CRISPR/Cas12a mediated genome editing to introduce amino acid substitutions into the mechanosensitive channel MscCG of *Corynebacterium glutamicum*

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ABSTRACT

Against the background of a growing demand for the implementation of environmentally friendly production processes, microorganisms are engineered for the large-scale biosynthesis of chemicals, fuels or food and feed additives from sustainable resources. Since strain development is expensive and time-consuming, continuous improvement of molecular tools for the genetic modification of the microbial production hosts is absolutely vital. Recently, the CRISPR/Cas12a technology for engineering of Corynebacterium glutamicum as important platform organism for industrial amino acid production has been introduced. Here, this system was advanced by designing an easy-to-construct crRNA delivery vector using simple oligonucleotides. In combination with a C. glutamicum strain engineered for the chromosomal expression of the β-galactosidase-encoding lacZ gene, this new plasmid was used to investigate CRISPR/Cas12a targeting and editing at various positions relative to the PAM site. Finally, we used this system to perform codon saturation mutagenesis at critical positions in the mechanosensitive channel MscCG to identify new gain-of-function mutations for increased L-glutamate export. The mutations obtained can be explained by particular demands of the channel on its immediate lipid environment to allow L-glutamate efflux.

KEYWORDS: CRISPR/Cas, recombineering, L-glutamate, Corynebacterium glutamicum, metabolic engineering, genome editing
Clustered regularly interspaced short palindromic repeats (CRISPR) together with CRISPR-associated (Cas) proteins constitute an immune system in bacteria and archaea that cut foreign DNA entering the cell\(^1\). DNA degradation is achieved by DNA cleavage at specific positions via the Cas-encoded nuclease activity, which is guided to the target site by a short CRISPR-RNA (crRNA) complementary to one of the strands of the target DNA. Simplicity and programmability of the CRISPR/Cas system, established for a broad range of different organisms, has enabled its straightforward application to specifically target any genomic location. Its applicability for engineering bacteria was first demonstrated in the context of recombineering in *Escherichia coli* where 65% of recombinant cells could be recovered by specifically killing of non-mutated cells\(^2\). Since then, this approach has been successfully used for manipulating numerous bacterial species, including amongst others *Streptococcus pneumoniae\(^2\), Lactobacillus reuteri\(^3\), Clostridium beijerinckii\(^4\), and Streptomyces coelicolor\(^5\) and more recently also *Corynebacterium glutamicum*\(^6\).\(^7\).

While Cas9 is currently the best-characterized and most widely used endonuclease among available CRISPR editing methods, Cas12a (previously named Cpf1) has recently emerged as an alternative for Cas9. This can be reasoned by the cytotoxicity of Cas9 for a number of bacteria\(^6,8\)-\(^10\), possible reasons could be the generation of blunt ends by Cas9 and lack or reduced activity of non-homologous end-joining (NHEJ) in bacteria\(^11\). In contrast, Cas12a generates staggered cut DNA with 5'-overhangs, which can be easier repaired by prokaryotes. Recently, Cas12a found additional applications in CRISPR genome editing methods for bacteria, such as *E. coli*\(^12\), *Mycobacterium smegmatis*\(^12\), *Cyanobacteria*\(^13\), *Pseudomonas putida*\(^14\) and *Clostridium difficile*\(^15\).

The first developed CRISPR/Cas method for bacteria taking advantage of Cas12a was developed for *C. glutamicum*, directly combined with single-stranded DNA (ssDNA) recombineering to relieve allosteric feedback inhibition of the γ-glutamyl kinase by L-proline\(^16\). *C. glutamicum*, generally recognized as safe (GRAS), is used for decades in industry for the million ton-scale production of monosodium L-glutamate\(^17\).\(^18\). Despite intensive efforts to understand L-glutamate synthesis in this organism in detail, it was only recently possible to
solve the mystery of L-glutamate export by identifying the responsible mechanosensitive channel MscCG^{19}. In addition, *C. glutamicum* has also been engineered for the production of organic acids^{20}, alcohols^{21,22}, polyphenols^{23,24}, and speciality chemicals^{25}.

Methodical milestones for the engineering of *C. glutamicum* include the availability of DNA transfer methods^{26}, a two-step homologous recombination method (the “sacB-system”)^{27} and, more recently, the introduction of RecT-mediated recombineering^{28}. The sacB-system, based on the conditionally lethal levansucrase-activity (SacB), is currently the method of choice for the site-specific introduction of genomic mutations. In principle, RecT-mediated recombineering with oligonucleotides (ssDNA) represents a true and less laborious alternative to the sacB-system, but applicability of this method requires a selection step since the frequency of obtaining recombinants in *C. glutamicum* is about 10^{-3}^{28}. Here, biosensors, enabling direct screening for mutated cells with increased metabolite formation via fluorescence-activated cell sorting (FACS) were successfully used to isolate recombineered variants^{29–32}. However, depending on the application in question, development of a biosensor-based screen is not an option. In such cases, CRISPR/Cas12a mediated killing of non-recombineered *C. glutamicum* cells is another option for quickly identifying variants with the desired genotype by selection, which holds the potential to decrease the experimental effort even further.

In this manuscript, we characterized the CRISPR/Cas12a system with regard to the introduction of point mutations into the genome of *C. glutamicum* and developed a simplified CRISPR/Cas12a protocol. Furthermore, we used this system to identify new gain-of-function mutations in the mechanosensitive channel MscCG leading to L-glutamate efflux.

