Tsetse flies: Genetics, evolution, and role as vectors

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Keywords
Glossina, Trypanosoma, Tsetse, Gene flow, African trypanosomiasis

Disciplines
Entomology | Evolution | Population Biology | Terrestrial and Aquatic Ecology

Comments
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Tsetse flies: Genetics, evolution, and role as vectors

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Abstract
Tsetse flies (Diptera: Glossinidae) are an ancient taxon of one genus, Glossina, and limited species diversity. All are exclusively haematophagous and confined to sub-Saharan Africa. The Glossina are the principal vectors of African trypanosomes Trypanosoma sp (Kinetoplastida: Trypanosomatidae) and as such, are of great medical and economic importance. Clearly tsetse flies and trypanosomes are coadapted and evolutionary interactions between them are manifest. Numerous clonally reproducing strains of Trypanosoma sp exist and their genetic diversities and spatial distributions are inadequately known. Here I review the breeding structures of the principle trypanosome vectors, G. morsitans s.l., G. pallidipes, G. palpalis s.l and G. fuscipes fuscipes. All show highly structured populations among which there is surprisingly little detectable gene flow. Rather less is known of the breeding structure of T. brucei sensu lato vis à vis their vector tsetse flies but many genetically differentiated strains exist in nature. Genetic recombination in Trypanosoma via meiosis has recently been demonstrated in the laboratory thereby furnishing a mechanism of strain differentiation in addition to that of simple mutation. Spatially and genetically representative sampling of both trypanosome species and strains and their Glossina vectors is a major barrier to a comprehensive understanding of their mutual relationships.

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INTRODUCTION
Drosophila melanogaster Mieigen (Diptera: Drosophilidae) is probably the best-known insect, even though its agricultural significance is small. The reasons for its eminence are clear: easy culturing, high reproductive rate, small size, rapid generation time, easily demonstrable polytene chromosomes, and much genetic variation. Drosophila melanogaster is the favored lab animal of many thousands of scientists all over the world. In contrast, tsetse flies (Diptera: Glossinidae) are difficult to culture, have long generation times and low reproductive rates. Only a handful of cultures exist and the numbers of scientists that now work directly with tsetse flies are probably no more than a hundred. Only one laboratory in North America now works with tsetse. Nevertheless, knowledge of tsetse biology is not trivial or insignificant. Tsetse systematics, ecology, physiology, and vector biology have been intensively studied for a hundred and fifty years because of their medical and economic significance.
Tsetse flies are most unusual insects because of their exclusively haematophagous feeding habits and reproductive biology. There are six major texts in English devoted principally to tsetse flies (Buxton, 1955; Glasgow, 1963; Nash, 1969; Ford, 1971; Jordan, 1986; Leak, 1998). These texts review a particularly rich literature on ecology, trypanosomiasis epidemiology and epizootiology, pest management, and history. Investigation of tsetse genetics and physiology, on the other hand, has been greatly inhibited by the difficulty and cost of culturing tsetse flies, their long generation times, low reproductive rates, and sampling of natural populations that has been geographically limited and mostly opportunistic. Earlier reviews of tsetse genetics include Gooding (1984) and Gooding and Krafsur (2004, 2005). Knowledge of trypanosome biology and genetics, on the other hand, is rather well developed because, apart from being medically and intrinsically interesting, trypanosome cultures can be propagated in the laboratory independently of tsetse flies. A thorough understanding of disease epidemiology and management, however, requires information that relates parasite distribution to that of its vectors (Tibayrenc and Ayala, 2002; De Meeus et al., 2007).

Interest in tsetse population genetics has recently developed, fueled by the notion that genetic methods can promote effective area-wide population management by (1) better defining taxonomic units (Gooding and Krafsur, 2005) and estimating degrees of isolation of populations of interest (e.g., Feldmann, 2004; Vreysen, 2006) or (2) modulating vector competence by genetic means (e.g., Aksoy et al., 2001). In these ways it is hoped that the well-known propensity of tsetse dispersal can be overcome and long-term tsetse population management can be achieved, thereby greatly reducing, or eliminating trypanosomiasis altogether.

In this article, I review tsetse fly evolution, trypanosome interactions, and genetics with an emphasis on population and ecological genetics as found in the more recent literature since Gooding and Krafsur’s review (2005).

Systematics and evolution of tsetse flies

TSETSE FLIES are classified into a single genus Glossina Weidemann 1830. There are three extant subgenera, Austenina Townsend, Nemorhina Robineau-Desvoidy, and Glossina Wiedemann that correspond to the Fusca, Palpalis, and Morsitans species groups respectively. A new subgenus, Machadomia Dias 1987, was established to incorporate the anomalous tsetse, G. austeni Newstead. There is an extinct sister group to extant Glossina known from the Florissant shale of Colorado, dating from 35 mya (Grimaldi, 1992) and Glossina-like fossils have recently been recovered from Oligocene strata in Germany (Grimaldi and Engel, 2005). It seems, therefore, that tsetse flies (to include the putative, unnamed, sister taxon) had nearly a worldwide distribution some 30 to 40 million years ago. Their origins, therefore, must predate continental separation in the Cretaceous, more than 100 MYA. The higher classification of Glossina is, following McAlpine (1989) and Grimaldi and Engel (2005): suborder Brachycera – infraorder Cyclorrhapha – Section Schizophora – Subsection Calyptrata – Superfamily Pupiparia (= Hippoboscoidea) – Family Glossinidae. An extinct sister taxon has not been agreed upon. The other families in the Pupiparia (Hippoboscidae, Streblidae, Nycteribiidae) are more highly specialized morphologically than Glossina and are true ectoparasites. All Pupiparia reproduce by adenotrophic viviparity, such that a zygote develops and hatches in the female’s reproductive tract and the larva feeds on “milk” produced by the female’s reproductive accessory glands until it completes its development. Hippoboscoidea phylogeny and evolution was recently addressed by Peterson et al. (2007).

Thirty-three extant taxa have been described of 22 species that include five species complexes (Gooding and Krafsur, 2004 for review). Leak (1999) lists 31 taxa. Thirty taxa evaluated by Rogers and Robinson (2004) have well-defined habitats in terms of satellite imagery derived from...
variables. The species complexes include *G. morsitans morsitans* Westwood 1850, *G. m. centralis* Machado 1970, *G. m. submorsitans* Newstead 1910 and *G. m. submorsitans ugandensis* Vanderplank 1949; *G. palpalis palpalis* Robineau-Desvoidy 1830 and *G. p. gambiense* Vanderplank 1949. The foregoing taxa are particularly important medically and economically. Other subspecies include *G. pallicera pallicera* Bigot 1891 and *G. p. newsteadi* Austen 1929; *G. nigrofusca nigrofusca* Newstead 1910 and *G. n. hopkinsi* Van Emden 1944; *G. fuscipes fuscipes* Newstead 1910, *G. f. quanzensis* Pires 1948, and *G. f. martini* Zumpt 1933. Taxa within each species complex are mostly allopatric but morphologically indistinguishable from their sisters, or very nearly so. Results of recent studies suggested incipient speciation in *G. p. palpalis* although alternative explanations also were offered (Gooding et al., 2004; Ravel et al., 2007). With the likely exception of *G. p. palpalis* in Bonon, Cote d’Ivoire, sibling species are allopatric, hybrid males are sterile, and the hybrid females vary in their fertilities. The *G. morsitans* subspecies share little microsatellite variation among them, are reproductively isolated intrinsically and extrinsically, and it was suggested they should be raised to specific rank (Krafsur and Endsley, 2006).

Extant tsetse flies are confined to sub-Saharan Africa, although *G. fuscipes fuscipes* and *G. morsitans submorsitans* have been recorded in the southwestern corner of the Arabian Peninsula (Elsen et al., 1990). They are relicts from earlier times when more equable climates prevailed.

Glossinidae, indeed, all Pupiparia, are exclusively haematophagous. Reproduction in tsetse is by adenotrophic viviparity: a larva develops to maturity in its mother over a temperature dependent 7 - 12 day interval. When mature, the mother deposits her larva in an appropriate microhabitat and the animal pupariates in the soil. Adult development requires about 27 days at 25°C. Females become inseminated by the end of their first week of adulthood. The youngest larvapositing female is at least 16 days old, so the minimum time necessary to produce two offspring is about 25 days. Generation time is c. 43 days at 25°C. Compensating for their slow reproduction rates are their adult survival rates, which typically exceed 97% per day (Rogers and Randolph, 1984a, b; 1985).

**Medical and economic significance**

The impact of tsetse flies is severe in much of sub-Saharan Africa because of their blood feeding habits and their role as vectors of pathogenic trypanosomes. Chief among these is *Trypanosoma (Trypanozoon) brucei brucei*, *T. b. rhodesiense*, and *T. b. gambiense*, two of which are highly pathogenic to man, the agents of ‘sleeping sickness’ or human African trypanosomiasis (HAT). It seems that *G. brucei* s.l. are not well-defined taxonomic units nor clinical entities, however (Hide and Tait, 2004; Gibson, 2005). *T. b. gambiense* infection causes a chronic, slow wasting disease that ultimately causes death if untreated. This form of HAT occurs in West and central Africa and is unknown east of the Rift Valley (Welburn et al., 2001). Its chief vectors are *Palpalis* group flies. Humans are the principal reservoir of *T. b. gambiense*. The parasite has been isolated from pigs, dogs, and sheep but the epidemiological significance of this remains opaque. *T. b. rhodesiense* is zoonotic and causes a much more acute disease in humans that is rapidly fatal if not properly treated. *T. b. rhodesiense* occurs east of the Rift valley. Its vectors are *Morsitans* group flies. Uganda is the only country where both forms of HAT are known to occur. They do so in separate foci and there is a concern that the two agents soon will overlap because transport of infected cattle can introduce *T. b. rhodesiense* to western Uganda (Picozzi et al., 2005). *T. b. brucei* occurs throughout sub-Saharan Africa wherever its vectors may be found. It is not infective to humans because it is rapidly lysed by human serum. The principal vectors of HAT include *G. palpalis* s.l., *G. morsitans* s.l., and *G. tachinoides* Westwood 1850. *G. fuscipes fuscipes* is the chief vector in Uganda and western Kenya. The principle foci of HAT seem to be static and ancient. The epidemiology of HAT has been comprehensively
treated by Welburn et al. (2004) and an epidemiological update is available from the WHO (2006). Trypanosome molecular epidemiology was reviewed by Hide and Tait (2004).

Trypanosomiasis in domestic animals is termed ‘nagana’, a chronic wasting disease that precludes most animal agriculture in much of sub-Saharan Africa but poultry are unaffected by trypanosomiasis. In addition to *T. b. brucei* and *T. b. rhodesiense*, tsetse-transmitted agents include *T. vivax, T. congolense, T. simiae, and T. godfreyi* (Stevens and Brisse, 2004). *T. evansi* is pathogenic but not transmitted by tsetse flies. Nagana is of much economic significance because where it is prevalent; meat, milk, dung, and draft power production are greatly reduced or lost altogether. The long-term consequence of greatly reducing Nagana is controversial, however, because it has been argued that its elimination will inexorably lead to habitat degradation: deforestation, overgrazing, and erosion (ERGO, 1999).

Tsetse population control methods include aerial application of ultra-low volume insecticide applications, the use of insecticide-laced targets and traps, and the sterile insect technique (Torr et al., 2006). Research efforts are well underway to find ways to employ transgenesis to render vector tsetse flies insusceptible to trypanosome infection and replace natural vector populations with refractory ones (Aksoy et al., 2001). Earlier control methods, no longer extensively applied, included ground spraying of persistent insecticides such as DDT, the clearing of brush, and extermination of native mammals that provide tsetse fly blood meals and act as trypanosome reservoirs.

