A gene expression panel for estimating age in males and females of the sleeping sickness vector Glossina morsitans.

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10 Abstract

11 Many vector-borne diseases are controlled by methods that kill the insect vectors responsible for disease transmission. Recording the age structure of vector populations provides 12 information on mortality rates and vectorial capacity, and should form part of the detailed 13 monitoring that occurs in the wake of control programmes, yet tools for obtaining estimates of 14 15 individual age remain limited. We investigate the potential of using markers of gene expression to predict age in tsetse flies, which are the vectors of deadly and economically 16 17 damaging African trypanosomiases. We use RNAseq to identify candidate expression 18 markers, and test these markers using qPCR in laboratory-reared *Glossina morstians* 19 morsitans of known age. Measuring the expression of six genes was sufficient to obtain a 20 prediction of age with root mean squared error of less than 8 days, while just two genes were 21 sufficient to classify flies into age categories of ≤ 15 and >15 days old. Further testing of these 22 markers in field-caught samples and in other species will determine the accuracy of these 23 markers in the field.

24 Keywords: tsetse flies, age prediction, transcriptomics, machine learning

25 **1** Introduction

26 Vector-borne diseases represent major threats to health and livelihood world-wide, being 27 directly responsible for 680,000 deaths annually (Roth et al. 2018), as well as causing huge 28 economic damage to livestock (Eisler et al. 2003, Shaw 2004). Control of the vectors that 29 transmit these diseases is an integral tool for reducing disease burden (Wilson et al. 2020). 30 The metric of success for these control programmes is a reduction in disease burden in the 31 host population. However, when vector control is accompanied by other interventions such as 32 screening and treating the host population for the disease, the contribution of vector control to 33 the subsequent reduction of disease can be hard to determine (World Health Organization 34 2012). Conversely, while the impact on the vector population may not bear a simple 35 relationship to disease burden, it is a direct outcome of vector control. Control efforts should 36 thus be accompanied by detailed monitoring of the targeted vector populations, to estimate 37 impact, to monitor population recovery and to understand the transmission dynamics of the 38 disease. Mostly, monitoring currently relies on counting the number of vectors caught in sentinel traps, which can be greatly affected by trapping method, effort and efficacy, and may 39 40 only partly reflect the ability of the vector population to transmit disease (Wilson et al. 2015).

41 One aspect of vector monitoring that has been particularly challenging is the quantification of 42 the age-distribution (demographics) of natural populations (Caragata et al. 2011, Cook et al. 43 2006, Sikulu et al. 2010). Estimating vector age is important for two reasons. First, it can 44 provide a measure of the effectiveness of vector control because increased adult mortality should lead to a younger population age structure. Importantly, this measure of control 45 46 effectiveness is independent of catch size and trapping effort because only the distribution of 47 age needs to be known. Second, in most cases, the probability that an individual vector is 48 infectious for a given disease increases with age (Dye 1992, Woolhouse & Hargrove 1998). 49 Before transmitting the disease, vectors first need to have taken an infected blood meal, and 50 there is then typically a delay between acquisition of infection and onward transmission due 51 to the need for the pathogen to replicate and/or mature. Age grading is therefore useful to 52 determine the proportion of individuals old enough to transmit disease.

Tsetse flies (genus *Glossina*) are the vectors of Human African Trypanosomiasis (HAT, or sleeping sickness) and Animal African Trypanosomiasis (AAT, or nagana). HAT is, without treatment, a fatal disease endemic to sub-Saharan Africa (Franco et al. 2014), while AAT presents a major economic burden to rural communities by affecting livestock (Eisler et al.

2003). Being a disease primarily of animals and with reservoirs across multiple species, AAT 57 58 cannot be controlled through treatment alone and is thus highly dependant on vector control 59 (Holmes 2013). G. morsitans morsitans is a major vector of AAT in East and Southern Africa 60 and can also transmit HAT (Dale et al. 1995). Catch rates of this species in the wake of vector 61 control can be extremely low (Kgori et al. 2006, Vale et al. 1988, Van den Bossche 1997), 62 making it particularly challenging to conduct ongoing monitoring of important populations. It 63 is therefore all the more important to extract as much information as possible from the limited 64 number of flies obtained.