**MATERIAL AND METHODS**

**Plasmid constructions.** In the used CRISPR/Cas12a system^{7}, vector pJYS2_crtYf is applied for crRNA delivery. Using this vector as a template, plasmid pJYScr was constructed comprising a *BamH*I/*BstBI* cloning site which facilitates the insertion of protospacer sequences into the delivery vector. Removal of the original crtYf protospacer sequence was
performed by DNA restriction using BamHI and Dral. The respective restriction sites for these enzymes are located at position 3196 and 3244. Since the available vector pJYS2_crtYf contains three additional Dral sites\textsuperscript{16}, the selected Dral restriction site was not suitable for cloning. However, to make this site available for cloning, the restriction site was replaced by a BstB1 site using a synthesized 313 bp XhoI-AflII fragment (Thermo Fisher Scientific, Waltham, MA, USA), comprising the parental sequence but replacing two nucleotides in the Dral site to create a BstB1 restriction site. For plasmid construction, the delivery vector pJYS2_crtYf was cut with XhoI/AflII. Subsequently, the resulting plasmid backbone fragment was purified and combined with the synthesized fragment. Since the vector backbone contains an additional BstB1 restriction site, site-directed mutagenesis was performed to introduce a C1165T mutation thereby eliminating this restriction site. The QuikChange II XL Site-Directed Mutagenesis Kit (Agilent, Santa Clara, CA, USA) was used for this purpose. However, we experienced that even after gel purification of vector DNA treated with BamHI/BstB1 still intact vector was obtained after ligation with oligonucleotides. Therefore, the 49 bp BamHI/BstB1 fragment was replaced by a 931 bp dummy fragment (nt 379847-383111 of BA000036) yielding vector pJYScr. This vector makes it easy to separate individual DNA fragments after BamHI/BstB1 restriction for further processing. The DNA sequence of pJYScr and the vector itself are deposited at Addgene (#132719). As control experiment, pJYScr:crtYf bearing the same crRNA as pJYS2_crtYf was constructed to evaluate, whether the replacement of restriction sites has an effect on the editing efficiency in comparison to pJYS2_crtYf. In these experiments, 21±7 cfu were obtained with the pJYScr:crtYf delivery vector and with the original vector pJYS2_crtYf 34±10 cfu as escaper cells could be counted. This indicated that the vector backbone modifications did not affect the formation of an active Cas12a/crRNA complex in C. glutamicum cells.

For construction of the individual crRNA delivery vectors, annealed oligonucleotides encoding loop region and spacer were used. For annealing, 10 µl of both oligonucleotides (200 nmol/ml H\textsubscript{2}O) were mixed, incubated for 10 minutes at 98°C, at room temperature for one hour, and subsequently stored on ice. The annealed oligonucleotides were then ligated.
with the 4332 bp BamHI/BstBI vector fragment of pJYScr prepared via gel purification. Then *E. coli* DH5α was transformed with the ligation mix. Spectinomycin resistant clones indicating a successful transformation event were identified on LB plates containing the antibiotic at a concentration of 100 mg/L. Oligonucleotides pJYS2-fw2 and pJYS2-rv2 to amplify a 995 bp PCR fragment and DNA sequencing were used to verify the presence the crRNA coding region in the vector. If desired, recombinant strains were cured from the pJYScr plasmids by cultivation at 34 °C without supplementation of antibiotics as described earlier7.

**Strain constructions.** *C. glutamicum* ATCC13032 was used throughout this study. Integration of the lacZ gene under control of the synthetic PH36 promoter was achieved by isolating a 3190 bp SmaI-fragment from the plasmid pMK-RQ_PH36_LacZ synthesized by invitrogen (Thermo Fisher Scientific, Waltham, MA, USA). This fragment was inserted into the pK18mobsacB vector carrying the two flanking regions of the chromosomal none-coding region at nucleotide 1.404.651 (Acc.-no. NC_006958) as previously described33. Using two homologous recombination events strain *C. glutamicum* WT::lacZ was obtained. DNA sequencing confirmed a correct chromosomal integration. *C. glutamicum* WT::lacZ colonies plated onto BHIS-X-gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) turned blue after three days, whereas the *C. glutamicum* wild type cells remained yellowish. The two constructed mscCG-mutants devoid of the carboxyterminal end were constructed by homologous recombination using pK19mobsacB-420stopΔ and pK19mobsacB-424stopΔ (Supplementary table S1), each carrying two flanking fragments of ~ 500 bp34.

**Transformation and recombination.** Preparation of electrocompetent *C. glutamicum* pJYScr cells providing inherent expression of RecT and Cas12a was performed as described previously with only minor modifications16. In short, a single colony from a BHI agar plate was used to inoculate a pre-culture of 4 mL BHISG-Kan and incubated overnight at 30°C and 220 rpm. Subsequently, 500 μl of the preculture was transferred into 50 ml of BHISG-Kan supplemented with 1 ml/L Tween 80 and 4 g/L glycine. The culture was harvested when the optical density at 600 nm (OD600) reached approx. 1. Cells were chilled on ice for 20 min and centrifuged at 2,600 g and 4 °C for 10 min. Subsequently, two washing steps in 50 ml of 10%
glycerol were performed. Competent cells were resuspended in 500 μl 10% glycerol, and aliquots of 90 μl were stored at −80 °C. For transformation, an aliquot was thawed on ice, mixed with 0.5 μg of the pJYScr derivative and 10 μg of PAGE purified oligonucleotide (Eurofins MWG Operon (Ebersberg, Germany), and transferred into 4°C pre-cooled electroporation cuvettes (width 2 mm). The oligonucleotides used were 59 bp in size and encoded a mutated PAM site, spacer region, and, if applicable, the mutation to be introduced. After electroporation, cells were immediately transferred to 900 μl of prewarmed BHISG medium and heat-shocked for 6 min at 46 °C. The cells were incubated to recover for 2 h at 30 °C with shaking at 170 rpm. Cells were plated on BHISG containing kanamycin and spectinomycin (BHISG-Kan-Spc-Xgal), and incubated for 3-4 days to determine the number of colony forming units and their respective color.

**Media and L-glutamate formation.** *E. coli* was grown on LB and *C. glutamicum* on BHI (Difco Laboratories, Detroit, USA). BHIS contained in addition 90 g/L sorbitol, BHISG moreover 10 g/L glucose, BHISG-Kan 15 mg/L kanamycin, BHISG-Kan-Spec 100 mg/L spectinomycin, and BHISG-Kan-Spc-Xgal 50 mg/L. For glutamate formation precultures were grown overnight in 20 mL BHI. Cells were harvested by centrifugation, washed with 20 mL 0.9% NaCl and inoculated to an OD_600 of in defined CGXII medium (Eggeling and Bott 2005) containing 50 μg/L biotin. L-Glutamate in supernatants was determined as its o-phthalaldialdehyde derivative via HPLC and fluorescence detection as previously described.

