1 Evidence of the absence of Human African Trypanosomiasis in northern Uganda: analyses of cattle, pigs

2 and tsetse flies for the presence of Trypanosoma brucei gambiense

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8 Abstract

9 Background

Large-scale control of sleeping sickness has led to a decline in the number of cases of Gambian human
 African trypanosomiasis (g-HAT) to <2000/year. However, achieving complete and lasting interruption of
 transmission may be difficult because animals may act as reservoir hosts for *T. b. gambiense*. Our study
 aims to update our understanding of *T. b. gambiense* in local vectors and domestic animals of N.W.
 Uganda.

15 Methods

- 16 We collected blood from 2896 cattle and 400 pigs and In addition, 6664 tsetse underwent microscopical
- 17 examination for the presence of trypanosomes. *Trypanosoma* species were identified in tsetse from a
- 18 subsample of 2184 using PCR. Primers specific for *T. brucei* s.l. and for *T. brucei* sub-species were used to
- 19 screen cattle, pig and tsetse samples.

20 Results

- 21 In total, 39/2,088 (1.9%; 95% CI=1.9-2.5) cattle, 25/400 (6.3%; 95% CI=4.1-9.1) pigs and 40/2,184 (1.8%;
- 22 95% CI=1.3-2.5) tsetse, were positive for *T. brucei* s.l.. Of these samples 24 cattle (61.5%), 15 pig (60%) and
- 25 tsetse (62.5%) samples had sufficient DNA to be screened using the *T. brucei* sub-species PCR. Further
- analysis found no cattle or pigs positive for *T. b. gambiense,* however, 17/40 of the tsetse samples
- 25 produced a band suggestive of *T. b. gambiense.* When three of these 17 PCR products were sequenced the
- sequences were markedly different to *T. b. gambiense*, indicating that these flies were not infected with *T.*
- 27 b. gambiense.

28 Conclusion

- 29 The absence of *T. b. gambiense* in cattle, pigs and tsetse accords with the low prevalence of g-HAT in the
- 30 human population. We found no evidence that livestock are acting as reservoir hosts. However, this study
- 31 highlights the limitations of current methods of detecting and identifying T. b. gambiense which relies on a
- 32 single copy-gene to discriminate between the different sub-species of *T. brucei* s.l.

33 Author Summary

34	The decline of annual cases of West-African sleeping sickness in Uganda raises the prospect that
35	elimination of the disease is achievable for the country. However, with the decrease in incidence and the
36	likely subsequent change in priorities there is a need to confirm that the disease is truly eliminated. One
37	unanswered question is the role that domestic animals play in maintaining transmission of the disease. The
38	potential of cryptic-animal reservoirs is a serious threat to successful and sustained elimination of the
39	disease. It is with the intent of resolving this question that we have carried out this study whereby we
40	examined 2088 cattle, 400 pigs and 2184 tsetse for Trypanosoma brucei gambiense, the parasite
41	responsible for the disease. Our study found <i>T. brucei</i> s.l. in local cattle, pigs and tsetse flies, with their
42	respective prevalences as follows, 1.9%, 6.3% and 1.8%. Further analysis to establish identity of these
43	positives to the sub-species level found that no cattle, pigs or tsetse were carrying the pathogen
44	responsible for Gambian sleeping sickness. Our work highlights the difficulty of establishing the absence of
45	a disease, especially in an extremely low endemic setting, and the limitations of some of the most
46	commonly used methods.

47 Introduction

The term "human African trypanosomiasis" (HAT) is used to describe two diseases that are clinically, geographically 48 49 and parasitological distinct. The majority of HAT cases (98%) occur in West and Central Africa and are referred to as 50 West African sleeping sickness or Gambian HAT (g-HAT) indicating the geographical range of the disease and the 51 protozoan parasites responsible, Trypanosoma brucei gambiense. Similarly, East African sleeping sickness or Rhodesian HAT (r-HAT) results from an infection caused by T. b. rhodesiense. While T. b. rhodesiense has long been 52 known to have a primarily zoonotic lifecycle, T. b. gambiense is considered to be largely anthropophilic with the 53 parasites largely circulating between tsetse and humans only. T. b. gambiense has been identified in domestic 54 55 animals such as pigs, sheep and goats (1, 2) as well as in a number of wild animals (3, 4). Similarly, a wide range of 56 animals have been experimentally infected with T. b. gambiense some of which were shown to be infective to tsetse. 57 These observations suggest that it may be possible for animals to act as reservoirs hosts for T. b. gambiense (5-7) 58 and play a role in transmission. Another study that supports the possibility of cryptic animal reservoirs are the 59 reports of tsetse infected with T. b. gambiense caught in areas without cases of g-HAT (8). The unresolved question, 60 of a zoonotic host, in the life history of T. b. gambiense has significant consequences if elimination by 2030 is to be 61 achieved (9). The importance of a successful elimination campaign that does not result in low prevalence pockets of 62 transmission is evident when one considers the history of HAT. Since the turn of the 20th century there have been 63 three major outbreaks of sleeping sickness resulting in hundreds of thousands of deaths. Crucially the third outbreak 64 occurred after intense control efforts had reduced the number of HAT cases to near-elimination levels (10). The 65 threat of resurgence will always be present and require continued pressure to keep HAT in check unless it is truly eradicated. 66

Despite the well documented reports of animals infected with *T. b. gambiense*, and the evidence that tsetse can become infected through animal hosts, it is not known if zoonotic cases of *T. b. gambiense* act as cryptic reservoirs that play a role in sustaining transmission of gHAT. Modelling studies (11-13) have shown that the success or failure of eliminating sleeping sickness depends on a number of parameters, one of which is the existence of a crypticanimal reservoir. The presence of an animal reservoir can also change the importance of the other parameters such as the importance of human migration to an area with a high tsetse biting rate in the context of heterogenous biting

73 (12).

