A transient CRISPR/Cas9 expression system for genome editing in *Trypanosoma brucei*

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**Abstract**

**Objective:** Generation of knockouts and in situ tagging of genes in *Trypanosoma brucei* has been greatly facilitated by using CRISPR/Cas9 as a genome editing tool. To date, this has entailed using a limited number of cell lines that are stably transformed to express Cas9 and T7 RNA polymerase (T7RNAP). It would be desirable, however, to be able to use CRISPR/Cas9 for any trypanosome cell line.

**Results:** We describe a sequential transfection expression system that enables transient expression of the two proteins, followed by delivery of PCR products for gRNAs and repair templates. This procedure can be used for genome editing without the need for stable integration of the Cas9 and T7RNAP genes.

**Keywords:** CRISPR/Cas9, Transient transfection, Trypanosome

**Introduction**

The establishment of genome editing by CRISPR/Cas9 in trypanosomatids has greatly increased the ease with which knockouts can be generated, as two copies of a non-essential gene can often be deleted in a single round of transfection. The system most widely used for *Trypanosoma brucei* entails creating cell lines which express (either constitutively or inducibly) both Cas9 to generate a DNA double-strand break and T7RNAP for the transcription of guide RNAs from a DNA template [1–4]. The original pTB011 plasmid generated for *T. brucei* [4] encodes Cas9 flanked by tubulin sequences, enabling the construct to be integrated into the corresponding multicycopy array. We modified this plasmid by replacing the α-tubulin sequence upstream of T7RNAP with 366 bp of the EP procyclin promoter and 5′ UTR [6]. To test the functionality of the plasmid, we used the LeishGEEdit software [4] to design a guide RNA that allows tagging of the C-terminus of phosphodiesterase B1 (PDEB1) with mNeonGreen (mNG). The repair template, in the form of a purified PCR product, including homology arms of 30 bp on each side, was amplified from pPOTv7 [7]. As a positive control we transfected a derivative of *T. brucei* Lister 427 procyclic forms that already

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expresses Cas9 and T7RNAP constitutively (Lister 427/Cas9) [5]. This gave rise to cells with a fluorescent flagellum as expected (tryptag.org) (Fig. 2a).

For genome editing by transient expression of Cas9 and T7RNAP, we tested two different transfection schemes. Initially, all components were transfected simultaneously: the circular plasmid pAi1C9 encoding Cas9 and T7RNAP, the DNA template for in vivo transcription of a guide RNA, and the DNA repair template harbouring mNG for C-terminal tagging and a hygromycin resistance gene. This procedure gave few hygromycin-resistant clones, only 10 clones in total, from 4 separate transfections. These clones either did not express mNG (4 clones), or expressed it as a cytoplasmic protein (2 clones) or expressed it correctly localised to the flagellum (4 clones). Examples are shown in Fig. 2b. We hypothesise that, due to insufficient expression of Cas9 and T7RNAP, the amounts of site-specific guide RNAs and double-strand breaks at the correct locus were too low to drive efficient integration by homology-directed repair.

We therefore tested sequential transfections in which we first used pAi1C9 to enable expression of Cas9 and T7RNAP. A second transfection 20 h later provided the templates for the gRNA and the mNG repair construct. The same electroporation conditions were used for the plasmid and the templates (see Methods and Fig. 3). This procedure yielded 20 clones. There were 16 clones in which mNG was correctly localised to the flagellum, one in which it was cytoplasmic, 2 clones with mixed populations of mNG localised either to the flagellum or the cytoplasm, and one that was negative (see Fig. 2b for representative examples). One clone with flagellar mNeonGreen had a fluorescence intensity twice that of the others (clone B1) and might have been tagged on both alleles.

We have also used this procedure of sequential transfections to simultaneously knock out both copies of trypanin or GPI-anchor transamidase subunit 8 (GPI8), which are non-essential genes in cultured procyclic form T. brucei [8, 9]. Genotyping data is provided in Additional file 1.

