Codon-Optimized Fluorescent Proteins Designed for Expression in Low-GC Gram-Positive Bacteria

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Fluorescent proteins have wide applications in biology. However, not all of these proteins are properly expressed in bacteria, especially if the codon usage and genomic GC content of the host organism are not ideal for high expression. In this study, we analyzed the DNA sequences of multiple fluorescent protein genes with respect to codons and GC content and compared them to a low-GC gram-positive bacterium, Bacillus anthracis. We found high discrepancies for cyan fluorescent protein (CFP), yellow fluorescent protein (YFP), and the photoactivatable green fluorescent protein (PAGFP), but not GFP, with regard to GC content and codon usage. Concomitantly, when the proteins were expressed in B. anthracis, CFP- and YFP-derived fluorescence was undetectable microscopically, a phenomenon caused not by lack of gene transcription or degradation of the proteins but by lack of protein expression. To improve expression in bacteria with low genomic GC contents, we synthesized a codon-optimized gfp and constructed optimized photoactivatable pagfp, cfp, and yfp, which were in contrast to nonoptimized genes highly expressed in B. anthracis and in another low-GC gram-positive bacterium, Staphylococcus aureus. Using optimized GFP as a reporter, we were able to monitor the activity of the protective antigen promoter of B. anthracis and confirm its dependence on bicarbonate and regulators present on virulence plasmid pX01.

Fluorescent proteins (FPs) based on the Aequoria green FP (GFP) are widely used for elucidating molecular mechanisms in cells and bacteria. Today, a large number of different FPs are available from different groups of cnidaria, some of which exhibit distinct absorption and emission spectra, superior folding, and improved activation at higher temperatures (14, 38, 50). Unlike other fluorescent reporters, the chromophore in the Aequoria GFP is intrinsic to the primary protein structure and consequently, in addition to oxygen needed for the activation of the chromophore, does not require substrates or other cofactors to fluoresce (8, 17, 46). FPs have found wide applications not only in eukaryotes but also in prokaryotes, including reporter systems to monitor protein expression or promoter activity or for analysis of protein localization within the cell (48). To elucidate bacterial protein expression, it would be desirable to have multiple fluorescent markers available for expression in the target bacterium; Suel et al. (42), for example, analyzed the regulatory circuit of Bacillus subtilis competence using multiple FPs. However, not all FPs are expressed properly in all bacteria. To overcome these problems, Veening and colleagues were able to increase FP expression in B. subtilis by 20 to 70% by adding the first eight amino acids of ComGA to the N-terminal sequences of cyan FP (CFP) and yellow FP (YFP), speculating that the overcoming of a slow translation initiation caused by the eukaryotic codon bias led to this expression improvement (49). In addition, codon optimizations have been successfully used to increase expression of extrinsic proteins in different cells (3, 27). Using this method, unfavorable or rare codons in an extrinsic gene are exchanged in favor of more abundant ones without affecting amino acid sequences. This approach not only may lead to higher expression yields of recombinant proteins in bacteria and mammalian cells (3, 27, 34, 43, 51) but may also lead to fewer mistranslations, therefore improving the quality of the protein (22).

Gram-positive bacteria can be divided into two distinct groups: those with an overall high GC content, including Mycobacteria and Streptomyces, and those with low genomic GC content, such as Bacillus, Lactococccaeae, and Clostridia (28). Bacillus anthracis, the causative agent of anthrax, belongs to the latter group. We hypothesized that by replacing rare GC-rich codons with more abundant, AT-rich ones, we could improve gene expression of FPs in a low-genomic-GC-content bacterium such as B. anthracis. Our ultimate aims were the acquisition of a variety of colors for investigating molecular mechanisms in these prokaryotes and the improvement of FP expression. Codon optimizations by gene synthesis are widely offered and present a cost-effective way to increase recombinant protein yield.

Here we show that by exchanging amino acids of just one codon-optimized protein, we were able to obtain three FPs: YFPopt, CFPopt, and PAGFPopt, a photoactivatable GFP (31). All FPs, including the codon-optimized GFP, GFPopt, are highly expressed not only in B. anthracis but also in Staphylococcus aureus and can be used to analyze promoter activity. The corresponding genes have a low GC content and may be suitable for expression in other low-GC gram-positive bacteria.

MATERIALS AND METHODS

Bacterial strains, media, and growth conditions. All strains used in this study are listed in Table 1. Escherichia coli strains used for cloning purposes were grown in Luria-Bertani (LB) broth or plates. B. anthracis strain Ames 33, a
Template DNA was subsequently digested with DpnI, and the residual plasmids, PCR was performed as recommended, with an extension time of 6 min 45 s. /TechToolbox) and were used to amplify the entire pUC18-GFPopt plasmid.

A607C, C608A, and G609T. All oligonucleotides designated with “SDM” (Table pagfpopt

codon is also present in cfp

valine at position 2 of the protein was incorporated into C198G, A437T, T458C, A692T, and C693A (in addition, a GTT coding for a

esis, the following nucleotides were subjected to exchange: (i) for cfpopt

as a template, the Stratagene QuikChange Multi Kit was used as recommended by the manufacturer. During the mutagen-

Site-directed mutagenesis.