Vector-parasite-mammalian co-adaptations

*Trypanosoma brucei* s.l. salivary gland infection rates in tsetse flies are low, typically < 1% in (Maudlin et al., 1998). Much higher infection rates may be expected given the great longevity of the flies (adult daily survival rates typically exceed 97% and half lives exceeding ~ 28 d) and prevalence of trypanosome-infected animals. A likely reason that tsetse infection rates are small is they have a robust innate immune system in which most ingested trypanosomes die in the fly midgut. Once established in the midgut, procyclic trypomastigotes must transform to epimastigotes and proceed via the proventriculus and foregut to the host salivary glands where they develop to infective metacyclic trypanosomes. Only a small fraction of gut infections succeed in developing into infective metacyclic forms lodged in the vector salivary glands. In laboratory studies, injection of *E. coli* into tsetse flies induced immune stimulation thereby reducing trypanosome-infective feeds and the anti-microbial peptides attacin, defensin, and diptericin were demonstrated in tsetse tissues (Hao et al., 2001). Attacin was shown to provide an important immune response that regulates trypanosome infection in some, but not all, *Morsitans* group flies (Hu and Aksoy, 2006; Nayduch and Aksoy, 2007; Roditi and Lehane, 2008). The prophenoloxidase cascade (Nigam et al. 1997) and phagocytosis of trypanosomes by haemocytes have also been suggested to play a role in vector competence. Another mechanism that may suppress trypanosome development in tsetse is the production of certain midgut lectins (reviews in Welburn and Maudlin, 1999; Rodite and Lehane, 2008). Lectins are proteins that bind specifically with carbohydrates to form glycoproteins. Lectin titers increase in post-teneral flies and help to explain the insusceptibility of older flies to new trypanosome infections. The foregoing picture, however, has been modified by recent experimental work that demonstrated the addition of antioxidants (e.g. ascorbic acid, uric acid, glutathione, N-acetyl-cysteine) to blood meals increased *T. b. brucei* midgut infection rates in *G. m. morsitans* – remarkably to 100% in some experiments. The mechanism of trypanosome suppression by lectins, therefore, is related to their properties as pro-oxidant free radicals induced during digestion of blood meals (MacLeod et al., 2007a). Maturation and migration of midgut forms to the salivary glands, however, also seems to require reactive oxygen species nitric oxide and/or L-cysteine (MacLeod et al., 2007b). Clearly, the pathway from ingestion of blood forms by tsetse flies to the production of infective metacyclic trypomastigotes in the
tsetse salivary glands is long and complex, with many critical steps that developing trypanosomes must overcome.

Susceptibility to midgut infection was reported to be under genetic control and inherited matrilinearly in *G. m. morsitans* (Maudlin, 1982) and refractory and susceptible lines have been established experimentally (Welburn and Maudlin, 1999). The mode of matrilinearly inheritance can be attributed to the commensal inter- and intracellular bacterium in tsetse flies, *Sodalis glossinidius* (Dale and Maudlin, 1999). Recent work of MacLeod et al. (2007a), however, raises the question, do the *Sodalis* flora of natural tsetse fly populations vary in their production of anti-oxidants and is this the underlying mechanism of susceptibility to trypanosome infection in tsetse flies? Could engineered *Sodalis* eventually be used in a scheme to replace vectorally competent natural tsetse populations with transgenic, insusceptible populations?

Salivary gland infected tsetse flies in the laboratory have a reduced expectation of life and males show greater infection rates than females (Maudlin et al., 1998; Welburn and Maudlin, 1999). Laboratory studies using *T. b. brucei* and *G. m. morsitans* have demonstrated that reproductive success by female tsetse can depress the proportion of midgut infections that mature, but midgut infections do not detectably depress tsetse reproductive success (MacLeod et al., 2007b). Co-infection of tsetse flies by more than one trypanosome species and strain is commonly observed in field samples (MacLeod et al., 1999; Lehane et al., 2000). Experimental co-infection and superinfection of *G. m. morsitans* by two fluorescently marked *T. b. brucei* strains demonstrated that salivary gland infection by one strain of trypanosomes did not inhibit the salivary gland establishment of another strain; the chief determinant was the insect’s susceptibility to a mature infection (Peacock et al., 2007). The finding is important because genetic recombination between trypanosome strains takes place in the tsetse salivary glands (Gibson and Stevens, 1999) as discussed below.

Are there locally co-adapted trypanosome-tsetse demes distributed throughout a tsetse belt? Genotyping of *T. brucei* isolates indicate both spatial and host differentiation (e.g., MacLeod et al., 2001). Probably more *T. brucei* s.l. strains will be uncovered with the development of additional genetic markers and improved techniques. The source of such differentiation is mutation and possibly genetic recombination, first demonstrated in laboratory experiments (e.g., Jenni et al., 1986). Indeed, genetic recombination has been shown to occur among the three *G. brucei* ‘subspecies’ but the extent of genetic recombination between trypanosome populations in the field remains controversial and largely unknown. It is believed that trypanosome mating and meiosis take place in the tsetse fly salivary glands, probably among epimastigotes (Tait et al., 2007; Gibson et al. 2008). Meiosis in trypanosomes provides an essential adaptive mechanism and may explain the numerous strains that have been described (Gibson, 2001)

Is trypanosome genetic differentiation adaptive, i.e., is it maintained by natural selection? In particular, is it maintained by adaptation to the locally prevalent tsetse vectors? Gene flow among natural *T. b. brucei* and *T. b. rhodesiense* populations is said to be absent or greatly limited; indeed, the two are said to be genetically isolated from each other with *T. b. brucei* showing epidemic population structure and *T. b. rhodesiense* showing clonal population structure (MacLeod et al., 2000). Estimating the strength and degree of substructuring among *T. brucei* spp in nature, however, seems a herculean task, and I know of no attempts to evaluate trypanosome population structure over large areas or to evaluate it with respect to its tsetse vectors. Indeed, most trypanosome research is conducted without reference to its vectors. *T. brucei* s.l. loses its ability to infect tsetse flies after multiple passages in laboratory mice thereby calling into question the epidemiological significance of some findings. Biased sampling of natural *T. brucei* s.l. populations is a formidable problem such that their population structures are poorly known; the bias arises from a requirement of obtaining large numbers of blood forms.
from infected hosts and amplifying their numbers in mice before genotyping them (Garcia et al., 2006).

To complete the triangle of co-adaptation, trypanosomes vary in their infectiousness to mammals. For example, *T. brucei brucei*, ubiquitous in mammals throughout sub-Saharan Africa, is rapidly lysed in human serum but *T. b. gambiense* and *T. b. rhodesiense* are not, and therefore are pathogenic to humans as well as many other mammals. A single gene, the serum resistance associated (SRA) gene, is sufficient to confer the trait of human infectivity on *T. brucei brucei* (Xong et al., 1998); all known strains of *T. b. rhodesiense* have this gene but *T. b. gambiense* does not, and therefore must have another mechanism of human serum resistance (Gibson, 2005).

Susceptibility of tsetse species to trypanosome infection in lab trials may not be strongly predictive of their vectorial capacity in nature, for a number of reasons. First, tests of susceptibility/refractoriness typically are based on a single genotype of trypanosome and a well-adapted laboratory strain of Glossina. As we have seen, susceptible and insusceptible lines of a common lab strain of *G. m. morsitans* have been developed, and it will be shown that tsetse field populations are typically genetically differentiated - often, highly so. A single *Glossina* sp lab strain is most unlikely to be representative of the species over much of its geographic range and multiple, independently derived lab strains from diverse natural populations are unavailable. Second, even though HAT epidemics are typically of one trypanosome genotype, i.e., clonal (but see Agbo et al., 2003 for contrary evidence), natural populations of *T. brucei* s.l. are genetically differentiated in ways that remain largely unknown. Indeed, an objective and scientifically adequate appreciation of vector-parasite compatibilities in the field is quite beyond reach at this time because the sampling problems seem insurmountable.

## Genetics and cytogenetics of tsetse flies

All tsetse examined have two pairs of metacentric autosomes and a sex bivalent: 2N = 4 + XY. *G. m. morsitans*, *G. m. submorsitans*, and *G. m. centralis* also have small, telocentric, heterochromatic supernumerary chromosomes that vary in number from zero to eight (Warnes and Maudlin, 1992). A lab culture of *G. austeni* Newstead 1912 had rather more supernumerary chromosomes that varied from 8 to 12 in number (Southern, 1980). Supernumararies vary from 12 to 22 in two taxa of *Fusca* group flies (review in Gooding and Krafsur, 2005). Sex chromosome karyotypes XXY, XO, XXY, XYY were found in Nigerian populations of *G. palpalis palpalis* (Maudlin, 1979). The high frequency of these polymorphisms suggested that they are maintained by some kind of balancing selection instead of high frequencies of spontaneous non-disjunction.

Well-banded polytene chromosomes can be obtained from pupal trichogen cells found in apical scutellar bristles (Southern and Pell, 1974). Gariou-Papalexioi et al. (2007) compared banding patterns among *G. pallidipes, G.m. morsitans, G. m. submorsitans*, and *G. austeni*. The karyotypes of these taxa differ by numerous paracentric inversions in the X chromosome and chromosome L1. *G. austeni* differs from the others by bearing a pericentric inversion on chromosome L2. *G. m. morsitans* and *G. m. submorsitans* are homosequential and differ only by a paracentric inversion on the X and another on the L2 chromosomes.

Evidence to date shows abundant chromosome diversity among *Morsitans* and *Palpalis* group flies. Chromosome diversity, however, has been examined inadequately among and within species; few tsetse cultures exist and sampling wild populations cytogenetically required rather more interest and support than available. Clearly there is much to be learned by further study, and it is to be hoped that surveys of natural populations can be initiated.
Formal genetics and genetic markers

Five classical morphological markers have been described and these were assigned to two linkage groups in *G. p. palpalis*, *G. m. morsitans*, and *G. m. submorsitans*. Each of thirteen biochemical markers was assigned to one of three linkage groups (Gooding and Rolseth, 1995; Gooding and Challoner, 1999). Composite maps are reviewed and shown in Gooding and Krafsur (2005). The X chromosome map was recently updated with two microsatellite loci in *G. palpalis* (Gooding et al., 2004). There are three linkage groups in each species examined, but they have not been associated with their physical chromosomes.

Population genetics

The forces of evolutionary change include mutation, selection, drift, and migration. When these forces do not operate there is random mating and no change in gene frequencies from one generation to the next. This is the Hardy-Weinberg rule that provides the basic null hypotheses with which to evaluate gene frequencies within and among demes. Natural populations, however, are of finite size and more or less discontinuously distributed so that chance variations in gene frequencies occur from place to place and time to time. Mutations arise and contribute to gene diversity. Matings are more likely within subdivided populations than among them. Selection may act locally and differentially on genotypes thereby contributing to spatial diversity. Migration of reproductives among demes tends to neutralize the forces of drift and local selection. A primer of population genetics applied to molecular epidemiology can be found in an earlier volume of this journal (De Meeus et al., 2007).