65 As is the case for all insect vectors, a means to accurately determine the age of tsetse flies is a valuable but elusive goal, and current methods have many shortcomings. Laborious ovary 66 67 dissections can be used to age females up to their fourth ovarian cycle (Hargrove 2012), but 68 this technique requires specialist dissection skills and cannot be applied to males, despite 69 males being at least as competent at transmission as females, and perhaps more so (Dale et al. 70 1995, Maudlin et al. 1990). Estimates of age based on wing damage (Hargrove 1990) or 71 analysis of pteridines have also been used (Langley et al. 1988, Lehane & Hargrove 1988), 72 but experience in practical applications has shown that measurements in the field vary 73 enormously (for example in mosquitoes: (Lardeux et al. 2000, Penilla et al. 2002)) and cannot 74 be used to reliably estimate age on an individual basis (Hargrove 2020).

75 Here we explore the value of using gene expression to estimate age in tsetse flies. This 76 method has previously been tested in mosquitoes (Caragata et al. 2011, Cook et al. 2006), 77 with encouraging results, but has yet to be applied in tsetse. We use laboratory-reared G. 78 morsitans as a proof of concept, and show that measuring the expression of just six genes can 79 estimate the age of both male and female tsetse flies with a root mean squared error of less 80 than 8 days. We also trained models to classify tsetse into those younger or older than 15 81 days, since flies younger than 15 days are unlikely to harbour a mature trypanosome infection 82 (Dale et al. 1995), and found that just two genes are sufficient for 95% accurate classification.

83 **2 Methods**

84 **2.1 Sample collection and RNA extraction**

6. *morsitans morsitans* individuals were collected from colonies maintained at the Liverpool School of Tropical Medicine. Colonies are kept in meshed boxes (cages) at $26^{\circ}C \pm 2 ^{\circ}C$ and 72 ± 4% humidity, with a 12hr light-dark photoperiod, and fed three times per week using 88 defibrinated horse blood (TCS Biosciences Ltd., Buckingham, UK) provided through silicon-89 membrane feeders. Pupae are regularly collected and allowed to emerge to form new cages. 90 Each fly cage contains flies which eclosed over a 2-3 day window, and thus the age of all flies 91 in the cage are known to a precision of either 2 or 3 days. The ages reported here are the 92 middle of the age range (eg: a fly aged 13-15 days or 13-16 days is reported as 14 days old). 93 The age of the samples ranged from 2 to 62 days. While reproductive status of females was 94 not measured precisely, we tried to include a range of physiological states (based on visual 95 inspection of the size of the abdomen) within each age group, so that genes could be identified 96 that are predictive of age in spite of variation caused by the ovarian cycle. Overall, 505 flies 97 were collected (301 female and 204 male, Supplementary Data S1).

98 For sample collection, fly cages were briefly transferred to a cold room (4 °C) where flies to 99 be collected were removed from the cage once quiescent and decapitated. Heads were placed 100 into RNAlater and stored at -20 °C. In case repeated exposure to the cold room created 101 alterations in gene expression, we minimised this exposure by never collecting flies from a 102 given cage more than three times over the course of the experiment. No more than two flies 103 were collected from a cage on a given day, for three reasons. Firstly, we wanted to make sure 104 that flies were obtained from a range of different cages in order to avoid issues of results 105 being confounded by cage of origin (such as an infection specific to one cage of flies). We 106 therefore never obtained more than six flies from a single cage over the course of the 107 experiment. Second, we wanted to minimise the time that samples spent at temperatures 108 above -20 °C after death, limiting the number of samples that could be collected in a single 109 sitting. Third, all flies were collected at the same approximate time of day (morning) to 110 minimise gene expression variation due to circadian cycles (Rund et al. 2011), limiting the number of collections that could be performed on the same day. 111

