.

9. The isolated eukaryotic cell according to claim 8, wherein

   a) the eukaryotic cell is a hamster cell and at least a portion of the telomeric region of chromosome 8 is deleted, wherein said deleted portion comprises gene FAM60A; or

   b) the eukaryotic cell is a mouse cell and at least a portion of the telomeric region of chromosome 6 is deleted, wherein said deleted portion comprises gene FAM60A.

10. The isolated eukaryotic cell according to claim 9, having one or more of the following characteristics:

   a) the deleted telomeric region comprises gene FAM60A and comprises one or more genes selected from Caprin2 and Lpo8;
b) the deletion is induced by chromosome breakage and the breakpoint is located centromeric of the lpo8 gene, preferably within the Tmtc1 gene;
c) at least a portion of the telomeric region is deleted in both chromosomes of the respective chromosome pair, wherein the deleted portions comprise gene FAM60A.

11. The isolated eukaryotic cell according to one or more of claims 1 to 10, having one or more of the following characteristics:

a) the eukaryotic cell is a rodent cell,
b) the eukaryotic cell is a hamster cell, preferably a CHO cell,
c) the eukaryotic cell expresses endogenously DHFR and a folate receptor, and/or
d) the eukaryotic cell is provided as cell clone or cell line.

12. The isolated eukaryotic cell according to one or more of claims 1 to 11, wherein the endogenous FAM60A protein shares at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% homology or identity to one or more of the amino acid sequences shown in SEQ ID NO: 1 to 8.

13. The isolated eukaryotic cell according to one or more of claims 1 to 12, wherein in said cell additionally the effect of the expression product of gene C12orf35 is impaired, preferably by reducing or eliminating functional expression of gene C12orf35.

14. The isolated eukaryotic cell according to claim 13, wherein the C12orf35 gene is a gene encoding a protein that shares at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% homology or identity to one or more of the amino acid sequences shown in SEQ ID NO: 10 to 16 or the protein encoded by SEQ ID NO: 17.

15. The isolated eukaryotic cell according to one or more of claims 1 to 14, wherein the eukaryotic cell comprises stably integrated into its genome at least one heterologous polynucleotide encoding a product of interest and at least one heterologous polynucleotide encoding a selectable marker or reporter polypeptide and wherein said heterologous polynucleotides are located on the same or different expression vectors.

16. The isolated eukaryotic cell according to one or more of claims 1 to 15, having one or more of the following characteristics:

a) the product of interest is a polypeptide;
b) the product of interest is a polypeptide and the eukaryotic cell secretes said polypeptide of interest into the cell culture medium; and/or
c) the product of interest is a polypeptide selected from a therapeutic polypeptide and a diagnostic polypeptide.
17. The isolated eukaryotic cell according to one or more of claims 1 to 16, originating from a population of respective cells, wherein on average at least 40%, preferably at least 50% of the cells originating from said population of respective cells do not lose more than 30%, preferably 25%, of their expression titer (volumetric) over a time period of 8 weeks, preferably 10 weeks, more preferably over a time period of 12 weeks.

18. A method for selecting a host cell which recombinantly expresses a product of interest, comprising

(a) providing eukaryotic cells according to one or more of claims 1 to 17 as host cells; and
(b) selecting one or more host cells expressing the product of interest.

19. The method according to claim 18, wherein stage (a) comprises stably transfecting eukaryotic cells, wherein the genome of said cells is altered so that the effect of protein FAM60A is impaired in said cells, with at least one polynucleotide encoding the product of interest to provide eukaryotic host cells which comprise stably integrated into the genome at least one heterologous polynucleotide encoding the product of interest.