**RESULTS**

**Simple construction of crRNA delivery vectors using oligonucleotides.**

According to the protocol currently used, amplification of the entire pJYS2_crtYf vector is necessary for inserting the desired crRNA-encoding spacers. Despite different PCR conditions and the use of megaprimers, this procedure resulted in a low yield and often a doubling of the oligonucleotides. With the original protocol we were able to construct only 1 of 5 desired vectors, which motivated us to develop an improved crRNA delivery vector expediting necessary constructions. The starting vector pJYS2_crtYf encodes the loop region
of the crRNA in combination with the specific crtYf-targeting sequence on a 49 bp BamHI-DraI-fragment. The single BamHI site is located between the strong promoter pJ23119 and the specific spacer, whereas one of the four DraI sites of the vector is located between the spacer and the T1rrnB terminator. Using a synthesized fragment and applying site-directed mutagenesis, we replaced the DraI site by a BstBI site and removed an existing BstBI site in the vector backbone. The resulting vector in principle allows replacing the crtYf-specific crRNA-encoding sequence on the 49 bp BamHI-BstBI fragment by any other sequence.

However, we substituted the crtYf-fragment by a larger fragment of 931 bp resulting in the vector pJYScr of 5.252 kb, which enables easy preparation of the 4.321 kb BamHI-BstBI vector fragment for the insertion of oligonucleotides encoding the desired crRNA (Figure S1). This new vector for genome editing has been made available through Addgene (#132719).

To validate the functionality of pJYScr in a CRISPR/Cas12a selection experiment, a crRNA targeting murE of C. glutamicum was cloned into the vector. The murE gene encodes the essential UDP-N-acetylmuramoylalanyl-D-glutamate 2,6-diaminopimelate ligase\textsuperscript{36}. The corresponding oligonucleotides comprising the PAM site 5´-GTTG-3´ (nt 255-258) within murE, were annealed and ligated with the 4.321 kb BamHI-BstBI fragment of pJYScr, resulting in pJYScr:murE. After transformation of C. glutamicum pJYS1-petFU (providing constitutive Cas12a-activity) with 1 µg of the pJYScr:murE plasmid, only 81±26 cfu as escaper cells were obtained demonstrating the formation of an active Cas12a/crRNA complex in the cells. A control experiment, in which C. glutamicum wild type cells without pJYS1-petFU were transformed with the same vector yielded 3.2x10\textsuperscript{5} cfu.

**Efficiency of CRISPR/Cas12a-assisted ssDNA-recombineering in C. glutamicum.**

With the aim to explore the capabilities of combining ssDNA-recombineering with CRISPR/Cas12a-mediated elimination of non-recombinant cells, we first constructed a C. glutamicum test strain carrying a chromosomal integration of the lacZ gene encoding for the β-galactosidase from *E. coli* (C. glutamicum WT::lacZ). In this particular strain, lacZ-expression is under control of the synthetic promoter pH36 \textsuperscript{37}, enabling easy identification of generated lacZ mutants on agar plates supplemented with 5-bromo-4-chloro-3-indolyl-β-D-
galactopyranoside (X-Gal) by blue-white screening. Subsequently, this strain was transformed with pJYS1-petFU providing constitutive Cas12a- as well as RecT-activity to this strain for the following recombineering-CRISPR/Cas12a experiments\textsuperscript{16}. When transformed with the empty pJYScr-vector with or without the recombineering oligonucleotide o-1574a, high transformation rates comparable to that with the standard pEKEx3 vector as a control were obtained (Figure 1). A high transformation rate is essential to obtain recombinants in the combined recombineering-CRISPR/Cas12a-killing approach. Noteworthy, the transformation rates obtained also confirmed the non-toxicity of Cas12a opposed to Cas9 as it has been already described for various organisms\textsuperscript{13} including \textit{C. glutamicum}\textsuperscript{16}. As crRNA delivery vector targeting the chromosomal \textit{lacZ} gene, we have built pJYScr-lacZ-1574 using the intragenic PAM site 5´-(T)TT(C)-3´ with a guide sequence of 21 nts. When \textit{C. glutamicum} WT::lacZ pJYS1-petFU was transformed with this vector, only 51±21 cells survived underlining the efficiency of Cas12a targeting in \textit{C. glutamicum} (Figure 1). To assay for DNA-editing plus targeting of unmutated cells, \textit{C. glutamicum} WT::lacZ pJYS1-petFU was transformed with pJYScr-lacZ-1574 in combination with the editing oligonucleotide o-1574a (all recombinogenic oligos are listed in supplementary Table S2). This oligonucleotide was designed to replace the PAM site and the adjacent codon TCG (5´-TTTCG-3´) by 5´-TATAG-3´, which results in an inactivated PAM site and the introduction of a stop codon within the open reading frame of \textit{lacZ}. This experiment yielded 1,500 ± 172 colonies of which more than 97% were white (Figure 1). Comparable numbers could be obtained when targeting a second PAM-site (5´-(T)TT(A)-3´) within \textit{lacZ} using the oligonucleotide o-1654a and the simultaneous introduction of a stop codon by substituting a single nucleotide (Figure 1).
Figure 1: Efficiency of genome targeting and editing in C. glutamicum. The transformation assays to transform C. glutamicum WT::lacZ pJYS1-petFU contained 500 ng plasmid and 10 µg oligonucleotide where indicated. The transformants were plated on X-Gal plates for blue-white screening. The results are averages from at least two independent experiments, and error bars depict standard deviations. In control experiments, C. glutamicum was either transformed with the cloning vector pEKEx3 (column 1), the empty vector pJYScr (column 2) inscribed as pcr, or empty vector plus oligo o-1574a targeting a PAM site in the chromosomal lacZ gene (column 3). When the crRNA was provided by vector pJYScr-lacZ-1574 (pcr-1574, column 4) or vector pJYScr_lacZ-1654 (pcr-1654, column 6), only few cells survived, whereas addition of the corresponding oligonucleotides (o-1574a, column 5 or o-1654a, column 7) for PAM site-inactivation and introduction of a stop codon resulted in a drastic increase of surviving colonies with >96% of having the desired mutation.