RNA was extracted from individual fly heads. Single heads contain enough material for RNA sequencing and can easily be removed without the need for precise dissection, providing a quick and convenient tissue for sampling. We avoided the abdomen because of the important effect that sex and the ovarian cycle would have on gene expression in these tissues. RNA extractions were performed using PicoPure kits (Arcturus), increasing the volume of extraction buffer and alcohol to 120µl. cDNA libraries were prepared using SuperScript III Reverse Transcriptase (Invitrogen).

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Figure 1: Number of samples used for RNAseq (top; total = 50) and qPCR (bottom; total = 498), split by age category (2 - 62 days old). Individual female and male flies shown as blue triangles and orange circles respectively.

119 2.2 Sequencing

120 cDNA libraries from 22 male and 28 female individual flies ranging in age from 2 to 62 days 121 post-eclosion (Fig. 1, Supplementary Data S1) were sent to the Liverpool Centre for Genomic 122 Research (CGR) for 150bp paired-end sequencing on an Illumina HiSeq 4000 sequencer. 123 Strand-specific library preparation was performed using NEBNext poly A selection and Ultra Directional RNA library preparation kits, producing an average of 23.8 million reads per 124 125 sample. Reads were then trimmed as part of the CGR's genomic pipeline using Cutadapt version 1.2.1 (Martin 2011) with option -O 3 to remove Illumina adapter sequences, and 126 127 Sickle version 1.2 (https://github.com/najoshi/sickle/releases/tag/v1.2) with a minimum window quality score of 20. Reads shorter than 20 bp after trimming were removed and 128 129 subsequently unpaired reads were excluded. Data were quality checked using FastOC 130 (Andrews 2010) before analysis.

131 **2.3 RNAseq analysis**

Trimmed reads were aligned to the GmorY1.9 genome using STAR aligner version 2.7.0
(Dobin et al. 2013) using the --quantMode GeneCounts option to obtain mapping counts for
each gene.

135 Differential expression analysis was performed using the R package *EdgeR* (Robinson et al. 136 2010), with library size normalisation performed using Trimmed Mean of M-values 137 (Robinson & Oshlack 2010) and dispersion calculated with trended and tag-wise estimates. 138 Genes with fewer than 10 reads across all 50 samples were excluded from the analysis. All plotting figures show expression measured as reads per million reads (RPM) from normalised 139 library sizes. Association of gene expression with age and sex was tested using generalised 140 linear modelling (glm) implemented in *edgeR*, with age coded as a continuous variable and 141 142 sex as a categorical variable. Preliminary analysis found little evidence of an important effect 143 of the number of times a colony was exposed to the cold room on gene expression, but there 144 was a significant effect of the number of days since flies had received a blood meal 145 (Supplementary Data S2). We therefore controlled for days since receiving a blood meal by 146 including it as a fixed continuous factor in the glm. False discovery rate control was set at 1% 147 using the R package *fdrtool* (Klaus & Strimmer 2015).

Gene clustering analysis was performed with the *WGCNA* package in R (Langfelder & Horvath 2008), using the normalised read counts generated by *edgeR* and keeping only the 5000 genes with the highest variance in expression. We used the hybrid module merging algorithm with a deep split value of 4, a minimum cluster size of 30 and a power parameter of 8, followed by module merging using the absolute value of the correlation coefficient between eigengenes as a distance matrix and a merging threshold of 0.2.

Prediction of age based on normalised read counts from the RNAseq data was performed using lasso regression implemented with the *glmnet* package in R (Friedman et al. 2010). As the aim was to find genes with consistently high predictive value for age, we explored a range of lasso parameters. This exploratory procedure is recorded in detail in the R script "02_lasso.r" provided on GitHub (https://github.com/EricRLucas/TsetseAgeMarkers).