Ecological background

Clearly the outcome of Hardy-Weinberg criteria applied to tsetse fly demes will depend on their distribution and abundance. *Morsitans* group flies are largely savanna and woodland inhabitants, although *G. pallidipes* may also be found in forests. *Morsitans* group taxa are adapted to drier habitats than the other two subgenera (Rogers and Robinson, 2004). *Palpalis* group flies tend to occur in riverine and lacustrine habitats. *Fusca* group flies largely inhabit moist forests of West Africa although *G. brevipalpis* occurs discontinuously in East Africa, Zaire, and Mozambique (Jordan, 1963; Leak, 1998). Indeed, maps show that many *Glossina* taxa are discontinuously distributed according to environmental criteria that include host blood meal sources and abiotic factors. Abiotic factors have been extensively evaluated in terms of satellite imagery (Robinson et al., 1997a, b; Rogers, 2000; Rogers and Robinson, 2004). The chief predictor variables include NDVI, the normalized density vegetation index, elevation, maximum and minimum temperature indices, rainfall, and saturation deficits. Different combinations of variables are necessary for each tsetse species and subspecies. These approaches have allowed a detailed mapping of suitable tsetse habitats in sub-Saharan Africa (Rogers and Robinson, 2004 and references therein). The distribution of tsetse flies spans many ecological zones, among which, different discriminant models and predictor variables best describe the distribution (Robinson et al., 1997a). More accurate predictions of tsetse distributions are possible if an area is first subdivided into ecological zones, and the distribution within each zone treated separately. This suggests the existence of population clusters (i.e., subpopulations or demes) in which the flies have adapted to their differing environments. It seems likely, therefore, that the flies’ adaptations may differ from one cluster to another.

Life history and dispersal

Male *G. m. centralis* are active only a mean 32 minutes per day (Bursell and Taylor, 1980). Presumably, males and females of the other species show similar activity patterns. Tsetse dispersal is a critical component of models constructed to evaluate efficacies of area-wide population management procedures and to evaluate historical biogeographical events (Hargrove, 1981; Williams et al., 1992; Hargrove, 2000; Vale and Torr, 2005). Dispersion rates
are said to be high, compromising otherwise effective control programmes (Williams et al., 1992; Brightwell et al., 1997; Hargrove, 2003). Rogers (1977) modeled dispersion as a random diffusion process in which the distance \( d \) moved \( \approx s \sqrt{t} \), where \( s \) is the step length (the mean daily displacement) and \( t \) is time in days. Rogers calculated daily displacement rates for 13 field studies, estimated over a scale of days to 17 years. The overall mean \( s = 252 \text{ m} \) for savannah species. This implies a frontal advance of 1.4 km a month and 4.7 km a year. Riverine taxa are said to disperse at higher rates. A more recent review offered by Hargrove (2000) supports Rogers’ analysis.

The foregoing citations predict significant rates of gene flow. There is a caveat, however. Dispersion is delayed because teneral and young flies are less likely than older flies to disperse because their flight muscles require, at tropical temperatures, 2 – 3 bloodmeals and 6 - 8 days to mature (Hargrove, 2003, 2005). Female flies become receptive at 3 days of age and virtually all are inseminated by 6 - 8 days. Under these conditions, most matings are likely to be between members of a deme or nearby demes. Exchange of flies among demes, therefore, critically depends on post-teneral dispersal rates and the reproductive success of emigrants in their ‘new’ environments.

Genetic variation in tsetse flies

Theory indicates a relationship between population size and genetic diversity (\( H_e \)): \[ H_e = \frac{4N_e \nu}{4N_e \nu + 1} \] where \( N_e \) is effective population size (typically a large number) and \( \nu \) is the mutation rate (a very small number, \( \sim 10^{-4} \) to \( 10^{-9} \)). Tsetse populations seem small when compared with Diptera such as *Drosophila*, Muscoid flies, mosquitoes, etc. Indeed, early studies suggested a paucity of genetic variation in tsetse flies. Morphological mutants were few and allozyme diversity low (Gooding, 1992), features that could be attributed to historically small population sizes. More recent, comprehensive surveys of allozyme variation confirmed the low diversity but disclosed sufficient variation to allow studies of gene flow in natural populations (Krafsur et al., 1997). Comparative genetic statistics for allozymes (Table 1) generally indicate lesser diversities (heterozygosities) than occur in other medically and economically important Diptera (Krafsur and Griffiths, 1997). The four taxa investigated in *Morsitans* group flies, moreover, showed closely similar heterozygosities although the proportion of polymorphic loci was less in *G. swynnertoni* Austen 1923. The homogeneous heterozygosities are notable because *G. swynnertoni* has a much smaller geographic range than the others and its populations have contracted in recent times; indeed, its effective population size is quite small, of tens to hundreds of flies (Marquez et al., 2006). Mitochondrial variation in *G. swynnertoni*, on the other hand, was less than in other *Morsitans* group taxa. A similar picture was obtained in Zimbabwe and Zambian populations of *G. pallidipes*, among which allozyme diversity was greater than that in Kenya populations, but mitochondrial and microsatellite variation was much reduced (Krafsur, 2002a). The conservation of allozyme variation in two tsetse species seems to be an example of balanced polymorphism in which selective advantage accrues to heterozygotes at structural gene loci. A similar phenomenon was observed in *Drosophila* (Hale and Singh, 1991).

Microsatellite variation

Simple sequence repeats (SSRs) occur throughout the nuclear genome. They are codominant and highly polymorphic. SSRs have been developed for the *Morsitans* group (Baker and Krafsur, 2001; Ouma et al., 2003, 2006). Some primers for *Morsitans* group tsetse also amplify orthologous loci in the *Fusca* and *Palpalis* groups. SSRs have also been developed specifically for *G. palpalis gambiensis* (Luna et al., 2001) and *G. f. fuscipes* (Abila et al., 2008).

Diversities in microsatellite loci vary among tsetse taxa and compare with those in house flies, which have very much larger effective populations sizes (Table 2). Rather more loci would be
helpful in furthering genetic understanding of tsetse flies. SSRs are expensive and time
consuming to identify and evaluate as useful markers. They are typically isolated from
untranscribed regions of the genome such that well conserved flanking regions that serve as
good priming sites are not abundant, leading to high frequencies of null alleles and poor
amplifications in related taxa. Expressed sequence tag libraries are available for tsetse (Lehane
et al., 2003) that could be used to find and develop new SSR loci thereby avoiding the problems
often associated with loci in unexpressed genomic regions (Kim et al., 2008).

Population structure: genetic differentiation and gene flow among demes

Estimators of genetic differentiation include $F_{ST}$ (Wright, 1978), $G_{ST}$ (Nei, 1987), and theta
($\theta$) (Weir & Cockerham, 1984). $F_{ST}$ and its analogues describe departures from random matings
among demes at hierarchical population levels. These estimators are based on different
assumptions, but each attempts to measure the same thing. $F_{ST}$ can be considered as the ratio
between within deme diversity $H_S$ and the diversity in the total population $H_T$ thus
($F_{ST} = 1 - H_S/H_T$), and as correlations between uniting alleles. In theory, $F_{ST}$ can take a value from $-1$
to 1, but that holds only for a single locus with two alleles. In practice, estimates cannot exceed
the mean level of homozygosity in subpopulations, $1 - H_S$ (Hedrick, 1999). Departures from
random mating within demes (= subpopulations) are estimated by $F_{IS}$ (individuals in
subpopulations).

The geographic scale of sampling tsetse flies varied from hundreds of meters to thousands of
kilometers. Often it was found that a nested design was appropriate, for example samples nested
in regions. High indices of genetic differentiation can indicate the occurrence of sibling species
and must be considered given the substantial number of sibling species (morphologically
identical species) that have been detected in other Diptera. Evaluating the question in the
Glossinidae is not a simple task, however, because sampling and establishing geographically
diverse cultures for genetic analysis is extraordinarily time consuming and expensive. Existing
lab cultures are few and longstanding. For example, the $G. fuscipes fuscipes$ culture dates from
1986 or earlier – documentation about its origins are now lost although it is thought the strain
originated in the Central African Republic (see Krafsur et al., 2008). $G. morsitans s.l.$ cultures
date from the 1960s. The final test of intrinsic reproductive isolation is to cross reciprocally
different strains and evaluate the fertilities of the matings and the fertilities of any progeny in
backcrosses. Nevertheless, a number of investigations have demonstrated sibling species. A
study in chromosome mapping has disclosed the putative existence of two mating types in $G.$
$palpalis palpalis$, one derived from Nigeria and the other from Zaire (Gooding et al., 2004).

Much earlier, $G. m. morsitans$, $G. m. submorsitans$, and $G. m. centralis$ were separated initially
on the basis of hybrid sterility and shown to be allopatric (Vanderplank, 1949a). More recent
studies on the phylectic relationships among $G. morsitans s.l.$ and the closely related $G.$
$swynnertoni$ have been reviewed (Gooding and Krafsur, 2004, 2005).

Population structure in subgenus Morsitans – the Morsitans group

$Morsitans$ group flies occur chiefly in the wooded savannahs of East and southern Africa where
they are medically and economically highly significant (Fig. 1). Distribution maps (Rogers and
Robinson, 2004) show that only $G. longipalpis$ Weidemann 1830 is exclusively West African;
$G. pallidipes$ Austen 1903, found in East and southern Africa, is morphologically and
ecologically similar (Vanderplank, 1949a; Leak, 1999) but no crosses between them seem to
have been reported. It would be most interesting to investigate their phylogenetic relationships.

As we have seen, allozyme, microsatellite, and mitochondrial variation have been characterized
in $G. morsitans s.l.$ and $G. pallidipes$. Summary statistics for the foregoing taxa are offered in
Tables 2 and 3. In making within-taxon comparisons, it should first be observed that
improvements in methods have a strong bearing on summary statistics. In $G. m. morsitans$, for

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example, the use of automated nucleotide-sequencing technologies and proprietary allele scoring software doubled the number of alleles detected when compared with silver-stained 4.5% acrylamide gels in the author’s lab. Estimates of diversity $H_e$ were not affected, but $F$ statistics differed. Use of automated sequencing methods for mitochondrial PCR amplicons provided greater numbers of haplotypes, hence greater diversities, than the SSCP technique. In particular, the number of singletons detected was greater, a demographically important index because a high frequency can indicate recent population expansion or the operation of natural selection.

G. morsitans sensu lato consist of subspecies principally because they are morphologically difficult to distinguish and intermate freely in cages (Vanderplank, 1949a). G. swynnertoni was considered to be only a variant of G. morsitans s.l. for the same reason. Machado (1970) regarded G. morsitans s.l. to be only ‘major geographical races’ even though Vanderplank had demonstrated hybrid sterility, i.e., reproductive isolation. Numerous studies have shown that hybrid males are sterile and the females sterile or semisterile (reviews in Gooding and Krafsur, 2004, 2005). Genetic studies, moreover, provide no tangible evidence of gene flow among natural populations of the three taxa. The subspecies are entirely allopatric. Clearly, these taxa have a common ancestor and share large regions of their genomes, but each has evolved different adaptations. For example, G. m. centralis Machado 1970 occurs in cooler, wetter conditions, compared with G.m. morsitans Westwood 1850, which occurs in hotter and drier areas (Robinson et al. 1997a, b). Krafsur and Endsley (2006) argued that G. morsitans subspecies should be elevated to species rank because genetic evidence indicated complete premating and postmating reproductive isolation among them.

G. m. morsitans is distributed in four broad belts. Two belts occur in Mozambique, one of which extends well into southern and coastal Tanzania. Another large belt occurs from northern Zimbabwe through eastern Zambia (Rogers and Robinson, 2004). This G. m. morsitans belt lies immediately east of two extensive G. m. centralis belts (Robinson et al., 1997a, b). Samples for genetic analysis were obtained from western Mozambique, Zimbabwe, Zambia, and Tanzania at distances of 12 to 917 km (Wohlford et al., 1999; Krafsur and Endsley, 2001; Ouma et al., 2007). Genotypes were obtained for seven microsatellite loci and mitochondrial locus Cox I. Microsatellite allelic diversities (22.6 alleles per locus) and heterozygosities ($H_e = 0.77$) were greater in four Tanzanian populations than in five southern African populations (18.7 and 0.71, respectively). Thirty-three Cox I haplotypes were recorded in 96 flies from seven populations (only three southern Africa populations could be amplified by PCR). Only one haplotype was shared among six populations and it occurred in 43 (45%) G.m. morsitans. Seventy haplotypes (73%) were singletons. Tanzania flies had the most private and singular haplotypes; diversity was 0.72 in Tanzania and 0.50 in southern Africa. An hypothesis of isolation by distance (IBD) was not rejected when based on microsatellite loci, but the slope was very shallow. IBD was rejected, however, when based on mitochondrial variation.