**Methods**

**Transfection of pAi1C9**

$4 \times 10^7$ T. b. brucei 427 procyclic forms were transfected with 10 µg pAi1C9 dissolved in 100 µl TbBSF transfection buffer [10], using an Amaxa Nucleofector IIB (Lonza), program X-014 [11]. Cells were transferred to 13 ml SDM79 medium [12] supplemented with 10% FBS and incubated at 27 °C and 2.5% CO$_2$ for 20 h.

**Transfection of pooled PCR products for repair constructs and sgRNA templates**

The entire culture from the first transfection was centrifuged at 1700 g for 5 min, the supernatant discarded and the cells resuspended in 100 µl TbBSF transfection buffer containing the pooled PCR products (see protocol for PCRs below). For tagging one sgRNA template (targeted to the 3’ end of the ORF) and one repair template (hygromycin resistance) were provided; to generate the knockouts, we provided two sgRNA templates (targeted to the 5’ and 3’ ends of the ORF) and two repair templates (hygromycin and neomycin resistance genes).

Transfection was performed as described above and the cells were transfected to 10 ml SDM79 supplemented with 10% FBS. The cells were diluted 1:5, 1:50

![Fig. 1 Plasmid maps of a pTB011_Cas9_T7RNAP_blast and b pAi1C9](image-url)
Fig. 2 Trypanosomes tagged with mNeonGreen at the C-terminus of phosphodiesterase B1 (PDEB1). a Tagging PDEB1 in Lister 427/Cas9. Scale bar: 10 microns. b Tagging PDEB1 in Lister 427. Fluorescence intensity of live cells was quantified with a benchtop flow cytometer (ACEA NovoCyte). To remove particles of subcellular size, a cut-off of $3 \times 10^4$ was applied to the forward scatter. A total of $10^4$ events were recorded and analysed using FlowJo software without gating. TF transfection
and 1:500 in conditioned medium (fresh medium + 20% supernatant of a log-phase culture) and distributed into 24-well plates (1 ml in each well). Transformants were selected using 25 µg ml\(^{-1}\) Hygromycin B and/or 15 µg ml\(^{-1}\) Geneticin; stable clones were obtained 2 weeks post selection.

**Polymerase chain reactions (PCR)**

Reactions were performed with reagents from New England Biolabs: Phusion High-Fidelity DNA polymerase (M0530S), 5x Phusion HF buffer (B0519S) and dNTP mix (N0447S). Cycling conditions are identical to those previously published [4, 13].

- 1st PCR: template for sgRNAs; two 20 µl reactions per target.
  
  1 µl G00 primer (0.5 µM)

- Program:
  1) 30'', 98 °C
  2) 10'', 98 °C
  3) 30'', 60 °C
  4) 15'', 72 °C
  5) go to step 2, 35 cycles in total
  6) 10', 72 °C
  7) hold at 10 °C

Fig. 3 Flow chart of sequential transfections. Procyclic form trypanosomes were transfected first with pAi1C9 to allow expression of Cas9 and T7RNAP. After 20 h, the pool of trypanosomes was transfected a second time to provide the templates for gRNA(s) and the repair template(s)/resistance cassette(s).
– 2nd PCR: template for resistance gene; two 40 µl reactions per resistance gene

2 µl 60 ng pPOTv7 mNG (hygro or G418) plasmid
8 µl 5 x Phusion HF buffer
1 µl dNTP mix (250 µM)
2 µl Upstream forward primer (0.5 µM)
2 µl Downstream reverse KO/TAG primer (0.5 µM)
0.4 µl Phusion High-Fidelity DNA polymerase (0.4U)
24.6 µl H2O

– Program:
1) 5′, 94 °C
2) 30 ′′, 94 °C
3) 30 ′′, 65 °C
4) 2′30′′, 72 °C
5) go to step 2, 40 cycles in total
6) 10′, 72 °C
7) hold at 10 °C

DNA purification after PCR
PCR reactions were pooled and extracted with 1 volume water-saturated phenol (pH 8), followed by extraction with 1 volume chloroform. DNA was precipitated from the aqueous phase by addition of 0.1 volume 3 M sodium acetate, pH 5.2, and 3 volumes ice-cold ethanol. The DNA was pelleted by centrifugation, washed twice with 1 ml 80% ethanol, air-dried at room temperature, dissolved in 40 µl Milli-Q-water and stored at −20 °C until transfection.