These results raised the question at which frequency a reliable introduction of mutations at different positions relative to the PAM site is possible with this system. To answer this, nine different oligonucleotides with a length of 59 nt were designed for introducing a stop codon over a stretch of 18 nt along with a mutation inactivating the PAM site. Oligonucleotides used for recombineering and introduction of the desired mutations in
the lacZ gene and the PAM site are given in Table S2. Experiments conducted with these oligonucleotides yielded no white colonies when oligonucleotide o-1574b was used, which would introduce two nucleotides substitutions upstream of the PAM-site (Figure 2). In this case, sequencing of eight randomly picked blue clones revealed indeed that only the PAM site was mutated. When performing recombineering-CRISPR/Cas12a with the editing oligonucleotides o-1574c, -d, -a, e, and –f, more than 96% of the surviving C. glutamicum variants were white in all cases indicating a successful introduction of the stop codon into the lacZ gene. This observation was supported by DNA sequencing of randomly selected clones, which confirmed correct incorporation of both mutations in all clones analyzed. However, when using the recombinogenic oligonucleotides o-1574g, o-1574h or o-1574i designed for introducing a stop codon further downstream of the PAM site, introduction of the desired mutations was still possible, but only at a reduced frequency ranging from 75% to 90% (blue bar), depending on the number of substitutions necessary to generate a stop codon. Interestingly, DNA-sequencing revealed that in a few cases only the stop codon was successfully introduced whereas the PAM site was still intact (given in Fig. 2 as “stop only”). We obtained a similar result of such partially successful editing events in experiments targeting a second PAM site in lacZ (Table S2). In these cases, possibly only a short sequence of the genomic DNA was exchanged for the respective oligonucleotide during recombineering. These clones escape Cas12a cleavage when only mismatches in the PAM-distal region of the DNA target are present. Collectively, these data suggest that site-directed mutagenesis using the combined recombineering/Cas12a selection strategy is possible over a stretch of four nucleotides upstream and ten nucleotides downstream of the PAM sequence.
Figure 2: Genome editing efficiency relative to the PAM site in *C. glutamicum* using the combined recombineering-CRISPR/Cas12a selection method. Shown is a portion of the chromosome-integrated *lacZ* gene with the PAM sequence highlighted in blue and the protospacer highlighted in yellow. The Cas12a-cleavage site yielding staggered ends is indicated by red triangles. The relevant DNA sequence of the oligonucleotides for site-directed mutagenesis is shown at the bottom with all mutations highlighted in red. The fraction of white colonies obtained, is given as percentage share. From each assay, the *lacZ* gene of randomly selected white clones was sequenced (n=4-16). Either, the PAM site was mutated and the stop codon was introduced (blue bars), only the PAM site was mutated (red bars), or only the stop codon was introduced (green bars). The respective numbers for each type of mutation (PAM+stop/stop only/PAM only) assessed via DNA sequencing, are also given as numbers below the colored bars. Detailed DNA sequencing results are given in supplementary table S1.

Inspired by these results, the mutagenic oligonucleotide o-1574fe-random was designed, which carries a continuous stretch of 18 random nucleotides spanning the previously targeted PAM sequence plus five nucleotides upstream and ten nucleotides...
downstream of that PAM site (Supplementary Table S3). After performing the recombineering-CRISPR/Cas12a selection, only eight white colonies were obtained. DNA sequencing of the lacZ gene of all clones revealed a mutated and thus non-functional PAM sequence in all cases with four clones carrying additional mutation(s) downstream of the targeted position (nucleotides +11 to +16, Supplementary Table S3). Apparently, when considering the low number of obtained colonies one can assume, that with the given oligonucleotide bearing such a long random sequence, recombineering does not work. Hence, this method cannot be used to exchange a longer stretch of DNA and is limited to the introduction of three consecutive nucleotide substitutions into the genome of C. glutamicum.

This conclusion is supported by the results of two additionally performed experiments using oligonucleotides o-1574e-mix and o-1574g-mix, which were both designed for the saturation mutagenesis of a selected codon (Supplementary Table S3). 24 white colonies obtained by recombineering-CRISPR/Cas12a selection using o-1574e-mix were randomly picked and sequenced. All variants were characterized by a mutated PAM site and 16 of them carried additional mutation(s) in the desired codon leading to twelve amino acid substitutions on the protein level without a discernable preference for specific substitutions (Supplementary Table S3). Similarly, targeted DNA sequencing of 24 white clones resulting from the genome editing experiment using oligonucleotide o-1574g-mix yielded 13 variants bearing mutations in the desired codon, leading to eight different amino acids at this particular position (Supplementary Table S3).

**Site-directed mutagenesis targeting mscCG in C. glutamicum**

An efficient L-glutamate export is an important prerequisite for achieving high L-glutamate titers with C. glutamicum. In case of this bacterium, the mechanosensitive channel MscCG (533 aa) is mainly responsible for L-glutamate efflux\(^{19}\). However, L-glutamate efflux only occurs in response to changes in the structure of this channel, induced either by growth under biotin limitation or addition of penicillin, ethambutol or fatty acid ester surfactants to the medium\(^{40}\). In addition, an altered lipid composition may cause efflux\(^{41,42}\). According to the current model, these cultivation/production conditions mentioned increase membrane tension...
and thus activate the MscCG channel\textsuperscript{43}. Interestingly, several gain-of-function (GOF) variants of MscCG are known causing constitutive L-glutamate efflux\textsuperscript{19,44}. These mutational hot spots are located in the pore-lining helix of the N-terminal domain (1-286 aa), which has been also shown to be responsible for L-glutamate excretion in response to the above mentioned treatments\textsuperscript{44}. Additionally, two additional GOF mutants, characterized by mutations in the C-terminal extracytoplasmic domain are known. In one case, insertion of an IS-element at position 419 was described, resulting in the introduction of a stop codon at position 424\textsuperscript{19}. The other GOF mutant bears an P424L-substitution\textsuperscript{19}. Finally, a third mutant devoid of the entire cytoplasmic domain (amino acids 420-533) has been described\textsuperscript{44}. We were interested in studying consequences of mutations in this particular region at positions 422 and 423 of MscCG and used the recombineering-CRISPR/Cas12a method to assay additional amino acid substitutions at these particular positions. Both targeted amino acid positions of MscCG are predicted to be located at the transition of the fourth transmembrane helix to the periplasmic space\textsuperscript{19}. Presumably, this site determines the overall structure of MscCG and is crucial for the interaction of the channel protein with other components of the cell wall.

