

Tracking zoonotic pathogens using bloodsucking flies as 'flying syringes'

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Abstract About 60% of emerging infectious diseases in humans are of zoonotic origin. Their increasing number requires the development of new methods for early detection and monitoring of infectious agents in wildlife. Here, we investigated whether blood meals from hematophagous flies could be used to identify the infectious agents circulating in wild vertebrates. To this aim, 1230 blood-engorged flies were caught in the forests of Gabon. Identified blood meals (30%) were from 20 vertebrate species including mammals, birds and reptiles. Among them, 9% were infected by different extant malaria parasites among which some belonged to known parasite species, others to new parasite species or to parasite lineages for which only the vector was known. This study demonstrates that using hematophagous flies as 'flying syringes' constitutes an interesting approach to investigate blood-borne pathogen diversity in wild vertebrates and could be used as an early detection tool of zoonotic pathogens.

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Introduction

Emerging and re-emerging human infectious diseases have increased in recent years. Around one-fourth of the 1415 pathogens known to infect humans appeared between 1940 and 2004 and their appearance has gradually increased since 1980 (*Taylor et al., 2001*; *Woolhouse and Gaunt, 2007*; *Jones et al., 2008*; *Daszak et al., 2004*). Today, seven new pathogens appear every year and this number should reach 15–20 by 2020 (*Woolhouse et al., 2008*), mostly due to the growth of human activities that increase contact with novel sources of pathogens and favor their spread worldwide (*Murray et al., 2015*). Emerging threats mainly concern viruses, such as HIV (*Sharp and Hahn, 2011*), SARS-CoV and MERS-CoV (*de Wit et al., 2016*), avian flu (*Alexander, 2007*) and more recently Ebola (*Baize et al., 2014*), chikungunya (*Burt et al., 2012*) and Zika (*Wikan and Smith, 2016*). However, disease emergence and re-emergence also concern bacteria (e.g. *Helicobacter*



eLife digest About 60% of new infectious diseases in humans come from animals. Their increasing number and rapid spread are linked to increasing levels of contact between humans and wildlife, as recently highlighted by the epidemics of Zika in Brazil or Ebola in West Africa. To anticipate and prevent similar outbreaks in the future, it would be ideal to develop new methods for the early detection and monitoring of infectious diseases in wild animals.

Currently, three methods are mainly used to screen wild animals for infectious disease, but these all have limitations. Analyses of bushmeat and game meat only investigate those animals that are eaten by humans. Testing the organs and tissues of trapped animals can be difficult and harmful for both the humans and animals involved. Collecting and examining samples of feces, urine or saliva cannot detect all diseases and can be difficult to do for some species.

Bitome-Essono et al. now demonstrate a new method for assessing the diseases carried by wild animals: using blood-sucking flies as 'flying syringes' to collect their blood. During several weeks of sampling in Gabon, Central Africa, Bitome-Essono et al. trapped thousands of these flies, about a third of which were engorged with blood. Analyses of these blood samples revealed that they had come from 20 different species, including birds, mammals and reptiles. Different malaria parasites could also be detected in the blood.

Although the study performed by Bitome-Essono et al. only focused on malaria parasites, in the future the technique could be extended to analyze a number of disease-causing microbes – including viruses, bacteria, protozoa and macroparasites – that are found in the blood of wild animals.

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pylori, Salmonella sp., etc.) and parasites (e.g. Plasmodium knowlesi in South-East Asia). Sixty per cent of diseases emerging in humans are zoonoses and wildlife plays a key role by providing a zoonotic pool from which previously unknown pathogens may emerge (Taylor et al., 2001; Woolhouse and Gaunt, 2007; Jones et al., 2008; Daszak et al., 2004). The case of P. knowlesi in South-East Asia is a good example. This parasite emerged in the human population after a transfer from Asian macagues. It is now considered as the fifth human malaria agent after Plasmodium falciparum, Plasmodium vivax, Plasmodium malariae and Plasmodium ovale (Singh and Daneshvar, 2013). Such emerging diseases constitute a massive public health issue that requires active monitoring for signs of outbreaks and rapid diagnosis of the involved pathogen. Therefore, it is crucial to anticipate and prevent potential epidemic and pandemic outbreaks by developing new methods for the early detection and monitoring of infectious agents in wild animal sources (Kuiken et al., 2005; Wolfe et al., 2005). However, in many cases, monitoring is limited or impossible due to our poor knowledge about the ecology of these pathogens (i.e. where, when and how these agents circulate in the wildlife). The case of the Ebola virus is quite exemplary. Indeed, the exact nature of its reservoir(s) remains uncertain, although thousands of animals have been screened during the last 40 years (e.g. [Marí Saéz et al., 2015]).

