

Red Fluorescent Proteins for Gene Expression and Protein Localization Studies in *Streptococcus pneumoniae* and Efficient Transformation with DNA Assembled via the Gibson Assembly Method

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During the last decades, a wide range of fluorescent proteins (FPs) have been developed and improved. This has had a great impact on the possibilities in biological imaging and the investigation of cellular processes at the single-cell level. Recently, we have benchmarked a set of green fluorescent proteins (GFPs) and generated a codon-optimized superfolder GFP for efficient use in the important human pathogen *Streptococcus pneumoniae* and other low-GC Gram-positive bacteria. In the present work, we constructed and compared four red fluorescent proteins (RFPs) in *S. pneumoniae*. Two orange-red variants, mOrange2 and TagRFP, and two far-red FPs, mKate2 and mCherry, were codon optimized and examined by fluorescence microscopy and plate reader assays. Notably, protein fusions of the RFPs to FtsZ were constructed by direct transformation of linear Gibson assembly (isothermal assembly) products, a method that speeds up the strain construction process significantly. Our data show that mCherry is the fastest-maturing RFP in *S. pneumoniae* and is best suited for studying gene expression, while mKate2 and TagRFP are more stable and are the preferred choices for protein localization studies. The RFPs described here will be useful for cell biology studies that require multicolor labeling in *S. pneumoniae* and related organisms.

S*treptococcus pneumoniae* is a human pathogen that annually kills about 1.5 million people worldwide (1). It causes harmful infections such as pneumonia, meningitis, and sepsis, especially in elderly people, young children, and immunosuppressed patients. The increase of antibiotic resistance and the lack of long-lasting vaccines against this organism make it crucial to better understand the cell biology of S. pneumoniae and gain insights into essential processes such as cell division, chromosome segregation, and cell wall synthesis. Therefore, the development of tools for cell biological studies in this organism is important.

Studies of protein localization and gene expression at the single cell level have brought new insights into the cell biology of bacteria. Nevertheless, most of our knowledge on bacterial cell biology is based on experiments performed in rod-shaped model organisms such as *Escherichia coli* and *Bacillus subtilis*, and only recently have researchers been able to examine these processes in coccoid bacteria such as *Staphylococcus aureus* and *S. pneumoniae* (2).

S. pneumoniae is a microaerophilic organism that cannot tolerate high levels of oxygen. Folding and maturation of fluorescent proteins often require sufficient concentrations of oxygen, and microaerophilic conditions might have a negative effect on maturation (3). More recently, molecular tools to study cell biology of living cells of *S. pneumoniae* have been published (4– 6), and a set of green fluorescent proteins (GFPs) that can be used in *S. pneumoniae* with adequate efficiency has been described (6).

The GFP from the jellyfish *Aequorea victoria* was the first FP to be described (7) and cloned (8, 9). Other FP variants, such as blue, cyan, and yellow FPs, have been engineered from the jellyfishderived GFP, whereas orange, red, and far-red variants were found in other sea animals (10–12). Most red fluorescent proteins (RFPs) naturally occur in tetrameric form, and monomerization

mutants often resulted in a great loss of photostability and fluorescence intensities. However, in recent years, improved variants have been developed (11, 13). The four different RFPs that we benchmarked in this study were codon optimized for low-GC Gram-positive bacteria. We chose four monomeric RFPs that can possibly be used in protein fusions. The orange-red fluorescent protein we tested, mOrange2, is a monomeric derivative of DsRed from the coral Dictiosoma sp. with improved photostability (11). TagRFP, a monomeric derivate of eqFP578 from the sea anemone *Entacmaea quadricolor* (13), is an orange-red FP for which high brightness but low photostability has been reported (13). For farred variants, we tested mCherry (10), another derivative of DsRed and one of the most widely used RFPs in bacterial research, and mKate2, derived from E. quadricolor and with excellent pH resistance and photostability (12). While the four selected RFPs share a high degree of sequence and structural similarity, mKate2 has the highest reported quantum yield when examined in vitro (12). To

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TABLE 1 Bacterial strains and plasmids