\textit{C. glutamicum} pJYS1petFU was transformed with pJYScr\_mscCG1269c providing the crRNA and with 10 \( \mu \)g of the mutagenic oligo o-mscCG1269-V422nnn for site saturation mutagenesis (Table S4). The assay yielded 43 clones of which 16 were randomly choosen and sequenced. In these clones, altogether seven different amino acid substitutions at amino acid position V422 of MscCG could be identified. Similarly, position E423 was also targeted, eventually yielding 27 clones surviving CRISPR/Cas12a selection after recombineering. In 19 clones, DNA sequencing revealed introduction of eight different amino acids and one stop codon. In addition, one variant with an undesired deletion of two nucleotides in the PAM sequence was found. This frameshift mutation inevitably resulted in a stop codon (E423AQstop). Recombinant strains were cured from the respective plasmids and subsequently assayed for L-glutamate accumulation in the culture supernatant.

As previous studies have shown, L-glutamate formation, even in described MscCG mutants, is very much dependent on biotin concentration, medium composition and sugar
concentration\textsuperscript{19,44,45}. For instance, the known MscCG mutant devoid of the extracytoplasmic domain\textsuperscript{44} excretes L-glutamate only at low biotin concentrations, whereas the mutant carrying the IS-element insertion at position 419 accumulates L-glutamate even in the presence of excess biotin\textsuperscript{19}. For assaying the L-glutamate accumulation capabilities of the newly constructed MscCG-variants obtained by recombineering-CRISPR/Cas12a selection, we chose stringent conditions in defined medium with biotin excess (50 µg/L) in combination with a low glucose concentration (40g/L). Under these conditions, the \textit{C. glutamicum} wild type does not excrete L-glutamate (Figure 3a). The MscCG variants V422A, V422C, V422T, V422L, and V422S did not accumulate L-glutamate. In contrast, in the supernatant of the MscCG-variants V422K and V422D, in which the valine residue was substituted by charged amino acids, a substantial L-glutamate accumulation of up to 13 mM could be detected (Figure 3a). However, variants E423K and E423D did not show L-glutamate accumulation, as did mutants E423N, E423T, E423A, and E423H (Figure 3b). In contrast, mutant E423P, E423S as well as the strain with the MscCG variant E423stop promoted L-glutamate export. Noteworthy, L-glutamate concentrations decreased with time due to reuptake and utilization of this amino acid as known from previous studies\textsuperscript{45}.

Interestingly, the MscCG variant E423AQstop did not cause L-glutamate accumulation as expected (data not shown). We also used the conventional sacB-system to delete two different portions of the extracytoplasmic domain of MscCG. In mutant M420stopΔ methionine in position 420 was replaced by a stop codon and the carboxyterminal end of 113 amino acids deleted. Similarly, the \textit{C. glutamicum} strain carrying the shortened MscCG variant P424stopΔ missing the carboxyterminal 109 amino acids was constructed. With variant M420stopΔ, L-glutamate concentrations of up to 27 mM could be detected, and with mutant P424stopΔ concentrations of up to 130 mM could be determined in the supernatant (Supplementary Table S5). These observations are in line with previous data also implying that presence of the carboxyterminal domain has a negative impact on L-glutamate channeling activity\textsuperscript{44}.
Figure 3: L-Glutamate accumulation in the supernatant of *C. glutamicum* variants with different mutations in the mechanosensitive channel MscCG gene at position V422 (top) and E423 (bottom). In addition, MscCG variants with stop codons at positions M420 (M420stop\(\Delta\)) and E423 (E423stop), respectively were also characterized with regard to the accumulation of L-glutamate. L-Glutamate concentrations for each variant were determined after 14, 18, 20 and 22 hours.

**DISCUSSION**

CRISPR/Cas selection is a powerful technology to perform genome editing in living cells. Based on this system, methods for inhibiting transcription (CRISPRi), activation of gene expression, or modification of nucleobases have been developed\(^{46}\). In bacteria, for the most part, methods introducing double-strand breaks using the crRNA/Cas-complex to eliminate
unaltered strain variants find an application. The main challenge during combined recombineering-selection experiments using RecT-recombineering and CRISPR/Cas represents balancing activities of all individual components including the recombinase, the crRNA and the endonuclease involved. The need to ensure the availability of the respective components in appropriate quantities at the right time initially led to the development CRISPR/Cas methods for *E. coli*, which require up to three vectors or three independent induction systems. However, these established Cas9-based systems were not suitable for applications in *C. glutamicum*, for the most part due to the cytotoxicity of the Cas9 endonuclease.

However, a CRISPR/Cas genome editing system based on the Cas12a endonuclease originating from *Francisella novicida* was introduced for *C. glutamicum*. This genome editing method is more suitable for this bacterium as (i) Cas12a is not toxic for *C. glutamicum*; (ii) induction of gene expression (to establish endonuclease or recombinase activity and crRNA formation) is not necessary due to a fortunate choice of constitutive promoters; and (iii) the crRNA, 43 nt in length, is about half the size of a typical Cas9 crDNA. However, the construction of the crRNA delivery vector for a distinct protospacer sequence remained laborious and time-consuming, lowering the overall applicability of this method. The reconstructed vector pJYScr presented here, overcomes this bottleneck as it allows for fast and simple vector assembly to enable crRNA delivery. With CRISPPathBrick, a similar strategy was developed by the Koffas group, which allows for the modular assembly of CRISPR-bricks to be used for CRISPR-Cas9 applications in *E. coli*. In this study, CRISPR/Cas12a-assisted recombineering with ssDNA oligonucleotides using this new vector was assessed by performing experiments to introduce point mutations up- and downstream of PAM-sites in a chromosomally integrated *lacZ* gene. Interestingly, the results are consistent with previous experiments in *Mycobacterium smegmatis*, showing that any targeting distal from the PAM site results in a drastically decreased recombineering efficiency. Nonetheless, the results obtained in this study confirm the potential of CRISPR/Cas12a-assisted recombineering using ssDNA oligonucleotides for the efficient and
precise introduction of nucleotide substitutions into the genome of \textit{C. glutamicum}. This was demonstrated when performing site-directed mutagenesis of the gene encoding the MscCG channel of \textit{C. glutamicum}. We focused on positions 422 and 423 of the MscCG protein located at the transition of the fourth transmembrane helix to the periplasmic space\textsuperscript{19}. It is striking that only substitution by charged amino acids residues resulted in an open-channel conformation at position 422. Given that the hydrophobic head groups of cardiolipin and phosphatidylglycerol are present in \textit{C. glutamicum} in this region\textsuperscript{41}, the direct local membrane environment may be altered, which in turn creates the necessary conditions for the amino-terminal part of MscCG to mediate the L-glutamate efflux\textsuperscript{44}. In contrast, amino acid substitutions E423P and E423S, presumably causing drastic structural alterations, support L-glutamate efflux at position 422. Against the background that complete deletion of the periplasmic domain also fosters L-glutamate efflux\textsuperscript{44}, there are two obvious possible explanations. Again, the immediate channel environment could be affected, which could result in a deformed channel. Alternatively, the extracytoplasmic domain could be forced away from the channel opening, thus enabling the efflux of L-glutamate. Such models where a periplasmic domain functions as an efflux-controlling lid and where this lid further interacts with the extracellular matrix have been described for eukaryotes\textsuperscript{51}.

