Title: Optimization of the DsRed fluorescent protein for use in Mycobacterium tuberculosis

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Running title: Optimized DsRed for M. tuberculosis
**ABSTRACT**

**Objective:** We have previously codon-optimized a number of red fluorescent proteins for use in *Mycobacterium tuberculosis* (mCherry, tdTomato, Turbo-635). We aimed to expand this repertoire to include DsRed, another widely used and flexible red fluorescent protein.

**Results:** We generated expression constructs with a full length DsRed under the control of one of three strong, constitutive promoters ($P_{hsp60}$, $P_{rpsA}$ or $P_{G13}$) for use in mycobacteria. We confirmed that full length DsRed (225 amino acids) was expressed and fluoresced brightly. In contrast to mCherry, truncated versions of DsRed lacking several amino acids at the N-terminus were not functional. Thus, we have expanded the repertoire of optimized fluorescent proteins for mycobacteria.

**Keywords:** fluorescent protein, mycobacteria, reporter system
INTRODUCTION

Fluorescent proteins (FPs) have become the work horses of molecular biology and microbiology, with numerous applications. A plethora of variants of *Aequorea victoria* green fluorescent protein (GFP) (1) and *Discosoma* sp red fluorescent protein (*DsRed*) (2) are available covering almost the whole light spectrum from green to infra-red (3). Mutant derivatives have been engineered with altered excitation and emission wavelengths, increased or decreased stability, resistance to photo bleaching, sensitivity to environmental stimuli and substrates, as well as time for fluorophore maturation, intrinsic brightness and multimeric formats (3, 4). We previously described the use of a range of red reporters, of which the brightest was mCherry (5). We wanted to expand our repertoire of proteins. Since *DsRed* has been widely used as a bright and stable reporter, we optimized constructs for its expression in *M. tuberculosis*.

MAIN TEXT

MATERIALS AND METHODS

*Bacterial culture*

*Escherichia coli* DH5α was cultured in LB medium or on LA agar. *M. tuberculosis* H37Rv was grown in Middlebrook 7H9 medium plus 10% v/v OADC (oleic acid, albumen, dextrose, catalase) supplement (Becton Dickinson) and 0.05% w/v Tween 80 or on Middlebrook 7H10 agar (Becton Dickinson) plus 10% v/v OADC. Hygromycin was used at 100 μg/ml where required.
**Construction of expression vectors**

The *DsRed* expression vectors were constructed as follows: a partial *DsRed* sequence was codon optimized for *M. tuberculosis*, synthesised and cloned into pUC57 (Genscript USA Inc.) to generate pRed1. The *DsRed* ORF was excised from pUC57 as a BamHI/HindIII fragment and cloned into pSMT3 (6) to generate pBlaze1. The *DsRed* ORF was extended three times by PCR to generate pRedA1, pRedB1 and pRedC1 using primers DsRed-F1 5’-GGA TCC ATG CGC TTC AAG GTG CGC ATG GAG GGC TCG GTG AAC-3’, DsRed-F2 5’-GGA TCC GAC GTG ATC AAG GAG TTC ATG CGC TTC AAG GTG CGC-3’ and DsRed-F3 5’-GGA TCC ATG GCC TCG TCG GAG GTG ATC AAG GAG TTC- together with the reverse primer DsRed-R 5’-AAG CTT TTA CAG GAA CAG GTG GTG CCG-3’. The restriction sites are underlined, potential start codons are in bold. The ORFs were excised and cloned into pSMT3 (6) as BamHI/HindIII fragments to generate pBlazeA1, pBlazeB1 and pBlazeC1 with *DsRed* under the control of the *hsp60* promoter. Plasmids pBlazeC8 and pBlazeC10 were generated by replacing P <sub>hsp60</sub> with P <sub>rpsA</sub> and P <sub>G13</sub> respectively. All three promoters should drive constitutive high level expression (5, 7, 8).

**Quantitation of fluorescence in whole cells**

*M. tuberculosis* was electroporated as described (9) and transformants selected with hygromycin. *M. tuberculosis* was grown to stationary phase, harvested, washed twice in 10 mM Tris pH 8.0 and resuspended in 10 mM Tris pH 8.0 to an OD<sub>580</sub> of 0.25, 0.10, 0.05 and 0.01 in 12 x 100 mm glass culture tubes. Fluorescence was measured on a
Shimadzu RF-1501 spectrofluorimeter (Shimadzu) with a detection range of 0-1015 relative fluorescent units at Ex/Em 558/583nm (5).