Primers
G00: aaaaagcaccgactcggtgccttattttactgtgatgtacggactagtcttttaacttgctatttctagctctaaaac
PDEB1 3′ sgRNA primer: gaattaatacactcatacgggaggatcagttgacagtgtgttttagagctagaaatagc
Trypanin 5′ sgRNA primer: gaattaatacactcactattgcaagggcagtaaaggggacacttatttttaacttgctatttctagctctaaaac
Trypanin 3′ sgRNA primer: gaattaatacactcactattgcaagggcagtaaaggggacacttatttttaacttgctatttctagctctaaaac
CCTAcgttttagcgtagcaaatagc
Trypanin 3′ sgRNA primer: gaattaatacactcactattgcaagggcagtaaaggggacacttatttttaacttgctatttctagctctaaaac
GTggttttagcgtagcaaatagc
GPI8 5′ sgRNA primer: gaattaatacactcactattgcaagggcagtaaaggggacacttatttttaacttgctatttctagctctaaaac
GPI8 3′ sgRNA primer: gaattaatacactcactattgcaagggcagtaaaggggacacttatttttaacttgctatttctagctctaaaac
GAggttttagcgtagcaaatagc
Trypanin upstream forward primer: TACATTTGACTGCTACGTGCGTACCCGTGATAATCGACGCTGC
Trypanin downstream reverse KO primer: CTGCAACAAAGCGTAGTAAGTTGGAACACGCA cgggaaccactcagaacc
PDEB1 downstream forward primer: ACGAGTTCTGGACAAACAGCAGTCATCTGTTggtctgttagtttccgg
PDEB1 downstream reverse TAG: ATACACCATTGACAAGACGTACATCTACCA ccaattttagcagacgctgc
GPI8 upstream forward primer: TTGGATCAGGCGCTTTGCAATAATATTTATTTCAGACgttttagagctagaaatagc
GPI8 downstream reverse KO primer: AGTTCAGGAAAGGTTGCTTTTTCCTCCTccggagaaccactcagaacc

Limitations
Transient transfection with pAi1C9 gives rise to fewer clones than cell lines that are stably transformed with the T7RNAP and Cas9 genes. It has the advantage, however, that it can be applied to any T. brucei cell line. Moreover, in contrast to stable integration of the CRISPR/Cas9 machinery, transient transfection does not require additional selectable markers and it has the added advantage that it circumvents possible Cas9 toxicity. In addition to using Cas9 for deletion, mutation, tagging or integration of ectopic copies, this procedure could also be adapted for nuclease inactive Cas9 variants for targeting RNAs or epigenetic modifications [14, 15].

Supplementary information
Supplementary information accompanies this paper at https://doi.org/10.1186/s13104-020-05089-z.

Additional file 1: Validation of knockout (KO) clones. A+B Trypanin-KO (Tb427.10.6350). A Clonal selection of stable transformants in 24-well plates. Dilutions from transfected pool cultures and number of cells seeded are indicated. Circles depict wells containing cells with successful (●) or unsuccessful (○) genome editing. B Assessment of editing by genotyping PCR and agarose gel electrophoresis. Genes amplified from genomic DNA and target loci are indicated. Lanes corresponding to validated KO clones are marked with green circles. C+D GPI8-KO (Tb427.10.13860) in a cell line allowing inducible expression of an ectopic copy of GPI8. C as in A. D as in B. E List of primers used for genotyping PCRs of Trypanin-KO (left panels) and GPI8-KO (right panels).

Abbreviations
mNG: mNeonGreen; PDEB1: Phosphodiesterase B1; T7RNAP: T7 RNA polymerase.

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Authors' contributions
Designed study: SS and IR, performed experiments: SS, SH and SK, analysed data: SS, SK and IR; wrote the paper: SS, SK and IR. All authors read and approved the final manuscript.

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Availability of data and materials
Plasmids are available on request from Isabel Roditi (Isabel.roditi@izb.unibe.ch). The nucleotide sequences of pTB011_Cas9_T7RNAP_blast and pAi1C9 are provided on this website: https://www.izb.unibe.ch/research/prof_dr_isabel_rodit/index_eng.html.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
We, the authors, declare no conflict of interests.

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