159 2.4 Primer design and qPCR

160 Based on the results of the RNAseq analysis, 16 genes were short-listed to be tested as qPCR markers of age in *G. morsitans*, with two further genes being identified as suitable 161 162 housekeeping genes for our purposes (i.e.: showed minimal variation in expression in the 163 conditions included in our study and no evidence of association with age). Primers were 164 designed for these genes based on the GmorY1.9 genome using NCBI Primer blast (Ye et al. 165 2012). Where possible, amplicons were designed to span exon junctions. Based on testing amplification efficiency using 1:3 serial dilutions, the 10 best primer pairs for age-predictive 166 genes, and the two primer pairs for housekeeping genes, were kept for use in the study and 167 168 applied to 499 samples (298 females and 201 males), including 44 of the samples used for

169 RNAseq (the remaining 6 samples had too little cDNA left to be included in the qPCR study).

170 One of the samples failed to produce a Ct value for several genes and was therefore excluded

171 from subsequent analysis, leaving 498 samples (Fig. 1). All primers used in this study are

172 listed in Supplementary Data S3.

qPCR was run on a AriaMX RealTime PCR instrument in a total volume of 20 μl, containing
10 μl of SYBR 2x MM, 1.2 μl of forward primer (5μM), 1.2 μl of reverse primer (5μM), 6.6
μl of nuclease-free water and 1 μl of genomic DNA. Reaction conditions: one cycle of 95°C
(3 minutes), 40 cycles of 95°C (10 seconds) and 60°C (10 seconds), one cycle of 95°C (1
minute), 55°C (30 seconds) and 95°C (30 seconds, 5 seconds soak time).

178 Missing raw Ct values for age-predictive genes (where the signal never reached the threshold 179 even after 40 cycles) were replaced with the maximum value of 40. Δ Ct values were calculated using the mean Ct of the two housekeeping genes. Where Ct values were missing 180 181 for either housekeeping gene, normalisation was impossible and the normalised aging gene 182 value was recorded as missing (NA). All samples were run in two technical replicates and the final Δ Ct was taken as the mean of the two replicates. Gene GMOY005321 consistently 183 184 showed variable ΔCt values between technical replicates, possibly due to low expression of 185 this gene, and these values were kept unchanged. For all other genes, any gene-sample 186 combinations whose Δ Ct differed by more than 1 between technical replicates were rerun for 187 a third technical replicate, along with both housekeeping genes, providing a third Δ Ct. In most 188 cases, this third Δ Ct was very close to one of the first two and very different from the other, 189 indicating which of the first two technical replicates was wrong. The final Δ Ct was thus taken 190 as the mean of the third replicate and whichever of the first two replicates it was closest to.

191 **2.5 Predicting tsetse age from qPCR data.**

192 Machine learning predictions of tsetse age from qPCR data were performed using the *caret* 193 package in R (https://cran.r-project.org/package=caret). The Δ Ct values for each of the 10 194 study genes were used as continuous predictor variables, and sex was included as a 195 categorical predictor variable since some of the genes showed sex-dependent expression. 196 Samples were randomly split into training set (75% of samples) and test set (25% of samples), 197 stratified by sex and age to ensure equal representation of these two variables in the two sets. 198 Due to rounding of sample numbers within each stratification layer, the final numbers in the 199 train and test sets were 380 (76%) and 118 (24%) samples respectively. Model training was 200 performed using three rounds of 10-fold cross-validation. For regression models, whose aim



Figure 2: Gene expression clusters primarily by age. Principal component analysis of RNAseq data, coloured by age (left) or days since blood meal (right).

is to estimate age as a continuous variable, partial least squares regression (PLS), random forest and extreme gradient boosting (XGB) models were all trained on the data and their predictive accuracies compared. Categorical models were trained to categorise individuals into \leq 15 and >15 days old. Simple decision tree, random forest and XGB models were compared for these categorical models.