G. m. centralis populations are geographically discontinuous. This tsetse occupies an area in central Angola small relative to its chief habitats in western Tanzania, Zambia, and southeastern Democratic Republic of Congo (Zaire). A separate population occurs also in Botswana, the Caprivi Strip of Namibia, and southeastern Angola. Genetic sampling of six populations was carried out in Zambia, Botswana, and Namibia (Krafsur and Griffiths, 1997). No samples were obtained, however, from the huge belt in Tanzania and northern Zambia. Mitochondrial and microsatellite diversities were very low in G. m. centralis, compared with diversities in other Morsitans group tsetse. For example, only seven mitochondrial haplotypes were detected over two loci. Allozyme diversity indices of average alleles per locus and mean heterozygosity, on the other hand, were homogeneous with other G. morsitans s.l. when measured over 31–45 putative loci. Diversity (heterozygosity) at only polymorphic allozyme loci, however, was less in G. m. centralis: 18.7% vs 31.9%. Genetic differentiation was pronounced among the six
populations sampled; $F_{ST}$ for mitochondrial variation was 0.87 and 0.19 for microsatellites. The difference in estimates can be attributed to high mutation rates, the much greater sensitivity of the mitochondrial genome to demographic change, and homoplasy at microsatellite loci. It is notable that 171 of 174 flies from Botswana and the Caprivi Strip, taken from more than 10 sampling locations, possessed the same haplotype, suggesting the sampled populations originated from an earlier bottlenecked population. Periodic insecticidal treatments were applied aerially to Botswana from 1973 to 1991 with institutional claims of success, but the genetic data are consistent with the ecological analysis of Hargrove (2003), viz, survival of the resident population well within the treated region. Clearly, *G. m. centralis* populations are strongly subdivided through southern Africa, a consequence of geographic isolation and genetic drift. It will be interesting to see if genetic diversities among Tanzanian populations are similar to those recorded in Zambian and Botswana populations. *G. m. centralis* disappeared from large areas of southern Africa (Ford, 1971) because of the rinderpest, and only slowly recovered from very low, undetectable, population numbers. Thus, the paucity of genetic variation was caused by genetic bottlenecks. The species has now been eliminated by aerial applications of the pyrethroid insecticide deltamethrin over 16,000 km$^2$ of the Okavango delta in Botswana in 2001-2002 (Kgori et al., 2006). No tsetse flies have been detected by continuous and intensive sampling since (P. Kgori, Veterinary Services, Botswana, personal communication, 16/5/08). It should be noted that conspecific populations nearby in Namibia and southern Zambia were not treated nor have they yet furnished detectable surviving immigrants to Botswana.

*G. m. submorsitans* Newstead 1910 occurs in Guinea Bissau east to Uganda and southwestern Ethiopia but its distribution is highly discontinuous. The largest discontinuities occur in Nigeria and southern Sudan (Rogers and Robertson, 2004). Examination of male genitalia led Vanderplank (1949b) to describe four morphologically distinct races, later supported by Machado (1970). The Ugandan forms were designated *G. m. submorsitans* var *ugandensis*. Crossing experiments have not adequately tested phylogenetic relationships among the four nominal races, a difficult task because representative laboratory cultures do not exist. Some work has been accomplished on population genetics, however.

Microsatellite and mitochondrial variation was used to investigate differentiation and gene flow in *G. m. submorsitans* sampled at the eastern and western extremes of its distribution, approximately 5450 km apart (Krafsur et al., 2000; Krafsur and Endsley, 2002). Mitochondrial diversities in five samples from The Gambia ($h_e = 0.27$) were only a third of those in two Ethiopian samples ($h_e = 0.84$). Here the SSCP method was used to estimate diversity. Genetic differentiation within countries was significant in The Gambia ($F_{ST} = 0.04$) but not in Ethiopia ($F_{ST} = 0.003$) where the two demes were only c. 10 km apart. $F_{ST}$ between countries, however, was 0.34. Nei’s (1987) unbiased genetic identity $I$ varies from zero when no variant is shared, to one, when all are shared in equal measure, was $I = 0.65$ between countries, $I = 0.96$ among Gambian subpopulations, and about $I = 0.985$ between the Ethiopian subpopulations. Microsatellite diversities were $h_e = 0.68$ in The Gambia and $h_e = 0.72$ in Ethiopia. Subpopulation allele frequencies differed significantly within each country. Indices of genetic differentiation for microsatellite loci were $F_{ST} = 0.34$ between The Gambia and Ethiopia and $F_{ST} = 0.35$ among all subpopulations. The low mitochondrial diversities among *G. m. submorsitans* in The Gambia are indicative of recent contractions in population numbers. Human population increase, loss of fly habitat, and progressive decrease in annual rainfall are likely to have caused the decline in tsetse populations (Krafsur et al., 2000). Genetic diversities among *G. m. submorsitans* in Ethiopia, on the other hand, seemed typical of a large, longstanding population. Hutchinson (1971) and colleagues found it to be abundant and widespread. Ugandan and Ethiopian *G. m. submorsitans* are allopatric (see Rogers and Robinson, 2004), so the morphological pattern of Ethiopian *G. m. submorsitans* could be quite
different from the Ugandan morphotype although it has been assumed by some that the
Ethiopian race is variety *ugandensis*. Morphological differentiation has genetic and strong
environmental components. Covariances and genotypic by environment interactions further
complicates unambiguous genetic inferences of morphological variation. Interpretation of
quantal morphological traits therefore requires a formal experimental design and thorough
sampling before tentative genetic conclusions may be drawn.

*G. swynnertoni* occupies only a small region relative to the other *Morsitans* group tsetse taxa.
Two principle belts in north central Tanzania are figured in Rogers and Robinson’s (2004)
map. Its preferred habitat is open woodland savannah; its range has contracted because of
earlier control programmes and loss of habitat (Marquez et al., 2006). It is now confined largely
to national parks and reserves where it remains a serious trypanosomiasis vector. An early
report suggested that *G. swynnertoni* might have experienced a genetic bottleneck (Gooding
et al., 1993). The magnitude of allozyme variation in *G. swynnertoni*, however, was similar to
that in other *Morsitans* group taxa and provided no evidence of a stringent, earlier bottleneck
(Table 1). As we shall see, however, allozyme variation is subject to balancing selection that
can maintain heterozygosities at high levels even in bottlenecked populations. Mitochondrial
variation is more sensitive to demographic phenomena and has been systematically assessed
in *G. swynnertoni* (Marquez et al., 2006). Only 18 haplotypes were detected by sequencing
two loci (*r16S2* and *Cox I*) totaling 668 bp among 8 populations totaling 149 flies. Only two
haplotypes accounted for 80% of the total number of flies sequenced. Ten of the 18 haplotypes
were singletons (56%), indicative of mutation-drift disequilibrium and consistent with the
notion that the *G. swynnertoni* populations examined had experienced an earlier bottleneck
and subsequent expansion. The estimate of genetic differentiation among the locations was
$G_{ST} = 0.04$, the lowest estimate of any *Morsitans* group species. Temporal differentiation,
however, was four times the magnitude of spatial variation. Drift is inversely proportional to
population numbers, so it seems that the sampled populations were quite small. Better
resolution of population structure may be afforded by survey of diploid loci. Nine putative
c microsatellite loci were identified (Baker and Krafsur, 2003; Ouma et al., 2006), but no survey
of diversity has yet been carried out.

The temporal flux in *G. swynnertoni* haplotype frequencies allowed estimates of effective
population size, $N_e$ (Table 4). The most reliable estimate was derived from villages in Tarangiri,
where successive samplings were made at an interval of approximately 54 generations. Here,
the estimate of $N_e = 66$ reproducing females per generation. The other estimates varied from
two to 41 reproducing females. These are not large numbers so it is well to bear in mind that
$N_e$ estimates represent harmonic means and so are weighted by the smallest numbers in the
sampling intervals.

*G. pallidipes* geographic distribution is extremely patchy according to Rogers and Robinson’s
(2004) map (Fig. 2). Sizable but discontinuous belts are figured in Ethiopia, Somalia, Kenya,
Uganda, Tanzania, Zaire, Mozambique, Zambia, and Zimbabwe. Probabilities of occurrence
are small in vast areas of Tanzania and Zambia even though many of these same regions can
support *G. morsitans*. Genetic sampling has been carried out in many of the principal belts.
Allozyme variation at eight loci was used to estimate genetic differentiation and gene flow
among 11 *G. pallidipes* populations from Kenya, Zambia, Zimbabwe, and Mozambique (Table
1). These data pointed to a high degree of population structure in which the average level of
gene flow $Nm$ was much less than anticipated by a reading of the ecological literature ($F_{ST} =
0.24, Nm \sim 0.8$). Mating was random within the sampled populations thereby falsifying an
hypothesis of two or more sibling species within populations. The indices of mean allozyme
diversity, $H_E$, and of genetic differentiation, $F_{ST}$, varied between the East African and southern
African populations. $H_E$ was greater in the southern populations but $F_{ST}$ was less. Further
investigation compared allozyme, microsatellite, and mitochondrial variation in the same 11

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populations (Krafsur, 2002a). Microsatellite diversity and mitochondrial diversity were less in southern Africa than in East Africa and both were strongly correlated with each other, but neither was correlated with allozyme diversity. In contrast to the allozyme diversities, mitochondrial and microsatellite variation was consistent with a severe and prolonged reduction in population sizes in southern Africa. Allozyme variation was conserved, however, and differentiation at allozyme loci among populations was much less than at microsatellite and mitochondrial loci. This is illustrated in Fig. 3, where pairwise estimates of mitochondrial, microsatellite, and allozyme genetic distances are regressed separately on geographic distances. Conservation of allozyme variation in southern African G. pallidipes that had experienced an earlier, severe bottleneck (Ford, 1971) is an example of balancing selection. Allozymes, it seems, are not selectively neutral in tsetse.

Because of its wide, but discontinuous distribution, high degree of subdivision, and economic importance, it was decided to investigate further the population structure of G. pallidipes. Microsatellite and mitochondrial variation was examined in detail at macrgeographic (Krafsur and Wohlford, 1999; Ouma et al., 2005) and microgeographic scales (Ouma et al., 2006). Single strand conformational polymorphisms (SSCPs) were used to characterize mitochondrial variation (later, direct sequencing was adopted to obtain better resolution). The picture afforded by these further studies was consistent with earlier inferences based upon allozyme variation. At the macrogeographic scale, 21 samples representing populations from Ethiopia south to Zimbabwe were placed into five geographically and ecologically homogeneous groups (Table 5). Microsatellite diversities, in terms of mean alleles per locus and heterozygosity, varied significantly among the grouped populations. The eastern populations in Kenya and Tanzania had twice the allelic diversity as the other groups and greater mean heterozygosities. The bottlenecked southern African populations showed similar microsatellite diversity indices as three other groups.

