Serological Responses and Biomarker Evaluation in Mice and Pigs Exposed to Tsetse Fly Bites



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Abstract

Background: Tsetse flies are obligate blood-feeding insects that transmit African trypanosomes responsible for human sleeping sickness and *nagana* in livestock. The tsetse salivary proteome contains a highly immunogenic family of the endonuclease-like Tsal proteins. In this study, a recombinant version of Tsal1 (rTsal1) was evaluated in an indirect ELISA to quantify the contact with total *Glossina morsitans morsitans* saliva, and thus the tsetse fly bite exposure.

Methodology/Principal Findings: Mice and pigs were experimentally exposed to different *G. m. morsitans* exposure regimens, followed by a long-term follow-up of the specific antibody responses against total tsetse fly saliva and rTsal1. In mice, a single tsetse fly bite was sufficient to induce detectable IgG antibody responses with an estimated half-life of 36–40 days. Specific antibody responses could be detected for more than a year after initial exposure, and a single bite was sufficient to boost anti-saliva immunity. Also, plasmas collected from tsetse-exposed pigs displayed increased anti-rTsal1 and anti-saliva IgG levels that correlated with the exposure intensity. A strong correlation between the detection of anti-rTsal1 and anti-saliva responses was recorded. The ELISA test performance and intra-laboratory repeatability was adequate in the two tested animal models. Cross-reactivity of the mouse IgGs induced by exposure to different *Glossina* species (*G. m. morsitans, G. pallidipes, G. palpalis gambiensis* and *G. fuscipes*) and other hematophagous insects (*Stomoxys calcitrans* and *Tabanus yao*) was evaluated.

Conclusion: This study illustrates the potential use of rTsal1 from *G. m. morsitans* as a sensitive biomarker of exposure to a broad range of *Glossina* species. We propose that the detection of anti-rTsal1 IgGs could be a promising serological indicator of tsetse fly presence that will be a valuable tool to monitor the impact of tsetse control efforts on the African continent.

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Introduction

Tsetse flies (*Glossina* spp.) are notorious transmitters of trypanosome parasites responsible for Human and Animal African Trypanosomiasis (HAT and AAT). Since 2009, the annual number of reported cases of HAT has dropped below 10000 (www.who.int; [1]) with the prospect and challenge of entering into the elimination phase of HAT in the near future [2,3]. Additionally, some 46 million cattle in sub-Saharan Africa are estimated to be at risk of contracting AAT making deep inroads in the socio-economical development of this continent [4]. Beside active HAT case detection and treatment of humans as well as prophylactic and curative treatment of animals with trypanocidal drugs, tsetse vector control represents an important component of trypanosomiasis control, which is mainly based on the use of insecticides through the sequential aerosol spraying technique (SAT), ground spraying, insecticide-treated targets or insecticidetreated animals [reviewed in [5,6,7]]. After a successful campaign as part of an area-wide integrated pest management on Unguja island (Zanzibar) [8], the sterile insect technique has been added to the vector control arsenal, with ongoing activities in Ethiopia, Senegal and Burkina Faso [9] under the auspices of the Pan African Tsetse and Trypanosomosis Eradication Campaign (PATTEC). However, beside laborious conventional

Author Summary

Salivary proteins of hematophagous disease vectors represent potential biomarkers of exposure and could be used in serological assays that are complementary to entomological surveys. We illustrate that a recombinant version of the highly immunogenic Tsal1 protein of the savannah tsetse fly (Glossina morsitans morsitans) is a sensitive immunological probe to detect contact with tsetse flies. Experimental exposure of mice and pigs to different regimens of tsetse fly bites combined with serological testing revealed that rTsal1 is a sensitive indicator that can differentiate the various degrees of exposure of animals. Tsetse-induced antibodies persisted relatively long, and an efficient boosting of immunity was observed upon re-exposure. Recombinant Tsal1 is a promising candidate to detect contact with various tsetse species, which would enable screening of populations or herds for exposure to tsetse flies in various areas on the African continent. This exposure indicator could be a valuable tool to monitor the impact of vector control programs and to detect re-invasion of cleared areas by tsetse flies.

entomological surveys, no sensitive rapid tests are yet available to provide a semi-quantitative measure of the evolution of tsetse fly densities in areas subjected to tsetse control interventions. Indeed, easy-to-use monitoring of tsetse fly exposure on a regular basis would be a highly valuable tool in the follow-up of the efficacy of the applied and/or ongoing tsetse fly control activities.

The obligatory blood feeding tsetse flies are the cyclical insect vectors of HAT and a majority of AAT infections are initiated by the bite of an infected tsetse fly. Although it can be assumed that all tsetse fly species could act as vector, a number of Glossina species of the Palpalis group (e.g. G. palpalis spp., G. fuscipes spp., G. tachinoides) and the Morsitans group (e.g. G. morsitans spp., G. pallidipes, G. swynnertoni) are implicated as major vectors for HAT and animal trypanosomiasis. Given that only a limited percentage of these tsetse flies acquire an infection with trypanosomes, vertebrate hosts living in the tsetse fly belt are predominantly exposed to the bites of uninfected flies. It has been demonstrated for a number of hematophagous insects that salivary proteins induce humoral immune responses that could represent attractive sero-epidemiological markers of exposure (reviewed in [10]). The saliva of Glossina morsitans morsitans tsetse flies was documented to contain over 200 protein constituents [11] from which some are implicated in manipulating the vertebrate hemostatic and inflammatory reactions [12,13,14]. In G. m. morsitans saliva, the most abundant proteins were shown to be highly immunogenic and to correspond to the 43-45 kDa tsetse salivary gland (Tsal) protein family [15]. The physiological role of these proteins remains elusive, but biochemical characterization revealed that they are nucleic acid binding proteins with low endonuclease activity [16]. Immunoglobulin responses to tsetse fly saliva have been detected in humans living in Uganda [15], Democratic Republic of Congo [17,18] and Guinea [19]. Also cattle experimentally exposed to tsetse fly bites displayed elevated levels of anti-saliva antibodies [20]. Immunoblotting studies using the immune plasmas have shown that salivary proteins of several tsetse fly species are recognized by the circulating antibodies [15,18,19]. The highly abundant Tsal proteins were commonly recognized by the human plasmas and an indirect ELISA using recombinant Tsal1 and Tsal2 proteins as antigens was clearly able to differentiate the tsetse-exposed Ugandan plasmas from control plasmas [15].