**Western analysis of fluorescent proteins**

Cell extracts were prepared from liquid cultures. Cells were harvested by centrifugation, washed twice in 10 mM Tris (pH 8.0), resuspended in 1 ml of 10 mM Tris (pH 8.0), and added to lysing matrix B tubes (QBiogene). Cells were disrupted using the Fastprep (QBiogene) set at speed 6.0 for 30 seconds. Samples were centrifuged for two min, and the supernatant was recovered and filter sterilized. Protein was quantified using a BCA kit (Pierce), and 10 μg of total protein was subjected to Western blot using a rabbit anti-body (Clonetech). The primary antibody was detected using horseradish peroxidase goat-anti-rabbit (Sigma), and activity was detected using an ECL kit (GE Healthcare).

**RESULTS**

We were interested in the use of FPs in *M. tuberculosis* and had previously used these as reporters of bacterial viability for *in vitro* and *in vivo* studies (5, 8). We were successful in obtaining high level expression by using codon-optimized versions of red fluorescent proteins driven by strong mycobacterial promoters (5).

**Optimization of DsRed expression**
We wanted to expand the range of reporters available for use to increase flexibility and allow dual reporter expression and monitoring. We selected *DsRed* for optimization, based on its Ex/Em wavelengths, and the fact that it is a well-characterized FP in wide use.

**Expression of DsRed uses a different translational start site than mCherry**

Our initial attempts to obtain expression of a codon-optimized *DsRed* were unsuccessful. We constructed a synthetic gene for *DsRed* using a similar approach as we used with another red fluorescent protein mCherry (Figure 1). We designed a codon-optimized version based on the *DsRed*-T3 protein previously used. We cloned the synthetic version into a mycobacterial expression vector and tested for fluorescence in *M. tuberculosis*. Surprisingly, we did not detect any fluorescence from this construct (Figure 1C).

mCherry is a variant of *DsRed* and we expected the two proteins would be similarly functional. Our previous work demonstrated that mCherry is expressed from a distal translational start site than the one annotated in the databases (10). Sequence alignment shows the few mutations which differ between the two (Figure 2A); these include loss of the translational start site we identified for mCherry, although there are still multiple translational start sites (Figure 1A). The version we used for the synthetic gene used a downstream translation start site and would produce a truncated version of *DsRed* as compared to mCherry. Therefore it was possible that we did not express the full protein (Figure 1B). In order to determine the functional start site for *DsRed* we used
a different approach in which we cloned several versions of the coding region into the expression vector under the control of the constitutive *hsp60* promoter (Figure 2B).

In order to test this, we used PCR amplification to extend the region sequentially. We extended the gene to incorporate both additional start sites and generate proteins of 214, 220 and 225 amino acids. These variants were cloned into the same mycobacterial expression system and tested. Plasmids were transformed into *M. tuberculosis* and fluorescence was monitored. In contrast to mCherry, expression of a functional fluorescent *DsRed* was not seen with any truncated versions of the protein. In fact fluorescence could only be detected when the full length amino acid sequence (as annotated) was cloned into the expression vector; high level fluorescence was seen with transformants carrying the plasmid pBlazeC1 (Figure 2B).

We constructed two alternative vectors with *DsRed* under the control of either *P*<sub>rpsA</sub> or *P*<sub>G13</sub> (pBlazeC8 and pBlazeC10 respectively); both of these constructs gave high level expression in *M. tuberculosis*. Western blotting using an anti-*DsRed* antibody in *E. coli* demonstrated that a protein of the expected size was only seen in bacteria carrying the full length construct (pBlazeC series), but not in the strains carrying the truncated version (Figure 2C).

**DISCUSSION**

We have determined that the functional translational start sites for two closely related FPs are different in *M. tuberculosis*. Although mCherry was functional even when a
truncated version was being expressed, DsRed was non-functional in a truncated form and only fluoresced when expressed as a full length protein (225 amino acids). Western blotting suggested that the lack of fluorescence was most likely due to a lack of protein expression, since proteins could not be detected in the plasmids carrying truncated forms. This difference may relate to protein stability, with the extended N-terminal portion of DsRed increasing stability or protein maturation; alternatively this could be attributed to the physical state of the active proteins, since mCherry functions as a monomer, whereas DsRed is a tetramer which might also affect protein degradation.