Strain or plasmid	Relevant properties	Source or reference	
E. coli DH5α	F ^{$-$} araD139 Δ (ara-leu)7696 Δ (lac)X74 galU galK hsdR2 mcrA mcrB1 rpsL	Invitrogen, CA	
S. pneumoniae			
D39	Encapsulated strain	25	
KB1-69	D39 Tet bgaA::P _{Zn} -mOrange2	This study	
KB1-70	D39 Tet bgaA::P _{Zn} -tagRFP	This study	
KB1-71	D39 Tet <i>bgaA</i> ::P _{Zn} - <i>mCherry</i>	This study	
KB1-72	D39 Tet bgaA::P _{Zn} -mKate2	This study	
MK119	D39 Cam hlpA hlpA-mKate2	18	
KB1-64	D39 Cam <i>hlpA-mCherry</i>	This study	
MK218	D39 Cam hlpA hlpA-mCherry	This study	
KB1-65	D39 Cam hlpA hlpA-tagRFP	This study	
KB04-03	D39 Tet bgaA::P _{ssbB} -mOrange2	This study	
KB04-04	D39 Tet <i>bgaA</i> ::P _{ssbB} -tagRFP	This study	
KB04-05	D39 Tet <i>bgaA</i> ::P _{ssbB} -mCherry	This study	
KB04-06	D39 Tet bgaA::P _{ssbB} -mKate2	This study	
DRR27	D39 Ery ftsZ-mCherry	This study	
DRR28	D39 Ery ftsZ-mKate2	This study	
DRR54	D39 Ery ftsZ-tagRFP	This study	
MK125	D39 Cam <i>hlpA hlpA-mKate2</i> Tet <i>bgaA</i> ::P _{Zn} - <i>gfp-stkP</i>	This study	
MK289	D39 Cam <i>hlpA hlpA-mCherry</i> Tet <i>bgaA</i> ::P _{Zn} - <i>gfp-stkP</i>	This study	
MK290	D39 Cam <i>hlpA</i> , <i>hlpA-tagRFP</i> Tet <i>bgaA</i> ::P _{Zn} - <i>gfp-stkP</i>	This study	
MK292	D39 Ery <i>ftsZ-mCherry</i> Tet <i>bgaA</i> ::P _{Zn} - <i>gfp-stkP</i>	This study	
MK293	D39 Ery ftsZ-mKate2 Tet bgaA::P _{Zn} -gfp-stkP	This study	
MK294	D39 Ery ftsZ-tagRFP Tet bgaA::P _{Zn} -gfp-stkP	This study	
Plasmids			
pKB01_sfgfp(Sp)	<i>bla tet bgaA</i> ::P _{Zn} -sfgfp(Sp)	6	
pKB01_mOrange2	bla tet bgaA::P _{zn} -mOrange2(Sp)	This study	
pKB01_tagRFP	bla tet bgaA::P _{Zn} -tagRFP(Sp)	This study	
pKB01_mCherry	<i>bla tet bgaA</i> ::P _{Zn} - <i>mCherry</i> (<i>Bs</i>)	This study	
pKB01_mKate2	bla tet bgaA::P _{zn} -mKate2(Sp)	This study	
pKB01-P _{ssbB} -mOrange2	bla tet bgaA P _{ssbB} -mOrange2(Sp)	This study	
pKB01-P _{ssbB} -tagRFP	bla tet bgaA P _{ssbB} -tagRFP(Sp)	This study	
pKB01-P _{ssbB} -mCherry	<i>bla tet bgaA</i> P _{ssbB} -mCherry(Sp)	This study	
pKB01-P _{ssbB} -mKate2	bla tet bgaA P _{ssbB} -mKate2(Sp)	This study	
pJWV25-stkP	pJWV25-stkP bla tet bgaA P _{Zn} -gfp-stkP		
pLA18	bla tet bgaA::P _{ssbB} -luc-gfp	26	

identify which of the four RFPs is the brightest and most rapidly maturing variant in vivo in S. pneumoniae, we placed the respective genes under control of inducible promoters PZn and PssbB and integrated them stably as a single copy within the S. pneumoniae genome. The speed of FP maturation is not always important, for instance, when the question is simply where the protein localizes. In such cases, brightness or functionality of the FP fusion might be more crucial. Therefore, we tested the properties of TagRFP, mCherry, and mKate2 when used as C-terminal protein fusions to the tubulin-like protein FtsZ and the histone-like protein HlpA. We successfully used Gibson's one-step isothermal (ISO) DNA assembly method (14) to generate several of our constructs. Importantly, using S. pneumoniae's natural competence, we show that pneumococcus can take up and integrate the linear assembly product even directly from the reaction mixture. The RFPs and the use of a fast DNA assembly method (Gibson assembly) for direct transformation presented in this study further expand the cell biological toolbox available for S. pneumoniae and related organisms.

MATERIALS AND METHODS

Bacterial strains, plasmids, media, and growth conditions. All strains and plasmids used in the study are listed in Table 1. *Streptococcus pneumoniae* was grown in liquid C+Y medium at 37°C (15). Blood agar plates were prepared from Columbia agar with addition of 3% defibrinated sheep blood (Johnny Rottier, Kloosterzande, The Netherlands). The P_{Zn} promoter was induced with addition of ZnCl₂ to a final concentration of 0.1 mM or 0.15 mM, and P_{ssbB} was induced by adding 100 ng/ml of competence-stimulating peptide (CSP). *Escherichia coli* DH5 α was used for cloning and grown in LB medium at 37°C with shaking or on LB medium solidified with 1.5% (wt/vol) agar. When required, 100 µg ml⁻¹ ampicillin (Amp) for *E. coli* or 1 µg ml⁻¹ tetracycline (Tet) and 4.5 µg ml⁻¹ chloramphenicol (Cm) for *S. pneumoniae* was used for selection.

Recombinant DNA techniques and oligonucleotides. Procedures for DNA isolation, restriction, ligation, agarose gel electrophoresis, and transformation of *E. coli* were performed as described by Sambrook and Russell (16). Plasmid DNA and PCR products were isolated and purified using the High Pure plasmid isolation kit (Roche Applied Science, Mannheim, Germany) according to the manufacturer's instructions. Enzymes were purchased from Fermentas (Vilnius, Lithuania) and used as described by the manufacturer. For PCR amplification, Phusion and *Taq*