TABLE 1. Strains and plasmids used in this study

<table>
<thead>
<tr>
<th>Strains</th>
<th>Relevant characteristic(s)</th>
<th>Source or reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>E. coli</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TOP10</td>
<td></td>
<td>Invitrogen</td>
</tr>
<tr>
<td>SCS110</td>
<td></td>
<td>Stratagene</td>
</tr>
<tr>
<td>B. anthracis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ames 33</td>
<td>pXO1− pXO2−</td>
<td>32</td>
</tr>
<tr>
<td>Ames 35</td>
<td>pXO1− pXO2−</td>
<td>32</td>
</tr>
<tr>
<td>A33(pSW4-GFPmut1)</td>
<td>Ames 33 electroporated with pSW4-GFPmut1, Km'</td>
<td>This study</td>
</tr>
<tr>
<td>A33(pSW4-GFPopt)</td>
<td>Ames 33 electroporated with pSW4-GFPopt, Km'</td>
<td>This study</td>
</tr>
<tr>
<td>A33(pSW4-CEP)</td>
<td>Ames 33 electroporated with pSW4-CEP, Km'</td>
<td>This study</td>
</tr>
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<td>This study</td>
</tr>
<tr>
<td>S. aureus</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RN4220</td>
<td>Hemolysin-defective S. aureus strain, Sp'</td>
<td>13</td>
</tr>
<tr>
<td>RN(pTetONGFPopt)</td>
<td>RN4220 electroporated with pTetONGFPopt, Sp'</td>
<td>This study</td>
</tr>
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<td>RN(pTetONCFPopt)</td>
<td>RN4220 electroporated with pTetONCFPopt, Sp'</td>
<td>This study</td>
</tr>
<tr>
<td>RN(pTetONYFPopt)</td>
<td>RN4220 electroporated with pTetONYFPopt, Sp'</td>
<td>This study</td>
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</tbody>
</table>

Plasmids

pCR2.1 TOPO                Cloning vector, Km' Ap' Invitrogen
pUC18-GFPopt              Vector harboring custom codon-optimized GFPmut1, Ap' BlueHeron
pS10-CFP                  pCR2.1 harboring CFP under the control of S10 promoter of S. aureus This study
pS10-CFPopt               pCR2.1 harboring CFPopt under the control of S10 promoter of S. aureus This study
pSW4                      Gram-positive/gram-negative shuttle vector with PA promoter, Km' Ap' 32
pSW4-GFPmut1              pSW4 plasmid containing 729-bp AseI/BamHI-cloned gfpmut1 This study
pSW4-GFPopt               pSW4 plasmid containing 729-bp AseI/BamHI-cloned gfpopt This study
pSW4-CFP                  pSW4 plasmid containing 732-bp AseI/BamHI-cloned cfp This study
pSW4-CFPopt               pSW4 plasmid containing 732-bp AseI/BamHI-cloned cfpopt This study
pSW4-YFP                  pSW4 plasmid containing 732-bp AseI/BamHI-cloned yfp This study
pSW4-YFPopt               pSW4 plasmid containing 732-bp AseI/BamHI-cloned yfpopt This study
pYJ3335                   Vector with tetracycline-inducible Pnov promoter 20
pJR3S12                   Vector containing spectinomycin cassette 36
pTetON                    pYJ3335 derivative with inactive tetracycline repressor promoter, Ap' Sp' This study
pTetONGFPopt              pTetON containing 729-bp gfpopt cloned via SmaI/SbfI This study
pTetONCFPopt              pTetON containing 729-bp cfpopt cloned via SmaI/SbfI This study
pTetONYFPopt              pTetON containing 729-bp yfpopt cloned via SmaI/SbfI This study

Abbreviations: Apr, ampicillin resistance; Km', kanamycin resistance; Sp', spectinomycin resistance.

pXO1− pXO2− derivative of strain Ames 34 (32), and strain Ames 35 (pXO1− / pXO2− ) were grown in LB broth or NBY broth, containing 0.8% (wt/vol) nutrient broth, 0.3% yeast extract, 0.5% glucose, 1% fetal bovine serum (FBS), and 0.9% sodium bicarbonate. Bacteria were grown at 37°C and 225 rpm in either air or air supplemented with CO2 regulated at 5% (vol/vol). Sporulation plates contained 0.92% nutrient broth, 0.001% MnSO4, 0.001% KH2PO4, and 1.8% agar. S. aureus strain RN4220, a gift from Michael Otto, was grown in tryptic soy broth (TSB) at 37°C and 225 rpm or on tryptic soy agar. When required, the following antibiotics (Sigma) were added: ampicillin (100 μg/ml), kanamycin (20 μg/ml for B. anthracis and 100 μg/ml for S. aureus), and specti-

nomycin (100 μg/ml).

Site-directed mutagenesis. gfpopt was synthesized by oligonucleotide assembly performed by BlueHeron. For generation of cfpopt, yfpopt, and pagfpopt by site-directed mutagenesis using gfpopt as a template, the Stratagene QuikChange Multi Kit was used as recommended by the manufacturer. During the mutagen-

esis, the following nucleotides were subjected to exchange: (i) for cfpopt, A197G, C198G, A437T, T458C, A692T, and C693A (in addition, a GTT coding for a valine at position 2 of the protein was incorporated into cfpopt, because this codon is also present in cfp); (ii) for yfpopt, A193G, C194G, G202C, T214G, A607C, C608A, and G609T; and (iii) for pagfpopt, A192T, A193T, T488C, A607C, C608A, and G609T. All oligonucleotides designated with “SDM” (Table 2) were designed by using the Stratagene web page (http://stratagene.com/ /TechToolbox) and were used to amplify the entire pUC18-GFPopt plasmid. PCR was performed as recommended, with an extension time of 6 min 45 s. Template DNA was subsequently digested with DpnI, and the residual plasmids, pUC18-CFPopt, pUC18-YFPopt, and pUC18-PAGFPopt, were transformed into XL10-Gold (Stratagene). Positive transformants were identified by PCR and verified by sequencing using M13 primers.