20. The method according to claim 18 or 19, having one or more of the following characteristics

a) the eukaryotic cells are mammalian cells;

b) said host cells provided in stage (a) additionally comprise at least one heterologous polynucleotide encoding a selectable marker and stage (b) comprises culturing said plurality of host cells under conditions selective for the selectable marker;

c) heterologous polynucleotides are introduced into the eukaryotic cells by transfecting one or more expression vectors;

d) the recombinant product of interest is a polypeptide;

e) stage (b) comprises multiple selection steps;

f) stage (b) comprises performing a flow cytometry based selection;

g) the selected host cells recombinantly express an immunoglobulin molecule;

h) said host cells provided in stage (a) comprise at least two heterologous polynucleotides each encoding a selectable marker wherein the first selectable marker is a folate receptor and wherein the second selectable marker is a DHFR and wherein stage (b) comprises culturing said plurality of host cells in a selective culture medium which comprises folate in a limiting concentration and a DHFR inhibitor.

21. A method for recombinantly producing a product of interest, comprising utilizing a eukaryotic cell according to one or more of claims 1 to 17 as host cell for recombinant expression of the product of interest.

22. The method according to claim 21, comprising
(a) culturing host cells according to one or more of claims 1 to 17 under conditions that allow for the expression of the product of interest;
(b) isolating the product of interest from the cell culture medium and/or from said host cells; and
(c) optionally processing the isolated product of interest.

23. The method according to claim 21 or 22, wherein for production a eukaryotic cell is used in which additionally, the effect of the expression product of gene C12orf35 is impaired, preferably by reducing or eliminating functional expression of gene C12orf35.

24. The method according to one or more of claims 21 to 23, having one or more of the following characteristics:

   a) the eukaryotic cell is a metazoan cell, a vertebrate cell or a mammalian cell;
   b) the eukaryotic cell is a mammalian cell, preferably a rodent cell;
   c) the product of interest is a polypeptide;
   d) the product of interest is a polypeptide and the host cell secretes the polypeptide of interest into the cell culture medium.

25. A method for producing a eukaryotic cell according to one or more of claims 1 to 17, comprising altering the genome of an eukaryotic cell to impair the effect of protein FAM60A in said cell and stably transfecting into said cell at least one expression vector comprising a polynucleotide encoding a product of interest.

26. The method according to claim 25, comprising reducing or eliminating the functional expression of gene FAM60A thereby impairing the effect of protein FAM60A in the eukaryotic cell.

27. A method for analyzing eukaryotic cells for their suitability as host cells for recombinant expression of a product of interest, comprising analyzing directly or indirectly whether the effect of FAM60A is impaired in said cells.

28. The method according to claim 27, wherein directly analyzing comprises analyzing whether the functional expression of gene FAM60A is reduced or eliminated in said cells.

29. The method according to claim 27 or 28, wherein prior to analysis, the eukaryotic cells are treated with an agent that induces chromosome breakage and wherein the analysis comprises analyzing whether treatment with said agent resulted in a deletion of a portion of a chromosome which includes gene FAM60A.

30. The method according to one or more of claims 27 to 29, wherein the cells are hamster cells and the method comprises analyzing whether expression of one or more genes located in the telomeric region of chromosome 8 and being selected from the group consisting of the Tmtc1 gene and genes located telomeric of the Tmtd gene is eliminated or reduced thereby
analyzing whether the functional expression of gene FAM60A is reduced or eliminated in said cells.

31. The method according to one or more of claims 27 to 30, comprising additionally analyzing directly or indirectly whether the effect of the expression product of endogenous gene C12orf35 is impaired in said cells.

32. The method according to one or more of claims 27 to 31, wherein prior to analysis, the eukaryotic cells are transfected with a heterologous polynucleotide encoding a product of interest and a heterologous polynucleotide encoding a selectable marker, and wherein prior to analysis at least one selection step is performed to identify successfully transfected host cells.

33. The method according to one or more of claims 27 to 32, wherein a plurality of cell clones is analyzed for discriminating between stable and unstable and optionally between high and low producing cell clones and wherein one or more cell clones wherein the functional expression of gene FAM60A is reduced or eliminated are selected as production clone.