TABLE 2 Oligonucleotides used in this study

Oligonucleotide	Sequence $(5' \rightarrow 3')^a$
mOrange-F+NotI	GCAG <u>GCGGCC</u> GCGGAGGAAAATTAATGTCTAAGGGAG
mOrange-R+SpeI	GCAG <u>ACTAGT</u> CTCATGAATTCTTTCTCGAGTTAGG
mKate2-F+XbaI	GCAG <u>TCTAGA</u> GGAGGAAAATTAATGTCAGAACTTATCAAGG
mKate2-R+SpeI	GCAG <u>ACTAGT</u> GCGAGATCTCATGAATTCTTTCTCG
tagRFP-F+XbaI	GCAG <u>TCTAGA</u> GGAGGAAAATTAATGTCAGAAC
tagRFP-R+SpeI	GCAG <u>ACTAGT</u> CATGAATTCTTTCTCGAGTTAGGATCC
hlpA-up-F	AACAAGTCAGCCACCTGTAG
hlpA-up-R+BamHI	CTGC <u>GGATCC</u> TTTAACAGCGTCTTTAAGAGCTTTACCAGC
camR-up-F+EcoRI	ACTCG <u>GAATTC</u> TAATGAGCACTAGTAGGAGGCATATC
hlpA-down-R	CGTGGCTGACGATAATGAGG
Linker-mCherry-F+BamHI	CGAT <u>GGATCC</u> GGATCTGGTGGAGAAGCTGCAGCTAAAGGAAGCAAAGGAAGAAGATAACATGGCAATCATC
mCherry-R+EcoRI	GCGC <u>GAATTC</u> TTATTATTTGTAAAGCTCATCCATTCCGCC
hlpA-F-rbs-SphI	CGTGCATGCTGGAGGAATCATTAACATGGCA
hlpA-R-SphI	CGCGCATGCAGACTGATTATTTAACAGCGTC
hlpA-linker-up-R+BglII	GCGCAGATCTTCCTTTAGCTGCAGCTTCTCC
mKate2-tagRFP-F+BamHI	CGCTGGATCCTCAGAACTTATCAAGG
tagRFP-R+EcoRI	GCATGAATTCTTATTAACGGTGTCCCAATTTACTAG
integration 1	CTTGATGAAACCTACATTTG
integration 2	GCTTCCATTAAGGATAGTTC
integration 5	GCTATCGCTGAGCGCCGG
integration 6	AGCTAGAGTTCCGCAATTGG
FtsK-down-RV	ATCAAAACCGAACTCACCCTGTTGAT
ftsZ-down_FW_eryOE	<i>GGGAGGAAATAAGCGGCCGATGA</i> ATGTAAAAGAAAATACAGAACTTG
ftsZ-up-FW	CCTGTTATTGCTCGTATCGCCAAA
ftsZ-up-RV-mCherryOE	<i>GCTTCTCCACCAGATCCGGA</i> ACGATTTTTGAAAAATGGAGGTG
ery-FW-mcherryOE	<i>GAATGGATGAGCTTTACAAATAATAAG</i> CCCGGAGGAATTTTCATATGAAC
ery-RV-ftszdOE	CAAGTTCTGTATTTTCTTTTACATTCATCGGCCGCTTATTTCCTCCC
ery_F_mKateOE	<i>GTAAATTGGGACACCGTTAATAA</i> CCCGGAGGAATTTTCATATGAAC
mKate_RV_eryOE	<i>GTTCATATGAAAATTCCTCC</i> GGGTTATTAACGGTGTCCCAATTTAC
tagRFP_RV_eryOE	<i>GTTCATATGAAAATTCCTCC</i> GGGCTAGTTTATTTATGTCCCAATTTAC
ery_F_tagRFPOE	<i>GTAAATTGGGACATAAATAA</i> ACTAGCCCGGAGGAATTTTCATATGAAC
FtsZ_checkRV	AGTGGTGCCGATATGG
FtsZ_checkFW	TGACAGCGAAGGCTAC
mCherry-RV-eryOE	GTTCATATGAAAATTCCTCCGGGCTTATTATTTGTAAAGCTCATCCATTC
mCherry-FW-ftszuOE	CACCTCCATTTTTCCAAAAATCGTTCCGGATCTGGTGGAGAAGC

^a Underlined sequences represent the endonuclease restriction sites; italic sequences represent the overlapping sequences for Gibson assembly.

DNA polymerases (Fermentas) were used. *S. pneumoniae* was transformed as described by Martin et al. (15). The oligonucleotides used in this study are listed in Table 2 and were purchased from Biolegio (Nijmegen, The Netherlands). Constructs were verified by sequencing.

Codon optimization. *mKate2, tagRFP,* and *mOrange2* were codon optimized for *S. pneumoniae* using OptimumGene (GenScript Inc., Piscataway, NJ, USA), resulting in genes *mKate2(Sp), tagRFP(Sp),* and *mOrange2(Sp),* while the sequence for a codon-harmonized variant of *mCherry* (based on the *Bacillus subtilis* codon usage table) was obtained from DSM Biotechnology Center (Delft, The Netherlands), resulting in gene *mCherry(Bs).* Genes were synthesized by GenScript [*mKate2(Sp), tagRFP(Sp), and mOrange2(Sp)*] or by GeneArt (Regensburg, Germany) [*mCherry(Bs)*]. Note that throughout this paper we refer to these codon-optimized genes as *mKate2, tagRFP, mOrange2, and mCherry.*

Construction of promoter fusion plasmids. Different RFP genes were first fused to the Zn²⁺-inducible promoter P_{Zn} , resulting in construct *pKB01-P_{Zn}-RFP. mKate2* and *tagRFP* were amplified from the synthetic genes using primer pairs mKate-F+XbaI/mKate-R-SpeI and tagRFP-F+XbaI/tagRFP-R-SpeI, respectively, and cut using XbaI and SpeI. *mCherry* was cut from the GeneArt plasmid pMA-*mCherry* using XbaI and SpeI and purified. Digested *mKate2*, *tagRFP*, and *mCherry* were ligated in XbaI- and SpeI-cut plasmid pJWV102 (6), resulting in plasmids pKB01-*mKate2*, pKB01-*tagRFP*, and pKB01-*mCherry*, respectively. *mOrange2* was amplified using mOrange-F+NotI/mOrange-R+SpeI, cut using NotI and SpeI, and ligated in similarly digested pJWV102. Fragments were ligated accordingly, resulting in plasmid pKB01-*mOrange2*.