Nowadays, pathogen circulation in wild animals is screened using mainly two methods: bushmeat analysis or direct trapping of animals for organ and tissue collection. These methods are pertinent in many cases, but present some weaknesses. Bushmeat represents only a fraction of the fauna (the one consumed by humans), whereas animal trapping can be difficult or dangerous. Moreover, such manipulation may be harmful for threatened and protected species. As a consequence, several methods were developed in the last years to study pathogen diversity from wild fauna without the need of direct contacts with animals, for example, by using fecal, urine or saliva samples (e.g. [Santiago et al., 2002; Prugnolle et al., 2010; Pesapane et al., 2013; Taberlet et al., 2012]). However, the value of these non-invasive methods remains limited because not all pathogens can be detected and not all reservoirs can be explored by these methods (for instance, it is difficult to collect feces or saliva of reptiles without trapping them). Therefore, new non-invasive methods are crucially needed to provide new opportunities for screening a larger range of hosts and pathogens.

The use of hematophagous flies as 'flying syringes' may constitute a new approach to track and survey blood-borne pathogens in the wild (*Calvignac-Spencer et al., 2013*). Nucleic acids (DNA or



RNA) of vertebrate hosts or of pathogens in arthropod blood meals are preserved and detectable for several days (Calvignac-Spencer et al., 2013; Kent, 2009; Muturi et al., 2011; Grubaugh et al., 2015; Lee et al., 2015). For example, HIV was detected 8 days and 10 to 14 days after blood ingestion by bugs and by ticks, respectively (Webb et al., 1989; Humphery-Smith et al., 1993). Recently, the H5N1 flu virus was found viable in mosquitoes (Barbazan et al., 2008), although its transmission by these insects is unproven (Sawabe et al., 2006). Grubaugh and colleagues (Grubaugh et al., 2015) applied such an idea (that they called 'xenosurveillance') using Anohpeles mosquitoes to estimate the diversity of viruses infecting human populations in remote areas. Nevertheless, bloodengorged mosquitoes are very difficult to collect in forest and often show strong host preferences (in particular for mammals). Arthropods with more generalist blood feeding patterns would be more useful to survey pathogens from a large range of vertebrates (including mammals, birds and reptiles) in these highly complex ecosystems.

Hematophagous flies (tsetse flies, stomoxids and tabanids) could be good candidates for this purpose since they are usually large Diptera (length comprised between 3 and 25 mm) and hematophagous in both sexes, with the exception of male tabanids (*Mullens, 2002*). They are easy to trap and some studies performed on tsetse flies and stomoxids showed that 20 to 40% of trapped flies are engorged with blood (*Mavoungou et al., 2008; Simo et al., 2012*). These flies feed on a large spectrum of vertebrate hosts, including birds, reptiles and mammals (*Muturi et al., 2011; Clausen et al., 1998; Muzari et al., 2010*). The omnipresence of hematophagous flies in certain habitats and their opportunistic blood-feeding behaviour (*Muturi et al., 2011; Muzari et al., 2010; Späth, 2000*) make of them compelling candidates to obtain blood meals from different vertebrate hosts for pathogen detection.

In the present study, we investigated the possibility of using hematophagous flies as 'flying syringes' to explore the diversity of extant malaria parasites (Haemosporida) infecting wild vertebrates living in the forests of Gabon (Central Africa).

Results

Host identification from blood meals

A total of 4099 hematophagous flies were caught in four national parks of Gabon during dry and rainy seasons over a cumulated sampling period of 16 weeks (*Figure 1a*). Among them, six tsetse fly species, six stomoxid species and six tabanid species were identified (*Table 1*).