Mitochondrial variation is more sensitive to demographic flux than diploid loci because of its uniparental inheritance and single copy number. Thus, of 39 SSCP haplotypes recorded, only four (10%) were found in southern Africa, compared with 23 (59%) in coastal Kenya and Tanzania. Mean diversity was 0.42 and \( F_{ST} = 0.51 \) over the 21 populations (Table 6). Here, diversity is the probability that two randomly chosen flies have different haplotypes and the estimate of mean gene flow among populations is approximately one reproductive female every two generations.

Sequencing r16S2 and Cox I (a sum of 670 bp) yielded 181 haplotypes among the 873 flies sequenced, only 12 (7%) of which occurred in southern Africa and 144 (80%) in Kenya and Tanzania (Marquez and Krafsur, unpublished data). Mean haplotype diversity was much greater at \( h_s = 0.73 \) than estimated by SSCP but \( F_{ST} \) was about the same at 0.47. There was a high frequency of singleton haplotypes − 64%. Singletons can be indicative of rapidly expanding populations (Slatkin and Hudson, 1991; Rogers and Harpending, 1992) or more complex demographic phenomena including selection against mitochondrial mutants. Detection of demographic changes by genetic means, however, is uncertain when expanding populations are subject to pronounced environmental and temporal heterogeneities (Wegmann et al., 2006). Nevertheless, the high frequencies of singular mitochondrial haplotypes is consistent with rapid population expansions and well supported by the historical record relating the African rinderpest epizootic to the disappearance of tsetse fly populations in southern Africa. This is illustrated in regional lineage through time plots (Fig. 4) that show abrupt increases in lineage numbers about 300 to 400 generations ago, corresponding to 41 – 55 years before sampling, in the years 1940 – 1954. The oldest lineages varied from c. 671 generations in Tanzania and southern African G. pallidipes (~ 92 y), 1864 generations in Kenya (~ 255 y), to c. 9200 generations in Ethiopia (~1260 y).
At the microgeographic level, *G. pallidipes* subpopulations were sampled at scales of 200 m to 14 km (Ouma et al., 2006). The chief locations were Lambwe in the Ruma National Park in western Kenya, and Nguruman, in southwestern Kenya, adjacent to Tanzania. Lambwe and Nguruman are 240 km apart and much of the area between is at least 2000 m elevation and, therefore, inhospitable to tsetse fies. The tsetse populations in Lambwe and Nguruman have been intensively studied (e.g., Turner and Brightwell, 1986; Williams et al., 1992; Brightwell et al., 1992, 1997). Trap catches in Nguruman showed a high degree of spatial aggregation (Odulaja et al., 2001) raising the question, does this aggregation represent a component of genetic differentiation among local demes? *G. pallidipes* populations closest to Nguruman and Lambwe also were sampled. $F_{ST}$ over samples in Lambwe was 0.05; in Nguruman, $F_{ST} = 0.04$, both significantly different from zero ($P < 0.0001$). Differentiation of allele frequencies among the three blocks in Lambwe was highly significant ($\chi^2(14) = 1067, P < 0.0001$). A similar result obtained among the seven blocks in Nguruman ($\chi^2(42) = 1160, P < 0.0001$). Even at distances of a few hundred meters, it seems that tsetse flies form demes among which genetic drift is stronger than gene flow. Genetic differentiation of Lambwe *G. pallidipes* from its nearest conspecific neighbour at Kodera was $F_{ST} = 0.15$. Similarly, the contrast between Nguruman and its neighbour Serengeti yields $F_{ST} = 0.16$. The foregoing estimates are nearly 40% of the maximum values they can take because levels of homozygosity at microsatellite loci were ~ 0.39. Sequencing *Cox I* and *r16S2* loci supported the view of little shared variation (Marquez and Krafsur, unpublished data). Only one of 187 Nguruman *G. pallidipes* shared its mitochondrial haplotype with one of 31 Serengeti flies.

Estimates of effective population numbers in Nguruman were approximately double those in Lambwe: the modal values were approximately 700 and 350 respectively (Table 4). Taken together with indices of genetic differentiation, the $N_e$ estimates suggest strongly that longstanding, substantial populations existed even in the face of sustained control efforts.

**Subgenus Machadomyia**

As discussed earlier, *G. austeni* cytogenetics have been examined (Southern et al., 1972; Gariou-Papalexiou et al., 2007) and chromosome interchanges induced and manipulated experimentally (Curtis et al., 1972). Two subspecies are allopatric and have been described based on differences in body size, color, habitats, and responses to traps (Dias, 1987). The chief systematic question concerning *G. austeni* is its phylogenetic relationships to the other *Glossina* (Gooding and Krafsur, 2004, 2005 for reviews). The distribution of *G. austeni* is largely coastal and highly discontinuous from Natal Province in South Africa north to the River Jubba in Somalia. Representatively sampling this species is difficult because it does not respond well to traps (Vreysen et al., 1998). Its population genetics are uninvestigated but RAPDs were used as markers to detect genetic differences between cultured flies and wild flies on the continent and Zanzibar; unquantified differences in polytene chromosome banding patterns were found among flies from the lab culture and field samples from Zanzibar, Tanga (Tanzania), and Zululand (Gariou-Papalexiou et al., 2000). Sequencing of the *r16S2* and *Cox I* mitochondrial genes in 32 specimens from Pretoria, South Africa and 15 specimens from a lab culture in Tanga, Tanzania yielded only eight haplotypes (Table 7). The Pretoria culture was monomorphic and its haplotype was unshared with the Tanga culture.

**Population structure in subgenus Nemorhina – the Palpalis group (Fig. 5)**

A more complex picture obtains for palpalis group than morsitans group tsetses. *G. p. gambiensis* is riparian and also occurs in humid savannah. Its range includes Senegal, Guinea, Sierra Leone, southwestern Mali, and Burkina Faso. It disperses along watercourses (Cuisance et al., 1985). *G. p. palpalis* inhabits forest and occurs in Cote d’Ivoire, Benin, Nigeria, Cameroon, Equatorial Guinea and south to Angola (Robinson and Rogers, 2004). Empirical evidence suggested that dispersion and emigration is density-dependent in *G. p. palpalis* in
Cote d’Ivoire (Rogers and Randolph, 1985). Such dispersion would promote gene flow and diminish the effects of genetic drift. No mitochondrial variation was detected by the SSCP technique in G. p. gambiensis from Senegal but nine haplotypes were detected among flies sampled from three river drainages in Mali (Marquez et al., 2004). The Senegal haplotype was detected in only two of 303 flies sampled in Mali. It had been claimed earlier that the Senegal population had been eradicated, but the genetic evidence supports the notion of an expanded, relict population at the time of genetic sampling. The paucity of detectable mitochondrial variation among Senegal tsetse may be indicative of earlier population bottlenecks or of inadequate genetic resolution afforded by SSCP. Sequencing r16S2 and Cox1 mitochondrial genes in 32 Senegalese flies, however, demonstrated 17 haplotypes, and 10 haplotypes among 28 flies from Bado, Mali (Marquez and Krafsur, unpublished data).

Examination of SSCP’s indicate that only 0.06% of the genetic variance among Mali G. p. gambiensis was attributed to differences among three river drainages, but GST = 0.17, indicative of significant genetic differentiation. Microsatellite variation at two X-linked loci in G. p. gambiensis showed strong differentiation (FST = 0.18) between a Burkina Faso population and a Senegal population, some 2,000 km distant (Solano et al., 1999). Especially notable was significant differentiation between two sampled populations on the River Koba in Burkina Faso only 15 km apart (FST = 0.12) and an unusually large deficit of heterozygotes (FIS = 0.33) in just one of the two sampled areas. A later report utilizing three microsatellite loci (Solano et al., 2000) confirmed genetic differentiation across sampled demes on the R. Koba (FST = 0.07) and the deficiency of heterozygotes (FIS = 0.20) in the same location as before. Further sampling of G. p. gambiensis in Burkina Faso along 216 km of the River Mouhoun provided a much lesser estimate of FST = 0.012. Of the four sampled demes, two showed significant departures from random matings (FIS = 0.15 and 0.16; Bouyer et al., 2007). Here the deficiencies of heterozygotes were attributed to null alleles. Consistently large heterozygote deficiencies were recorded at each of five loci in G. p. palpalis in Bonon, Cote d’Ivoire (Ravel et al., 2007). Estimates of FIS varied from 0.12 to 0.44 and a mean over loci and samples of 0.34. Ravel et al. (2007) ruled out linkage disequilibrium and null alleles as the primary causes of the extraordinary departures from random mating within all sampled populations (i.e., FIS). Neither gene frequencies nor estimates of genetic differentiation were provided but application of a Bayesian clustering method showed graphically that samples could be reduced to four highly branched clusters. Ravel et al. (2007) concluded that the Wahlund effect explained the extraordinary FIS. Further analysis using Bayesian methods seemed to confirm that individual samples of G. palpalis s.l. included representatives of two or more demes - up to nine, in fact (de Meeus et al., 2007). Thus, it was suggested that even individual fly traps had sampled multiple demes of only one species. As suggested by the authors, heterozygote deficits recorded in G. p. gambiensis and G. p. palpalis could have been examples of the Wahlund effect (the mixing of conspecific demes with differing allele frequencies).

I find the foregoing explanation unlikely, however, and suggest the data indicate sympatric sibling species: Annual cycling of seasonal range expansions and contractions could, in principle, result in the establishment of demes differentiated by founder effects and genetic drift during the favourable wet season, followed by concentration in moist refugia during the dry season. Initial dry season populations could then show the Wahlund effect but subsequent matings, if random, would lead to a population at Hardy-Weinberg equilibrium and FIS would approach zero. Tsetse generation times are of the order of 50 days, so the timing of sampling is important. A full understanding the relative magnitudes of FST and FIS under the foregoing scenario, however, requires sophisticated simulation studies and estimates of deme sizes. Why do only Palpalis group flies in West Africa show extraordinary FIS estimates? Wahlund effects have not been detected in G. f. fuscipes (see below) and in five taxa of morsitans group tsetse.
An alternative hypothesis to mixing of genetically differentiated, conspecific demes in *G. palpalis* s.l. at some sites but not others is the presence of morphologically identical sibling species, an explanation supported by reciprocal crosses between strains of *G. p. palpalis* (Gooding et al., 2004). Significant microgeographic genetic differentiation in *G. p. gambiensis* (i.e., $F_{ST} \gg 0$) was manifest particularly on the R. Koba, but not the R. Mouhoun (Solano et al., 2000). The uneven spatial distribution in samples of two (or more) unrecognized sibling species would have the effect of inflating both $F_{IS}$ and $F_{ST}$. Colonization of geographic strains of the foregoing taxa followed by cross-matings and backcrosses of surviving progenies need to be carried out to help clarify the picture. Cytological examination of polytene chromosomes would also be helpful in resolving what seems likely, in my opinion, to be two instances of sibling species.

**Population structure in *G. fuscipes fuscipes*—**Krafsur et al. (2008) examined the spatial patterns of mitochondrial cytochrome oxidase I ($Cox$ I) and $r16S2$ variants in two Kenya, one Uganda, and a laboratory population derived from the Central African Republic sometime before 1986. Like many other *Nemorhina*, *G. f. fuscipes* is riverine and lacustrine. The East African sampling sites were within 55 km of each other but they shared only four of 21 variants and were genetically differentiated; pairwise $F_{ST}$s varied from 0 to 0.28. None of its 21 mitochondrial variants was shared between the lab culture and East Africa samples; pairwise $F_{ST}$ varied from 0.46 to 0.58. In a geographically more extensive study, Abila et al. (2008) examined microsatellite and mitochondrial variation at cytochrome oxidase II and cytochrome B in nine Uganda populations and in a sample from Sudan. Pairwise $F_{ST}$ based on five microsatellite loci varied from 0.02 to 0.39 (mean 0.22) among eight sampled populations. $F_{ST}$ based on mitochondrial variation in ten samples varied from 0.01 to 0.91 (mean 0.59). No correlation was observed between tsetse allele frequencies and the distribution of *T. b. rhodesiense* and *T. b. gambiense*. Thus, no evidence was obtained to support an hypothesis of local co-adaptation between vector and parasite strains that would reduce gene flow among tsetse nor help to explain the disjunct distribution of *T. brucei* subspecies.