Subsequently, the zinc-inducible promoter was replaced by the competence-induced promoter P_{sbB} (17), resulting in plasmid *pKB01-P*_{ssbB}-*RFP*, which was excised from pLA18 using SphI and EcoRI and introduced into pKB01-RFP plasmids cut with the same restriction enzymes.

Construction of strains with P_{Zn}- and P_{ssbB}-controlled RFP gene expression. S. pneumoniae strains KB1-69, KB1-70, KB1-71, and KB1-72, expressing different RFP variants under control of the Zn²⁺-inducible promoter (P_{Zn}), were obtained by transformation of wild-type strain D39 with pKB01 derivatives. Correct integration by double crossover in the *bgaA* locus was tested by colony PCR using primer pairs integration 1/integration 2 and integration 5/integration 6.

Similarly, strains KB04-03, KB04-04, KB04-05, KB04-06, expressing RFPs from the competence-activated promoter P_{sbb} , were obtained by transforming the corresponding plasmids into D39. Correct integration in the *bgaA* locus was checked as described above.

Construction of strains expressing translational HlpA-RFP fusions. The construction of strain MK119 (*hlpA hlpA-mKate2*; Cam^r), containing an *hlpA-mKate2* fusion (in which *hlpA* and *mKate2* is separated by a flexible linker) and a chloramphenicol resistance gene on the same transcriptional unit as *hlpA*, is described elsewhere (18).

To construct strain KB1-64 (D39 *hlpA-mCherry*; Cam^r), the *hlpA*-up fragment (containing the *hlpA* gene and its promoter) was amplified from

D39 chromosomal DNA using hlpA-up-F/hlpA-up-R+BamHI, and the hlpA-down fragment (containing ~1,000 bp immediately downstream of the hlpA stop codon, including a chloramphenicol resistance marker) was amplified from strain MK119 (D39 hlpA hlpA-mKate2; Cam^r) using primer pair camR-up-F+EcoRI/hlpA-down-R. *mCherry* including a flex-ible linker was amplified from pMA-*mCherry* using linker-mCherry-F+BamHI/mCherry-R+EcoRI. Fragments were purified and cut using BamHI (fragment hlpA-up), BamHI and EcoRI (linker-*mCherry*), or EcoRI (hlpA-down). Cut and purified fragments were ligated. *S. pneumoniae* D39 was transformed with the ligation mix, and transformants were selected on plates containing chloramphenicol. A single transformant was sequence verified, and this strain was named KB1-64.

To construct strain MK218 (D39 *hlpA hlpA-mCherry*; Cam^r), an *hlpA*-up fragment (containing *hlpA* and the upstream sequence) was amplified from genomic DNA of D39 using primers hlpA-up-F and hlpA-R. SphI. The *hlpA-mCherry*; Cam^r fragment (containing the *hlpA-mCherry* fusion with a downstream; Cam^r resistance gene and the *hlpA* downstream sequence) was amplified from strain KB1-64 using primers hlpA-F-rbs-SphI and hlpA-down-R. The two fragments were digested with SphI and ligated. The ligation product was transformed into D39, and transformants were selected on plates containing chloramphenicol. Transformants were verified by PCR and sequencing.

To construct strain KB1-65 (D39 *hlpA hlpA-tagRFP*; Cam^r), the *hlpA*linker-up fragment, containing an extra copy of *hlpA* including a flexible linker downstream of *hlpA*, was amplified from MK119 using primer pair hlpA-up-F/hlpA-linker-R+BgIII, and the *hlpA*-down fragment was amplified as described for KB1-64. *tagRFP* was amplified from pUC57*tagRFP* using primers mKate2-tagRFP-F+BamHI/tagRFP-R+EcoRI. Fragments were purified and cut with BgIII (*hlpA*-linker-up), EcoRI (*hlpA*-down), and BamHI and EcoRI (*tagRFP*). The fragments were ligated and directly transformed into *S. pneumoniae* D39. Positive transformants were selected on plates containing chloramphenicol. A single transformant was sequence verified, and the strain was named KB1-65.

Gibson assembly for construction of FtsZ-RFP strains. Strains expressing the different FtsZ-RFP fusions were constructed by direct transformation of linear Gibson assembly products. The parts for strain DRR27 were assembled using Gibson's one-step ISO assembly of overlapping double-stranded DNA (dsDNA) (14). The ftsZ-up and ftsZ-down fragments were amplified from chromosomal DNA using ftsZ-up-FW and ftsZ-up-RV-mCherryOE for ftsZ-up and ftsZ-down_RV and ftsZ-down_FW_eryOE for ftsZ-down. The fragment encoding erythromycin resistance was amplified from plasmid JWV502 using ery-FW-mcherryOE and ery-RV-ftszdOE. mCherry including a flexible linker was amplified from strain KB1-64 using primers mcherry-RV-eryOE and mcherry-FW-ftszuOE. All four parts included a sequence of around 20 bp (Table 2) that overlapped with the next part. Gibson one-step ISO assembly was done as described by Gibson et al. (14), and 10 µl of the reaction mixture was directly transformed into *Streptococcus pneumoniae* D39.