Recombinant DNA techniques. Vectors pSW4-GFPmut1, pSW4-CFP, pSW4-CFPopt, pSW4-YFP, pSW4-YFPopt, and pSW4-PAGFPopt (Table 2) were constructed as follows. Fluorescent genes were amplified with proofreading Phusion polymerase (New England Biolabs) from pUC18 with oligonucleotide FPAscf and FPBamr for codon-optimized constructs, CFPf in combination with CFPfw for cfp, and YFPf in combination with YFPfw for yfp, incorporating a BamHI restriction site at the 3' end and an Asel restriction site at the 5' end. Templates for yfp and cfp amplification were the vectors pIYFP and pICFP, respectively (49), which we received from the Bacillus Genetic Stock Center. PCR fragments were subsequently cloned into TOPO pCR2.1 (Invitro-
gen), and positive TOP10 clones were sequenced with M13 primers to verify the fidelity of the insert. pSW4 was restricted with Ndel and BamHI, and FP genes were ligated into the vector via BamHI and Asel sites; the latter site has single-

strand extensions compatible with Ndel products. Positive XLA2Blue clones were verified by restriction analysis and sequencing, using oligonucleotide pSW4seq, which anneals in pSW4. To generate CFP fusions with the S10 promoter of S. aureus, the S10 promoter was amplified using oligonucleotide SI06/Asec in combination with SI0CFPrv with genomic DNA of strain RN4420 as a template. cfp was amplified with oligonucleotide CFPSOElw in combination with CFPSOElw, resulting in an overlap with the S10 promoter at the 5' end of the FP gene, as well as incorporation of a BamHI site at the 3' end. The S10 promoter and FP gene fragments
<table>
<thead>
<tr>
<th>Primer name</th>
<th>Sequence (5' to 3')</th>
<th>Restriction site</th>
<th>Purpose</th>
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<tr>
<td>FPAsefw</td>
<td>GGATTATGCTAAAAAGGAGAAGAAGATATTACAG</td>
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<td>Amplification of fluorescent protein genes</td>
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<td>FPBamrv</td>
<td>GGATCCCTATTATATAATTCAATCTATTTCCTGGT</td>
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</tr>
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<td>GFPAsfw</td>
<td>GGATTAAATGAGTGAAGAGAGAGAGAACTTCTTACCTG</td>
<td>AseI</td>
<td>Amplification of non-codon-optimized gfpopt1</td>
</tr>
<tr>
<td>GFPBamrv</td>
<td>GGATCCGTATTATAGATCTCTCTATTCTATGCC</td>
<td>BamHI</td>
<td></td>
</tr>
<tr>
<td>CFPAsefw</td>
<td>GGATTAAATGCTGAGCAAGAACGCGGA</td>
<td>AseI</td>
<td>Amplification of non-codon-optimized gfpopt</td>
</tr>
<tr>
<td>CFPBamrv</td>
<td>GGATCCCTATTATAGATCTCTCTATTCTATGCC</td>
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</tr>
<tr>
<td>YFPAsfw</td>
<td>GGATTAAATGCTGAGCAAGAACGCGGA</td>
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<td>Amplification of non-codon-optimized gfpopt</td>
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<tr>
<td>YFPBamrv</td>
<td>GGATCCCTATTATAGATCTCTCTATTCTATGCC</td>
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<tr>
<td>SDMCFP2</td>
<td>TATACAAAAAGGAGAAGCAATAATGTTTCAAAAGGAGAAGAATTATTTACAG</td>
<td>None</td>
<td>Site-directed mutagenesis of gfpopt to introduce valine-2 (CFPopt)</td>
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<tr>
<td>SDMCFPH-L</td>
<td>TTTGTAACAGCAGCAAGGAATTACACTAGGAATGGATGAATATTATAT</td>
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<td>SDMCFPSY-TW</td>
<td>GGCCCACACTTGTGACTCTTTTAAACTACGAGATACATGAGATTATTTTC</td>
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<td>Site-directed mutagenesis of gfpopt for exchanges S65T and Y66W (CFPopt)</td>
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<td>SDMCFPN-I</td>
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<td>SDMYFPSV-GL</td>
<td>ATGGCCCCACACTTGTGACACTCTTTAAGAGATACAGACTACAAATGTGTTT</td>
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<td>SDMYFPT-Y</td>
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<td>SDMPAGFPS-A</td>
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<td>pSW4Seq</td>
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<td>M13f</td>
<td>GTAAAGACGGGCCAGT</td>
<td>None</td>
<td>Sequencing oligonucleotide for verification of pCR2.1 inserts</td>
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<tr>
<td>M13r</td>
<td>AACAGCTATGACCAGT</td>
<td>None</td>
<td>Sequencing oligonucleotide for verification of pCR2.1 inserts</td>
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<tr>
<td>S10fwaSe</td>
<td>GGATTATATTTACACCCACCCTTTTACTGACTA</td>
<td>AseI</td>
<td>Forward primer to amplify S10 promoter from S. aureus</td>
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<td>S10CFPrv</td>
<td>CTTGCTACATATTTCCCTCTTATTCGCTTA</td>
<td>None</td>
<td>Reverse primer for S10 promoter with cfp overlap</td>
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<tr>
<td>S10CFPoptrv</td>
<td>CTTTGGAAACCATAATTCCCTCTCTATTCGCTTA</td>
<td>None</td>
<td>Reverse primer for S10 promoter with cfpopt overlap</td>
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<tr>
<td>CFPoptBelfw</td>
<td>ATTTGATCTCATAAAAAGGAGAAGCAATATGTTGAGCAAGGCGGA</td>
<td>BclI</td>
<td>Amplification of cfpopt with ribosomal binding site for pTetON cloning</td>
</tr>
</tbody>
</table>

Continued on following page
were fused by overlap extension PCR (18) and cloned into pCR2.1 (Invitrogen), and single colonies were directly analyzed for CFP fluorescence by microscopy.