Recently, a peptide (amino acids 18–43) derived from the *G. m. morsitans* adenosine deaminase-related TSGF1 protein was evaluated using a panel of human plasmas from West Africa, revealing that obtained ELISA signals correlated with the anticipated levels of tsetse exposure of the tested populations [21]. Allergic and anaphylactic reactions against tsetse fly bites have also been reported, in which IgE antibodies directed against an Antigen5-related salivary allergen are implicated [22,23,24]. Efforts to develop a serological test based on a TAg5-derived peptide were not yet successful [21].

We here provide experimental evidence that anti-rTsall and anti-G. m. morsitans saliva antibodies can mark exposure of mice and pigs to tsetse flies. Although the anti-tsetse saliva ELISA exhibits a better average test performance, we propose that a serological assay based on the individual recombinant Tsall protein could be an alternative to assess the exposure of populations or herds to tsetse fly challenge and hence could be used for tsetse fly epidemiological studies, for prioritizing tsetse fly control, for monitoring and evaluating tsetse fly control schemes and for risk assessment of trypanosome transmission in endemic regions.

Methods

Ethics statement

The experiments, maintenance and care of animals complied with the guidelines of the European Convention for the Protection of Vertebrate Animals used for Experimental and other Scientific Purposes (CETS n° 123). Rodent care and experimental procedures were performed under approval from the Animal Ethical Committee of the Institute of Tropical Medicine (Permit Nrs. PAR013-MC-M-Tryp and PAR014-MC-K-Tryp). Breeding and experimental work with tsetse flies was approved by the Scientific Institute Public Health department Biosafety and Biotechnology (SBB 219.2007/1410). Pig experiments were approved by the ITM Animal Ethical Committee (Permit Nr. PAR-021) and the Ethical Committee of the Faculty of Veterinary Medicine of Ghent University (Permit Nr. EC2010/030) and were performed in the Ghent university stables under the supervision of a veterinary doctor.

Salivary antigens and recombinant proteins

Saliva of Glossina m. morsitans from the ITMA tsetse fly colony was collected as outflow from the salivary glands as described elsewhere [16]. Saliva of Tabanus yao was kindly provided by Prof. Ren Lai (Kunming Institute of Zoology, Yunnan, China), Stomoxys calcitrans saliva was harvested from flies received from Dr. Christopher J. Geden (United States Department of Agriculture, Gainesville, US). Recombinant Tsall was purified as described elsewhere from inclusion bodies of IPTG-induced Top10F' Escherichia coli host cells harboring a pQE60:Tsal1 plasmid [15]. Tsal1 was resolubilized in 6M guanidinium hydrochloride, enriched by Ni-NTA affinity chromatography (Qiagen) and further purified on a Superdex 200 size exclusion column connected to an Akta Explorer (GE Healthcare) in 6 M ureum 50 mM Tris pH 8.0 and 600 mM NaCl. Protein concentrations were determined using Nanodrop spectrophotometry and samples were stored in aliquots at -20° C. E. coli soluble extract was prepared as an additive for the porcine ELISA assay diluent as described for other porcine assays [25]. The extract was made from a 1L overnight culture of Top10F' E. coli in Terrific broth. The bacterial pellet was resuspended in 12 ml PBS supplemented with a Complete protease inhibitor cocktail tablet (Roche), followed by 5 rounds of 1 minute sonication. The soluble fraction was harvested as the supernatant after 30 minutes centrifugation at $20.000 \times g$. *E. coli* soluble extract was stored in aliquots at -20° C.

Experimental study animals and immunization

Groups of eight female outbred mice (NMRI, Charles River) were subjected to different intensities of exposure to G. m. morsitans bites, followed by regular blood sampling and evaluation of the antibody responses in ELISA: (i) once exposed to a single fly, (i)once exposed to 10 flies, (iii) 3 times per week for 3 weeks exposed to a single fly, (iv) 3 times per week for 3 weeks exposed to 10 flies and (v) not exposed to tsetse fly bites. One mouse that underwent multiple exposures to a single fly succumbed by day 28 after initiation of exposure. Plasmas were collected over a period of 390 days. After this >1 year non-exposure period, six mice that were exposed to single or multiple (10) flies were selected based on their physical appearance and behavior and were re-exposed to a single G. m. morsitans tsetse fly followed by weekly plasma collection for up to 42 days after re-challenge. Two mice that were previously exposed to the multiple fly bites succumbed within 4 weeks after the boosting.

A *Glossina* species cross-reactivity study was performed by exposing five mice (OF1, Charles River) per group every 3 days for 6 weeks to 10 flies of either *G. m. morsitans, G. p. gambiensis, G. pallidipes* (kindly provided by Peter Takac, Institute of Zoology, Bratislava, Slovakia) or *G. f. fuscipes* (kindly provided by the International Centre of Insect Physiology and Ecology, Mbita Point, Kenya). Immune plasma was harvested 10 days after the last exposure. Five non-exposed mice served as negative controls. A cross-reactivity study between salivary antigens of hematophagous insects was performed by intrapinna exposure of 6 OF1 mice per group at 3-weekly interval to decreasing amounts of saliva (5, 2, 1 and 1 µg) harvested from *Glossina morsitans, Stomoxys calcitrans* and *Tabanus yao.* Plasma samples were collected 10 days after the last immunization.