Among the 4099 caught flies, 1230 (30%) were engorged with blood. These were mostly tsetse flies (n = 1218; 99%), particularly *Glossina palpalis palpalis* (n = 662; 54%) and *G. fuscipes fuscipes* (n = 214; 18%) specimens. The blood meal origin was successfully identified in 33% and 43% of these flies, respectively (*Table 1*).

Overall, the blood meal origin was successfully identified in 428 fly samples (35%) using a PCR system amplifying long fragments of *Cytb* (450 bp) or *COI* genes (330 bp or 660 bp). Specifically, blood meals were from 20 vertebrate species, including 12 families and 8 orders (*Figure 1b* and *Tables 2* and *3*).

A trial study using a PCR system amplifying a shorter fragment (150 bp of the gene 16S) to deal with potential DNA degradation in the blood meal showed a high gain of sensitivity in the determination of the origin of the blood meal. Thus, out of 89 previously unidentified blood meals, the host was identified for 76% (n = 68) of them. The list of newly identified hosts is given in **Figure 2**. This shows a high gain of sensitivity with the new PCR system.

Pathogen identification from blood meals

Extant malaria parasites were detected in 37 (8.7%) of the 428 identified blood meals (*Figure 1c*, red isolates). Phylogenetic analyses revealed that 29.7% of these parasites belonged to *Plasmodium falciparum* (n = 11, *Figure 1c*; group 1), 8.1% to *Plasmodium adleri* (n = 3, *Figure 1c*; group 2), and 8.1% to a recently described lineage of parasites infecting wild ungulates (n = 4, *Figure 1c*; group 3) (*Boundenga et al., 2016*). For all blood meals, the identified host represented the known natural host (or one of the hosts) of such parasites. Sequences of unknown parasite lineages or of parasites for which the hosts were not known were also obtained. For instance, one sequence (*Figure 1c*; group 4) detected in a blood meal originating from an ungulate was related to parasites previously



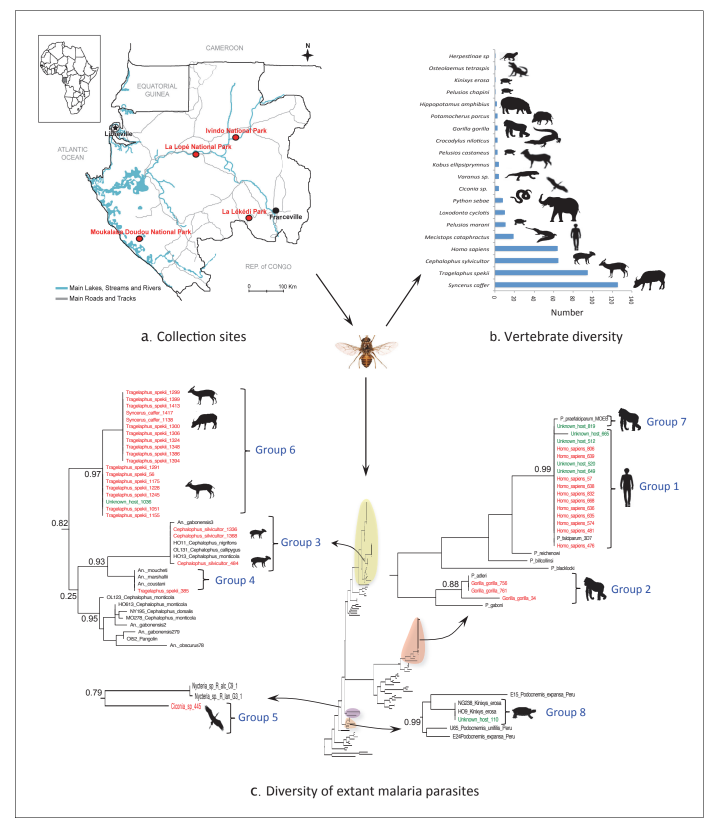


Figure 1. Monitoring vertebrate haemosporidian diversity using haematophagous flies. (a) Localization of the sampling sites (red dots) in Gabon (Central Africa). (b) Number of blood meals originating from the different vertebrate species. (c) Position within the *Cytb* phylogeny of the haemosporidian *Cytb* sequences PCR-amplified from the blood meals of engorged flies with identified hosts (red isolates) and unidentified hosts (green isolates). Black isolates: references (*Table 4*). Bootstrap values at important nodes are shown.