206 The minimum number of expression markers (genes) required to obtain accurate predictions 207 of age was determined by training the models with different numbers of loci. For each of the random forest and XGB models, the ten genes were ranked according to their variable 208 209 importance in the full model training described above (sex was found to have a variable importance of 0 in both cases, and was therefore excluded from these models). The models 210 211 were then trained with all ten genes, the top nine genes, the top eight genes, and so on. For 212 each set of genes, 20 models were trained with a different random split of training and test 213 sets, to account for stochastic variation in model accuracy.

214 All statistical analysis was conducted in R version 3.4.4 (R Core Team 2015). Analysis 215 scripts. qPCR raw data and RNAseq read counts are available on GitHub 216 (https://github.com/EricRLucas/TsetseAgeMarkers). Raw sequencing will be submitted to 217 ENA shotgun sequencing archive upon final acceptance of the paper for publication.

218 **3 Results**

We collected 301 female and 204 male *G. morsitans* flies of known age from laboratory colonies, ranging in age from 2 to 62 days old. An initial RNAseq analysis of 28 female and 221 22 male samples showed that gene expression in these samples was primarily affected by age, rather than sex or days since last blood meal (Fig. 2, Supplementary Fig. S1), although this

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Figure 3: Expression of ten age-related genes and two housekeeping genes from RNAseq data, ordered according to the variable importance in the XGB model (Fig. 4). Very strong early-age expression changes in some genes (eg: GMOY005321, GMOY002920) allow good discrimination among young individuals, but show little change in later life. Genes with continuous changes (eg: GMOY003371, GMOY000749) are more gradual and offer more consistent, but less powerful, discrimination at all ages.

- 223 was primarily due to the strong changes in gene expression found during the first 15 days of
- life, with older individuals clustering primarily by sex (Supplementary Fig. S2).

225 We identified a set of genes that was likely to provide strong age prediction by looking for genes that: 1. Were strongly correlated with age, or 2. consistently performed well in 226 227 prediction of age using lasso regression and 3. where possible, belonged to different gene 228 clusters as defined by weighted gene network clustering analysis. We particularly looked for genes showing strong expression changes in older individuals by identifying the genes most 229 230 differentially expressed when considering only individuals older than 15 days, but even these 231 showed relatively slight changes with age compared to some of the changes seen in the first 232 15 days of life (Fig. 3, Supplementary Fig. S3). Using our criteria, and after testing qPCR 233 primer efficient, we manually picked 10 genes associated with age, and 2 genes with very 234 little variation across samples to serve as housekeeping genes (Figs. 3 and 4).

Gene	Description	Top Drosophila BLAST hit	XGB import.	RF import.	XGB class. import.
GMOY005321	Cuticular protein 49Aa	tr A8DRW0 A8DRW0_DROME Cuticular protein 49Aa	100.0	60.1	3.9
GMOY002920	Cuticular protein 92F	tr Q9VDJ8 Q9VDJ8_DROME Cuticular protein 92F	71.4	72.0	100.0
GMOY003090	Porin	sp Q94920 VDAC_DROME Voltage-dependent anion-selective channel	70.4	68.6	2.6
GMOY003588		tr]Q7K188 Q7K188_DROME Protein quiver	67.9	100.0	11.7
GMOY001603	friend of echinoid	tr A0A023GPK8 A0A023GPK8_DROME Friend of echinoid, isoform H	26.5	78.4	2.0
GMOY003371	Elongation factor 1-alpha	sp P05303 EF1A2_DROME Elongation factor 1-alpha 2	18.5	84.3	0.3
GMOY000749		sp P05303 EF1A2_DROME Elongation factor 1-alpha 2	16.7	46.3	0.1
GMOY011979	Vacuolar H+-ATPase v1 sector subunit E	sp P54611 VATE_DROME V-type proton ATPase subunit E	7.9	38.7	0.1
GMOY005053		tr Q9VG81 Q9VG81_DROME RH49330p	6.8	29.8	6.4
GMOY009908		tr Q9VLZ6 Q9VLZ6_DROME FI24007p1	5.5	38.2	54.8
GMOY003952*	nuclear pore complex component	tr Q7K2X8 Q7K2X8_DROME Nucleoporin at 44A, isoform A	NA	NA	NA
GMOY010976*		sp Q9W123 POF_DROME Protein painting of fourth	NA	NA	NA