A total of 11 female pigs (Seghers, Belgium), hybrids of Belgian Landrace, Large White and a specific company line were used for experimental exposure to three different G. m. morsitans exposure regimens: (i) 5 pigs were weekly exposed for 7 weeks to 30 tsetse flies (high exposure), (ii) 5 pigs were exposed 2-weekly for 6 weeks to 3 tsetse flies (low exposure, 1 pig died at day 7) and (iii) 1 pig was not exposed to tsetse (negative control). The experimental pigs were 6 weeks old at the start of the experiment following a 4-days acclimatisation period in the university stables (Faculty of Veterinary Medicine, UGhent). For anaesthesia of the pigs, intramuscular injection of midazolam (0.5 mg/kg), ketamine (10 mg/kg) and morphine (0.1 mg/kg) was used. Blood was collected every week for 11 successive weeks from the jugular vein using Vacutainer EDTA tubes (BD) a few minutes after the pigs had been anesthetised and prior to exposure to the tsetse fly bites. After blood collection, the blood was centrifuged at 3000 rpm for 15 minutes and plasma stored at -20° C. Due to the unexpected death of 1 pig, no tsetse challenge and blood sampling was performed on day 7 for all animals.

Due to limited housing capacity in the university stables, only two pigs from the low exposure group could be kept for an additional 2-month period of non-exposure, followed by a rechallenge by the bites of 10 *G. m. morsitans* flies and a weekly plasma collection over a period of 6 weeks.

Serological analyses

IgG responses in the exposed animals were analyzed by indirect ELISA against rTsal1 and saliva from different hematophagous insects. For this purpose, polystyrene 96 well plates (Thermo

Scientific NUNC MaxiSorp Surface) were coated overnight at 4°C with 200 ng antigen (G. morsitans, S. calcitrans or T. yao saliva or rTsall) per well in 0.1 M NaHCO₃ (pH 8.3). Plates were overcoated for 1 h with 10% fetal bovine serum (FBS) at ambient temperature. Serial half plasma dilutions starting from 1:100 in assay diluent (PBS/10%FBS) were applied for 2 h to antigen and FBS-coated wells. For the analysis of porcine plasma samples, 20% Top10F' E. coli soluble extract was added to the assay diluent to reduce unspecific binding to the antigenic coat. Based on the plasma dilution experiments, a 1:1600 dilution was chosen for the time course analyses. Specific immunoglobulin detection was achieved using horseradish peroxidase conjugated detection antibodies. For the detection of mouse and porcine IgGs respectively a 1:1000 diluted rabbit F(ab)2 anti-mouse IgG (STAR13B, Serotec) and a rabbit anti-pig IgG conjugate (A5670, Sigma) 1:4000 diluted in PBS/10%FBS were used. Detection was with TMB substrate (3,3',5,5'-Tetramethylbenzidine, Sigma) and reactions were stopped by the 1:3 addition of 1N H₂SO₄. Optical density (O.D.) was measured using a Multiskan Ascent plate reader (Thermo) at a 450 nm wavelength.

Data analysis

Antigen-specific responses were expressed as the $\Delta O.D.$ between antigen and non-antigen-coated wells. Statistical analyses were performed in SAS Version 9.3 (SAS Institute Inc., Cary, NC, USA). The effect of tsetse fly exposure intensity and boosting on anti-G. m. morsitans saliva and anti-rTsal1 immune responses in mice and pigs was analysed over the entire sampling period based on a mixed model with animal as random effect and challenge, time and their interaction as categorical fixed effects and F-tests were performed at the 5% significance level. Pairwise comparisons were performed using Tukey's multiple comparisons technique to adjust the significance level. Cross-reactivity of immune responses induced by different Glossina species and other hematophagous insects was analysed using a linear fixed effects model using the 1:100 or 1:1600 dilution with challenge as categorical fixed effect. Pairwise comparisons were performed with control using Dunnett's multiple comparisons technique to adjust the significance level. Intra-laboratory repeatability of the ELISA tests was assessed by the non-parametric Spearman correlation test. Sensitivity and specificity of the assays were assessed by receiver operating characteristic (ROC) curve analysis of the Δ O.D. values of exposed and non-exposed animals, starting from 3 weeks after the initial exposure. The area under the ROC curve (AUC) was used as a global index of diagnostic accuracy. Kinetics of the IgG clearance in mice was assessed in five mice from the multiple exposure schemes with sufficient remaining plasma for the different timepoints and using the plasma sample with the highest antibody titer (set as 100%) to prepare a standard curve. Percent decrease in antibody titers over time was assessed by a two-phase non-linear regression allowing the estimation of the antibody halflife.

Results

Serological responses in mice exposed to *Glossina* morsitans

Induction of specific antibody responses was assessed by indirect ELISA in mice following different regimens of exposure to *Glossina* morsitans morsitans bites. Anti-*G. m. morsitans* saliva and anti-rTsal1 IgGs were detectable within 7 days after the initial exposure and remained persistently detectable up to 390 days (Figure 1). The detected IgG titers against both antigens correlated with the different intensities of tsetse fly exposure. Differences in number of