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Table 1. Number and proportion of specimens captured per fly species. The number of engorged specimens and blood meals identified in each fly species are also indicated.

Fly species	Number of collected specimens	Proportion (%)	Number of engorged specimens	Number of identified blood meals
Glossinidae	2252	54.94	1218	423
Glossina caliginea	144	3.51	87	33
G. fusca congolensis	210	5.12	104	42
G. fuscipes fuscipes	290	7.07	214	93
G. pallicera newsteadi	157	3.83	97	37
G. palpalis palpalis	1372	33.47	662	218
G. tabaniformis	79	1.93	54	0
Muscidae	1362	33.23	9	4
Stomoxys calcitrans	245	5.98	5	2
S. inornatus	334	8.14	0	0
S. niger niger	253	6.17	4	2
S. niger bilineatus	224	5.46	0	0
S. omega omega	197	4.81	0	0
S. transvittatus	109	2.66	0	0
Tabanidae	485	11.83	3	1
Ancala sp	41	1	0	0
Atylotus sp	104	2.53	0	0
Chrysops sp	156	3.81	3	1
Haematopota sp	13	0.31	0	0
Tabanus par	52	1.27	0	0
Tabanus taeniola	120	2.93	0	0
Total	4099	100	1230	428

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isolated only from Anopheles mosquitoes (**Boundenga et al., 2016**). One sequence detected in a blood meal originating from a bird was related to bat Haemosporida (*Nycteria*), (**Figure 1c**; group 5). Finally, 18 sequences (**Figure 1c**; group 6) that were amplified from blood meals originating from ungulates formed an independent and never described lineage related to groups 3 and 4.

In addition, 100 additional samples for which identification of the blood meal failed were randomly chosen for malarial parasite screening. This analysis showed that 7% were infected with P. falciparum (n = 4, group 1), P. praefalciparum (n = 1, group 7), malaria parasites of antelopes from group 6 (n = 1) and parasites of tortoises (group 8, n = 1) (**Figure 1c**, green isolates).

For the parasite, the use of a shorter PCR system led to less conclusive results than those obtained for the host identification. Out of the 91 blood meals that were negative to *Plasmodium* with a PCR system amplifying a long *Cytb* fragment, only one was found positive with the new system. The positive individual corresponded to a *Tragelaphus spekii* and was infected with a parasite belonging to group 3 (*Figure 1c*).

Discussion

In this study, we tested whether hematophagous flies could be used as 'flying syringes' to identify blood-borne pathogens circulating in the wild vertebrate fauna of Gabon. Our results show that the blood meals of the captured engorged flies can be successfully used to analyze the diversity of extant malaria parasites. Despite a limited sampling effort (a total of 4 weeks of sampling for each park), we could screen the diversity of haemosporidian parasites from a large range of vertebrate hosts, including mammals, birds and reptiles. Parasites were detected in more than 8% of the analyzed samples. These malaria parasites belonged to already known, but also to never previously

Table 2. Number and origin of blood meals according to the fly species (Fsp), park and climatic season.

Number of identified blood meals by fly species (Fsp)