**Population genetics in subgenus Austenina - the Fusca group (Fig. 6)**

It seems that little fieldwork has been reported for any member of the group. Allozyme variation was evaluated in lab stocks of *G. brevipalpis* Newstead 1910 and *G. longipennis* Corti 1895 (reviewed in Gooding, 1996). *G. brevipalpis* is a trypanosome vector and widely but discontinuously distributed from eastern South Africa north to the river deltas in Somalia (Rogers and Robinson, 2004). The largest patches occur in Mozambique, Zaire, northern Tanzania and southeast Kenya. Its population structure would seem to be of interest in anticipation of area-wide control measures. Mitochondrial nucleotide sequences indicated 14 haplotypes and mean diversity $H_s = 0.65$ in *G. brevipalpis*; 7 haplotypes and $H_s = 0.55$ were recorded in *G. longipennis* (Table 7).

**General comments**

Estimates of genetic differentiation ($F_{ST}$, $G_{ST}$) were less when estimated at nuclear loci than mitochondrial loci. The mitochondrial genome is single copy and inherited maternally. Moreover, there is no recombination and mutation rates are generally greater than in the nuclear genome. Mitochondrial variation is more sensitive to demographic phenomena. Genetic data based on microsatellite loci tend to underestimate genetic differentiation because $F_{ST}$, $G_{ST}$, and their analogues cannot take values greater than the level of homozygosity (Hedrick, 1999). Microsatellites are also subject to homoplasy (Estoup et al., 1995; Balloux et al., 2000), where two or more alleles become identical in state by convergence from different ancestors. Such alleles cannot be distinguished on gels because they have the same molecular weights thus leading to reductions in estimated diversity. In addition, high frequencies of private and singular alleles bias downward indices of genetic differentiation. A high degree of
genetic differentiation was detected at allozyme loci (Krafsur et al., 1997). Balancing selection may occur at some allozyme loci and it tends to minimize estimates of $F_{ST}$ (Schierup et al., 2000). In sum, microsatellites and allozymes underestimate genetic differentiation thereby overestimating magnitudes of gene flow (Nagylaki, 1998; Balloux et al., 2000).

Tests for isolation by distance, in which pairwise estimates of $F_{ST}/(1 - F_{ST})$ are regressed on log pairwise geographic distances, have generally shown shallow slopes indicating weak relationships between genetic and geographic distances. The results are in keeping with the principal assumptions of the island model of population structure, in which selectively neutral genetic variation is distributed among a very large number of subpopulations of equal size that demonstrate equal rates of gene flow among them. Of course, few, if any, natural populations fit such simplifying assumptions, but the picture in the savannah species is largely consistent with a large number of inhabited patches that exchange reproductive flies at low rates of gene flow.

Because *G. pallidipes* is considered the most dispersive of all tsetse species (e.g., Vale et al., 1984; Williams et al., 1992; Jordan, 1993; Hargrove, 2003), the high degree of genetic differentiation among its populations is remarkable and calls for further investigation. A full appraisal of its population genetics must include sampling and genetic analysis of heretofore unsampled populations in the DRC (Zaire), Mozambique, southern Somalia, Uganda, northwestern Zambia, and Tanzania. Because many fly belts and geographically small patches throughout its range remain unsampled, it is a valid question to ask if *G. pallidipes* constitutes a species complex. Perhaps some patches support different mating types of a species complex. It should be noted, however, that no sampled population showed a significant deficiency of heterozygotes, thereby suggesting only one taxonomic unit in each. The suggestion that the West African *G. longipalpis* may be conspecific with *G. pallidipes* should be followed up by comparing their polytene chromosome karyotypes, mitochondrial loci, and patterns of microsatellite variation. A compelling test would be to attempt reciprocal crosses and testing the fertilities of any progeny. Earlier attempts to obtain cage matings of *G. longipalpis*, however, were unsuccessful (Roubaud, 1935).

**Effective population sizes (numbers) in Glossina**

Effective population size $N_e$ is the harmonic mean number of reproducing individuals in an ideal population that have the same allele frequencies as an actual population under study. $N_e$ determines the rate of genetic drift – the less the $N_e$, the greater the magnitude of drift. An ideal population has the same rate of inbreeding (i.e., loss of heterozygosity) as the study population and is one in which all Hardy-Weinberg assumptions apply. Clearly, population effective number is bound to be less than the corresponding census number because reproductive success varies enormously among individuals and because of deviations from Hardy-Weinberg assumptions, particularly that of infinite population size.

Estimation of census size is problematic. Accurately estimating $N_e$ is also problematic but for different reasons (Wang, 2005). Moreover, there are a number of estimation methods each of which usually provides numerically quite different values. Short term estimates are based on temporal variance in allele frequencies. The principal assumptions include selective neutrality and a negligible mutation rate in a population closed to immigration. Long-term estimates are based on genetic diversity estimates with the chief assumption of mutation-drift equilibrium and a good estimation of the prevailing mutation rate.

Estimates of mean effective population sizes in *G. pallidipes* in Lambwe varied from c. 200 to 520, depending principally on the interval between successive samplings (Table 4). At the geographically larger Nguruman site, estimates varied from c. 250 to 1000 (Ouma et al., 2006). Estimates of $N_e$ varied from 18 to 66 in *G. swynnertoni* (Marquez et al., 2006). A much
different approach was used to estimate $N_e$ in *G. f. fuscipes* (Krafsur et al. 2008). *G. f. fuscipes* in Kenya and Uganda were estimated to be 403 and 2258 respectively compared with $N_e = 1,060$ in the International Atomic Energy Agency’s (IAEA) lab culture (Krafsur et al., 2008). It is well to recall that $N_e$ is a harmonic mean value, and is strongly weighted by the least values over numerous generations. The estimates were based on mitochondrial variation, reference to an estimated number of breeders in IAEA’s large lab culture, and estimates of a mutation rate.

The foregoing estimates indicate that tsetse demes are quite small. Especially small effective population numbers seem to characterize *G. swynnertoni* although observed population densities can be quite high locally (Ndegwa and Mihok, 1999). Estimates of $N_e$ for the genetically isolated *G. pallidipes* in the Lambwe valley and Nguruman in Kenya suggest longstanding, substantial populations even though intensive control measures were carried out there.

In contrast to the estimates for tsetse flies, consider the effective population size of Leopards (*Pantheria pardus*) in Tanzania of 38–48 thousand (Spong et al., 2000). Five island populations of *Anopheles gambiae* among which gene flow was severely restricted varied from 7 to 25 thousand when estimated by the stepwise mutation model and 3.3 to 8.5 thousand when estimated on the basis of the infinite allele model (Pinto et al., 2002). $N_e$ was estimated in two widely separated *Anopheles gambiae* populations in Kenya: a population near Lake Victoria was 6.3 or 22.6 thousand, depending on the method of estimation. Another population on the coast, 700 km away, was 4.3 or 7.9 thousand (Lehmann et al., 1998). Thus, the estimation model used can give widely divergent estimates. A further example of $N_e$ is provided by Michel et al. (2006) for *Anopheles funestus*. Based on mitochondrial variation, effective population sizes for this mosquito in Burkina Faso were estimated at 93.7 to 98.8 thousand and 340 to 311 thousand, depending on the precise long-term model employed. Microsatellite-based estimates in Burkina Faso varied from 174 to 263 thousand. What is clear is that African *Anopheles* has large effective population numbers even when isolated on islands and surviving annual dry season declines.

Even though $N_e$ varies greatly with the methods used in its calculation, it is plain that tsetse estimates are very much less than other African Diptera and even leopards. Two reasons can be offered. First, tsetse gene frequencies varied greatly from one generation to another. Because genetic variance is inversely proportional to population size $N$ (binomial variance = $p(1−p)/2N$), a large estimate of variance is indicative of small $N$. Second, high $F_{ST}$ estimates indicate that gene flow among demes is restricted. $N_e$ is directly proportional to the variance in density and spatial dispersion of a population of interest. As we have seen, in tsetse flies spatial dispersion distances and population densities are small. Forces of genetic drift in *Morsitans* group (and probably *Palpalis* group) tsetse clearly overwhelm the forces of dispersion.

**Range expansions after the rinderpest epizootic**

Range expansions of tsetse flies during the colonial era are discussed in Buxton (1955), Ford (1971), Jordan (1986), and Rogers (1995), based on published and colonial administration records of small refugia after the rinderpest epizootics. Southern African populations of *Morsitans* group flies showed lesser genetic diversities than conspecific populations from East Africa, consistent with much earlier bottlenecks in population numbers. Do the geographically small but widely distributed populations of *G. pallidipes* correspond to relict survivors from more extensive, pre-rinderpest populations, founders from earlier surviving populations, or merely the inhabitants of environmentally suitable islands in the midst of unwelcome habitats? No genetic evidence was obtained to support earlier bottlenecks or founder effects in East Africa and Ethiopia because the sampled populations showed abundant variation. Hypotheses
of local physiological adaptation via natural selection to prevailing climate offer a mechanism that could explain low rates of gene flow. The question of adaptation was investigated in *G. pallidipes*.

**Physiological adaptation to locally prevailing climates**

In principle, adaptation to locally prevailing climates could help to explain the lack of gene flow indicated by spatial patterns of genetic variation, i.e., there may be spatial heterogeneity in natural selection (Lenormand, 2002). Thus, reproductive success of immigrant flies may be poor even though dispersion rates are high. Tests of relevant hypotheses were performed in *G. pallidipes* in five ecologically diverse sites in Kenya and Zambia where the prevailing climates substantially varied. Hypotheses tested included homogeneity among populations’ thermal tolerances, desiccation tolerances, body water amounts, lipid amounts, and activity levels. Critical thermal limits among the sampled populations were narrow for high temperatures but much less so for low temperatures. Seasonal variation was detected in *G. pallidipes* in Lambwe and Nguruman and this was attributed to acclimation. Indeed, most among-population variation was explained by phenotypic acclimation rather than genetic differences. Little potential for evolutionary change was detected. Thus, phenotypic plasticity, not genetically determined, heritable, variation explained differences in physiological responses (Terblanche and Chown, 2006; Terblanche et al., 2006).

Cuticular hydrocarbons and lipids are important in water conservation and sex recognition; they were measured quantitatively and qualitatively in the four disparate *G. pallidipes* populations (Jurenka et al., 2007). Cuticular lipid quantities in flies varied among the sampled populations but did not correlate with environmental measures of normalized density vegetation index or prevailing temperatures and humidities. Desiccation resistance is thought to depend on cuticular lipids. Desiccation rates varied significantly among populations but were uncorrelated with cuticular lipid amounts. Another mechanism that might rationalize high dispersion rates with low rates of gene flow is mate recognition, which is conferred by contact pheromones (Carlson et al., 1984). The chief sex pheromone in *G. pallidipes* is 12, 23-dimethylpentatricontane and this did not differ among flies from Ethiopia, Zimbabwe, and five independently derived lab cultures (Carlson et al., 2000) nor in four diverse Kenya populations (Jurenka et al., 2007).