For DRR28 and DRR54, ery-ftsZdown was amplified from strain DRR27 using ery-F-mKateOE and ftsZ-down-RV for DRR28 and ery-F-tagRFPOE and ftsZ-down-RV for DRR54. mKate2 was amplified from strain MK119 using mcherry-FW-ftszuOE and mKate_RV_eryOE, and TagRFP was amplified from strain KB1-65 using mcherry-FW-ftszuOE and tagRFP_RV_eryOE. FtsZ-up was amplified the same way as for DRR27. For DRR28, ery-ftsZdown (with overlap with mKate2), ftsZ-down, and mKate2 and for DRR54, ery-ftsZdown (with overlap with TagRFP), ftsZ-down, and tagRFP were combined in equimolar amounts for Gibson assembly and directly transformed into *S. pneumoniae* D39. Integration via double crossover was checked in all three strains using primers FtsZ-checkRV and FtsZ-checkFW.

Construction of RFP/GFP-double-labeled strains. Double-labeled strains GFP-StkP and FtsZ-RFP were made by transformation of plasmid pJWV25-stkP (19) into the FtsZ-RFP fusion strains DRR27, DRR28, and DRR54, to generate strains MK292 (FtsZ-mCherry, GFP-StkP), MK293 (FtsZ-mKate2, GFP-StkP), and MK293 (FtsZ-TagRFP, GFP-StkP). Dou-

ble-labeled strains GFP-StkP and HlpA-RFP were made in a similar manner by transformation of plasmid pJWV25-stkP into HlpA-RFP fusion strains to generate MK125 (HlpA-mKate2, GFP-StkP), MK289 (HlpA-mCherry, GFP-StkP), and MK290 (HlpA-TagRFP, GFP-StkP).

Fluorescence measurements using a plate reader. *S. pneumoniae* RFP gene-expressing strains and the D39 wild-type strain were grown at 37°C in liquid C+Y medium to an optical density at 600 nm (OD₆₀₀) of 0.1 and diluted 1:100 with fresh C+Y containing 0.15 mM ZnCl₂ for induction of P_{Zn} . Cells were then further grown and harvested at an OD₆₀₀ of 0.15. For plate reader assays, cells were first collected by centrifugation and washed with phosphate-buffered saline (PBS). Fluorescence was measured with the following equipment and settings: Infinite 200 Pro plate reader (Tecan Group Ltd.) with I-control 1.7.1.12 software (Tecan Group Ltd.), RFP filter set (excitation at 590 nm and emission at 635 nm with a 50% dichroic mirror), and RFP signals collected as top readings with a gain setting of 100. RFP fluorescence values were corrected for background fluorescence, OD₆₀₀, and negative controls (values for the wild-type strains). The OD₆₀₀ levels used were corrected for the background value of the corresponding buffer.

Fluorescence microscopy. Cells of S. pneumoniae were grown at 37°C in liquid C+Y medium to an OD₆₀₀ of 0.1, diluted 1:100 with fresh C+Y, and further grown to an OD_{600} of 0.05 before 0.15 mM ZnCl_2 was added to the cultures for induction of P_{Zn} . Cells were then further grown and collected at different time points. For microscopy, 0.4 µl of the cell suspension was directly spotted onto a microscope slide carrying a thin layer of 1.2% agarose in PBS covered with a coverslip. Images were taken with an Olympus IX71 microscope (Personal DV, Applied Precision; assembled by Imsol [Preston, United Kingdom] and Chromaphor [Münster, Germany]) using a CoolSNAP HQ2 camera (Princeton Instruments, Trenton, NJ, USA) with a $100 \times$ phase-contrast objective and Trulight (Deltavision, GE Healthcare, USA) illumination. The following standard fluorescence filter sets from Chroma Technology Corporation (Bellows Falls, VT, USA) were used to visualize RFPs: for mCherry, excitation at 575/25 nm, emission at 632/60 nm, and polychroic mirror QUAD2 (580 to 630 nm); for tetramethyl rhodamine isocyanate (TRITC), excitation at 542/27 nm, emission at 594/45 nm, and polychroic QUAD1 (550 to 630 nm). (Note that the second number indicates the bandwidth around the wavelength; e.g., 575/25 indicates excitation with light from 562.5 to 587.5 nm). For the Zn²⁺-inducible constructs, the exposure time was 1.4 s with 100% excitation for all RFPs. To visualize the RFP fusions to HlpA, an exposure time of 0.9 s was used. Softworx 3.6.0 (Applied Precision, Issaquah, WA, USA) software was used for image capturing. Phase-contrast images were segmented automatically, and fluorescence signal intensities were extracted using Microbetracker (20). Fluorescence levels were corrected for background fluorescence of the medium.