34. Use of an isolated eukaryotic cell for recombinantly expressing a product of interest, wherein the genome of the eukaryotic cell is altered so that the effect of protein FAM60A is impaired in said cell.

35. The use according to claim 34, having one or more of the following characteristics:

a) the eukaryotic cell is a metazoan, a vertebrate or mammalian cell;

b) the eukaryotic cell is a mammalian cell;

c) the genome of the eukaryotic cell is altered as defined in one or more of claims 2 to 5 and 8;

d) in said eukaryotic cell, additionally the effect of the expression product of gene C12orf35 is impaired, preferably by reducing or eliminating functional expression of endogenous gene C12orf35;

e) the product of interest is a polypeptide;

f) the product of interest is a polypeptide that upon expression in the eukaryotic cell is secreted into the cell culture medium;

g) the use comprises stably transflecting the eukaryotic cell with an expression vector comprising a polynucleotide encoding a polypeptide of interest, and/or

h) the eukaryotic cell is a cell as defined in one or more of claims 1 to 17.
1 = coiled-coil domain-containing protein 91
2 = hypothetical
3 = fatty acyl-CoA reductase 1 isoform 1
4 = endoplasmic reticulum-Golgi intermediate compartment protein 2
5 = 40S ribosomal protein S4, X isoform like
6 = transmembrane and TPR repeat-containing protein 1
7 = zinc finger HIT domain-containing protein 1
8 = importin-8-like protein
9 = caprin-2-like protein
10 = protein* FAM60A-like
11 = DENN domain-containing protein 5B-like protein
12 = methyltransferase-like protein 20
13 = putative protein AMN1 like protein
14 = hypothetical protein
15 = Retroviral nucleocapsid protein Gag containing protein
16 = opioid growth factor receptor-like protein
17 = hypothetical protein
18 = uncharacterized protein C12orf35 homolog*
19 = putative protein bicaudal D
20 = 6-phosphofructokinase type C-like protein
21 = zinc finger protein
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

INV. C07K16/00

ADD.

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

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

C07K C12N

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 practicable, search terms used)

EPO-Internal, WPI Data, BIOSIS, EMBASE, Sequence Search

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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<td>I. M. MUNOZ ET AL: &quot;Family with Sequence Similarity 60A (FAM60A) Protein is a Cell Cycl e-fluctuating Regulator of the SIN3-HDAC1 Histone Deacetylase Complex&quot;, JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 287, no. 39, 21 September 2012 (2012-09-21), pages 32346-32353, XP055184093, ISSN: 0021-9258, DOI: 10.1074/jbc.M112.382499</td>
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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 application or patent but published on or after the international filing date

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"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

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"Z" document member of the same patent family

Date of the actual completion of the international search

20 April 2015

Date of mailing of the international search report

29/04/2015

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

Gresinger, Iri na
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<td>Figure 2, 4 &amp; Karen T. Smith: &quot;Human Family with Sequence Similarity 60 Member A (FAM60A) Protei n&quot;, Molecular and Cellular Proteomics, 14 September 2012 (2012-09-14), pages SL-SII, XP055184114, Retrieved from the Internet: URL: <a href="http://www.mcponline.org/content/suppl/2012/09/14/M112.020255.DCl/mcp.M112.020255-1.doc">http://www.mcponline.org/content/suppl/2012/09/14/M112.020255.DCl/mcp.M112.020255-1.doc</a> [retrieved on 2015-04-20]</td>
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<td>H Kobayashi ET AL: &quot;Fluorescence in situ hybridization on mapping of translocations and deletions involving the short arm of human chromosome 12 in malignant hematologic diseases&quot;, Blood, 15 November 1994 (1994-11-15), pages 3473-3482, XP055173050, UNITED STATES Retrieved from the Internet: URL: <a href="http://bl">http://bl</a> oodjournal .hemat olog ylibrary .org/cgi/content/abstract/84/10/3473</td>
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