For expression of CFPot and YFPot in \textit{S. aureus}, both FP genes were amplified using oligonucleotide CFPotBclfw in combination with CFPot-Bamrv and oligonucleotide YFPotBclfw with YFPotBamrv, respectively. After subcloning into pCR2.1 (Invitrogen), sequencing to verify the fidelity of the construct, and transformation of plasmids into \textit{E. coli} strain SC110 to receive nonmethylated DNA, fragments were excised with BclI and BamHI and ligated into vector pTetON, which is a derivative of plasmid pYJ335 (20) to which the following changes have been made: (i) the erythromycin resistance cassette was replaced by a spectinomycin cassette originating from vector pJR312 (30); (ii) the tetracycline repressor gene was exchanged with an improved repressor called tetR(B/D), which has superior repression characteristics (37); and (iii) the \textit{tetr} promoter was replaced by an \textit{nrl} promoter from \textit{Bacillus megaterium} (21), which is nonfunctional in \textit{S. aureus}, and consequently, in the absence of tetR(B/D), any gene placed under the control of the (normally) tetracycline-inducible promoter will be constitutively expressed. All oligonucleotides used in this study are listed in Table 2.

\textbf{Transformation and analysis of recombinant bacteria.} \textit{B. anthracis} was transformed as described previously (30) and plated onto selective LB agar.

\textit{S. aureus} competent bacteria were prepared as described by Fitzgerald (15). Briefly, bacteria were grown in 25 ml TSB until an optical density of 1.0 at 600 nm was reached, washed multiple times in 10% glycerol (vol/vol), and resuspended in the same buffer, and then 1 \mu g of plasmid DNA was added. Bacteria were electrotransformed at 1.75 kV, 100 \mu F, and 25 m\AA. TSB was added, and the suspension was incubated at 37°C for 1 h before aliquots were plated on tryptic soy agar containing the appropriate antibiotic.

To verify the presence of plasmids, colonies were boiled for 2 min at 96°C in either Tris-EDTA buffer (for \textit{B. anthracis}) or lysis buffer (for \textit{S. aureus}), consisting of 1\% Triton X-100, 0.5\% Tween 20, 10 mM Tris-HCl (pH 8.0), and 1 mM EDTA (41). Bacterial debris were centrifuged for 1 min at 12,000 \times g, and supernatants were analyzed by PCR.

\textbf{Microscopy and fluorimetric assays of bacteria.} For microscopic evaluation of fluorescence, bacteria grown overnight were diluted in phosphate-buffered saline (PBS) and directly analyzed by microscopy, using a Nikon Eclipse TE2000 U microscope with the appropriate filters for detection of CFP (excitation at 405 ± 10 nm and emission at 430 ± 25 nm) or GFP or YFP (excitation at 470 ± 20 nm and emission at 515 nm). For analysis of spores, bacteria were streaked out on TSA plates and incubated for 1 week at 37°C in air or 5\% CO\textsubscript{2}. Fluorimetric measurements of bacteria grown to different optical densities were performed in a Victor3 reader (Perkin-Elmer) using a GFP filter set (excitation at 485 nm and emission at 535 nm). Fluorescence intensities were normalized against the nonfluorescent wild type.

\textbf{Flow cytometry.} For fluorescence-activated cell sorter (FACS) analysis, bacteria grown overnight were centrifuged, washed, and resuspended at 1/20 in PBS. Samples were analyzed using a FACS Aria cell sorting system (Becton Dickinson) with a 100-nm W, 488-nm-kr tone sapphire solid state laser for GFP and YFP and a 20-mW-kr tone violet solid state laser for CFP. The fluorescence intensity of 50,000 unengaged events was measured in FL-1 and detected at a 515- to 545-nm range with a fluorescence isothiocyanate filter (GFP/YFP) or at 430- to 470-nm range with a Pacific Blue filter (CFP). \textit{B. anthracis} strain Ames 33 not carrying FP genes served as negative control. Data were analyzed using FACS Diva (Becton Dickinson).

\textbf{Western blot and protein analysis.} Bacteria were grown in 10 ml LB broth until the mid-logarithmic (Ames 33 expressing different FPS) or stationary (for ppa4 promoter analysis) growth phase. Cultures were centrifuged, washed twice with PBS, and resuspended in 500 \mu l PBS, and bacteria were lysed in a FastPrep system (MP Biomedical) using the FastProtein kit (MP Biomedical) as recommended by the manufacturer. Lysed suspensions were centrifuged briefly, and concentrations of protein lysates were determined using a bichromonic acid protein assay kit (Pierce). Equal amounts of cellular proteins were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis on a 4 to 20\% Tris-glycine gel and blotted onto a nitrocellulose membrane. FPS were subsequently identified using a rabbit anti-GFP polyclonal antibody (Rockland) as the primary antibody and a polyclonal goat anti-rabbit IR800 conjugate (Rockland) as the secondary antibody. The Western blot was developed on an Odyssey infrared scanner (Licer), and bands were quantified using ImageJ software version 1.40g. Protein alignments were performed with the Lasergene MegAlign program (version 7.1.0) using the Clustal W method.