74	A limiting factor to analysing the role and importance of non-human hosts is the type of diagnostic method used to
75	detect the presence of trypanosomes. Classically, microscopic detection of parasites in blood of a human is regarded
76	as evidence of infection. However, for animal hosts this method is unable to distinguish human-infective T. b.
77	gambiense from animal-infective T. b. brucei (14). Molecular methods can reliably distinguish the different
78	trypanosome species, with a high degree of sensitivity and specificity (15). However, differentiation of the T. brucei
79	sub-species, although possible, has limited sensitivity, as single copy regions of the genome must be targeted (16).
80	Ideally samples positive for <i>T. brucei</i> s.l. will need to be assessed to verify there is enough DNA present to undergo
81	the less sensitive sub-species PCR assay (17). To date the application of these molecular methods has not been fully
82	applied to the N.W. of Uganda, although an animal survey was conducted from 2004-2008, this study did not
83	validate the samples suitability of single copy gene amplification (18). It is likely that a potion of those samples
84	identified as being positive for <i>T. brucei</i> s.l. lack sufficient DNA to undergo the sub-species-specific detection assay.
85	An alternative to molecular methods are serological techniques including the card agglutination test for
86	trypanosomiasis (CATT) and the trypanolysis (TL) test, however the CATT can produce false positives due to malaria
87	and transient trypanosome infections (19, 20). The sensitivity of the two methods also varies between geographical
88	locations (21). The unreliability of these methods can vary across different geographical areas due to the heterogenic
89	distribution of the markers in wild trypanosome populations, making their reliability variable (22).
90	Aim
91	Here we use currently available molecular assays to determine the presence or absence of <i>T. b. gambiense</i> in NW
92	Uganda by screening the vector population and two potential, animal-reservoir populations, cattle and pigs, in a
93	large-scale xenomonitoring campaign, using molecular methods to first identify cases of <i>T. brucei</i> s.l. and

- 94 subsequently the sub-species of *T. brucei* with PCR assays.
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- 96
- 97 Methods
- 98 Study site

99 The North West of Uganda has nine districts, Nebbi, Arua, Koboko, Yumbe, Moyo, Adjumani, Maracha, Amuru and 100 Gulu, of which Arua, Koboko, Yumbe, Moyo, Maracha, Amuru and Adjumani have historic sleeping sickness foci (23). Records from 1905-1920 show deaths from HAT in the West Nile region to be 1-100 per 10,000 (24). Recent surveys 101 from 2000-2015 show that this area of Uganda still has foci of disease, (25) recently there has been a decrease in 102 new cases of HAT being reported and only three new cases in 2016. In 2018, 10,000 people from 28 villages across 103 four districts (Arua, Maracha, Koboko and Yumbe) of N.W. Uganda were screened using the card agglutination test 104 for trypanosomiasis (CATT), with any positives being followed up by CTC microscopy and trypanolysis. Out of the 105 10,000 individuals screened none were found to have a current T. b. gambiense infection, although three 106 participnats did test positive for having had T. b. gambiense in the past (26). In this study tsetse were caught in the 107 108 district of Koboko and screened for trypanosomes and in both Koboko and Arua cattle were sampled and screened 109 for trypanosomes. No sampling for tsetse was carried out in Arua due to a vector control programme being carried out in this district (27). Pigs were sampled from Arua but not from Koboko as there are few pigs there due to it being 110 111 predominantly Muslim and hence domestic pigs are scant.

112 Tsetse catches

113 A total of 12,152 tsetse flies were caught along the Kochi River in the district of Koboko over a period of 16 months

from April 2013 to July 2014. To catch tsetse, pyramidal traps (28) were deployed at four locations (383200N-

283715E, 381611N-287545E, 383674N-280855E, 383550N-283841E). The traps were located <1m from the river and
tsetse were collected twice a day, at ~07:30 h and ~15:30 h. Tsetse were transported from the trap sites to the field
laboratory in cool boxes containing a damp towel to reduce heat and maintain humidity to reduce mortality of tsetse
prior to dissection.

Of the tsetse caught, 6,664 were dissected at the field laboratory (333842N-269418E) to screen for trypanosomes in their mouthparts, salivary glands and midguts (29). Following their dissection, the three tissue types (mouthparts, salivary glands and midguts) were screened for trypanosomes at x400 using a compound-microscope with a darkfield filter. Both negative and positive tissues were then preserved, individually, in 100% ethanol, on a 96 well plate,

123 for further molecular analysis. Of the 6,664-tsetse preserved in this manner 2,184 were processed using the

124 molecular methods described below. The sub-sample of tsetse was selected based on the season they were caught,

125 with samples from the wet and dry season of Septemeber2013- February 2014 being chosen.

127

128 Tsetse DNA extraction

129	After transportation to the Liverpool School of Tropical Medicine, at room temperature, samples were stored at 4°C
130	until undergoing DNA extraction, after which the samples were stored, in a separate fridge, at 4°C. Each individually-
131	preserved tsetse tissue underwent DNA extraction, previously described in Cunningham etal. (30). Briefly, ethanol
132	was evaporated by incubating the samples at 56°C for 3 hours and to the sample $135\mu l$ of extraction buffer was
133	added (1% Proteinase K, 5% TE Chelex suspension). Finally, the samples underwent incubation at 56°C for one hour
134	followed by incubation at 93°C for 30 minutes to halt the enzymatic activity of the proteinase K.
135	Cattle and Pig sampling
136	From August 2011 to December 2013, 2,896 cattle blood samples were collected across seven sites in N.W. Uganda,
137	as part of an impact assessment study following the deployment of tiny targets to control the local tsetse population
138	(27). Of the 2,896 cattle samples taken 2088 were screened for <i>T. brucei</i> s.l. using the FIND-TBR primers (27).
139	Alongside the cattle sampling, in August 2013, 400 pigs were sampled from seven sites across in Arua (Table 1).
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148	
149	Table 1. Tsetse, cattle and pig sampling sites with corresponding numbers of animals sampled

Animal	Site	Number sampled	Dates	Northing	Easting
Tsetse	Koboko (KO16)	217	September	383200	283715