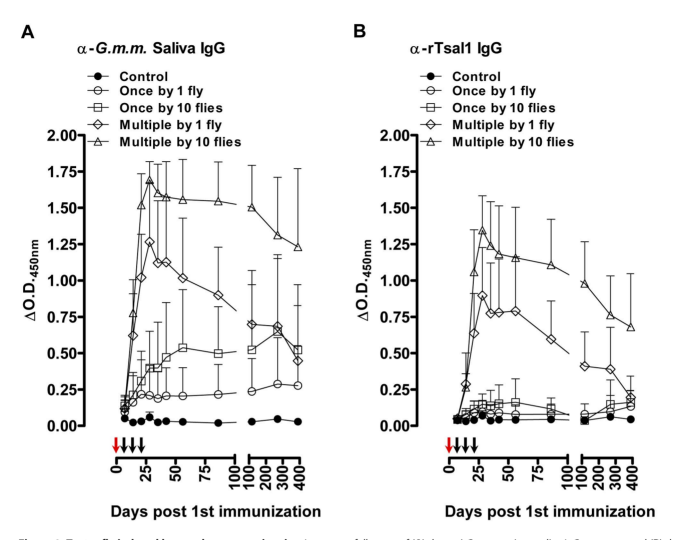


Figure 1. Tsetse fly induced humoral responses in mice. Long-term follow-up of (**A**) the anti-*G. m. morsitans* saliva IgG responses and (**B**) the anti-rTsal1 IgG responses in mice (n = 8/group) exposed to 4 different tsetse fly biting intensities (1 versus 10 flies) and frequencies (single versus weekly exposure for 3 weeks). Arrows indicate the tsetse exposure regimen for single exposed (red arrows) and multiple exposed mice (red and black arrows). Presented data are the mean Δ O.D._{450 nm} values and the 95% CI obtained with a 1:1600 dilution of the individual plasmas. doi:10.1371/journal.pntd.0002911.g001

flies (1 versus 10 flies) as well as differences in frequency (single versus weekly exposure over 3 weeks) were detected by ELISA. Exposure to a single tsetse fly bite was sufficient to induce slightly elevated levels of antibodies against G. m. morsitans saliva and rTsal1 (Figure 1), although the recorded differences with control plasmas were not significant (p = 0.9843 and p = 0.6146 for the anti-rTsal1 and anti-G. m. morsitans saliva tests respectively). Single exposure of mice to 10 flies resulted in significantly increased reactivity against G. m. morsitans saliva (p = 0.0160) but not against rTsal1 (p = 0.8618). With both antigens, statistically significant differences were recorded considering the entire sampling period between control mice and mice subjected to the repeated exposure to 1 and 10 flies ($p \le 0.0001$). Both the anti-rTsal1 and the anti-G. m. morsitans saliva ELISA test were able to differentiate between these two repeated tsetse exposure schemes ($p \le 0.0001$ and p = 0.0012 respectively). Based on a standard curve generated using the plasma sample with the highest IgG titer (i.e. day 28, mouse exposed to the most intense biting regimen), antibody halflives in five mice exposed multiple times to a single fly or 10 flies were determined by two-phase decay regressions (Figure 2). The average half-lives $(T_{1/2}^{2})$ of the anti-saliva and anti-rTsal IgGs in a

second phase of decay after a first phase immediately after cessation of tsetse exposure were respectively 36 and 40 days (Figure 2). There was a significant individual variation within the different exposure groups with up to a 10-fold difference between the strongest and weakest responder. On average, a 3 to 4-fold difference in peak IgG titer was recorded between mice exposed to a low (multiple bites by a single fly) and high exposure regimen (multiple bites by 10 flies). The antibodies appeared relatively persistent over an evaluation period of more than 1 year. Following a long period of non-exposure, the bite of a single tsetse fly was sufficient to boost the anti-saliva and anti-rTsal1 IgG titers. This boosting appeared independent of the previous exposure intensity (Figure 3) as statistical analysis over the 6-week sampling period revealed the inability of the saliva and rTsal1based tests to differentiate mice subjected to a low or high initial exposure regimen (p = 0.4141 and p = 0.9609 respectively). Antibody titers against both antigens reached a plateau within 7 days after re-challenge, while in naive animals the responses were slightly lower (p = 0.0833 and p = 0.1925 for the anti-saliva and anti-rTsal1 IgG levels respectively) and only reached peak titers within 4 weeks after initial exposure (Figure 3).

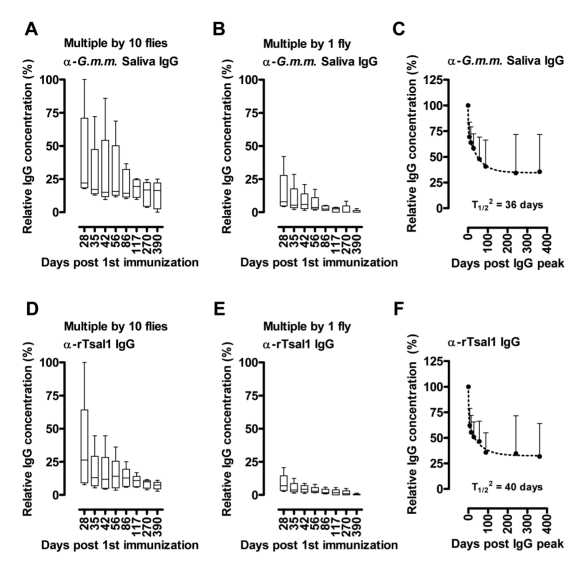


Figure 2. Antibody kinetics in tsetse exposed mice. Evolution over time in the absence of tsetse fly challenge (from day 28 onwards) of the anti-*G. m. morsitans* saliva IgG concentrations (**A**, **B**, **C**) and the anti-rTsal1 IgG concentrations (**D**, **E**, **F**) in mice (n = 5/group) exposed multiple times to a single fly (**B**, **E**) or 10 flies (**A**, **D**). Presented data are the IgG concentrations relative to the highest responder plasma and expressed as percent based on a standard dilution series. Box plots represent the means, with minimum and maximum and the 75-25% percentiles. The average antibody half-life ($T_{1/2}^2$, **C**, **F**) over the 2 exposure groups (n = 10) was determined by a two-phase non-linear decay regression (dotted line) of the declining antibody concentrations relative to the concentration recorded after cessation of the tsetse exposure (day 28 set as starting concentration of 100%). Scatter plots indicate the mean concentrations with the 95% CI. doi:10.1371/journal.pntd.0002911.g002

A strong correlation between anti-saliva and anti-rTsal1 ELISA results was recorded with a Spearman correlation coefficient r of 0.92. Intra-laboratory repeatability of the anti-saliva and anti-rTsal1 detection assays was excellent (Spearman r=0.98). A comparison of the diagnostic value of the anti-saliva and rTsal1 indirect ELISA was conducted using ROC curve analysis. Comparison of the area under the curve (AUC), based on the mouse plasmas from the experiment presented in Figure 1 and collected at least 3 weeks after the initial tsetse fly exposure, indicated that the anti-tsetse saliva ELISA has a better test performance than the rTsal1-based assay (AUC 0.94 versus 0.82, Figure S1).