												ľ											
			Mouk	alaba	Moukalaba-Doudou	nc	1						Lopé				1						
			Rainy season	seas	n		Δ	Dry season	Son			<u> </u>	Rainy season	eason			בֿ	Dry season	on				
Taxonomic group/ Order/Family	Host species	N° Identified	Fsp1 Fsp2	Fsp2	Fsp3 F	Fsp4 Fs	Fsp5 Fs	Fsp1 Fs	Fsp2 Fsp3	3 Fsp4	1 Fsp5	Fsp8 F	Fsp1 F	Fsp2 Fs	Fsp3 Fs	Fsp4 Fsp5		Fsp1 Fsp2	2 Fsp3	3 Fsp4	. Fsp5	Fsp6	Fsp7
Mammals																							
Artiodactyla		295		_	4 3	ω	_	-	7	_	14		2	_		2	m	က	4	-	က		
Bovidae	Cephalophus silvicultor	92																					
	kobus ellipsiprymnus	4				co co					-												
	Syncerus caffer	126	ю		2	7	_	_	m	2	6	2	_	10	2	7	-	2	8	-	9		-
	Tragelaphus spekii	95							_		9	4	_	2	_	ĸ	7	9	6	4	12	_	
Hippopotamidae	Hippopotamus amphibius	2				_					-												
Suidae	Potamochoerus porcus	ю				_					2												
Carnivora		-																					
Herpestidae	Herpestinae sp	-				-																	
Primates		29																					
Hominidae	Gorilla gorilla	ю				2					-												
	Homo sapiens	64			1	13			2		22	—	m		-	2			-		4		_
Proboscidae		10																					
Elephantidae	Loxodonta cyclotis	10									7			_		2							
Reptiles																							
Crocodilia		23									က												
Crocodylidae	Crocodylus niloticus	က																					
	Mecistops cataphractus	19			_	_					9												
	Osteolaemus tetraspis	_									-												
Squamata		12																					
Pythonidae	Python sebae	80									2												
Varanidae	Varanus sp	4							2		-			_									
Testudines		16																					
Testunidae	Kinixys erosa	_									—												
Pelomedusidae	Pelusios castaneus	8							_	_	-												
	Pelusios chapini	1									_												
	Pelusios marani	11				3					8												
Table 2 continued on next page	on next page																						

Table 2 continued

			Mot	ukalab	Moukalaba-Doudou	nop							Lopé)é									
			Rainy	Se	ason			Dry season	eason				Rai	Rainy season	son			Dry season	ason				
Birds																							
Ciconiformes		4																					
Ciconiidae Ciconia sp	Ciconia sp	4					_				2												
8 orders/12 families 20 species	20 species	428	3	-	11 4	4 41	41	2 ;	2	6 4	2 2 16 4 89 1 6 7 18	-	9	7	18	7	7 16 11 11 22 6	11 1	11	22 6	25	-	2

Number of identified blood meals by fly species (Fsp)

Fsp1 = Glossina caliginea; Fsp2 = G. fusca congolensis; Fsp3 = G. fuscipes fuscipes; Fsp4 = G. pallicera newsteadi; Fsp5 = G. palpalis palpalis; Fsp6 = Stomoxys calcitrans; Fsp7 = S. niger niger; Fsp8 = Chrysops sp.

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Table 3. Number and origin of blood meals according to the fly species (Fsp), park and climatic season.

			Nump	er of ide	entified	Number of identified blood meals by fly species (Fsp)	eals by	fly spe	cies (Fs	(d						
			La Lékédi	édi									lvindo			
			Rainy season	season			_	Dry season	lson				Dry season	ason		
Taxonomic group/Order/Family	Host species	N° Identified	Fsp1	Fsp3	Fsp4	Fsp5 F	Fsp6 F	Fsp1	Fsp2	Fsp3	Fsp4	Fsp5	Fsp1	Fsp2	Fsp3	Fsp5
Mammals																
Artiodactyla		295		2	~	е										
Bovidae	Cephalophus silvicultor	92														
	kobus ellipsiprymnus	4														
	Syncerus caffer	126	2	_		_		9	10	ω	8	13			_	-
	Tragelaphus spekii	95		—		3	_	2	8	2	2	6		~		
Hippopotamidae	Hippopotamus amphibius	2														
Suidae	Potamochoerus porcus	₆														
Carnivora		_														
Herpestidae	Herpestinae sp	_														
Primates		29														
Hominidae	Gorilla gorilla	3														
	Homo sapiens	64				4				_		2			_	_
Proboscidae		10														
Elephantidae	Loxodonta cyclotis	10														
Reptiles																
Crocodilia		23														
Crocodylidae	Crocodylus niloticus	3														
	Mecistops cataphractus	19							2	4	2	8				
	Osteolaemus tetraspis	_														
Squamata		12														
Pythonidae	Python sebae	80				2				2		2				
Varanidae	Varanus sp	4														
Testudines		16														
Testunidae	Kinixys erosa	1														
Pelomedusidae	Pelusios castaneus	m														
	Pelusios chapini	1														
	Pelusios marani	11														
Birds																