Taken together, the crRNA delivery vector pJYScr introduced here simplifies plasmid constructions and expands the number of CRISPR/Cas12a applications in \textit{C. glutamicum}. In particular in combination with recombineering, CRISPR/Cas12a is a powerful tool when performing site-directed mutagenesis in the genome of \textit{C. glutamicum} by selectively eliminating unedited cells as presented in this study.

**ASSOCIATED CONTENTS**

**SUPPORTING INFORMATION**

The supporting information is available free of charge on the ACS Publication website:

**Supplementary figure**

Figure S1: Plasmid map of the pJYScr vector
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Table S1: Plasmids and oligonucleotides used for plasmid constructions.

Table S2: Recombinogenic oligos and obtained mutations at two *lacZ* locations.

Table S3. Recombinogenic oligos for the insertion of a random stretch of nucleotides and codon saturation mutagenesis.

Table S4. Recombinogenic oligonucleotides used for introducing mutations in *mscCG* and the mutations obtained.

Table S5: L-Glutamate accumulation in the culture supernatant of constructed *mscCG* deletion mutants

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**Author Contributions**

L.E. designed the experiments, K.K and C.K.S. conducted the experiments. L.E. and J.M. wrote the manuscript.

**NOTES**

The authors declare no competing financial interest.
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CRISPR/Cas12a mediated genome editing to introduce amino acid substitutions into the mechanosensitive channel MscCG of Corynebacterium glutamicum

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Supplementary Figure S1: Plasmid map of the pJYScr vector constructed in this study for easy incorporation of crRNA encoding oligonucleotides. The 931 BamHI-BstBI-“dummy” fragment with the relevant restrictions sites for easy subcloning is highlighted.
Supplementary Table S1. Plasmids and oligonucleotides used for plasmid constructions.

| Plasmids | Characteristics | Source |
|----------|-----------------|--------|
| pK19mobsacB | Integrative vector, Km' mob sacB | (1) |
| pEKEx3 | Shuttle vector, Spc' Ptac lacI | (2) |
| pJYS2_crtYf | Shuttle vector based on pMB1, Spc', Pj23119-crRNA targeting crtYf | (3) |
| pJYS1 | Shuttle vector based on pBL1, Km', RecT, FnCpf1, identical with pJYS1-petFu | (3) |
| pMK-RQ_PH36_LacZ | Vector providing lacZ under constant high expression of PH36 | invitrogen Thermo Fisher Scientific |
| pMA-RQ_crRNA_crtYf | Vector providing a 313 bp fragment in which a DraI site of pJYS2_crtYf was replaced by a BstBI site | invitrogen Thermo Fisher Scientific |

| Oligonucleotides | Sequence (5´-3´) | Purpose |
|------------------|------------------|---------|
| cr:lacZ 1574f _fw | CTGTTGTAGATGCTACCTGGAGAGACGC GCCCATTCGAAATAAAAACGAAAAGGCTC | pJYScr-lacZ-1574f |
| cr:lacZ 1574f _rv | CGTTTTATTTCGAATGGGCGCTCTCT CAGTAGCATCTACAACAGTAGAAATTCC G | pJYScr-lacZ-1574f |
| cr:lacZ1656c _fw | CTGTTGTAGATGCGAAACCGCCAAGACT GTTAATTCGAAATAAAAACGAAAAGGCTC | pJYScr-lacZ-1654c |
| cr:lacZ1656c _rev | CGTTTTATTTCGAATTAACAGTCTTGGCG GTTTGCATCTACAACAGTAGAAATTCC G | pJYScr-lacZ-1654c |
| cr:murE-2 _BamHI | GATCCGAATTTCATCACTGTGTAGATAGC ACCTCAAGTCAGCTGCGTTT | pJYScr:murE |
| cr:murE-2 _BstBI | CGAA ACGCAGCTGGACTTGAGTGCTATCTAC AACAGTAGAAATTCC | pJYScr:murE |
| cr:mscCG_1269c _fw | GATCCGAATTTCATCACTGTGTAGATAC AGTCATGACCTTAAATAGTTT | pJYScr_mscCG1269c |
| cr:mscCG_1269c _rv | CGAAAC TATTTAAGGTCATGACTGTGAT CTACAACAGTAGAAATTCC | pJYScr_mscCG1269c |
| m420stop∆-1fwd | tgcatgcctgcaggtcgactGGCGGATCGACCAC GGCT | pK19mobsacB-420stop∆ |
| m420stop∆-2rev | cagcgtcctgaGACCTTAAATAGTGACAGCAA | pK19mobsacB- |
| m420stopΔ-3fwd   | atttaaggtcTAGGACGCTGATTACAGAC | pK19mobsacB-420stopΔ |
|-----------------|--------------------------------|----------------------|
| m420stopΔ-4rev  | ttgtaaaacgacggcagtgGTAGCCGCTTTCTTGAAC | pK19mobsacB-420stopΔ |
| p424stopΔ-1fwd  | cctgcaggtcgactctagagCCGGCAATCAATGGACTGG | pK19mobsacB-424stopΔ |
| p424stopΔ-2rev  | AATCAGCGTCTTTATTATTCCACAGTCATGACCTAAATAGTG | pK19mobsacB-424stopΔ |
| p424stopΔ-3fwd  | GGAATAATAGACGCTGATTACAGACGTG | pK19mobsacB-424stopΔ |
| p424stopΔ-4rev  | ttgtaaaacgacggcagtgGTAGCCGCTTTCTTGAAC | pK19mobsacB-424stopΔ |
| pJYS2_fw2       | GTGTCAGTGAAAGGCAGCATCC | Sequencing primer |
| pJYS2_rv2       | TCGCCACCTCTGACTTGAGCG | Sequencing primer |
**Supplementary Table S2.** Recombinogenic oligos and obtained mutations at two *lacZ* locations. Shown on top is the *lacZ* sequence targeted for mutations using recombineering in combination with CRISPR/Cas12. The PAM sequence is underlined. Used recombinogenic oligonucleotides (o-1574x or o-1654x) have a grey background with the nucleotides deviating from the *lacZ* sequence in red. The DNA sequencing results for individual clones are given below the oligonucleotide sequences.