Serological responses of mice exposed to various *Glossina* species

Mice were repeatedly exposed to the bites of various *Glossina* species (*G. m. morsitans, G. p. gambiensis, G. pallidipes* and *G. f. fuscipes*).

It was noted that the feeding performance of G. fuscipes on mice was inferior to those of the other species, resulting in a high mortality of these flies when maintained on mice. G. m. morsitans saliva and rTsal1 were next used as antigens in indirect ELISA to detect the elicited antibodies. A marked cross-reactivity of the antisaliva IgGs elicited by three Glossina species (G. m. morsitans, G. p. gambiensis and G. pallidipes, p < 0.0001 at the 1:100 plasma dilution) and a relatively weaker cross-reactivity of G. f. fuscipes (p = 0.0053) was detected with G. m. morsitans saliva as antigen (Figure 4A&B). Similarly, antibodies induced by exposure to G. m. morsitans, G. p. gambiensis and G. pallidipes significantly reacted with rTsall (respectively $p \le 0.0001$, p = 0.0157 and p = 0.0002), but no crossreaction with rTsal1 was detected in mice exposed to G. f. fuscipes (p = 0.9883). This indicated that a number of antigens are sufficiently conserved in the saliva of several Glossina species to allow multi-species detection of exposure using G. m. morsitans saliva. The rTsal1 also enabled detection of exposure of mice to

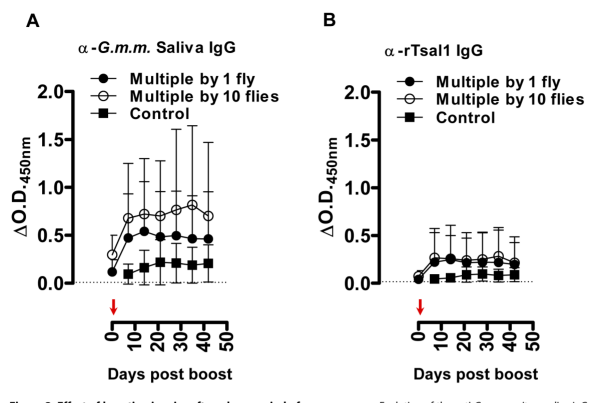


Figure 3. Effect of boosting in mice after a long period of non-exposure. Evolution of the anti-*G. m. morsitans* saliva IgG responses (**A**) and the anti-rTsal1 IgG repsonses (**B**) was evaluated in mice (n = 6/group) originally exposed to the bites of a single or 10 flies and re-exposed to a single tsetse fly bite after a 1-year period of non-exposure (indicated by an arrow). Controls are the naive mice exposed to a single bite as depicted in Figure 1. The original control animals (n = 3) were not exposed and served as negative controls (dotted line). Presented data are mean $\Delta O.D._{450 \text{ nm}}$ values with the 95% CI obtained with a 1:1600 dilution of the individual plasmas. doi:10.1371/journal.pntd.0002911.g003

various tested tsetse fly species, except for *G. fuscipes* (Palpalis subgenus).

Serological responses of mice exposed to various hematophagous insect species

Mice were experimentally immunized by 4 intradermal injections with the saliva of stable flies (Stomoxys calcitrans) or horse flies (Tabanus yao) and compared with control mice and mice that were immunized with tsetse fly saliva following the same immunization protocol. Immunization with Tabanus saliva did not result in IgGs that reacted with rTsal1 and total G. morsitans saliva as coating antigens (p = 0.7882 and p = 0.7637 at the 1:100 plasma dilution, Figure 4D&E). However, a slight increase that did not reach statistical significance was observed in responsiveness of the Stomoxys exposed plasmas in the rTsal1 and saliva-based assays (p=0.0837 and p=0.0639 respectively). At a standard 1:1600 dilution, no cross-reaction of the Stomoxys exposed plasmas with rTsal1 and saliva was detected (p = 0.9920 and p = 0.9915respectively). Analysis of the published salivary transcriptome of Stomoxys calcitrans suggested the presence of a Tsall homologue from which a truncated sequence was published (GenBank Accession N°: ACN69159, [26]). Within this region, only 36% identity in amino acid sequence was found, which could explain the slightly elevated anti-rTsall reactivity of Stomoxys exposed mice.

Serological responses in tsetse fly exposed pigs

Pigs were experimentally exposed to a low or a high tsetse fly bite regimen, followed by assessment of the antibody production. As compared to the mouse indirect ELISA, we have modified the porcine assay by including E. coli soluble extract in the sample diluent as described elsewhere [25] to reduce the background that was observed particularly onto the rTsall antigen which was produced in a bacterial expression system. Under the used assay conditions, anti-rTsal1 and anti-saliva IgGs were detected from day 21 after the first tsetse exposure onwards (Figure 5). The specific IgG titers in pigs correlated with the intensity of tsetse exposure. Anti-rTsal1 and anti-tsetse saliva IgGs were elevated in tsetse exposed groups as compared to the non-exposed control pig and pre-immune plasmas, with significant differences for the high exposure group (p = 0.0216 and p = 0.0030 respectively). The repeated exposure of pigs to 30 flies resulted in higher anti-rTsal1 and anti-tsetse saliva IgG titers as compared to the pigs exposed to a low tsetse challenge by 3 flies (p = 0.0014 and p = 0.0024respectively). However, exposure to the low exposure scheme did not result in significantly elevated responses in both the rTsal1 and saliva-based ELISA (p = 0.8844 and p = 0.4261 respectively, Figure 5). Boosting of 2 pigs from the low exposure group after a 2-month non-exposure period by the bites of 10 flies resulted in elevated anti-saliva IgG titers but was only weakly detectable using the rTsall-based ELISA and with higher individual variation (Figure 6). In general, a good correlation between anti-saliva and anti-rTsal1 ELISA results, obtained for the samples from the tsetse fly exposure experiment presented in Figure 5, was recorded with a Spearman correlation coefficient r of 0.79. Intra-laboratory repeatability of the anti-saliva and anti-rTsal1 detection assays was adequate (Spearman r = 0.94). Comparison of the area under the curve (AUC) for the anti-tsetse saliva and anti-rTsal1 ELISA ROC curve indicated that the assay based on total G. morsitans saliva has