Koboko (KO20) 651 2014 383674 28085 Koboko (KO21) 873 383550 28384 Total tsetse 2184 361283 28402 Arua 311 341589 26718 Ayi 301 August 2011- 364188 26880 Cattle Inve 312 December 351002 26138 Aiivu 280 2013 354092 28189 Koboko West 619 383176 27794 Koboko East 683 381647 28394						
Koboko (KO21) 873 3635074 26663 Koboko (KO21) 873 383550 28384 Total tsetse 2184 361283 28402 Arua 311 341589 26718 Ayi 301 August 2011- 364188 26880 Cattle Inve 312 December 351002 26138 Koboko West 619 383176 27794 Koboko East 683 381647 28394		boko (KO19)	443	2013- February	381611	287545
Total tsetse 2184 Kubala 390 361283 28402 Arua 311 341589 26718 Ayi 301 August 2011- 364188 26880 Cattle Inve 312 December 351002 26138 Aiivu 280 2013 354092 28189 Koboko West 619 383176 27794 Koboko East 683 381647 28394		boko (KO20)	651	2014	383674	280855
Kubala 390 361283 28402 Arua 311 341589 26718 Ayi 301 August 2011- 364188 26880 Ayi 301 August 2011- 364188 26880 Ayi 301 December 351002 26138 Aiivu 280 2013 354092 28189 Koboko West 619 383176 27794 Koboko East 683 381647 28394		boko (KO21)	873		383550	283841
Arua 311 341589 26718 Ayi 301 August 2011- 364188 26880 Cattle Inve 312 December 351002 26138 Aiivu 280 2013 354092 28189 Koboko West 619 383176 27794 Koboko East 683 381647 28394	Total tsetse		2184			
Ayi 301 August 2011- 364188 26880 Cattle Inve 312 December 351002 26138 Aiivu 280 2013 354092 28189 Koboko West 619 383176 27794 Koboko East 683 381647 28394		Kubala	390		361283	284022
Cattle Inve 312 December 351002 26138 Aiivu 280 2013 354092 28189 Koboko West 619 383176 27794 Koboko East 683 381647 28394		Arua	311		341589	267182
Aiivu 280 2013 354092 28189 Koboko West 619 383176 27794 Koboko East 683 381647 28394		Ayi	301	August 2011-	364188	268809
Koboko West 619 383176 27794 Koboko East 683 381647 28394	Cattle	Inve	312	December	351002	261388
Koboko East 683 381647 28394		Aiivu	280	2013	354092	281893
		oboko West	619		383176	277940
		loboko East	683		381647	283948
Total cattle 2896	Total cattle		2896			
Wiliffi 67 364383 28804		Wiliffi	67		364383	288042
Duku 45 361141 29238		Duku	45		361141	292384
Tondolo 25 354060 29008		Tondolo	25		354060	290083
Pig Ngalabia 68 August 2013 351690 28672	Pig	Ngalabia	68	August 2013	351690	286723
Muttee 51 360741 29093		Muttee	51		360741	290934
Drimveni 100 362178 28993		Drimveni	100		362178	289931
Inia 44 364383 28804		Inia	44		364383	288042
Total Pig 400	Total Pig		400			

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Both the cattle and pigs were sampled in the following manner, the animal was restrained, and a disposable lancet was used to puncture a pineal (ear) vein. Blood was collected with three 50mm heparinised capillary tube which collected 35µl of blood. Two tubes were centrifuged at 8,000 rpm for three minutes and the buffy coat layer examined as a wet preparation at x400 magnification using a compound-microscope with a dark-field filter. The contents of the third capillary tube was transferred to a Whatman FTA card (GE Health Care, Little Chalfont) and left to air dry before it was heat sealed in a foil pouch with a packet of silica gel to ensure the sample remained desiccated.

To extract the DNA from the FTA card, a modified version of the method described by Ahmed was carried out as follows, 10 2mm hole-punches were taken from each bloodspot, using a Harris micro-punch. The punches were washed three times in 1ml of distilled water and then 135µl of a 1% Proteinase K/10% Chelex TE suspension was added to each batch of 10-hole punches. These were then incubated at 56°C for an hour followed by 93°C for 30 minutes. In total 14 sampling sites (Fig. 1) were used to gather a total of 2896 samples from cattle and 400 from pigs across the N.W. of Uganda.

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<insert fig 1 here>

Fig. 1 Map of sampling sites for tsetse, cattle and pigs from N.W. Uganda. Intervention zone denotes the area that was under tsetse control during the collection of samples described in this paper (27). The map was created by authors for this publication using Gnu Image Manipulation Software (31).

170 Primer design

171 The tsetse and livestock samples underwent different PCR assays for the detection of T. brucei s.l.. Tsetse were processed with a nested multiplex primer set that targeted T. brucei s.l., T. congolense and T. vivax whereas cattle 172 and pig samples were screened with the FIND-TBR primers (30). The nested multiplex primers were a modified 173 version of the generic ITS primers designed by Adams etal. (32). These primers were used as part of a larger study to 174 identify tsetse positive for T. congolense and T. vivax as well as T. brucei. The outer nest utilised the previously 175 published forward and reverse primers that target the internal transcribed spacer region (ITS) of the trypanosome 176 genome (32). New primers were designed to amplify species-specific regions from the first amplicon generated. This 177 was achieved by aligning the ITS sequence for T. congolense Kilifi (accession number U22317), T. congolense Forest 178 179 (accession number U22319), T. congolense Savannah (accession number U22315), T. brucei s.l. (accession number JX910373), T. vivax (accession number U22316), T. godfreyi (accession number JN673383) and T. simiae (accession 180 number AB742533). A new universal forward primer and three species-specific reverse primers were designed and 181 182 used in a multiplex. The location of the new primers are shown in a diagrammatic form in relation to the universal primers designed by Adams etal. (Fig 2). 183

184

<Insert Fig 2 here>

Fig 2. Diagrammatic representation of the location of the multiplex ITS primers in relation to the Universal primers
designed by Adams et. al. (2006) on the ribosomal DNA.

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The resulting products vary in size based on the species of trypanosome that was amplified with the largest product
 belonging to *T. congolense* s.p. measuring 392bp (*T. congolense* Kilifi) to 433bp (*T. congolense* Savannah and *T. congolense* Forest). The products for *T. brucei* s.l. and *T. vivax* measure 342bp and 139bp, respectively (Fig. 3).