| lacZ | CC ATC AAA AAA TGG CTT CTA CCT GCT GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
|------|-----------------------------------------------------------------------------|
| o-1574a | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 1 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 2 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 3 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 4 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 5 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 6 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 7 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 8 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 9 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 10 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 11 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 12 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 13 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 14 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 15 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 16 | CC ATC AAA AAA TGG CTA TAG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| o-1574b | CC ATC AAA TAG TGG CTA TCG CTA CCT GCT GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 1 | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 2 | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 3 | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 4 | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 5 | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 6 | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 7 | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 8 | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| o-1574c | CC ATC AAA AAA TAG CTA TCG CTA CCT GCT GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 1 | CC ATC AAA AAA TAG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 2 | CC ATC AAA AAA TAG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 3 | CC ATC AAA AAA TAG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 4 | CC ATC AAA AAA TAG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 5 | CC ATC AAA AAA TAG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 6 | CC ATC AAA AAA TAG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
| clone 7 | CC ATC AAA AAA TAG CTA TCG CTA CCT GGA GAG ACG CGG CGG CTG ATC CTT TGC GAA TAC |
clone 8  TAC GCC CAC GCG ATG GGT AAC AGT CTT GGC TTC TGA TAA TAC TGG CAG GCG TTT CG
o-1654cb  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGC TAA TAC TGG CAG GCG TTT CG
clone 1  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGC TAA TAC TGG CAG GCG TTT CG
clone 2  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGC TAA TAC TGG CAG GCG TTT CG
clone 3  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC GCT TAA TAC TGG CAG GCG TTT CG
clone 4  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGC TAA TAC TGG CAG GCG TTT CG
clone 5  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC GCT TAA TAC TGG CAG GCG TTT CG
clone 6  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC GCT TAA TAC TGG CAG GCG TTT CG
clone 7  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGC TAA TAC TGG CAG GCG TTT CG
clone 8  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC GCT TAA TAC TGG CAG GCG TTT CG

o-1654cc  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 1  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 2  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 3  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 4  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 5  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 6  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 7  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 8  TAC GCC CAC GCG ATG GGT AAC AGT CTT GCC TGA TTC GCT TAA TAC TGG CAG GCG TTT CG

o-1654cd  TAC GCC CAC GCG ATG GGT AAC AGT CTT TGA GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 1  TAC GCC CAC GCG ATG GGT AAC AGT CTT TGA GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 2  TAC GCC CAC GCG ATG GGT AAC AGT CTT TGA GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 3  TAC GCC CAC GCG ATG GGT AAC AGT CTT TGA GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 4  TAC GCC CAC GCG ATG GGT AAC AGT CTT TGA GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 5  TAC GCC CAC GCG ATG GGT AAC AGT CTT TGA GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 6  TAC GCC CAC GCG ATG GGT AAC AGT CTT TGA GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 7  TAC GCC CAC GCG ATG GGT AAC AGT CTT GGC GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
clone 8  TAC GCC CAC GCG ATG GGT AAC AGT CTT TGA GGT TTC GCT TAA TAC TGG CAG GCG TTT CG
**Supplementary Table S3.** Recombinogenic oligos for the insertion of a random stretch of nucleotides and codon saturation mutagenesis. On top always the lacZ sequence is shown, targeted by recombineering in combination with CRISPR/Cas12a. The PAM sequence is underlined. Used recombinogenic oligonucleotides have a grey background with the nucleotides deviating from the lacZ sequence in red. The DNA sequencing results for individual clones are given below the oligonucleotide sequences. With the oligonucleotide o-1574fe-random carrying a stretch of 18 random nucleotides, only few white colonies were obtained (see results section in the main text). This was also the case with an additional, independently synthesized oligonucleotide with identical sequence. Noteworthy, clones 7 and 8 obtained during experiments with this oligonucleotide appeared white, but do not carry any additional mutation near the PAM site. It is possible that these two clones carry additional unwanted mutations in the lacZ gene. However, the nucleotide sequence nt 1188 to nt 1902 of these two clones was not at variance from the wild type sequence. Results obtained during codon saturation mutagenesis (oligonucleotides o-1574e-mix and o-1574g-mix), the amino acid substitution obtained is indicated in addition to the nucleotide sequence.

|   | lacZ |   | o-lacZ_1574fe-random |   | clone 1 |   | clone 2 |   | clone 3 |   | clone 4 |   | clone 5 |   | clone 6 |   | clone 7 |   | clone 8 |
|---|------|---|----------------------|---|---------|---|---------|---|---------|---|---------|---|---------|---|---------|---|---------|---|---------|
|   |      |   |                      |   | CC ATC AAA AAA TGG CTA TCG TAA CCA CCT GGA GAG ACG CGG CCG CTG ATC TTT TGC GAA TAC |   | CC ATC AAA AAA TGG CTA TCG TCA CCT GGA GAG TAA CCG CCG CTG ATC TTT TGC GAA TAC |   | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG TAG CCG CCG CTG ATC TTT TGC GAA TAC |   | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG TGG CCG CCG CTG ATC TTT TGC GAA TAC |   | CC ATC AAA AAA TGG CTA TCG CTA CCT GGA GAG AGG CCA CCG CTG ATC TTT TGC GAA TAC |   | CC ATC AAA AAA TGG CTA TCG TCG TAA CCA CCT GGA GAG ACG CGG CCG CTG ATC TTT TGC GAA TAC |   |
|   | amino acid |   |                      |   | clone 1 |   | clone 2 |   | clone 3 |   | clone 4 |   | clone 5 |   | clone 6 |   | clone 7 |   | clone 8 |
|   |      |   |                      |   | L527A    | L527R    | L527N    | L527Y    | L527S    | L527I    | L527stop | L527T    | L527S    | L527G    | L527C    | L527Q    | L527A    |
| Clone 19 | CC ATC AAA AAA TGG CTA TG C GTG ATC CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 20 | CC ATC AAA AAA TGG CTA TG C CTG ATC CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 21 | CC ATC AAA AAA TGG CTA TG C CTA CTT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 22 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 23 | CC ATC AAA AAA TGG CTA TG C TAT CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 24 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |

| Clone 19 | L527I |
| Clone 20 | L527Y |

| Clone 1 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGC GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 2 | CC ATC AAA AAA TGG CTA TG C CTA CCT AAA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 3 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 4 | CC ATC AAA AAA TGG CTA TG C CTA CCT AGC GAG ACG CGG CCG CTG CTG ATC CTT TGC GAA TAC |
| Clone 5 | CC ATC AAA AAA TGG CTA TG C CTA CCT GAA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 6 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 7 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 8 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 9 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 10 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 11 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 12 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 13 | CC ATC AAA AAA TGG CTA TG C CTA CCT TTA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 14 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGC GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 15 | CC ATC AAA AAA TGG CTA TG C CTA CCT AGC GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 16 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 17 | CC ATC AAA AAA TGG CTA TG C CTA CCT TAG GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 18 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 19 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 20 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 21 | CC ATC AAA AAA TGG CTA TG C CTA CCT TGG GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 22 | CC ATC AAA AAA TGG CTA TG C CTA CCT TAA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 23 | CC ATC AAA AAA TGG CTA TG C CTA CCT GGA GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
| Clone 24 | CC ATC AAA AAA TGG CTA TG C CTA CCT TGG GAG ACG CGG CCG CTG ATC CTT TGC GAA TAC |
**Supplementary Table S4.** Recombinogenic oligonucleotides used for introducing mutations in *mscCG* and the mutations obtained. Shown on top is the targeted *mscCG* sequence with the PAM sequence underlined (PAM sequence is on the opposite strand). The recombinogenic oligonucleotides o-*mscCG1269*-V422nnn and o-*mscCG1xxx*-E423 used are highlighted by a grey background. The DNA sequencing results for all individual clones with nucleotides deviating from the wild type sequence are presented in red. In the right column the relevant amino acid sequences (amino acids 420 - 426) are shown.

| mscCG | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG GAA CCA AGT GAG AAT TGG CAA | - Met Thr Val Glu Pro Ser Glu - |
|-------|-------------------------------------------------------------------------------------------------|----------------------------------|
| o-*mscCG1269*-V422nnn | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT tTa GAG CCA AGT GAG AAT TGG CAA | - Met Thr Leu Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT tCT GAG CCA AGT GAG AAT TGG CAA | - Met Thr Ala Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT AAA GAG CCA AGT GAG AAT TGG CAA | - Met Thr Lys Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT TGC GAG CCA AGT GAG AAT TGG CAA | - Met Thr Cys Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT ACT GAG CCA AGT GAG AAT TGG CAA | - Met Thr Thr Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT ACT GAG CCA AGT GAG AAT TGG CAA | - Met Thr Thr Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT ACT GAG CCA AGT GAG AAT TGG CAA | - Met Thr Thr Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT ACT GAG CCA AGT GAG AAT TGG CAA | - Met Thr Thr Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT ACT GAG CCA AGT GAG AAT TGG CAA | - Met Thr Thr Glu Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT ACT GAG CCA AGT GAG AAT TGG CAA | - Met Thr Thr Glu Pro Ser Glu - |

| mscCG | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG GAA CCA AGT GAG AAT TGG CAA | - Met Thr Val Glu Pro Ser Glu - |
|-------|-------------------------------------------------------------------------------------------------|----------------------------------|
| o-*mscCG1269*-E423nnn | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG GAA CCA AGT GAG AAT TGG CAA | - Met Thr Val Pro Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG CCA AGT GAG AAT TGG CAA | - Met Thr Val Asn Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG AAC CCA AGT GAG AAT TGG CAA | - Met Thr Val Thr Pro Ser Glu - |
|       | CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG TCC CCA AGT GAG AAT TGG CAA | - Met Thr Val Ser Pro Ser Glu - |

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CG CTG CTC TTG CTG CTA CTA TTT AAG GTC ATG ACT GTG GAC CCA AGT GAG AAT TGG CAA - Met Thr Val Asp Pro Ser Glu -
CG CTG CTC TTG CTG CTA CTA TTT AAG GTC ATG ACT GTG TAA CCA AGT GAG AAT TGG CAA - Met Thr Val STOP
CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG TAA CCA AGT GAG AAT TGG CAA - Met Thr Val Pro Ser Glu -
CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG TCA CCA AGT GAG AAT TGG CAA - Met Thr Val Ser Pro Ser Glu -
CG CTG CTC TTG CTG TCA CTA TTT AAG GTC ATG ACT GTG TAA CCA AGT GAG AAT TGG CAA - Met Thr Val Ala Gln STOP
### Supplementary Table S5. L-Glutamate accumulation in the culture supernatant of constructed mscCG deletion mutants.

| C. glutamicum variant | Sampling time [h] and L-glutamate concentration [mM] |
|-----------------------|-----------------------------------------------------|
|                       | 14 h  | 16 h  | 18 h  | 20 h  | 22 h  | 24 h  |
| wild type             | 0     | 0     | 0     | 0     | 0     | 0     |
| M420stopΔ Clone 1     | 27.1   | 24    | 24.6  | 2.7   |       |       |
| M420stopΔ Clone 2     | 35.2   | 45.3  | 32.2  | 15.1  | 5.7   | 0.3   |
| P424stopΔ Clone 1     | 42.4   | 89.1  | 82.6  | 61.4  | 53.6  | 59.5  |
| P424stopΔ Clone 2     | 57.5   | 135.1 | 103.8 | 67.8  | 74.3  | 85.2  |
| P424stopΔ Clone 3     | 62.1   | 97    | 91.1  | 47.7  | 57.4  | 33    |

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