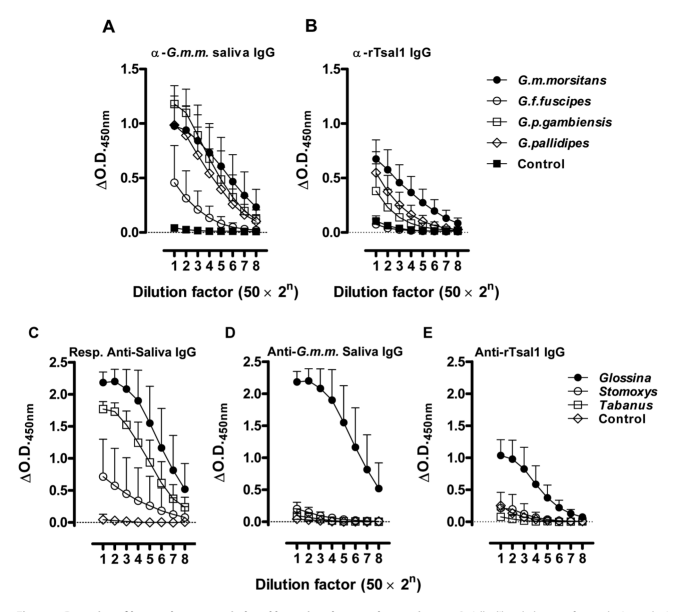


Figure 4. Detection of humoral responses induced by various hematophagous insects. Serially diluted plasmas of control mice and mice repeatedly exposed to the bites of *G. morsitans morsitans, G. fuscipes fuscipes, G. palpalis gambiensis* and *G. pallidipes* (n = 5/group) were tested in the *G. m. morsitans* saliva-based (**A**) and rTsal1-based IgG detection ELISA (**B**). Serially diluted plasmas of control mice and mice experimentally immunized with harvested *G. m. morsitans, Stomoxys calcitrans* and *Tabanus yao* saliva (n = 6/group) were analysed for responses against the respective saliva extracts (**C**) and were tested in the *G. m. morsitans* saliva-based (**D**) and rTsal1-based IgG detection ELISA (**E**). Presented data are the mean Δ O.D._{450 nm} values with the 95% CI. doi:10.1371/journal.pntd.0002911.q004

a better average performance than the anti-rTsall IgG detection test (AUC 0.96 versus 0.83, Figure S2).

Discussion

Assessment of exposure of populations at risk to the bites of tsetse fly vectors could be an important step towards improved control of both HAT and AAT. Given that novel innovative vector control strategies are being developed (e.g. miniaturized insecticide-treated targets [27], the release of sterile male insects [8,9]) and deployed on increasingly large scales on the African continent, monitoring the impact of these new as well as conventional interventions on actual exposure to tsetse fly bites is a logical

follow-up. The tsetse fly species implicated in the transmission of trypanosomes are not uniform throughout the African continent as more than 30 tsetse species and subspecies exist that play differential roles in parasite transmission, have preferences for specific biotopes and display specific host feeding preferences. A number of species are strongly implicated in HAT transmission, such as flies of the Morsitans subgenus (e.g. used in this study: *G. morsitans morsitans, G. pallidipes*) and flies of the Palpalis group (e.g. used in this study: *G. palpalis gambiensis, G. fuscipes fuscipes*). In case of AAT, a large panel of tsetse species and other sympatric hematophagous arthropods are respectively involved in biological and mechanical transmission. Especially stable flies (*Stomoxys* sp.) and horse flies (*Tabanus* sp.) play a role in mechanical transmission

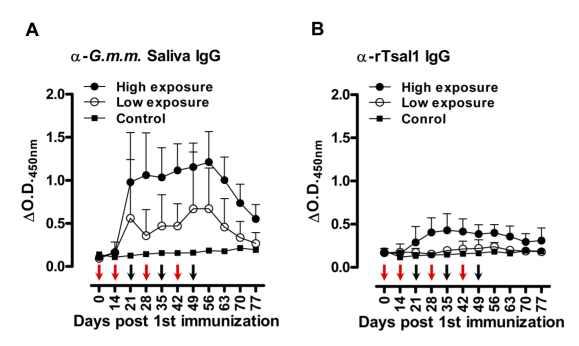


Figure 5. Tsetse fly induced humoral responses in pigs. Follow-up of the anti-G. m. morsitans saliva IgG responses (A) and the anti-rTsal1 IgG responses (B) in pigs exposed to 2 different tsetse fly biting intensities (two-weekly exposure for 6 weeks to 3 flies (low exposure group, n = 4) or weekly exposure for 7 weeks to 30 flies (high exposure group, n = 5)). One non-exposed animal was included as a negative control. Arrows indicate the tsetse exposure regimen for the low exposure (red arrows) and high exposure group (red and black arrows). Presented data are the mean Δ O.D._{450 nm} values and the 95% CI obtained with a 1:1600 dilution of the individual plasmas. doi:10.1371/journal.pntd.0002911.g005

of animal trypanosomes such as *T.vivax*. However, these insects are not able to biologically transmit trypanosomes as the parasite cannot complete its life cycle in these insects.