Fig. 3 Image showing the relative sizes of the mITS PCR reaction for *T. congolense* Savannah (1), *T. vivax* (2) and *T. b. brucei* (3), extraction negative control (4), reagent negative control (5) and a second *T. b. brucei* positive control (6)

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196 T. brucei s.l. PCR reaction set-up

197 The PCR reactions were carried out at a final volume of 25µL each containing the following reagents: 2.5µl of 10X 198 PCR Buffer (Bioline, London, UK), 200µM of each of the deoxynucleotide triphosphates (dNTPs) (Thermo Fisher Scientific, Leicstershire, UK), 1.2mM of MgCl2 (Bioline, London, UK), 0.4µM of both the forward and reverse primers 199 and 10µL of BIOTAQ Red DNA Polymerase (Bioline). The first reaction of the two nested PCRs and standard PCR used 200 5µL of DNA template. For the nested PCRs second reaction 1µL of the PCR product from the first reaction was used 201 202 as the template. The primers used to detect T. brucei s.l. positive samples are listed in Table 2. FIND-TBR primers were used for the cattle and pig samples and the novel multiplex ITS primers were used to screen the tsetse 203 samples. The different strategies to screen for T. brucei s.l. reflects the different objectives of the animal and tsetse 204 sampling with the focus on the animal samples being the identification of T. brucei positives while it was desirable to 205 206 confirm presence of different species of Trypanosoma to support the tsetse microscopic examination of different tissues (midgut, mouthparts, salivary glands) from tsetse. 207

208 Table 2. Primers used for detection of *T. brucei* s.l.

	Primer	Target Species	Primer Sequence 5'-3'	Expected product	Reference
	name	Target opened	Think Sequence 5 5	size	Kelerence
	FIND-TBR F		TGCGCAGTTAACGCTATTATACA		
FIND-TBR	FIND-TBR R	Trypanozoon	AAAGAACAGCGTTGCAAACTT	117	(Cunningham et al. 2016)
	Tryp 1		AAGCCAAGTCATCCATCG		
ITS Primers	Tryp 2	Trypanosomatidae	TAGAGGAAGCAAAAG	220-642	(Adams et al. 2006)
	mITSF	Trypanosomatidae	TAGCTGTAGGTGAACCTGCAGC	-	
mITS	mITS_TcR	T. congolense	GCGTCAGGCGGCRWAAGAA	392, 433*	Unpublished
	MpMkTbR	T .brucei sl	ATGCGAGGTTGATATACACATAGCA	342	
	MpMkTvR	T .vivax	GCCGTGCTCCACCTG	139	

209 *The size range of the T. congolense products varied depending on the strain type with Kilifi producing a smaller

210 product of 392bp in size compared to both the Savannah and Forest strains producing a product of 433bp in size.

212 However as there is a difference in the copy number being targeted by the different primers not all those samples 213 initially identified as T. brucei s.l. will be identified down to a sub-species level due to insufficient DNA. Following the 214 detection of *T. brucei* s.l. positive samples using either the multiplex ITS primers or the FIND TBR-PCR primers (30), 215 the positive samples were screened with the Picozzi multiplex primers (17). The FIND-TBR primer targets a copy region of several thousand (33) whereas the Picozzi primers and T. b. gambiense species specific primers (TgsGP) 216 (34), target a single copy region of the genome. Therefore, it will not be possible to identify a proportion (\sim 27%) of 217 the *T. brucei s.l.* positive samples down to the sub-species level as the species-specific primers are less sensitive than 218 the TBR-PCR primers (35, 36). The following methodology was used to clarify how many TgsGP negatives were 219 negative due to either the absence of T. b. gambiense or insufficient genomic material. 220

221 Identification of single copy gene and *T. brucei* sub-species

Having identified which samples are positive for T. brucei s.l. there is a need to determine what sub-species of T. 222 223 brucei the sample belongs, be it T. b. brucei, T. b. gambiense or T. b. rhodesiense due to the significance of the presence of a human infective sub-species. The PCR assays for positive identification of T. b. rhodesiense and T. b. 224 225 *gambiense* rely on the amplification of a single copy gene unique to either the West or East African parasite. If 226 neither Gambian or Rhodesian HAT is detected, then it is assumed the organism present is *T. b. brucei*. However due to the difficulty of reliably amplifying a single copy gene, and therefore the low sensitivity of the two sub-species 227 228 specific assays, there is a chance that a negative result occurs due to insufficient target DNA. To help increase the confidence of a negative result it is possible to determine if there is sufficient quantity of DNA by running an assay 229 that amplifies a single copy gene ubiquitous to all three sub-species. The multiplex designed by Picozzi et al (17) is 230 231 capable of assessing whether there is sufficient DNA for single copy gene amplification and to also screen for T. b. 232 rhodesiense. The multiplex consists of universal Trypanozoon primers that target the T. brucei s.l. single copy gene, phospholipase C (PLC), as well as primers that target the serum resistance associated gene (SRA) for T. b. 233 rhodesiense. Among the variable surface glycoprotein (VSG) genes there are regions with some sequence identity to 234 the SRA gene. Between the VSG and SRA genes there is an internal deletion within the SRA genes that allows it to be 235 distinguished between any VSG amplification and SRA amplification. Two pairs of primers were designed that 236 237 amplify across this deleted region to allow for clear size distinction between a SRA PCR product and a VSG PCR product. The combination of primers results in the amplification of a 324 bp product for all *Trypanozoon* species, a 238 239 669 bp product for *T. b. rhodesiense* and a >1 Kb for any *VSG* products amplified (Table 3).

240

241

242 Table 3. Primers for single copy gene detection and sub-species-specific analysis

Primer name	Target species	Primer sequence 5'-3'	Multiplex product size (bp)	Reference
PLC 657	T	CTT TGT TGA GGA GCT GCA	224	
PLC 658	Trypanozoon	CAC CGC AAA GTC GTT ATT	324	
SRA 02		AGC CAA AAC CAG TGG GCA		
SRA 03	T. b. rhodesiense	TAG CGC TGT CCT GTA GAC GCT	669	
VSG 651		GAA GAG CCC GTC AAG AAG GTT TG		
VSG 651	Trypanozoon	TTT TGA GCC TTC CAC AAG CTT GGG	>1Kb	
Primer name	Target species	Primer sequence 5'-3'	Singleplex product size (bp)	Reference
TgsGP forward		GCT GCT GTG TTC GGA GAG C		
TgsGP reverse	T. b. gambiense	GCC ATC GTG CTT GCC GCT C	308	

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244 Ethics Statement

- 245 Ethics to sample domestic animals from Uganda was granted by the Ugandan National Council for Science and
- 246 Technology ethics board, which approved the following protocol "Targeting tsetse: use of targets to eliminate African
- 247 sleeping sickness" Ref. Number HS 939. The protocol followed the guidelines set out in the Ag Guide (37) and

248 permission was granted by the animals owners for their involvement in the study.