Figure 4: Ten age-related genes and two housekeeping genes (denoted with *) were used for qPCR analysis. Gene descriptions are taken from the Contig names in the GmorY1.9 proteome. Top Drosophila BLAST hits obtained by blasting the GmorY1.9 proteome against the D. melanogaster swissprot proteome. Variable importance of each gene shown for XGB, random forest (RF) and XGB classifier models trained with all predictor variables.



Figure 5: Prediction accuracy of the XGB model was highest (RMSE lowest) for individuals under 15 days old (2.59), and highest when all individuals were considered (6.81). Females are shown as blue triangles and males as orange circles. Purple line shows idealised perfect prediction.

We obtained qPCR measurements of expression for these genes from 297 females and 201 males (Fig. 1). As expected, expression of all 10 age-related genes was strongly correlated with age (Supplementary Fig. S4) and with the RNAseq data (Supplementary Fig. S5). Principal component analysis of these age-related genes showed that age dominated the first principal component of the data. In particular, samples clustered strongly into those younger and older than 15 days (Supplementary Fig. S6)

The qPCR expression data produced strong overall predictions of age, with predictions being 241 242 much more accurate in young flies (15 days or younger) compared to older flies. For regression models, PLS provided the poorest predictions of age, while random forest and 243 XGB models performed equally well (Fig. 5, Supplementary Fig. S7). Taking the XGB model 244 245 as an example, the overall root mean squared error (RMSE) for the final model was 6.74 days, 246 but was 2.96 for individuals ≤15 days old. Variable importance for each gene in the random forest and XGB models are shown in Fig. 4. Training the model separately for males and 247 248 females did not improve prediction accuracy (Supplementary Fig. S8).

249 Models also performed well at classifying samples into age categories of ≤15 and >15 days
250 old (Supplementary Fig. S9). The XGB model performed best in this task, accurately
251 classifying 117 out of 118 samples in the test set.

252 For both the random forest and XGB regression models, prediction accuracy showed little decrease when the variables of least importance were dropped from the models (Fig. 6). In 253 254 both cases, accuracy remained comparable to that with all 10 genes when only 6 genes were 255 included, with RMSE changing from 7.3 to 7.7 (random forest) or from 7.3 to 7.8 (XGB). In 256 contrast, when moving to 5 genes instead of 6, RMSE changed from 7.7 to 8.4 (random 257 forest) or from 7.8 to 9.3 (XGB). Interestingly, the same 6 genes proved to be sufficient for 258 both model types (GMOY005321, GMOY002920, GMOY003090, GMOY003588, 259 GMOY001603, GMOY003371). For the classification models, even fewer genes were needed 260 (Fig. 6), with just two genes being sufficient for XGB classification accuracy consistently 261 better than 95% (GMOY002920, GMOY009908).

262 **4 Discussion**

We have identified a set of gene expression markers that can be used to predict the age of *G*. *morsitans* tsetse flies in the laboratory. Importantly, this method can be applied to both males and females, providing accurate estimates of age in male tsetse. This is particularly important



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Figure 6: Predictive power of XGB and random forest models plateaus after the top 6 genes are included in the models (left). Accuracy of classification models plateaus after top 3 genes are included, with >95% accuracy achievable with only two genes (right). Small points show models run on independent test-train splits of the data (20 replicates per gene number); large points show the mean for each category. Points are jittered on the x axis to show overlapping data.