\textbf{RNA isolation and reverse transcriptase PCR (RT-PCR).} Ten milliliters of mid-logarithmic-phase culture was centrifuged and washed once with PBS, and RNA was isolated using a FastRNA Pro Kit (MP Biomedical) as recommended by the manufacturer. RNA was ethanol precipitated and resuspended in 100 \mu l diethyl pyrocarbonate-treated water, and concentrations were determined using an ND-100 spectrophotometer (NanoDrop).

For RT-PCR, purified RNA was treated with DNase (Ambion) and screened for absence of contaminating DNA by PCR, and 2.5 \mu g of DNA-treated RNA was subjected to reverse transcription using the Superscript First Strand kit (Invitrogen). To amplify FP genes, the following oligonucleotides were used: FPAsfw and FPBamrv for codon-optimized genes, YFPAsfw and YFPBamrv for YFP, CFPAsfw and CFPBamrv for GFP, and CFPXasfw in combination with FPAServ for gfpmut1. As an internal control, reactions were also performed with primers GyraFw and GyraRv, which were designed to amplify a 819-bp fragment of the gyrA gene (accession no. YP_016609). All primers are listed in Table 2. As negative control, RNA isolated from the Ames 33 wild-type strain was used. Band intensities were quantified using ImageJ version 1.40g.

\textbf{Statistical analysis.} For statistical analysis of differences in mean fluorescence intensities (MFIs), the unpaired \textit{t} test was performed using GraphPad Prism (5.01).

\textbf{Nucleotide sequence accession numbers.} The DNA sequences of all four codon-optimized FP genes have been submitted to GenBank and are available under...
TABLE 3. GC content and unfavorable codons in fluorescent protein genes and endogenous genes

<table>
<thead>
<tr>
<th>Gene</th>
<th>% GC</th>
<th>Unfavorable codons</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>No./total</td>
</tr>
<tr>
<td>gfpmut1</td>
<td>38.77</td>
<td>96/239</td>
</tr>
<tr>
<td>cfp</td>
<td>61.94</td>
<td>226/240</td>
</tr>
<tr>
<td>yfp</td>
<td>61.53</td>
<td>213/240</td>
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<td>pagfp</td>
<td>61.53</td>
<td>224/240</td>
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</tr>
<tr>
<td>yfp</td>
<td>33.47</td>
<td>56/240</td>
</tr>
<tr>
<td>cfp</td>
<td>33.19</td>
<td>55/239</td>
</tr>
<tr>
<td>gfpmut1</td>
<td>37.36</td>
<td>76/265</td>
</tr>
<tr>
<td>spo0A</td>
<td>37.36</td>
<td>76/265</td>
</tr>
</tbody>
</table>

RESULTS

Sequence analysis of gfpmut1, cfp, yfp, and pagfp. In comparison to other bacteria, such as Mycobacterium tuberculosis or Corynebacterium diphtheriae, B. anthracis has an unusually low GC content of approximately 35% (http://cmr.tigr.org/tigr-scripts/CMR/GenomePage.cgi?org=gba). Consequently, for the expression of proteins derived from heterologous organisms having a higher GC content, a low availability of GC-rich codons in B. anthracis could be a limiting factor for protein synthesis and a negative influence on protein expression. In an initial in silico analysis, we considered whether B. anthracis could be expected to support expression of the Aequorea victoria-derived gfpmut1 gene (11), the cfp and yfp genes, encoding CFP and YFP, respectively (14), and the pagfp gene, encoding a photoactivatable FP (31). The main difference between the last three genes and gfpmut1, beyond the obvious differences in fluorescence excitation and emission spectra, is that cfp, yfp, and pagfp are derivatives of Clontech’s egfp, a gene with high GC content optimized for expression in mammalian cells. pagfp represents a photoactivatable GFP, first described by Patterson and Lippincott-Schwartz (31), that is nonfluorescent unless activated with a 413-nm laser. We analyzed the DNA sequences of all four genes with respect to GC content and percentage of unfavorable codons for expression in B. anthracis. A codon was deemed unfavorable if it was at least 50% less frequent than the most commonly used codon for a given amino acid in B. anthracis. Additionally, for comparison purposes, four randomly picked genes from B. anthracis were subjected to the same analysis: (i) sporulation gene spo0A, (ii) gyrase subunit A gene gyrA, (iii) PA gene pagA, and (iv) sortase A gene srtA. The results of this analysis are depicted in Table 3 and show that all nonoptimized FP genes exhibit a higher GC content than endogenous genes of B. anthracis, gfpmut1 has the lowest GC content of all four (39%) and the lowest percentage of unfavorable codons. In contrast, cfp, yfp, and pagfp exhibit a substantially higher GC content than intrinsic B. anthracis genes. Concomitantly, the percentages of unfavorable codons for B. anthracis are 94% (cfp), 89% (yfp), and 94% (pagfp), values greatly elevated in comparison to those for endogenous genes. In comparison, B. anthracis endogenous genes have an overall GC content of 33 to 37% and an unfavorable codon percentage of 22 to 29%. Taken together, these data imply that pagfp, cfp, and yfp are more likely to benefit from codon optimization than is gfpmut1.