249

250 Results

- 251 Microscopic examination
- 252 Of the 6,664 tsetse examined microscopically, 180 tsetse organs from 158 (2.4%; 95% CI=2-2.8) tsetse were
- 253 positive for trypanosomes comprising 73 single midgut infections, 69 single mouthpart infections, nine
- 254 mixed mouthpart-midgut infections, a single salivary gland-midgut infections and six cases where all three
- tissues were infected with trypanosomes. Of the 2,877 blood films examined from cattle, trypanosomes

- were identified in 568 (19.7%; 95% CI=18.3-21.2) samples, however, it was not possible to identify down to
- the species level using the microscopy methods.
- 258
- 259 Molecular Screening for *T. brucei* s.l.
- 260 In total 38/2,877 (1.3%; 95% CI=0.9-1.8) cattle and 25/400 (6.3%; 95% CI=4.1-9.1) pigs examined using the
- 261 FIND-TBR primers were positive for *T. brucei s.l.*. The number of tsetse positive for the three Salivarian
- 262 species of trypanosomes were as follows: *T. vivax* 46/2184 (2.1%; 95% CI=1.5-2.8), *T. brucei* s.l. 40/2184
- 263 (1.8%; 95% CI=1.3-2.5) and T. congolense 58/2184 (2.7%; 95% CI=2.0-3.4). Of the T. brucei positive tsetse,
- seven had a single positive mouthpart, nine were single salivary gland positives, 20 were single midgut
- 265 positives, two were mixed salivary gland and midgut positive and the remaining two had all three tissues
- 266 positive. The presence of *T. brucei* s.l. in the mouthparts deviates from the accepted life cycle of this
- 267 trypanosome and is either due to a transient presence of trypanosomes that have passed along the
- 268 proboscis during a feed or the result of cross contamination between tissues during dissection (Table 4).
- 269
- Table 4. The prevalence of PCR *T. brucei* s.l. positives across all samples and districts.

	Animal	District	T. brucei s.l. positives	Numbers sampled	Percent Positive	CI
	Tsetse	Koboko	40	2184	1.8	1.31-2.48
	Cattle	Koboko	8	1280	0.6	0.27-1.21
Cattle		Arua	30	1597	1.9	1.27-2.67
	Pigs	Arua	25	400	6.3	4.10-9.10

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β

- 272 Microscopy and Molecular comparison
- Of the 2,184 samples screened with multiplex ITS primers, trypanosomes were observed by microscopy in 62 samples (49 flies), comprising 30 midguts, four salivary glands and 28 mouthparts. Of the microscopy positives the molecular assay identified 48 of 62 positive tissues, (Table 5).
- 276 Table 5. Species identification by multiplex ITS primers of microscopy positive tsetse tissues

	Midgut	Salivary glands	Mouth parts	
Microscopy positive	30	4	28	
T. brucei s.l.	4	2	0	

	T. congolense	10	1	7
	T. vivax	0	0	15
LTS Its	T. brucei s.l. +T. vivax	0	0	1
ltiplex resul	T. brucei s.l.+T. congolense	6	1	1
lti r	PCR Neg	10	0	4

277

278

279 T. brucei s.l. sub-species screening

The 103 T. brucei s.l. positive cattle, pig and tsetse samples were processed using the T. brucei s.l. multiplex 280 assay to screen for T. b. gambiense, T. b. rhodesiense and the number of samples with enough DNA to 281 282 detect down to a single-copy gene. While not expecting to identify any cases of T. b. rhodesiense the inclusion of a positive control for the East African form of the disease acts as another quality control (Fig 4). 283 None of the samples tested positive for the SRA gene, confirming our expectation that T. b. rhodesiense 284 was not present. However of the cattle, pigs and tsetse, 24 (63%), 15 (60%) and 25 (56%) were positive for 285 the PLC gene, respectively, suggesting that sufficient DNA was present to detect DNA from T. b. 286 rhodesiense or T. b. gambiense if it were present. 287

- 288
- 289

<Insert Fig 4 here>

Fig 4. Example of gel run showing the results of the sub-species *Trypanozoon* multiplex PCR. The 324bp PLC product can be seen in the three positive controls and in sample number 2. The 669bp SRA product can be seen in the *T. b. rhodesiense* positive control but is absent from the other *T. brucei* sub-species. A >1 kb VSG product is visible in sample 2 and is faintly visible in the *T. b. rhodesiense* positive control. A fourth product of ~700 bp can also be seen in the *T. b. gambiense* and *T. b. rhodesiense* positive controls as well as sample 2.

The samples that proved to have enough genetic material for the amplification of the single copy PLC gene were then screened using the sub-species-specific *T. b. gambiense* primer, TgsGP. Among the cattle and pig samples there was no amplification of a 308 bp product however 17 tsetse samples produced a band approximately 308 bp in size.

Of the 17 bands, a subsample of three were sent for sequencing to determine the specific product size and 300 sequence. The samples were sent to SourceBioscience using both forward and reverse primers. The results 301 of the sequencing showed that the three bands sent were identical and that the product was 281 bp 302 (inclusive of primers) in length. The sequence when aligned against reference sequences for T. b. 303 *aambiense* using the NCBI database resulted in only a 90% identity and a query cover of 16% Fig 5. 304 305 <Insert Fig 5 here> Fig 5. Results of the sequence alignment of the three samples sent for sequencing (3923, 3861 and 4280) 306 against the T. b. gambiense positive control (Pos ctrl) and reference sequence (FN555990) from the NCBI 307 308 database. Image generated using MultAlin (38). 309 310 Discussion The aim of this paper was to determine the prevalence of T. b. gambiense in local tsetse, cattle and pig populations 311 312 from N.W. Uganda. The successful identification of T. b. gambiense would suggest that transmission of sleeping sickness in the area was continuing and the identification of the disease in either cattle or pigs would help resolve 313 the role of animal reservoirs in the transmission of the disease. 314 315 Tsetse 316 317 Of the 40 tsetse samples identified as T. brucei s.l. positive by PCR 56% were found to have enough DNA for the amplification of a single-copy gene region. Sixteen produced faint bands of approximately 300bp, comparable in size 318 to the expected band size for T. b. gambiense. However sequencing results showed that the size of the product 319 generated by the samples was smaller than the expected size at only 281bp compared to the expected 308bp sized 320 321 product. There was also significant variation in the 281bp sized sequences compared to *T. b. aambiense* sequence. Based on the sequencing results these positive samples cannot be unequivocally identified as T. b. gambiense. 322 323 Three conclusions arise from the tsetse survey, the first is that despite screening 2,184 tsetse, no tsetse were found to be positive for T. brucei gambiense, this could be due to T. b. gambiense no longer being transmitted in the area 324