To recapitulate, no evidence was obtained that would support the notion that the genetic differentiation of tsetse fly populations is maintained by sex recognition mechanisms or genetically based physiological adaptation to locally prevailing temperatures, saturation deficits, and other measured environmental variables. Instead, phenotypic plasticity and acclimation accounts for tsetse fly responses to short-term environmental variations. Bioclimatic models, however, imply that very long-term responses to environmental heterogeneity include speciation such as that seen among the *Morsitans* group tsetse.

**Summing up**

The epidemiological significance of tsetse fly population structures vis a vis population structures of trypanosomes is virtually unknown and requires both laboratory and field research. These endeavours will not be easy, however.

Clearly, demes in two (of nine) *Palpalis* group taxa, like those in the *Morsitans* group, are localized, with little detectable gene flow among demes. The extent of genetic sampling in *Palpalis* group species, however, is limited taxonomically and geographically. Clearly, gene flow in *G. p. gambiensis* should be investigated further by using markers that afford higher resolution than our use of SSCP's. A number of *G. palpalis* and *G. fusca* subspecies, *G.
tachinoides and G. pallicera s.l., remain to be studied. Our knowledge of Fusca group genetics is virtually nil.

Sampling of Morsitans group flies, while genetically and geographically much greater than Palpalis and Fusca taxa, has many deficiencies. In particular, G. longipalpis and G. austeni require investigation. For Morsitans group taxa, cytotgenetical work at a population level should prove highly interesting given the polymorphisms already detected in G. morsitans s.l. and G. pallidipes.

In terms of population biology, G. pallidipes is probably the most thoroughly investigated tsetse fly. The most recent distribution map of G. pallidipes shows highly fragmented populations (Rogers and Robinson, 2004) and compelling genetic evidence indicates little gene flow among them. Yet gene flow among G. pallidipes demes within environmentally suitable patches also seems restricted, as genetic sampling in Ethiopia, Zambia, Tanzania, and Zimbabwe has shown. Distribution maps of the other genetically sampled morsitans group and palpalis group show much less fragmentation than that for G. pallidipes. Indeed, G. morsitans s.l., G. palpalis s.l. and G. fuscipes s.l. distributions are more or less continuous throughout their ranges (Rogers and Robinson, 2004) yet the patterns of genetic differentiation in these taxa do not seem greatly different than G. pallidipes. Tsetse fly demes seem structured much like island populations. To be sure, there must be gene flow, at least historically, but its magnitude seems small. Even populations of species that recovered from earlier, severe bottlenecks are highly structured – for example, G. morsitans s.l. and G. pallidipes in southern Africa.

What may account for this structure? And how well does the hypothesis of little gene flow agree with ecological knowledge of this important group of vector insects? As we have seen earlier in this essay, tsetse dispersal has been measured and thoroughly analyzed (e.g., Rogers, 1977; Williams et al., 1992; Hargrove, 1981, 2000). Based on earlier direct estimates of movement, tsetse dispersal has been simulated (Hargrove, 2000; Vale and Torr, 2004). All the foregoing evidence strongly suggests there should be abundant gene flow among tsetse demes. Are conclusions drawn from the ecological research false? Or do the simplifying assumptions of population genetic theory render false conclusions when applied to tsetse populations?

Of course not. Ecologists and geneticists have measured different things at different scales (Krafsur, 2002b). Ecological estimates are based on scales of one to tens of kilometers measured over short time intervals. The genetic scales are tens to thousands of kilometers and represent demographic phenomena integrated over intervals of many thousands of generations. Let’s go back to the underlying model against which geneticists test hypotheses about gene flow. The model is an ideal population established by Hardy-Weinberg assumptions, viz., populations of infinite size, discrete generations, of randomly mating individuals, and uninfluenced by natural selection. Few taxa, if any, conform to such a model. Genetic inferences about tsetse population structure include small effective population sizes ($N_e$) of continuously overlapping generations and geographically fragmented populations. Genetic drift is inversely proportional to $N_e$ so is an especially potent force in tsetse population structure, accounting for most, if not all, of the measured differentiation. Also contributing to the genetic differentiation must be numerous and widespread inhospitable patches, in which the precise biotic and abiotic criteria vary greatly (review in Rogers and Randolph, 1985). There is a strong historical component to the picture, that of the rinderpest epizootic beginning in 1887 that killed out 90% of more of the mammalian fauna upon which tsetse feed. Although the epizootic occurred throughout much of sub-Saharan Africa, its effects on tsetse flies seemed to have been particularly severe in southern Africa, as documented by stockmen, entomologists, and veterinarians during the colonial era (Buxton, 1955; Ford, 1971; Jordan, 1986). Morsitans group flies were diminished to only a few, relict refugia. It is important to note that resurgence

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of *G. morsitans* s.l. and *G. pallidipes* there required many years to return to economically significant levels. A more recent example is afforded by results of the Okavango project in which *G. m. centralis* has failed to reappear after the six years since control operations ceased. Their slow recovery should be an important consideration in examining cost-benefit analysis of thorough, area-wide tsetse control schemes. As pointed out by Rogers and Randolph (2002), tsetse populations are extremely resilient. Adult female average daily survival probabilities are of the order of 98% or more. The half-life of an adult female cohort born simultaneously on day zero is approximately 35 days. Tsetse populations are able to recover from as few as 16 inseminated females, according to Hargrove’s (2005) estimates. Whether that is actually 10 or 100 inseminated females is moot because such small tsetse populations are not likely to be detected even with intensive sampling. Genetic studies support the existence of highly resilient *G. pallidipes* populations in two ecologically well-studied sites, Lambwe and Nguruman in Kenya. It was shown that populations there shared little genetic variation with the populations nearby. Hypotheses of massive immigration following control measures or seasonal declines were invoked to explain the reappearance of *G. pallidipes* in Lambwe and Nguruman (Turner and Brightwell, 1986; Williams et al., 1992). Based on genetic evidence, however, it seems those hypotheses are likely to be false.

Genetic models of population structure are gross oversimplifications of natural demographic phenomena, for example, $F_{ST}$ and $N_e$ (Whitlock and McCauley, 1999). Simulation modeling has shown that large variances in deme size, caused by environmental or temporal heterogeneity, can produce high values of $F_{ST}$ even when dispersion rates are large (Wegmann et al., 2006). It is clear that estimates of $N_e m$ are no substitute for estimates based on mark, release, and recapture trials. Based on the oversimplified genetic models, all we can really say about tsetse dispersion is that the subpopulations sampled contain numerically very few individuals that originated in other subpopulations and genetic drift is the dominant force in population structure. A scenario of massive exchange of individuals among any of the populations sampled to date, however, seems to be untenable when based on the genetic evidence.

**Acknowledgements**

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Figure 1.
Distribution of *Morsitans* group tsetse flies predicted from satellite imagery (courtesy of ERGO Ltd and TALA, Oxford, with permission. Available from <http://ergodd.zoo.ox.ac.uk/tseweb/distributions.htm>.)
Figure 2.
Predicted (grey) and observed (hatched) distribution of *G. pallidipes*, from Rogers and Robinson (2004) with permission.
Figure 3.
Pair-wise Cavalli-Sforza and Edwards chord genetic distances among *G. pallidipes* populations regressed on corresponding geographic distances. Note the slope for allozyme loci is much shallower than for mitochondrial and microsatellite loci (from Krafsur, 2002).
Figure 4.
Lineage (haplotype) through time plots showing temporal increases in mitochondrial diversities. Note the small number of lineages (12) in Zimbabwe and Zambia and the rapid increases in diversities 300 – 400 generations ago in each region.
Figure 5.
Distribution of *Palpalis* group tsetse flies predicted from satellite imagery (courtesy of ERGO Ltd and TALA, Oxford, with permission. Available from <http://ergodd.zoo.ox.ac.uk/tseweb/distributions.htm>.)
Figure 6.
Distribution of *Fusca* group tsetse flies predicted from satellite imagery (courtesy of ERGO Ltd and TALA, Oxford, with permission. Available from <http://ergodd.zoo.ox.ac.uk/tseweb/distributions.htm>.

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Table 1
Allozyme variation in tsetse flies compared with other Diptera

<table>
<thead>
<tr>
<th>Taxa</th>
<th>No. loci</th>
<th>Percentage polymorphic</th>
<th>Mean alleles per locus</th>
<th>Mean diversity, $H_e$</th>
<th>Diversity at polymorphic loci</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>G. m. morsitans</em></td>
<td>45</td>
<td>20.0</td>
<td>1.4</td>
<td>6.6</td>
<td>29.9</td>
</tr>
<tr>
<td><em>G. m. centralis</em></td>
<td>31</td>
<td>32.3</td>
<td>1.4</td>
<td>6.0</td>
<td>18.7</td>
</tr>
<tr>
<td><em>G. swynnertoni</em></td>
<td>34</td>
<td>17.6</td>
<td>1.2</td>
<td>7.2</td>
<td>40.5</td>
</tr>
<tr>
<td><em>G. pallidipes</em></td>
<td>38</td>
<td>26.3</td>
<td>1.5</td>
<td>6.8</td>
<td>25.4</td>
</tr>
<tr>
<td><em>Musca domestica</em></td>
<td>68</td>
<td>53.4</td>
<td>2.5</td>
<td>18.3</td>
<td>36.7</td>
</tr>
<tr>
<td><em>Musca autumnalis</em></td>
<td>50</td>
<td>62.0</td>
<td>2.4</td>
<td>18.6</td>
<td>30.5</td>
</tr>
<tr>
<td><em>Stomoxys calcitrans</em></td>
<td>38</td>
<td>52.6</td>
<td>1.8</td>
<td>9.6</td>
<td>18.2</td>
</tr>
</tbody>
</table>


Table 2
Microsatellite diversities and tests for random matings $F$ in *Glossina* spp and the house fly

<table>
<thead>
<tr>
<th>Species</th>
<th>No. Populations</th>
<th>No. loci</th>
<th>Alleles per locus</th>
<th>Diversity $H_e$</th>
<th>Within demes $F_{IS}$</th>
<th>Among demes $F_{ST}$</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>G. m. morsitans</em>&lt;sup&gt;a&lt;/sup&gt;</td>
<td>6</td>
<td>6</td>
<td>11.0 ± 5.6</td>
<td>0.73 ± 0.06</td>
<td>0.03 ± 0.03</td>
<td>0.19 ± 0.05</td>
</tr>
<tr>
<td><em>G. m. morsitans</em>&lt;sup&gt;b&lt;/sup&gt;</td>
<td>9</td>
<td>7</td>
<td>28.6 ± 4.5</td>
<td>0.74 ± 0.05</td>
<td>0.17 ± 0.07</td>
<td>0.13 ± 0.01</td>
</tr>
<tr>
<td><em>G. m. centralis</em>&lt;sup&gt;c&lt;/sup&gt;</td>
<td>7</td>
<td>7</td>
<td>8.8 ± 3.7</td>
<td>0.70 ± 0.09</td>
<td>-0.12 ± 0.04</td>
<td>0.19 ± 0.04</td>
</tr>
<tr>
<td><em>G. m. submorsitans</em>&lt;sup&gt;d&lt;/sup&gt;</td>
<td>7</td>
<td>7</td>
<td>12.7 ± 6.2</td>
<td>0.81 ± 0.04</td>
<td>0.03 ± 0.03</td>
<td>0.17 ± 0.07</td>
</tr>
<tr>
<td><em>G. pallidipes</em>&lt;sup&gt;e&lt;/sup&gt;</td>
<td>21</td>
<td>8</td>
<td>26.8 ± 8.7</td>
<td>0.80 ± 0.03</td>
<td>0.07 ± 0.03</td>
<td>0.18 ± 0.02</td>
</tr>
<tr>
<td><em>G. f. fuscipes</em>&lt;sup&gt;f&lt;/sup&gt;</td>
<td>8</td>
<td>5</td>
<td>8.2 ± 3.6</td>
<td>0.43 ± 0.07</td>
<td>0.11 ± 0.05</td>
<td>0.22 ± 0.07</td>
</tr>
<tr>
<td><em>Musca domestica</em>&lt;sup&gt;f&lt;/sup&gt;</td>
<td>14</td>
<td>7</td>
<td>7.9 ± 1.1</td>
<td>0.86 ± 0.02</td>
<td>0.08 ± 0.02</td>
<td>0.13 ± 0.02</td>
</tr>
</tbody>
</table>

<sup>a</sup>Krafsur and Endsley, 2002. Genotypes scored on acrylamide gels

<sup>b</sup>Ouma et al., 2007. Genotypes scored on ABI 3100 using GeneScan 3.1 and Genotyper 2.5

<sup>c</sup>Krafsur et al., 2001. Genotypes scored on acrylamide gels

<sup>d</sup>Ouma et al., 2005. Genotypes scored on ABI 3100 using GeneScan 3.1 and Genotyper 2.5

<sup>e</sup>Abila et al., 2008

<sup>f</sup>Krafsur et al., 2005
## Table 3
Mitochondrial diversities and genetic differentiation in wild *Glossina* species.