Western blot analysis and immunodetection. Cells were grown in C+Y liquid medium at 37°C to an OD₆₀₀ of 0.1, diluted 1:100 in C+Y containing 0.15 mM ZnCl₂ for induction, and harvested at an OD₆₀₀ of 0.15. Cells were harvested by centrifugation at 8,000 rpm for 10 min. For lysis, the pellet was resuspended in 100 μ l of SEDS lysis buffer (150 mM NaCl, 15 mM EDTA, 0.02% SDS, 0.01% deoxycholate) and was incubated for 5 min at 37°C. Lysates were diluted in 100 μ l 2× SDS loading buffer and boiled for 5 min. After separation by SDS-polyacrylamide gel electrophoresis (SDS-PAGE), proteins were transferred to a polyvinylidene difluoride (PVDF) membrane by Western blotting. Red fluorescent proteins were detected with polyclonal anti-RFP antibody (Molecular Probes) and rabbit anti-IgG–horseradish peroxidase (HRP) antibody (GE Healthcare) according to the manufacturer's instructions. Note that the polyclonal anti-RFP antibodies recognize all four RFPs due to the high structural homology.

Nucleotide sequence accession numbers. The sequences of the *mKate2, tagRFP, mOrange2*, and *mCherry* genes have been deposited in GenBank under accession numbers KJ908191, KJ908188, KJ908189, and KJ908190, respectively.

TABLE 3 Red fluorescent proteins

Fluorescent protein	$Maximum\lambda_{ex}(nm)$	$Maximum \ \lambda_{em} \ (nm)$	Color	Brightness ^a	Reference
mOrange2	549	565	Orange	35	11
TagRFP	555	584	Red	40	13
mCherry	587	610	Far red	16	11
mKate2	588	633	Far red	25	12

^{*a*} Brightness was determined as follows: (extinction coefficient \times quantum yield)/1,000.

RESULTS AND DISCUSSION

Expression of codon-optimized RFPs in S. pneumoniae. New variants of monomeric red fluorescent protein have been discovered and developed in recent years. They are important tools for use in protein fusions to study protein localization and dynamics and for multicolor labeling, for instance, in combination with GFP, cyan fluorescent protein (CFP), or yellow fluorescent protein (YFP). In the past, the brightness of monomeric RFPs has been problematic, but more recently, improved variants have been described. We selected four of these "next-generation" variants of red fluorescent proteins and codon optimized the genes for low-GC Gram-positive bacteria (Table 3), including those for mOrange2 and mCherry from the "mFruits" palette (11), as well as TagRFP (13) and mKate2 (12). mCherry is one of the most commonly used RFPs, with a reported short maturation time (varying between 15 min [11] and 40 min [13]) but a relatively poor photostability (11). mOrange2 is an orange-red-emitting FP variant and was reported to have a high photostability and high intrinsic brightness but a relatively long maturation time (\sim 4.5 h) compared to the doubling time of S. pneumoniae (~30 min under our experimental conditions) (11). TagRFP is a bright orange-red fluorescent protein with a reported maturation time of 100 min and high brightness (Table 3) (11, 13). mKate2 was also reported to have a short maturation time (<20 min) and good photostability (Table 3) (12). mKate2 and mCherry are far-red variants and therefore highly suitable for the use in combination with green and yellow fluorescent proteins. All variants are reported to be monomeric which makes them suitable for use in protein fusions.

In order to directly compare the four RFPs (mOrange2, TagRFP, mCherry, and mKate2), we used plasmid pJWV102 (6), a pPP2 derivative (21) containing the Zn^{2+} -inducible promoter P_{Zn} (4) or competence-inducible promoter P_{ssbB} (17), that integrates in the chromosome by double crossover at the nonessential *bgaA* locus. To exclude translational differences, the same ribosome binding site (RBS) was used for all RFPs. This allows for evaluation and unbiased comparison of the fluorescence intensities and folding times of the individual RFPs. A schematic presentation of chromosomally recombined RFP gene-carrying vectors is shown in Fig. 1.

Comparison of RFP intensities at the population and singlecell levels. The intensities of the different RFPs were analyzed at the population level using a plate reader and at the single-cell level using epifluorescence microscopy. First, to check whether all RFPs were expressed under our experimental conditions, we performed immunodetection on whole-cell lysates using an anti-RFP polyclonal antibody (Molecular Probes, Life Technologies). Cells were grown to exponential phase in the presence of Zn^{2+} for induction, and immunodetection was performed on whole-cell extracts, which allows for only relative quantification of protein levels. All fluorescent proteins were efficiently produced in *S. pneumoniae*, and only mCherry showed signs of protein degradation (Fig. 2A).

Fluorescence measurements with the plate reader were also performed on Zn^{2+} -induced cells in exponential phase which were resuspended in PBS. The plate reader we used contains a standard RFP filter set (excitation $[\lambda_{ex}]$, 590 nm; emission $[\lambda_{em}]$, 635 nm) which is best suitable for the far-red FPs. Not surprisingly, the signals for cells expressing TagRFP and mOrange2 hardly appeared above background. Interestingly, mCherry had the highest signal-to-noise ratio, with approximately 6-fold higher fluorescence than mKate2 (Fig. 2B), while mKate2 was reported to be brighter *in vitro* (12).