325 or that our sample size was too small to detect T. b. gambiense. Despite being understood as the sole vector of gHAT 326 (39) the prevalence of the disease amongst wild tsetse population is often extremely low (1, 40, 41) and attempts to infect tsetse with T. b. gambiense under laboratory conditions have often proven unsuccessful (42). Studies have 327 suggested that the prevalence of T. b. gambiense may be as low as 1 in 4,000 flies (11). However, this number is 328 329 based on microscopy methods, whereas PCR methods should be more sensitive and could identify immature and 330 transient infections reducing the number of tsetse needed to be screened (8). Second despite no T. b. gambiense being found, the tsetse population studied were actively transmitting T. b. brucei. Third, the TgsGP primers cross-331 reacted with DNA from an unidentified source and produced a band, similar in size, to T. b. gambiense, this raises 332 concerns about the specificity of the TgsGP primers are and the potential for erroneously reporting the presence of 333 T. b. gambiense.

335

334

336 Table 4 shows that the positive midguts were identified as either T. brucei or T. congolense similarly of the positive mouthparts all were infected with either T. congolense or T. vivax with no mono infections of T. brucei s.l. identified, 337 338 although in cases of mixed infections *T. brucei* s.l. was detected in the mouthparts.

339 The infected salivary glands were predominantly positive for *T. brucei* s.l. however there was one instance of a single T. congolense infection. The presence of a mITS positive does not indicate an infection of a specific tissue by the 340 341 trypanosome detected but merely the DNA, which could be a transient trypanosome, free floating DNA or DNA introduced during the dissection step; previous studies have found similar results (43). Although, overall, the 342 343 comparison between the mITS results and those of the dissection correlate closely with the traditional methods used 344 to identify trypanosome species based on their location in different tsetse tissues (29), however these methods 345 cannot distinguish between species easily and certainly not between sub-species.

346

Cattle and pigs 347

No animal samples (pig or cattle) produced a band of either 308bp or 281bp when screened with the TgsGP primers, 348 indicating that there were no zoonotic T. b. gambiense infections nor where there any cases similar to those found in 349 the fly samples, where non-target DNA was amplified. However, cases of T. b. brucei were found in both animal 350 351 populations with pigs proving to have the higher prevalence of T. b. brucei infection. This is typical of trypanosome

- epidemiology which has been shown to be highly localised in other studies (44). The lack of any positive *T. b.*
- 353 *gambiense* in the two animal populations sampled correlates with a previous study carried out in the same area (18),
- indicating that it is unlikely either cattle or pigs are acting as cryptic reservoirs of disease.
- 355

356 Diagnostics for *T. b. gambiense*

357	The diagnostic methods used in this paper involved both microscopy and PCR, of which only PCR has the potential to
358	discriminate sub-species of <i>T. b. gambiense</i> (17, 34). There are few diagnostic methods that are capable of
359	accurately distinguishing between the <i>T. brucei</i> sub-species (16, 17, 34). The molecular methods available for the
360	detection of <i>T. b. gambiense</i> are limited due to the practical aspect of conducting these assays in the field and the
361	limited diagnostic markers available. As mentioned previously the sensitivity of <i>T. b. gambiense</i> specific PCR is
362	limited to detecting a single copy gene. Some molecular assays attempt to overcome this problem by relying on the
363	human serums ability to lyse all salivarian trypanosomes (except for T. b. gambiense and T. b. rhodesiense) therefore
364	any T. brucei s.l. identified in a human sample would be one of the two HAT species (45). This allows for the targeting
365	of a higher copy region specific to the <i>T. brucei</i> species group. Using this method any positives would have to be one
366	of the two HAT species, however as the treatment of the two diseases differs and the only option to try and identify
367	if it was an East or West African sleeping sickness infection would be to try and determine the geographical location
368	of where the individual was infected. This approach would also only be limited to humans and could not be used in
369	either xenodiagnoses or screening animals, as all three <i>T. brucei</i> sub-species could be present in the vector or animal
370	populations. To put these differences of sensitivity in perspective, we can look at the limit of detection (LoD) of the
371	number of trypanosome per mL, between multiple diagnostic methods (Table 6.).

372 Table 6. Comparison of LoD between different diagnostic techniques

Туре	Method	LoD (tryps per mL)	Ref
Microscopy	CTC*	500	(AC)
Microscopy	mAECT**	30	(46)
	TBR	10	(30)
Molecular	LAMP	<1	(50)
	TgsGP	~100000	Unpublished