Generation of codon-optimized gfpmut1, cfp, and yfp for expression in B. anthracis. Initially, we designed a gfpmut1 gene with a lower GC content, hoping that it might show superior expression in B. anthracis. Using an in silico approach, we exchanged 110 nucleotides in the gfpmut1 gene, reducing the total number of unfavorable codons from 40% to 23%, which resembles the percentage in endogenous genes (Table 3). The codon-optimized gene, gfpopt, was then synthesized and inserted into pUC18 (BlueHeron). This plasmid served as a PCR template for generation of codon-optimized cfpopt, yfpopt, and pagpopt as outlined in Materials and Methods. An alignment of all codon-optimized FPs with CFPopt is given in Fig. 1. The translated GFPopt had the following exchanges in comparison to GFPopt: Y66W, N146I, M153T, V163A, and H231L, as well as an additional valine at position 2 of the mature protein (Fig. 1). The last two amino acid changes were incorporated to match the protein sequence of CFPopt exactly to that of the Clontech-derived CFP. Consequently, in comparison to the unmodified cfp, a total of 410 nucleotides were exchanged in cfpopt, leading to a lowered GC content of 33.5% and a decrease in the percentage of unfavorable codons to 23% (Table 3).

FIG. 1. Clustal W sequence alignment of all four codon-optimized FPs. Amino acid changes performed with respect to GFPopt are highlighted in gray.
For generation of YFPopt, we performed the following amino acid exchanges in GFPOpt: T65G, V68L, S72A, and T203Y (Fig. 1). Although we incorporated all amino acids crucial for a red-shifted fluorescence, our yfpopt is based on gfpopt, therefore resulting in an aberrant amino acid sequence that is not identical to that of Clontech's YFP. By codon optimization, we were able to reduce the number of unfavorable codons in yfpopt to 22% (Table 3), in comparison to 89% in yfp.

To design the photoactivatable PAGFP, we initially converted the enhanced GFPOpt, having F64L and S65T exchanges that cause loss of the major 397-nm peak in the GFPMut1 emission spectrum (11), back to the original wild-type GFP. A V163A exchange needed for high expression at 37°C (40) was incorporated, as well as the crucial exchange required for photoactivation, T203H (31) (Fig. 1). As for yfpopt, the valine at position 2 of the PAGFP protein was not included in the codon-optimized gene. However, in comparison to the nonoptimized pagfp, we were able to lower the GC content from 94% to 22% in pagfpopt (Table 3), incorporating 225 nucleotide exchanges with respect to gfpmut1.

Cloning and expression of FPs in B. anthracis. For evaluating protein expression of the unmodified and codon-optimized FPs, we cloned gfpmut1, gfpopt, cfp, cfpopt, yfpopt, and pagfpopt into the gram-positive E. coli shuttle vector pSW4 (32), a derivative of pSJ115 that allows expression of proteins under the control of the pagA promoter of B. anthracis. PA of B. anthracis is the moiety of the anthrax lethal toxin and edema toxin which binds to the eukaryotic cell membrane, using receptors CMG2 and TEM8 (6, 39). This bacterial protein is highly expressed in the E. coli FP shuttle vector pSW4 (32). We found that when using LB broth, expression occurs even in the absence of bicarbonate. The resulting plasmids pSW4-GFPMut1, etc. (Table 1), were partially sequenced to verify accuracy of the genes and promoter regions and electroporated into B. anthracis Ames 33, a strain devoid of both virulence plasmids, pXO1 and pXO2 (32).

We then analyzed positive clones grown in LB broth in the absence of bicarbonate to stationary phase for expression of FPs by microscopy (Fig. 2). MFIs can be assessed by flow cytometric analysis and show levels of FP expression in single bacteria. MFIs of bacteria grown in air corroborated the results obtained by microscopy. Thus, bacteria expressed both GFPOpt and GFPMut1 to a similar high level, with MFIs of 10,948 and 13,243, although expression of GFPMut1 was significantly better (P = 0.0309, GFP versus GFPOpt; P < 0.0001, CFP versus CFPOpt; P = 0.0005, YFP versus YFPopt). Clear differences were also seen between single bacteria expressing CFP and CFPOpt, with MFIs of 1,235 and 2,758, respectively, and, more strikingly, between YFP- and YFPopt-positive bacteria, having MFIs of 787 and 10,224, respectively. All differences were significant (P < 0.05). A comparison of MFIs of differently colored FPs could not be performed since different lasers and filters were used for excitation and detection. Observed variances in MFI can be attributed to the near absence of a fluorescent signal in CFP- and YFP-expressing bacteria, as FACS plots show (Fig. 3B). These results are consistent with the observations made by microscopy and confirm the superior expression of codon-optimized CFPOpt and YFPopt.

Expression and transcription of FP genes. To test whether low protein expression of yfp and cfp and higher fluorescence of gfpmut1 were caused by low production of mRNA, we performed semiquantitative RT-PCR experiments to compare transcription levels of the codon-optimized versus original FP genes. B. anthracis harboring plasmid pSW4 with FP genes was grown to mid-logarithmic growth phase in LB broth, allowing expression of genes in the absence of bicarbonate, and RNA was extracted and processed as outlined in Materials and Methods. Figure 4A shows that mRNA quantities of all genes were similar; however, in comparison to the internal gyrA control, transcription levels of nonoptimized CFP and YFP were slightly lower than for the codon-optimized counterparts. This might explain a weaker expression of these FPs but not a total lack of expression.