Research on a number of hematophagous arthropod vectors has resulted in the concept of exploiting salivary components as specific biomarkers of exposure [15,28,29,30,31,32,33]. An advantage of this approach is that relatively simple serological tests could provide information on actual exposure to the bites of disease vectors and provide a risk indicator of contracting a vectortransmitted disease without the need of strenuous entomological surveys. Whole salivary gland extracts of sand flies [33,34,35], triatomine bugs [30,31] and various mosquito species [36,37,38] can be used to assess biting exposure. It has been suggested that this type of serological approach would enable the detection of very low levels of exposure that could remain undetected by entomological trappings [39]. Also differences in exposure level

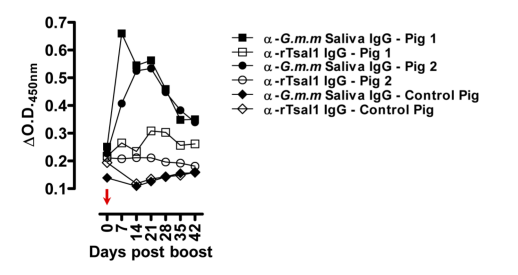


Figure 6. Effect of boosting in pigs after a period of non-exposure. Evolution of the anti-G. m. morsitans saliva IqG responses and the antirTsal1 IgG repsonses was evaluated in pigs (n = 2) originally exposed two-weekly for 6 weeks to the bites of 3 flies (= low exposure group) and reexposed to the bites of 10 flies (indicated by an arrow) after a 2-month period of non-exposure. One non-exposed animal sampled during the priming experiment (Figure 5) served as a negative control. Presented data are mean $\Delta O.D_{.450 nm}$ values with the 95% CI obtained with a 1:1600 dilution of the individual plasmas. doi:10.1371/journal.pntd.0002911.g006

due to vector control interventions (e.g. insecticide treated bednets) could be elucidated on the basis of salivary proteins as immunological probes [29,34,40,41]. While total salivary extracts can be used in various ELISA and immunoblot formats, some studies have moved towards the use of recombinant proteins or peptides which could lead to the development of more standardized immune assays [29,42,43,44,45,46,47,48,49]. Several strategies could be envisaged either aiming at species-specific or pan-species exposure detection.

Studies using various tsetse fly species have shown that salivary components are immunogenic in mice, rabbits, cattle and humans [15,17,18,19,20,23]. Immunoblotting of salivary proteins separated on 1D or 2D protein gels have highlighted immunogenic proteins corresponding to several protein families including endonuclease (Tsal), adenosine deaminase (TSGF), 5'nucleotidase (5'Nuc) and Antigen 5 (Ag5) related proteins [19]. In addition, the immunogenic nature of G. m. morsitans sgp1, sgp2 and sgp3 resulted in their identification on the basis of immune screening of a phage cDNA expression library [50]. We have previously observed by Western blot analysis that human plasma samples collected in Tororo (Uganda, [51]) where G.fuscipes fuscipes is the predominant tsetse fly, commonly recognized the 43–45 kDa Tsal proteins in G. m. morsitans saliva [15]. This suggested that G. m. morsitans Tsalbased immune screening of these Ugandan samples cross-detected exposure to G. fuscipes fuscipes, responsible for Trypanosoma brucei rhodesiense transmission in that area [52]. Plasma samples from tsetse fly exposed individuals in Guinean HAT foci also displayed strong reactivity against the highly abundant Tsal proteins in G. palpalis gambiensis saliva in immunoblots [19]. As the Tsal protein band is commonly recognized in samples from Glossina-exposed humans [15,19], Tsal proteins could have the potential of a pan-Glossina species exposure marker. However, reactivity with Tsal1 and Tsal2 was also observed with a pool of human control plasmas from Bobo-Dioulasso (Burkina Faso) [19]. Based on our current study we anticipate that the inclusion criteria for these urban unexposed plasma donors (not having traveled outside of the city for at least three months, [21]) might have been insufficiently stringent to exclude circulating anti-Tsal antibodies. Indeed, tsetse flies are present at sites outside the city and given the high immunogenicity and anticipated long antibody persistence, this could explain the documented positive reactions in the immunoblots.

Production of a recombinant version of the Tsal1 protein allowed evaluating its effectiveness for detecting tsetse fly exposure in an antibody-detection ELISA. By experimentally exposing mice and pigs to different tsetse fly biting regimens, the high immunogenicity of total G. m. morsitans saliva and Tsal1 as a major constituent was confirmed. In mice, a single bite was proven sufficient to induce a detectable immune response in naive animals and to boost antibody levels in previously exposed animals. Kinetics of the antibody clearance was assessed in mice over a >1year period, which revealed a long persistence with an average half-life of 36–40 days. In pigs, the apparent antibody clearance rate was faster and although the time of sampling and the number of experimental animals was limited, the anti-saliva IgG half-life could be estimated to be 15 days (data not shown). This clearance rate was consistent with the fast decline of anti-G. m. submorsitans saliva IgG levels observed in cattle within 10 weeks after cessation of tsetse exposure [20]. Both in mice and pigs, boosting of prior induced anti-saliva immunity was observed within 7 days by the anti-G. m. morsitans saliva and anti-rTsal1 ELISA. However, with the used sample sizes and experimental conditions, both the rTsal1 and saliva-based ELISA were unable to statistically differentiate the control pig from pigs exposed to a low tsetse fly challenge.