- since not only do both male and female tsetse flies transmit trypanosomes, but males appear 266 267 to be more likely to develop transmissible infections (Dale et al. 1995, Maudlin et al. 1990). 268 Our genetic markers were also unaffected by time since an individual's last blood meal, 269 making them more robust for use on wild-caught individuals, where such factors cannot be controlled. Further work is nevertheless required to test the applicability of these markers in 270 field conditions, since other environmental variables may still affect expression. For example, 271 272 temperature and humidity were constant in our rearing conditions, and all samples were 273 collected around the same time of day, leaving the possibility that these factors may yet 274 influence the expression of our markers.
- Like other methods for estimating the age of vectors, prediction accuracy decreases at older
 ages (Brei et al. 2004, Cook et al. 2006, Cook & Sinkins 2010, Gerade et al. 2004, Liebman et

277 al. 2015, Penilla et al. 2002, Sikulu et al. 2010). In our data, this was because the change in 278 expression with age was much greater in younger compared to older individuals, suggesting 279 that the overall physiology of tsetse changes slowly after a certain life stage, and that there is thus little to detect that can be used for age grading. While we found genes that continued to 280 change in older ages, the rate of change relative to the variance within age groups was not 281 282 sufficient to achieve the same prediction accuracies as found in younger individuals. While it 283 is likely that more accurate old-age predictions would be achievable using whole-284 transcriptome methods such as RNAseq, this is too costly to be applied at the scales required 285 for training predictive models. In mosquitoes, spectroscopy-based methods used to estimate 286 age initially suffered from a similar loss of precision at older ages (Liebman et al. 2015, 287 Mayagaya et al. 2009, Sikulu et al. 2010, Sikulu-Lord et al. 2016), but recent studies using 288 machine learning prediction methods have improved prediction accuracies (Lambert et al. 289 2018, Milali et al. 2019). Whether similar performance can be achieved with tsetse should be 290 explored.

291 While we used ten genes in our study, we found that using only the six genes most predictive 292 of age still provided high prediction accuracy, and only two genes were needed for classifying 293 individuals into age groups of ≤15 and >15 days old. By removing four genes from the 294 analysis, qPCR time and costs can be reduced by 1/3 (eight qPCR reactions per sample 295 instead of twelve), while removing eight genes will reduce costs by 2/3. We thus suggest that 296 further studies testing the applicability of these markers in the field restrict themselves to 297 either six or two genes, depending on how precisely age needs to be estimated. Such studies 298 are needed to determine the applicability of these markers in the field, but it would also be 299 interesting to measure the expression of these genes in age-controlled samples of other species 300 of tsetse to determine whether these markers have widespread applicability. Once the field applicability of these markers is confirmed, the technique can be rolled out in the context of 301 302 monitoring of tsetse control campaigns by comparing the age distribution before and after 303 interventions to confirm that a resulting shift in the population age distribution is observed. In 304 particular, in the wake of a 100% effective campaign, no flies older than the start of the 305 campaign should be found. The resulting data on age structure both before and after control 306 campaigns can then also be used to inform epidemiological models of trypanosomiasis 307 transmission.

308 In conclusion, our study provides a new method for estimating the age of tsetse flies which 309 does not require specialist dissection skills and can be applied to males. The problem remains

310 of finding methods for more accurately estimating age in older individuals. This may involve

311 identifying senescent changes whose rate is steady and consistent enough to be generalisable

312 to any individual in the population.

313 Acknowledgements

314 We are grateful to Rob Leyland for assistance and tuition for the insectary work, and to Tom

315 Churcher and Ben Lambert for valuable discussion on analytical approach. This work was

316 supported by a Liverpool School of Tropical Medicine internal grant (Director's Catalyst

317 Fund) to ERL and a Medical Research Council, UK (MR/T001070/1) grant to MJD and ERL.

318 SJT received support from the UK's Biotechnology and Biological Sciences Research Council

319 (grant numbers: BB/S01375X/1, BB/S00243X/1, BB/P005888/1) and the Bill and Melinda

320 Gates Foundation (INV-001785, OPP1155293).

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