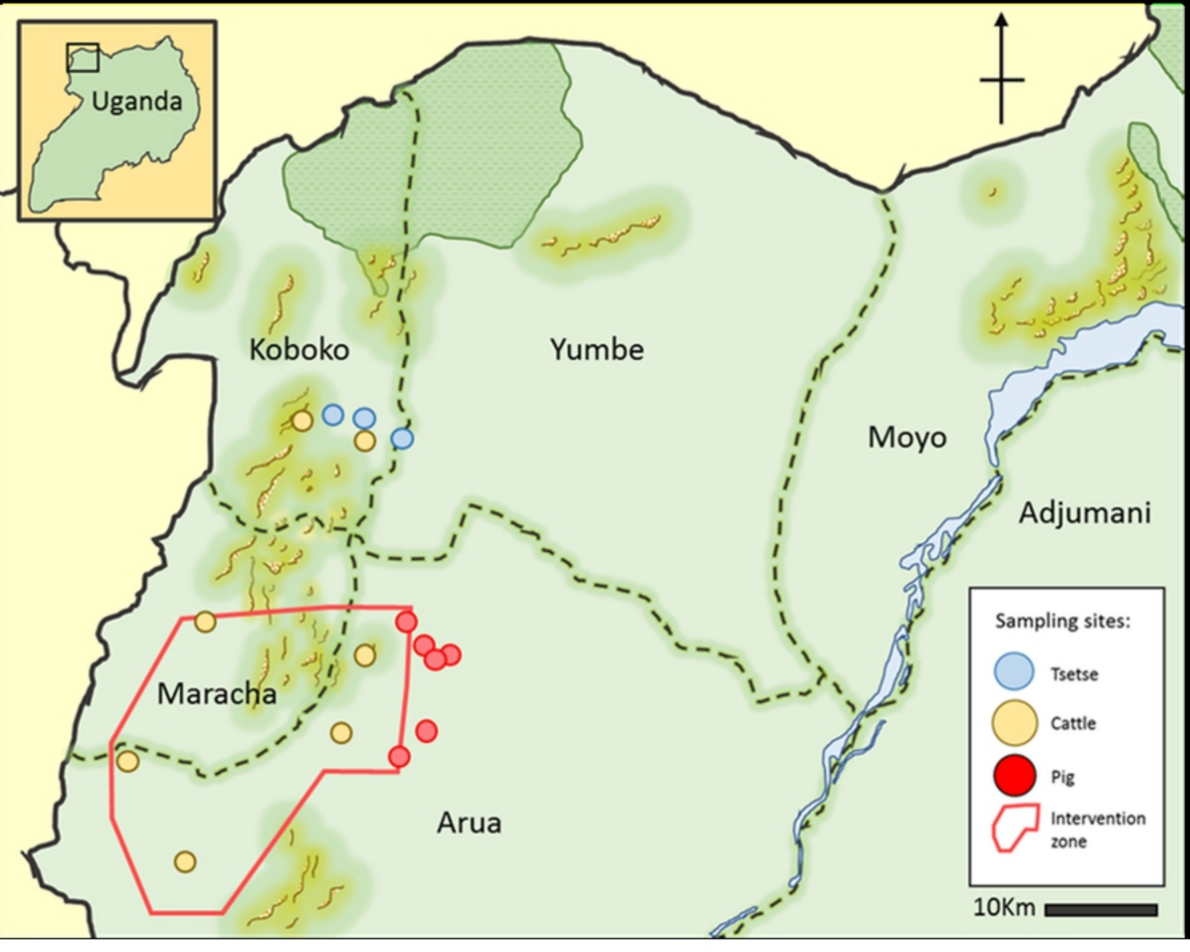
373

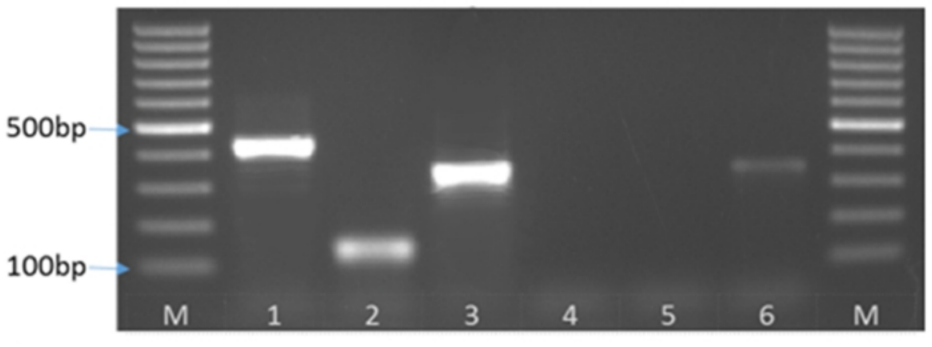
*Capillary tube centrifugation, **mini Anion Exchange Centrifugation Technique