For fluorescence microscopy, cells were grown to an OD_{600} of 0.05 before expression of RFPs was induced with 0.15 mM ZnCl₂. Cells were then harvested for microscopy at different time points. We used filter sets that are standard in most fluorescence microscopes: mCherry (λ_{ex} , 575/25 nm; λ_{em} , 632/60 nm; QUAD 2 polychroic [580 to 630 nm]) and TRITC (λ_{ex} , 542/27 nm; λ_{em} , 594/45 nm; QUAD 1 polychroic [550 to 630 nm]). The same light output and exposure times were used for all samples. Using the mCherry filter set, signals from mCherry and mKate2 were already visible after 30 min of induction (Fig. 2C). After 120 min, TagRFP also showed good fluorescence, similar in strength to that of mKate2 but still approximately 6 times less bright than mCherry (Fig. 2C). With these imaging conditions, signals for mOrange2 could not be detected. When the TRITC filter set was applied, which should be better suited for mOrange2 and TagRFP than for the far-red RFPs, mCherry still outperformed the other RFPs (Fig. 2D). With this filter set, TagRFP gave a good signal above background, with ap-



FIG 1 Inducible RFP expression in *Streptococcus pneumoniae*. The scheme for RFP gene expression strains after integration of the P_{Zn} -*rfp* promoter fusion at the nonessential *bgaA* locus by double crossover is shown. Strains were constructed by transformation with plasmids pKB01- P_{Zn} -*rfp* that carry P_{Zn} -*rfp* (*mCherry*, *mOrange2*, *tagRFP*, or *mKate2*) and a tetracycline resistance marker between the homologous recombination sites (shown in gray; '*gatC* and '*bgaA*). Lollipops depict transcriptional terminators. Similar constructs where the P_{Zn} promoter was replaced with the P_{ssbB} promoter were also made (see Fig. S1 in the supplemental material).



FIG 2 Fluorescence intensities of the different red fluorescent proteins. (A) Western blot analysis of expression levels of RFPs in whole-cell extracts detected using anti-RFP antibody. (B) Mean fluorescence on the population level determined using a plate reader equipped with a standard RFP filter set. See Materials and Methods for details of the filters used. (C and D)Average fluorescence intensities on the single-cell level measured by fluorescence microscopy using an mCherry filter set and polychroic mirror QUAD2 (C) or a TRITC filter set and polychroic mirror QUAD1 (D). Values are in arbitrary units (A.U.). Single-cell fluorescence levels were determined using MicrobeTracker (20). Error bars depict the standard error of the mean. mOrange2 values are not shown in the plots since signals above background level could not be detected.

proximately half of that of mCherry for the different time points, while the signal from mKate2 was consistently very low, and mOrange2 still gave no signals above background levels after 3.5 h.

To rule out that fluorescence intensity differences depended specifically on the Zn²⁺ regulation, we swapped the zinc-inducible promoter P_{Zn} with the competence-induced promoter P_{ssbB} (17) and measured fluorescence intensities after 1.5 h of induction with the competence-stimulating peptide (CSP) using the same filter sets (see Fig. S1 in the supplemental material). The results are consistent with the ones we obtained for the zinc induction after 1.5 h. It should be noted that the TRITC filter set and the QUAD1 polychroic mirror might block a large amount of the emitted photons from mOrange2. Also, the slow fluorophore maturation time of mOrange2 (11) might be a probable explanation for the absence of signal and thus makes this FP unsuitable for imaging in S. pneumoniae. TagRFP gives appreciable signals with a standard TRITC filter set and is a good orange-red RFP for use in S. pneumoniae (Fig. 2D), but it might be less suitable in multicolor experiments that include yellow fluorescent proteins due to possible spectral overlap. In total, these results show that mCherry is the fastestmaturing RFP in S. pneumoniae with the highest fluorescence signal when used in promoter fusions. This is probably not simply because of more efficient protein production, since all four RFPs seem to be produced to approximate similar levels using qualitative Western blotting (Fig. 2A).

Fast assembly of protein fusion constructs by Gibson assembly. In previous studies, mCherry has been used to tag *S. pneumoniae* FtsZ (22) and the capsule protein Wze (CpsD) (5), while mKate2 has been used to tag the eukaryotic-type serine/threonine kinase StkP (19) and also FtsZ (23). However, a systematic comparison of how different RFPs perform when fused to proteins for localization studies has not been made.

In order to compare the performances of different RFPs in protein fusions, the far-red and fast-folding mCherry and mKate2, as well as the red-orange TagRFP, were all fused to the C termini of FtsZ and HlpA (Hup, SPD_0997). TagRFP has been suggested as a suitable FP for protein fusions but, unlike mCherry and mKate2, has not yet been used for this purpose in *S. pneumoniae.* mOrange was not included in this part of the study, since we did not observe any signal when this FP was expressed alone (Fig. 2).

While fusions to *hlpA* were constructed by conventional methods (see Materials and Methods), the FtsZ-RFP constructs were all made using Gibson's one-step ISO assembly of overlapping ds-DNA (Gibson assembly) (14) (Fig. 3A). With this method, the four different parts of the construct were amplified with 40-bp

FIG 3 Fast generation of RFP fusions to FtsZ in *S. pneumoniae*. (A) Gibson assembly procedure (14) for construction of FtsZ-RFP-expressing strains. (B) Images show localization of FtsZ fused to TagRFP, mKate2, and mCherry. Phase-contrast (PC), RFP, and overlay (merge) images are shown. Scale bar, 2 µm. (C) Immunodetection of FtsZ-RFP fusion proteins in whole-cell extract by anti-RFP antibody. Asterisks indicate degradation products.

overlaps, and the different parts could then be fused without any restriction or ligation steps. Since no restriction sites are needed, it is possible to choose the optimal sequence for protein linkage. Importantly, as *S. pneumoniae* readily takes up linear DNA fragments via the competence pathway, we wondered whether we could directly use the Gibson assembly product, which is assembled in 1 h, to transform *S. pneumoniae* without any intermediate cleanup or amplification steps. Indeed, transformation with unpurified Gibson mixes readily yielded correct transformants. Thus, this direct transformation of assembly products is a fast and efficient method for strain construction and is now the preferred construction method for linear fragments in our laboratory.