In conclusion, we have codon-optimized DsRed for use in M. tuberculosis and demonstrated its high level fluorescence in that species from three different promoters of slightly varying strength (hsp60, rpsA, and G13). These vectors extend our current repertoire of functional fluorescent proteins for mycobacteria. They will be useful for generating fluorescent strains of M. tuberculosis for use in multiple studies, such as monitoring drug efficacy in vitro and in vivo (5, 8, 11, 12) and will allow for detection of multiple reporters simultaneously.

Limitations

- We have monitored the expression of DsRed under aerobic conditions only.
- We have not monitored long term stability of expression in the absence of antibiotic selection to maintain the plasmid.
- We have not monitored stability of expression in vivo.
Declarations

Ethics approval and consent to participate
Not applicable

Availability of data and materials
All data generated or analysed during this study are included in this published article.

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Consent for publication
Not applicable

Competing interests
The author(s) declare(s) that they have no competing interests.

Author’s contributions
Experimental design: PC, JM, TP
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Reviewing manuscript: PC, JM, TP
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Abbreviations
FP- fluorescent protein; OADC - oleic acid, albumin, D-glucose, catalase.

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Figure Legends

Figure 1. Expression of non-functional DsRed. (A) DsRed Protein sequence. Three potential translational start sites (methioinine) are indicated in bold. The valine which corresponds to the methionine start site of mCherry in *M. tuberculosis* is also indicated in bold. (B) DNA sequence of DsRed. The 5’ end of the synthetic gene designed to codon-optimize DsRed for *M. tuberculosis* is boxed. Potential starts sites are indicated in bold. The Shine Delgarno sequence is underlined. (C) *M. tuberculosis* was resuspended in 10 mM Tris pH 8.0 to an OD$_{580}$ of 0.25, 0.10, 0.05 and 0.01 in 12 x 100 mm glass culture tubes. Fluorescence was measured at Ex/Em 558/583nm. WT – wild-type (no plasmid). pBlaze1 – recombinant strain carrying DsRed 208aa. Data are the average ± SD of three cultures.

Figure 2. Expression of functional DsRed. (A) Sequence alignment of mCherry and DsRed. Protein sequences were aligned using Clustal (13). (B) Activity of full length DsRed expressed from mycobacterial promoters. pBlazeC1 – P$_{hsp60}$; pBlazeC8 - P$_{rpsA}$; pBlazeC10 - P$_{G13}$. Recombinant *M. tuberculosis* was resuspended in 10 mM Tris pH 8.0 to an OD$_{580}$ of 0.25, 0.10, 0.05 and 0.01 in 12 x 100 mm glass culture tubes. Fluorescence was measured at Ex/Em 558/583nm. Data are the average ± SD of three cultures. (C) Plasmids were transformed into *E. coli* and cell-free extracts analyzed by Western blotting; 10 µg protein were subjected to SDS-PAGE, blotted onto PVDF membrane and visualized with anti-DsRed antibody. Lane 1s and 11 – empty; Lane 2 – *E. coli* (no plasmid); Lane 3 – pRed1; Lane 4 – pRedA1; Lane 5 – pRedB1; Lane 6 –
pRedC1; Lanes 7 and 8 – pBlazeC1; Lane 9 - pBlazeC8; Lane 10 – pBlazeC10. The arrow indicates the size of the DsRed protein.
### Tables