- 374 The lack of a highly specific, sensitive and field-friendly assay that is capable of screening for *T. b. gambiense* in both
- the human, vector and local animal populations is sorely needed if the hope of eliminating sleeping sickness by 2020
- is to be achieved.
- 377
- 378
- 379 Conclusion
- 380 This lack of positive samples reflects the overall low prevalence of the disease and the continued decrease in the
- number of cases in Uganda (47). This study also highlights the lack of highly sensitive diagnostics that can
- discriminate between the different sub-species of *T. brucei* s.l.. Despite not finding *T. b. gambiense* in the tsetse
- population of Koboko vector control has been calculated to being essential to reach the elimination goal of 2030 (48)
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- 387
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	1	10	20	30	40	50	60	70	80	90
FN555990	GCTGCTO	GTGTTCGGAG	AGCTCAGACA	GGGCTGTAATA						ACG
Pos_ctrl				GGGCTGTAATA						
3923	GCTGCTO	GTGTTCGGAG	AGCGACA	GGGCCGCTAGT	GCGCGAA	CTAACGTCGC	TTGTT <mark>CA</mark> TCTG	AGTTAAAAG1	TTTCGGTCGTG	TCG
3861	GCTGCTO	GTGTTCGGAG	AGCGACA	GGGCCGCTAGT	GCGCGAA	CTAACGTCGC	TTGTT <mark>CA</mark> TCT <mark>G</mark>	AGTTAAAAG1	ITTCGGTCGTG	TCG
4280	GCTGCT(GTGTTCGGAG	AGCGACA	GGGCCGCTAGT	GCGCGAA	CTA <mark>ac</mark> gtcg <mark>c</mark>	TTGTT <mark>CA</mark> TCT <mark>G</mark>	a <mark>gttaaaag</mark> t	ITTCGGTCGTG	TCA
Consensus	GCTGCT(GTGTTCGGAG	AGCtcaGACA	GGGCtGtaAta	GCaaGCaAg	agcACaaaaC	cacagCAggcG	gGaTcAggGo	cggCccTgGcG	.Cg
						-				_
	91	100	110		130	140		160		180
	I	+	+	+	+	+	+	+	+	1
FN555990				GGACAACGGCT						
Pos_ctrl				GGACAACGGCT						
3923				ATACAGCCGAT						
3861				ATACAGCCGAT						
4280				ATACAGCCGAT						
Consensus	AtaAGga	agCaaaTTCC	aaaTtgaCGG	ggACAaCgGcTa	atctAGgaA.	GGTACgAc	aCcgaCGGcaa	CtgcaCagga	aACgGcGcCag	Gcg
	181	190	200	210	220	230	240	250	260	270
			T					+		
FN555990				CGGCACCAACA						
Pos_ctrl						The second se				
2002									CAACGGCCGCA	
3923	GAATCT	CGTAGACCG	TTGCCGGCG-	-GAATGCATCT	CGTGAACCG	ATGAGGAGGA	TTAGGGG	AATTTCACGO	CTTAGTA	GTT
3861	GAATCT(GAATCT(C <mark>CGTAGA</mark> CCG CCGTAGACCG	TTGCCGGCG- TTGCCGGCG-	-GAATGCATCT	CGTGAACCG	atgaggagga Atgaggagga	TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	CTTA <mark>gTa</mark> CTTA <mark>gTa</mark>	GTT GTT
3861 4280	GAATCT(GAATCT(GAATCT)	C <mark>GTAGA</mark> CCG CGTAGACCG CGTAGACCG	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG-	-GAATGCATCT -GAATGCATCT -GAATGCATCT	CGTGAACCG CGTGAACCG CGTGAACCG	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861	GAATCT(GAATCT(GAATCT)	C <mark>GTAGA</mark> CCG CGTAGACCG CGTAGACCG	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG-	-GAATGCATCT	CGTGAACCG CGTGAACCG CGTGAACCG	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861 4280	GAATCTO GAATCTO GAATCTO GLgTCTO	CGTAGACCG CGTAGACCG CGTAGACCG CGTAGACCG CGTLaAata	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG- T.GCCGGCCa	-GAATGCATCT -GAATGCATCT -GAATGCATCT cGgcacCAaCa	CGTGAACCG CGTGAACCG CGTGAACCG CtgGgAaCG	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861 4280	GAATCTO GAATCTO GAATCTO GLgTCTo 271	CGTAGACCG CGTAGACCG CGTAGACCG CGTAGACCG CGTLaAata 280	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG- T.GCCGGCCa 290	-GAATGCATCT -GAATGCATCT -GAATGCATCT -GAATGCATCT CGgcacCAaCa 300	CGTGAACCG CGTGAACCG CGTGAACCG CtgGgAaCG 311	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861 4280 Consensus	GAATCTO GAATCTO GAATCTO GL gTCTo 271	CGTAGACCG CGTAGACCG CGTAGACCG CGTAGACCG CGTLaAata 280	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG- T.GCCGGCCa 290	-GAATGCATCT -GAATGCATCT -GAATGCATCT -GAATGCATCT -GgcacCAaCa 300	CGTGAACCG CGTGAACCG CGTGAACCG CtgGgAaCG 311 +I	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861 4280 Consensus FN555990	GAATCTO GAATCTO GAATCTO GLgTCTO 271 I GCAGCAG	CGTAGACCG CGTAGACCG CGTAGACCG CGTLaAata 280 TTGAGGCAG	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG- T.GCCGGCCa 290 GAGCCCGAGC	-GAATGCATCT -GAATGCATCT -GAATGCATCT -GAATGCATCT -GgcacCAaCa 300 	CGTGAACCG CGTGAACCG CGTGAACCG CtgGgAaCG 311 +1 TGGC	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861 4280 Consensus FN555990 Pos_ctrl	GAATCTO GAATCTO GAATCTO GLgTCTo 271 I GCAGCAG GCAGCAG	CGTAGACCG CGTAGACCG CGTAGACCG CGTAGACCG CGTLaAata 280 TTGAGGCAG	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG- T.GCCGGCCa 290 GAGCCCGAGC	-GAATGCATCT -GAATGCATCT -GAATGCATCT -GAATGCATCT -GgcacCAaCa 300 	CGTGAACCG CGTGAACCG CGTGAACCG CtgGgAaCG 311 + TGGC TGGC	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861 4280 Consensus FN555990 Pos_ctr1 3923	GAATCTO GAATCTO GAATCTO GLgTCTO 271 I GCAGCAO GCAGCAO GCAGCAO GCA	CGTAGACCG CGTAGACCG CGTAGACCG CGTAGACCG CGTLaAata 280 TTGAGGCAG TTGAGGCAG	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG- T.GCCGGCCa 290 GAGCCCGAGC GAGCCCGAGC TAGGTAGAGC	-GAATGCATCT -GAATGCATCT -GAATGCATCT -GAATGCATCT -GgcacCAaCa 300 	CGTGAACCG CGTGAACCG CGTGAACCG CtgGgAaCG 311 +I TGGC TGGC TGGC	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861 4280 Consensus FN555990 Pos_ctrl 3923 3861	GAATCTO GAATCTO GAATCTO GLgTCTO 271 I GCAGCAO GCAGCAO GCA GCA	CGTAGACCG CGTAGACCG CGTAGACCG CGTAGACCG CGTLaAata 280 TTGAGGCAG TTGAGGCAG TTGAGGCAG TTGAGGCAG TTGAGGCAG TTGAGGCAG	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG- T.GCCGGCCa 290 GAGCCCGAGC GAGCCCGAGC TAGGTAGAGC	-GAATGCATCT -GAATGCATCT -GAATGCATCT -GAATGCATCT -GAATGCATCT -GgcacCAaCa 300 	CGTGAACCG CGTGAACCG CGTGAACCG CGTGAACCG CtgGgAaCG CtgGgAaCG 311 +1 TGGC TGGC TGGC TGGC	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT
3861 4280 Consensus FN555990 Pos_ctr1 3923	GAATCTO GAATCTO GAATCTO GLgTCTO 271 I GCAGCAO GCAGCAO GCA GCA GCA	CGTAGACCG CGTAGACCG CGTAGACCG CGTAGACCG CGTLaAata 280 TTGAGGCAG TTGAGGCAG TTGAGGCAG TTGAGGCAG TTGAGGCAG TTGAGGCAG TTGAGGCAG	TTGCCGGCG- TTGCCGGCG- TTGCCGGCG- T.GCCGGCCa 290 GAGCCCGAGC GAGCCCGAGC TAGGTAGAGC TAGGTAGAGC	-GAATGCATCT -GAATGCATCT -GAATGCATCT -GAATGCATCT -GgcacCAaCa 300 	CGTGAACCG CGTGAACCG CGTGAACCG CGTGAACCG CtgGgAaCG CtgGgAaCG 311 +1 TGGC TGGC TGGC TGGC TGGC	ATGAGGAGGA Atgaggagga Atgaggagga	TTAGGGG TTAGGGG TTAGGGG	AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark> AATTT <mark>CACGO</mark>	TTA <mark>gta</mark> Ttagta Ttagta	GTT GTT GTT