Consequently, we also investigated whether the lack of CFP and YFP fluorescence was due to rapid degradation of these FPs in B. anthracis. Therefore, we performed Western blot analysis of B. anthracis protein lysates and evaluated expression of FPs with a polyclonal GFP antibody. The results depicted in Fig. 4B show that in comparison to CFPOpt and YFPopt, the original CFP and YFP were poorly or not at all expressed, and we were unable to detect any degradation.
Interestingly, even though there was no visible fluorescence when analyzed microscopically and quantitatively, low protein expression was noticeable for CFP.

These results show that the dissimilarities observed in FP expression are unlikely to be caused by differences in mRNA expression or by degradation of FPs but must result from poor expression due to tRNA unavailability. Therefore, although expressed in higher-GC-content bacteria such as *E. coli*, non-codon-optimized YFP and CFP are not suitable for expression in *B. anthracis*.

Analysis of PagA promoter activity using codon-optimized GFPopt. Next, we wanted to test whether our improved codon-optimized FPs are suitable for analysis of promoter activity in *B. anthracis* by analyzing GFPopt production under the control of the pagA promoter. In *B. anthracis*, the expression of PA is influenced by two regulators, AtxA and PagR, present on virulence plasmid pXO1, and their activity is determined by the presence of carbon dioxide (12, 24, 47). Whereas AtxA up-regulates expression of PA in the presence of CO₂ (24), PagR, which is cotranscribed with PA, negatively influences transcrip-
tion of *atxA*, therefore causing low PA expression (19). To evaluate whether our optimized FPs are suitable for analysis of promoter activity in *B. anthracis*, we used fluorimetric assays to assess the influence of these factors on *pagA* promoter-driven GFPopt expression. We compared $P_{pagA}$-dependent GFPopt expression in an AtxA/PagR-harboring strain (Ames 35, pXO1 positive) with that in a regulator-negative strain (Ames 33, pXO1 negative) in the presence and absence of bicarbonate.
and carbon dioxide. For this analysis, we used NBY broth (45), which is a low-nutrient broth shown to promote efficient toxin production in \textit{B. anthracis} (10). Figure 5A shows growth curves and fluorescence over time for both strains and their GFP\textsuperscript{opt}-negative counterparts. High GFP\textsuperscript{opt} expression could be observed only for strain A35 in the presence of carbon dioxide, whereas there was little expression in regulator-negative strains, even in the presence of CO\textsubscript{2}. Fluorescence was highest at late exponential and stationary growth phases, showing that the \textit{pagA} promoter is more active during these phases. Western blot analysis of lysates harvested from bacteria at the end of the experiment after approximately 8 h of growth (Fig. 5B) confirms observations made in fluorimetric assays. In comparison to the pXO1-containing strain Ames 35 grown in carbon dioxide, all strains show lower GFP\textsuperscript{opt} expression. These results show the suitability of codon-optimized FPs, in our application GFP\textsuperscript{opt}, for the analysis of promoter activity in \textit{B. anthracis}.

\textbf{Expression of GFP\textsuperscript{opt}, CFP\textsuperscript{opt}, and YFP\textsuperscript{opt} in \textit{S. aureus}.} \textit{S. aureus} is another low-GC-content bacterium for which, to our knowledge, only GFP and YFP have been described as FP reporters (4, 9, 33). Analysis of the original YFP and CFP showed that as in \textit{B. anthracis}, 213 or 226 out of 240 codons can be deemed unfavorable for expression in \textit{S. aureus}. We therefore hypothesized that our newly developed codon-optimized FPs might be suitable for expression in \textit{S. aureus}. Initially, we attempted to utilize the \textit{pagA} promoter of \textit{B. anthracis} to control FP\textsuperscript{opt} expression; however, we failed due to its inactivity in \textit{S. aureus} (data not shown), a result which was to be expected. Consequently, we used a previously published tetracycline-inducible promoter that has been shown to function in \textit{S. aureus} (20). We cloned all nonactivatable codon-optimized FPs into plasmid pTetON under the control of the tetracycline-inducible promoter \textit{P}_{\text{xylo}}/\text{tetO}. This plasmid is a derivative of pYJ335 (20), as outlined in Materials and Methods. Due to lack of tetracycline repressor expression on this plasmid, \textit{P}_{\text{xylo}}/\text{tetO} is constitutively active. Ligation of \textit{yfp\textsuperscript{opt}} and \textit{cfp\textsuperscript{opt}} into pTetON and transformation of \textit{S. aureus} RN4220 resulted in high expression of all FPs (Fig. 6). These results demonstrate that our codon-optimized FP genes may be highly suitable as reporter genes not only in \textit{B. anthracis} but also in other low-GC bacteria.

\textbf{DISCUSSION} 

FPs have wide applications in imaging bacterial gene expression, promoter activity, and localization of proteins. In comparison to enzymatic reporters, a clear advantage is the ability to monitor gene activity in intact bacteria, a method that stands in contrast to the use of enzymatic reporters, such as \textit{β}-lactamase or chloramphenicol acetyltransferase, which require lysis of cells and release of the cytosol in order to assess reporter activity. A disadvantage of FPs is that not all bacteria are able to express them at high levels. Here, we were able to show superior expression of two codon-optimized FPs, YFP\textsuperscript{opt} and CFP\textsuperscript{opt}, in the low-genomic-GC-content pathogens \textit{B. anthracis} and \textit{S. aureus}.