Characterization of RFPs in a fusion to FtsZ and HlpA. The highly conserved tubulin-like protein FtsZ polymerizes at the septum during early cell division. The localization pattern of the FtsZ ring is well studied in bacteria, making it an excellent candidate for comparing the performances of the three red fluorescent proteins. HlpA is the only known histone-like protein encoded in the *S. pneumoniae* D39 genome (24) and binds unspecifically to DNA and thus localizes to the nucleoid in a fashion similar to that for the intercalating DNA dye DAPI (4',6'-diamidino-2-phenylindole) (18). C-terminal fusions of FtsZ to tagRFP, mCherry, and mKate2 were obtained by Gibson assembly replacing wild-type FtsZ. We also attempted to construct C-terminal fusions of these fluorescent proteins to HlpA by integrating the fusion gene into the native locus. Interestingly, we obtained the *hlpA-mCherry* fu-

sion strain but were unable make strains expressing the HlpA-mKate2 and HlpA-TagRFP fusions. In our attempts to make these strains, we always observed mutations in the linker sequence generating premature stop codons or deletion of (parts of) the RFP genes. Therefore, we instead constructed merodiploid strains where the hlpA fusions are present as an extra copy immediately downstream of the native hlpA gene and thus also under the control of the endogenous hlpA promoter. These strains could be readily made (Table 1) (see Materials and Methods).

Fluorescence microscopy of cells growing in mid-exponential phase showed that the total amount of fluorescence of both FtsZ and HlpA fused to TagRFP was comparable to that of the fusions with mKate2, whereas signals of the mCherry fusion proteins were lower (data not shown). Signals for the FtsZ-mKate2 and FtsZ-TagRFP strains appeared as sharp defined Z rings at midcell even before deconvolution (Fig. 3B and 4A), whereas FtsZ-mCherry demonstrated a slightly more diffuse and heterogeneous signal (Fig. 3B and 4A). The same was the case for the HlpA-fusions; the meroploid TagRFP and mKate2 fusions appeared as sharp defined nucleoid-localized spots in the cytoplasm (Fig. 4B; see Fig. S2 in the supplemental material), whereas both HlpA-mCherry strains frequently demonstrated a more diffuse and heterogeneous signal (Fig. 4B; see Fig. S2 in the supplemental material).

Western blot analysis of both proteins showed the presence of degradation products in the FtsZ-mCherry strain and in both HlpA-mCherry strains, whereas for fusions with mKate2 and

FIG 4 Double labeling with RFP and GFP in *S. pneumoniae*. (A) Localization of FtsZ-RFP (different RFP variants in the different panels) and GFP-StkP. (B) Localization of HlpA-RFP (different RFP variants in the different panels) and GFP-StkP. Phase-contrast (PC), RFP, and GFP images are shown, in addition to an overlay of GFP and RFP (merge). Scale bars, 2 µm.

TagRFP, no or only marginal degradation was observed (Fig. 3C; see Fig. S2B in the supplemental material). Degradation of mCherry might lead to a pool of untagged HlpA, which might be the reason why the single copy HlpA-mCherry could be made whereas this was not possible with mKate2 and TagRFP. In total, these results show that mCherry originating from our codon-op-timized gene is more readily degraded than mKate2 and TagRFP in *S. pneumoniae* (Fig. 2A and 3C), making the latter RFPs more suitable for protein fusions. Note that protein functionality always needs to be tested, since FP fusions may not always be fully functional, as probably is the case for the HlpA-mKate2 and HlpA-TagRFP fusions.

Far-red fluorescent proteins are particularly suitable for double-labeling experiments in combination with GFP or YFP due to nonoverlapping excitation and emission spectra and thus limited chance for cross talk. We show that all three RFPs can be successfully used in combination with GFP using our standard filter sets, as exemplified by double-labeled strains showing the localization of two cell division proteins, FtsZ and StkP (Fig. 4A), or localization of the division site relative to the nucleoid (Fig. 4B).

Final remarks. Here we benchmarked four commonly used RFPs (mOrange2, TagRFP, mCherry, and mKate2) in the important human pathogen *S. pneumoniae*. The genes encoding these RFPs were codon optimized, and Western blotting showed that all

proteins are efficiently produced in S. pneumoniae. Under our experimental conditions, no fluorescent signals were detected with mOrange2, likely because of its long maturation time. The three other RFPs were all reported to be fast folding. However, mCherry clearly was the fastest-folding variant in our assays and generated the highest fluorescence signals, and it is thus the best RFP to use in promoter fusions. On the other hand, mCherry is also partially degraded, making it less suitable in protein fusions. It would be interesting to identify the degradation site in mCherry and mutate it to generate a superior mCherry. However, for the time being, we recommend the use of mKate2 or TagRFP as a red fluorescent localization marker protein in S. pneumoniae. Further, we showed that mKate2, TagRFP, and mCherry are suitable for dual-color imaging using standard filter sets. Finally, we demonstrated that direct transformation of Gibson assembly products could be used as a fast and straightforward cloning method in S. pneumoniae.

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