| Plasmid   | Description                                      | Promoter | Fluorescent protein | Source         |
|-----------|--------------------------------------------------|----------|---------------------|----------------|
| pSMT3     | Shuttle vector, $P_{\text{hsp60}}$, HygR         |          |                     | (6)            |
|           | Codon-optimized $Ds\text{Red}$ in                |          |                     |                |
| pRED1     | pUC57                                            | None     | $Ds\text{Red208}$   | This study     |
| pBlaze1   | $Ds\text{Red}$ expression vector. HygR          | Hsp60    | $Ds\text{Red208}$   | This study     |
| pBlazeA1  | $Ds\text{Red}$ expression vector. HygR          | Hsp60    | $Ds\text{Red214}$   | This study     |
| pBlazeB1  | $Ds\text{Red}$ expression vector. HygR          | Hsp60    | $Ds\text{Red220}$   | This study     |
| pBlazeC1  | $Ds\text{Red}$ expression vector. HygR          | Hsp60    | $Ds\text{Red225}$   | This study     |
| pBlazeC8  | $Ds\text{Red}$ expression vector. HygR          | RpsA     | $Ds\text{Red225}$   | This study     |
| pBlazeC10 | $Ds\text{Red}$ expression vector. HygR          | G13      | $Ds\text{Red225}$   | This study     |

Table 1. Plasmids used in this study.
Figure 1

(A)

MASSEDVKEFMRFKVMGSGVNGHEFEIEEGEGRRPETFQTDKTKVTKGGLP
FAYDILSPQPQYGSKVVKHPADIPDYKLYSSLPSPEGMFKWVRVNMFDGGVVTQD
SSLQDGCIYKVKIFIGNFSDGPVMQKKTGMWEPSTERLYPRDGVKIMGHKL
KLKDGCHYLFKSMAKKPVQLPGYYVDSKLDITSHNEYTVTVEQYERTEGG
HNLFL

(B)

ATGCCTCCTCGGAGGCATGATCAAGGAGTTTATGCCTTTCAAGGTGGCAGATGG
AGGGCTCGGTGAACGCGCCACGAGTCTCGAGATCGAGGCAGGCGGAGGGCGGCGCC
GCTACGGACCCGAGACCCGAGAAGCTGAAGGTCAACAGGACGGGCGGCGCCGCGCC
TTCACTGCGGACATCCTGTGCACGCACTTCCAGTTGACAGCAAGGTGTACGTCA
AGCCTCCGGCCGACATCGCCGACTACAGAAGCTGAGCTTCTCCGGAGAGGCTTCAA
GTGGGACGCGGTGATGAACTTCTGAGACGGCGGCTGCTGACCGTACCCGACCGAC
TGGACCTGGACGACGCTGTCTACATCTCTACAGGAGTTGATCAGTTGCTCGACGCTCAACT
TCCGGAGCCACGCCCTGGATGCAGAAGAAGACCCAGGGCTGGGACGCCTGACG
CGAGCCTCTGTACCACCGGAGCGCTGCTCGAGAGGGCGGCTCCACACACCAGCTG
AGCTGAGAAGCGCGCCACCTGCTGTACCCGCCTGGACGACCCGACGACCTGCCTGACG
AGAAGCCGCGGCAGCTCAGCTCAGTCACGCTACAGCTGCTCAGCAGAGGCTAG
CAGGCACCAACGAGGACTACACCATCGCTCGACGCTACGAGCGCAGGGCCG
CACCACTTCTGCTGTAA

(C)
Figure 3

(A) DaRed
nCherry
MKLASSKEN---VTIEFNSRFKVRMEGTVHGMHFEIEIGEGEGRFYEGRHNYTVKLKV(TG2SPLP 57
MVSKSEEEDMMATKRERNSRFKVRMEGTVHGMHFEIEIGEGEGRFYEGRQTAlKL60

DaRed
nCherry
FAEMILSOPQQGVSQGTVKIHDPQDIYWGEPFPEKQVMNSHEDGTVATQD117
FAEMILSOPQQGVSQGTVKIHDPQDIYWGEPFPEKQVMNSHEDGTVATQD120

DaRed
nCherry
GCCFVKVYTFIGVYFQGIFQKINGWEASTERLYFA6GVLKGEKALKLKE6DVHVLV 177

DaRed
nCherry
EFHSIAQAQKFCQLPGYYDVDAKLDITSEHEDYTIVEQHITAEKSHKFLFRS6APPFF6L 237
nCherry
EVRTYKAGKFCQLPGAYNVNIKLIDISSEHEDYTIVEQHITAEKSHKFLFRS6APPFF6L 175

DaRed
nCherry
T 238
nCherry
K 176

(B) Graph showing fluorescence levels for WT, pBlazeC1, pBlazeC8, and pBlazeC10.

(C) Image showing protein bands for WT, pBlazeC1, pBlazeC8, and pBlazeC10.