Initial sequence comparison between the original FPs and codon-optimized ones with regard to codon usage showed high discrepancies compared to intrinsic \textit{B. anthracis} genes, making them good candidates for codon optimization. By lowering the GC content substantially and adapting the codon usages of fluorescent genes \textit{gfp\textsuperscript{opt}}, \textit{cfp\textsuperscript{opt}}, and \textit{yfp\textsuperscript{opt}} to those preferred
by *B. anthracis*, we were able to achieve high expression in this bacterium. Strikingly, no expression was observed for the original, nonoptimized genes *cfp* and *yfp*. This phenomenon of better expression after codon optimization has been described previously (3, 27). Surprisingly, although it was not microscopically obvious, we were able to observe differences in fluorescence intensities between the codon-optimized GFPopt and the original GFPmut1 by FACS; the latter was significantly brighter as determined by MFI analysis. We did not expect major differences in the fluorescence of these two proteins, since the GC content and codon utilization are not strikingly dissimilar. Nonetheless, our expectations were to observe a higher fluorescence for the codon-optimized version of the protein. An explanation for this phenomenon might be differential folding and structural properties of the mRNA, since the primary sequence of optimized and nonoptimized *gfp* has been altered, therefore leading to a differential translation rate. Another reason might be better protein folding properties of the original GFPmut1. To further investigate the reasons for expression differences between codon-optimized and original CFPs and YFPs, we performed transcriptional analysis by RT-PCR to quantify mRNA levels. The results showed the presence of all transcripts, with a slightly smaller amount for *cfp* and *yfp*, which could be explained by either mRNA instability or a lower transcription rate; the latter seems unlikely since all FPs were expressed from the same promoter. The fact that mRNAs of genes exhibit different stabilities has been shown for many bacteria and can have multiple reasons, such as secondary structures (hairpins), internal secondary structures marking them for RNase III degradation, or RNase recognition sequences (7). The lower mRNA levels observed for YFP might explain a lower protein expression but not the total absence of protein that was observed by Western blot analysis. We therefore conclude that a better availability of the cognate tRNAs is the more likely explanation for high expression of CFPopt and YFPopt in comparison to the nonoptimized counterparts.

We were able to demonstrate that our codon-optimized FPs can be used as reporters for promoter activity, as shown by the analysis of the *pagA* promoter in response to carbon dioxide, bicarbonate, and the presence of pXO1, the larger virulence plasmid of *B. anthracis* harboring PA regulators AtxA and PagR. The results showed a clear correlation between the presence of CO₂ and promoter activity, measured in GFP fluorescence. Furthermore, the presence of pXO1 greatly influenced PA promoter activity as observed by high fluores-

FIG. 5. *pagA* promoter analysis using GFPopt as a reporter. (A) Growth curves (broken lines) and fluorescence intensities (solid lines) of *B. anthracis* expressing GFPopt under the control of the *pagA* promoter. GFPopt-harboring bacteria containing virulence plasmid pXO1 (strain A35) or plasmid free (strain A33) were grown in NBY broth supplemented with 10% FBS in the presence or absence of sodium bicarbonate (0.9%, wt/vol) and a 5% CO₂ atmosphere. At different time points, samples were analyzed for fluorescence in a fluorimeter. CPS, counts per second. Error bars indicate standard deviations. (B) Western blotting of bacteria grown to stationary phase. Lysates were processed as described for Fig. 4.
cence, especially during stationary phase. To some degree, the higher fluorescence at later time points could also be caused by accumulation of GFP and amplification of the fluorescent signal within the cells. GFP has been shown to have an extremely long half-life of 24 h or more (25), making it less suitable for promoter reporter activity. Therefore, it would be desirable to have less stable variants for reporter purposes. For some bacteria, such as Staphylococcus epidermidis (16) or Mycobacterium (5), GFP variants with alternative C termini and substantially shorter half-lives have been described. By varying the C terminus, bacterial proteins can be marked for faster degradation by housekeeping proteases, lowering the half-lives of GFP variants to as low as to 40 min to several hours, depending on the bacterial species (2). Studies to improve our optimized proteins with respect to stability are under way, since the use of a stable FP might prove difficult for true quantitative determination of promoter activities during bacterial growth.

To evaluate whether high expression of our codon-optimized FPs is limited to just B. anthracis or whether it could also be applied to other low-GC bacteria, we cloned GFPopt, YFPopt, and CFPopt under the control of a strong promoter for expression in S. aureus. Fluorescent microscopic images confirmed that all proteins are also highly expressed in S. aureus. Even though YFP has been used to visualize biofilms in S. aureus (4), this is to our knowledge the first report showing CFP expression in this bacterium.

We were also able to construct a codon-optimized photoactivatable GFP suited for expression in low-GC bacteria. Photoactivatable proteins find application in mammalian cells, e.g., for monitoring organelle or molecule dynamics within cells (23, 29, 31). By photoactivation, highlighted subpopulations of proteins, organelles, or cells can be monitored temporally, and new synthesis of PAGFP-labeled proteins does not influence observations, as these molecules will not be fluorescent (26). No use of this protein in bacteria has been reported, but applications paralleling those in mammalian cells can be imagined.

Finally, another important observation made was that B. anthracis spores derived from bacteria expressing FPs showed high fluorescence which remained stable even after heat inactivation and for a period of at least 14 days. Stable maintenance of GFP within the spore has been reported before (35). These results show that fluorescent spores could be suitable for infection purposes and for monitoring localization of spores within, e.g., mammalian cells; fluorescence intensities of GFP-positive bacteria or spores in host cells may even be used as indicator for multiplicities of infection, as previously shown for Salmonella by Thone et al. (44). There are many possible applications for FPs in bacteria; extending the variety of color spectra available for expression in different species of bacteria may help us to understand basic pathogenic mechanisms and expression profiles of virulence factors in vitro and in vivo.

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