<table>
<thead>
<tr>
<th>Species</th>
<th>Method</th>
<th>No. populations</th>
<th>No. flies</th>
<th>No. haplotypes</th>
<th>Haplotype diversity, $H_S$</th>
<th>$F_{ST}/G_{ST}$</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>G. m. morsitansa</em></td>
<td>SSCP</td>
<td>5</td>
<td>111</td>
<td>25</td>
<td>0.81 ± 0.04</td>
<td>0.09 ± 0.02</td>
</tr>
<tr>
<td><em>G. m. morsitans b</em></td>
<td>ABI 3730</td>
<td>7</td>
<td>96</td>
<td>33</td>
<td>0.81</td>
<td>0.40 ± 0.08</td>
</tr>
<tr>
<td><em>G. m. centralis c</em></td>
<td>SSCP</td>
<td>6</td>
<td>265</td>
<td>7</td>
<td>0.54 ± 0.16</td>
<td>0.81 ± 0.07</td>
</tr>
<tr>
<td><em>G. m. submorsitans d</em></td>
<td>SSCP</td>
<td>7</td>
<td>282</td>
<td>26</td>
<td>0.51 ± 0.12</td>
<td>0.35</td>
</tr>
<tr>
<td><em>G. pallidipes e</em></td>
<td>SSCP</td>
<td>21</td>
<td>624</td>
<td>39</td>
<td>0.42 ± 0.02</td>
<td>0.52 ± 0.001</td>
</tr>
<tr>
<td><em>G. pallidipes f</em></td>
<td>ABI 3730</td>
<td>23</td>
<td>873</td>
<td>181</td>
<td>0.73 ± 0.09</td>
<td>0.47 ± 0.07</td>
</tr>
<tr>
<td><em>G. p. gambiensis g</em></td>
<td>SSCP</td>
<td>13</td>
<td>372</td>
<td>9</td>
<td>0.18</td>
<td>0.68</td>
</tr>
<tr>
<td><em>G. s. swynnertoni h</em></td>
<td>ABI 3730</td>
<td>8</td>
<td>149</td>
<td>18</td>
<td>0.59 ± 0.10</td>
<td>0.04</td>
</tr>
<tr>
<td><em>G. f. fuscipes i</em></td>
<td>ABI 3730</td>
<td>3</td>
<td>79</td>
<td>21</td>
<td>0.84 ± 0.05</td>
<td>0.28</td>
</tr>
<tr>
<td><em>G. f. fuscipes j</em></td>
<td>ABI 3730</td>
<td>9</td>
<td>202</td>
<td>37</td>
<td>0.93</td>
<td>0.60 ± 0.07</td>
</tr>
</tbody>
</table>

*a* Krafsur and Endsley, 2002  
*b* Ouma et al., 2007  
*c* Krafsur et al., 2001  
*d* Krafsur et al., 2000  
*e* Ouma et al., 2005  
*f* Marquez and Krafsur, unpublished data  
*g* Marquez et al., 2004  
*h* Marquez et al., 2006  
*i* Krafsur et al., 2008  
*j* Abila et al., 2008
### Table 4

Effective population numbers ($N_e$) in *Glossina* spp

<table>
<thead>
<tr>
<th>Location</th>
<th>Years</th>
<th>Interval in generations</th>
<th>$N_e$</th>
<th>95% C. L.</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>G. swynnertoni</em>&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Tarangiri</td>
<td>1996 - 99</td>
<td>22</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>Tarangiri</td>
<td>1999 - 03</td>
<td>35</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>Tarangiri</td>
<td>1996 - 04</td>
<td>54</td>
<td>66</td>
</tr>
<tr>
<td><em>G. pallidipes</em>&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Nguruman</td>
<td>2000 - 01</td>
<td>12</td>
<td>333</td>
</tr>
<tr>
<td></td>
<td>Nguruman</td>
<td>2000 - 03</td>
<td>26</td>
<td>679</td>
</tr>
<tr>
<td></td>
<td>Nguruman</td>
<td>2001 - 03</td>
<td>17</td>
<td>250</td>
</tr>
<tr>
<td></td>
<td>Nguruman</td>
<td>2000 - 03</td>
<td>29</td>
<td>1026</td>
</tr>
<tr>
<td></td>
<td>Lambwe</td>
<td>2000 - 02</td>
<td>20</td>
<td>202</td>
</tr>
<tr>
<td></td>
<td>Lambwe</td>
<td>2000 - 03</td>
<td>28</td>
<td>522</td>
</tr>
<tr>
<td><em>G. f. fuscipes</em>&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Kenya</td>
<td>-</td>
<td>-</td>
<td>403</td>
</tr>
<tr>
<td></td>
<td>Uganda</td>
<td>-</td>
<td>-</td>
<td>2258</td>
</tr>
</tbody>
</table>

<sup>a</sup>Marquez et al., 2006. $N_e$ estimated from temporal variances in gene frequencies

<sup>b</sup>Ouma et al., 2006. $N_e$ estimated from temporal variances in gene frequencies

<sup>c</sup>Krafsur et al., 2008. $N_e$ estimated from pairwise differences among mitochondrial haplotypes
Table 5
Microsatellite variation in regionally grouped *G. pallidipes* populations. Means are over 8 loci

<table>
<thead>
<tr>
<th>Grouping</th>
<th>No. populations</th>
<th>No. flies</th>
<th>Avg. no. alleles</th>
<th>Diversity</th>
<th>F$_{ST}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethiopia</td>
<td>2</td>
<td>48</td>
<td>10.0 ± 4.8</td>
<td>0.69 ± 0.05</td>
<td>0.072</td>
</tr>
<tr>
<td>Western Kenya</td>
<td>4</td>
<td>144</td>
<td>11.5 ± 5.4</td>
<td>0.64 ± 0.09</td>
<td>0.203</td>
</tr>
<tr>
<td>Southwest Kenya</td>
<td>7</td>
<td>176</td>
<td>9.9 ± 4.4</td>
<td>0.63 ± 0.01</td>
<td>0.081</td>
</tr>
<tr>
<td>East Kenya, Tanzania</td>
<td>5</td>
<td>240</td>
<td>22.4 ± 8.2</td>
<td>0.82 ± 0.04</td>
<td>0.110</td>
</tr>
<tr>
<td>Southern Africa</td>
<td>3</td>
<td>96</td>
<td>12.8 ± 5.4</td>
<td>0.73 ± 0.06</td>
<td>0.096</td>
</tr>
<tr>
<td>Totals and means</td>
<td>21</td>
<td>720</td>
<td>26.8 ± 8.7</td>
<td>0.80 ± 0.03</td>
<td>0.213</td>
</tr>
</tbody>
</table>

Kruskal-Wallis test of mean alleles per locus among groups $\chi^2_4 = 66.1, \ P < 0.001$

Kruskal-Wallis test of homogeneity of diversity among groups $\chi^2_4 = 19.97, \ P = 0.005$
Table 6
Mitochondrial variation estimated by single strand conformational polymorphisms in regionally grouped *G. pallidipes* populations.

<table>
<thead>
<tr>
<th>Grouping</th>
<th>No. populations</th>
<th>No. flies</th>
<th>No. haplotypes</th>
<th>Diversity, (H_S)</th>
<th>(F_{ST})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethiopia</td>
<td>2</td>
<td>48</td>
<td>7</td>
<td>0.44 ± 0.10</td>
<td>0.07</td>
</tr>
<tr>
<td>West Kenya</td>
<td>4</td>
<td>160</td>
<td>16</td>
<td>0.60 ± 0.06</td>
<td>0.27</td>
</tr>
<tr>
<td>Southwest Kenya</td>
<td>7</td>
<td>152</td>
<td>11</td>
<td>0.35 ± 0.09</td>
<td>0.41</td>
</tr>
<tr>
<td>East Kenya, Tanzania</td>
<td>5</td>
<td>168</td>
<td>23</td>
<td>0.59 ± 0.08</td>
<td>0.23</td>
</tr>
<tr>
<td>Southern Africa</td>
<td>3</td>
<td>96</td>
<td>4</td>
<td>0.04 ± 0.03</td>
<td>0.48</td>
</tr>
<tr>
<td>Totals and means</td>
<td>21</td>
<td>624</td>
<td>39</td>
<td>0.42 ± 0.02</td>
<td>0.51</td>
</tr>
</tbody>
</table>

Kruskal-Wallis test of homogeneity of diversities among groups \(\chi^2_4 = 10.97, P = 0.03\).
Table 7
Sequence variation at mitochondrial *r16S2* (197 bp) and *CoxI* (321 bp) among *Glossina* species. The sequences were aligned over species and concatenated.

<table>
<thead>
<tr>
<th>Species</th>
<th>No. samples</th>
<th>No. flies</th>
<th>No. haplotypes</th>
<th>Diversity $H_s$</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. swynnertoni</td>
<td>4</td>
<td>150</td>
<td>13</td>
<td>0.271</td>
</tr>
<tr>
<td>G. pallidipes</td>
<td>23</td>
<td>873</td>
<td>149</td>
<td>0.666</td>
</tr>
<tr>
<td>G. m. morsitans</td>
<td>3</td>
<td>88</td>
<td>21</td>
<td>0.685</td>
</tr>
<tr>
<td>G. m. centralis</td>
<td>1</td>
<td>47</td>
<td>15</td>
<td>0.684</td>
</tr>
<tr>
<td>G. m. submorsitans</td>
<td>3</td>
<td>87</td>
<td>30</td>
<td>0.777</td>
</tr>
<tr>
<td>G. austeni (lab culture)</td>
<td>2</td>
<td>47</td>
<td>8</td>
<td>0.362</td>
</tr>
<tr>
<td>G. fuscipes</td>
<td>4</td>
<td>127</td>
<td>41</td>
<td>0.863</td>
</tr>
<tr>
<td>G. p. gambiensis</td>
<td>2</td>
<td>60</td>
<td>23</td>
<td>0.822</td>
</tr>
<tr>
<td>G. longipennis</td>
<td>1</td>
<td>31</td>
<td>7</td>
<td>0.559</td>
</tr>
<tr>
<td>G. brevipalpis</td>
<td>2</td>
<td>72</td>
<td>13</td>
<td>0.652</td>
</tr>
</tbody>
</table>

* Kruskal-Wallis test of homogeneity in haplotype numbers: $\chi^2_9 = 9.40, P = 0.39$

** Kruskal-Wallis test of homogeneity in diversities: $\chi^2_9 = 14.4, P = 0.12$