Interestingly, the boosting of immunity in mice seemed independent of the previous exposure intensity. As such, surveys based on rTsall or total G. m. morsitans saliva in tsetse fly control intervention zones would give information on the actual tsetse exposure intensity rather than history and could provide a tool to monitor re-invasion. Consistent with this, serological results for cattle in South-West Burkina Faso based on G. palpalis gambiensis whole saliva extract related to the seasonality which directly impacts the intensity of host/vector contact [20]. Also humans living in HAT endemic areas display significant differences in anti-G. p. gambiensis saliva IgGs depending on the study site [19]. Similar results were obtained with a peptide derived from the adenosine deaminaserelated protein TSGF1, although obtained ELISA signals seemed very low [21]. In both the cattle and human ELISA tests, large inter-individual differences were observed [15,19,20,21]. It should be noted that in our study the individual variation in antibody titers in the different exposure groups also varied significantly. In controlled experimental conditions with outbred mice, we recorded up to 10-fold differences in anti-saliva and anti-rTsal1 antibody concentrations between the strongest and weakest responders. Consequently, using the tsetse salivary antigens as immunological probes has the limitation of inherent variability due to individual differences in immunological responsiveness. Nevertheless, we observed a clear overall difference in anti-rTsal1 and anti-saliva IgG levels between the different exposure groups of mice and pigs, with an average 3-4 fold difference in specific antibody concentration between a low exposure (repeated exposure to 1 fly for mice or to 3 flies for pigs) and a high exposure regimen (repeated exposure to 10 flies for mice or to 30 flies for pigs). This is in line with observations made for cattle that were experimentally exposed to different biting intensities [20]. Beside population-level studies, individual serological follow-up of selected animals or sentinel animals using the recombinant Tsal1 or the total G. morsitans saliva could provide a measure for tsetse prevalence, provide evidence for the impact of an intervention strategy or could be a sensitive tool to detect re-invasion of previously cleared areas or the efficacy of a barrier protecting a cleared area. Here, rTsall and total G. morsitans saliva seem to provide an indication of exposure to a broad range of Glossina species. Only for G. f. fuscipes, detection of induced IgGs is hampered which could relate to the genetic distance of this species as member of the Palpalis group (although G. palpalis gambiensis was efficiently cross-detected). Nevertheless, serological responses against rTsal1 were detected in humans from Uganda principally exposed to G. fuscipes, which suggests that sensitivity and specificity parameters related to a single antigen may vary considerably depending on the host species. It also remains to be evaluated whether the intensity and persistence of the anti-Tsall IgG responses would not result in a saturation in populations persistently exposed to tsetse bites. This could possibly limit application of the rTsal1-based assay to examining subjects in tsetse-free areas that are at risk of invasion or to monitoring sentinel animals in order to evaluate tsetse presence in endemic regions.

Exposure to the saliva of other hematophagous insects (*Tabanus* and *Stomoxys* sp.) that may be abundant in areas where tsetse flies are present, revealed that *G. m. morsitans* saliva did not yield false positive signals. Observations of *Stomoxys* and *Tabanus* exposed cattle revealed that also *G. m. submorsitans* saliva shares this feature of specificity, while *Glossina tachinoides* and *G. palpalis gambiensis* saliva yielded unspecific reactions with *Tabanus* exposed plasmas [20]. When using rTsal1 as an antigen, a weak unspecific signal was observed with the 1:100 to 1:800 diluted plasmas of mice exposed to *Stomoxys calcitrans* saliva which could be due to the

presence of a homologue (GenBank Accession N°: ACN69159, [26]) with relatively limited degree of identity to Tsal1. Given the strong immunogenicity of the Tsal proteins and based on our ELISA analyses with the experimentally exposed pigs and mice we propose to use a 1:1600 plasma dilution in favor of an increased specificity and an overall reduced risk of detecting cross-reactive antibodies. For the porcine ELISA, *E. coli* soluble extract was added to the sample diluent to increase the specificity similar to what was proposed for other pig serological tests [25]. Under these stringent experimental conditions, the recombinant Tsal1 was able to detect the high levels of exposure to tsetse fly bites and with a good correlation with the data obtained for total *G. m. morsitans* saliva as an antigen.

Collectively and combined with our previous analyses using a large panel of East-African human plasmas [15], this study indicates that a recombinant version of the *G. m. morsitans* Tsal1 fulfills the criteria of a candidate exposure biomarker for a broad range of tsetse fly species. We believe that the high sensitivity of the rTsal antigen and the broad species recognition could be an added value to the immunoassays that are tested in the framework of other studies based on tsetse salivary peptides.

Supporting Information

Figure S1 Repeatability and specificity/sensitivity analysis of the antibody detection test in mice. Scatter plot analysis of the anti-*G. m. morsitans* saliva IgG responses (**A**) and the anti-rTsall IgG repsonses (**B**) (Δ O.D._{450 nm}) in two separate tests performed on a panel of 1:1600 diluted mouse plasma samples (*n* = 352). Test repeatability was analyzed by the non-parametric Spearman correlation test. Sensitivity and specificity of the two

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assays were assessed by receiver operating characteristic (ROC) curve analysis of the Δ O.D. values of exposed and non-exposed mice (**C**). The area under the ROC curve (AUC) is reported as a measure for the test performance. (TIF)

Figure S2 Repeatability and specificity/sensitivity analysis of the anti-G. m. morsitans saliva IgG responses (**A**) and the anti-rTsall IgG repsonses (**B**) (Δ O.D._{450 nm}) in two separate tests performed on a panel of 1:1600 diluted porcine plasma samples (n=80). Test repeatability was analyzed by the non-parametric Spearman correlation test. Sensitivity and specificity of the two assays were assessed by receiver operating characteristic (ROC) curve analysis of the Δ O.D. values of exposed and non-exposed mice (**C**). The area under the ROC curve (AUC) is reported as a measure for the test performance.

(TIF)

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Author Contributions

Conceived and designed the experiments: GC RDD LD FG JVDA. Performed the experiments: GC RD RDD SS JVDA. Analyzed the data: GC RD LD JVDA. Contributed reagents/materials/analysis tools: GC RDD LD FG JVDA. Wrote the paper: GC RD RDD SS LD JVDA.

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