Table 3 continued on next page

Table 3 continued

			Num	ber of id	Number of identified blood meals by fly species (Fsp)	ood mea	ls by fly s	pecies (F	(d:					
			La Lé	Lékédi							_	lvindo		
			Rainy	/ season			Dry :	season				Dry season		
Ciconiformes		4												
Ciconiidae	Ciconia sp	4									_			
8 orders/12 families	20 species	428	2	4	_	3 1	∞	20	20	15	32 1	_	2	7

Fsp1 = Glossina caliginea; Fsp2 = G. fusca congolensis; Fsp3 = G. fuscipes fuscipes; Fsp4 = G. pallicera newsteadi; Fsp5 = G. palpalis palpalis; Fsp6 = Stomoxys calcitrans; Fsp7 = S. niger niger, Fsp8 = Chrysops sp.

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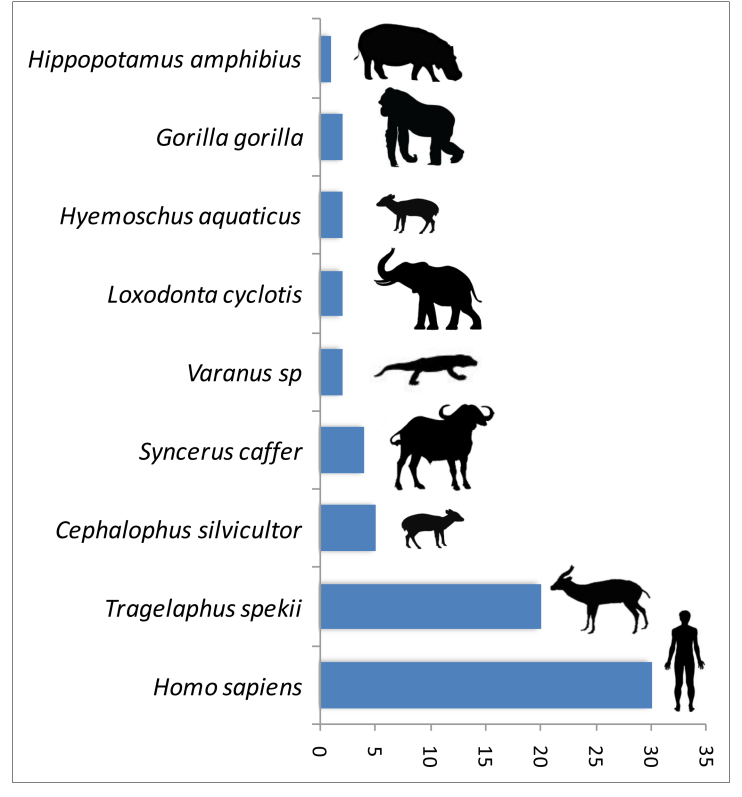


Figure 2. Number of blood meals identified using the shorter PCR system of **Boessenkool et al. (2012)** out of the previously unidentified 89 blood meals. DOI: 10.7554/eLife.22069.007

described lineages. In addition, the method allowed identifying the natural hosts of parasites for which only the vectors were known.



Table 4. Cytb sequences of parasites recovered in this study and of those used as references for phylogenetic analyses and their Genbank accession numbers.

Isolate	Accession number
Anopheles coustani	KT367855
An. gabonensis2	KT367852
An. gabonensis279	KT367853
An. gabonensis3	KT367861
An. marshallii	KT367857
An. moucheti	KT367864
An. obscurus2	KT367846
An. obscurus78	KT367849
Cephalophus_silvicultor_1336	KY631949
Cephalophus_silvicultor_1368	KY631947
Cephalophus_silvicultor_484	KY631963
Ciconia_sp_445	KY631985
E15_Podocnemis_expansa_Peru	KF049492
E24_Podocnemis_expansa_Peru	KF049495
Gorilla_gorilla_34	KY631983
Gorilla_gorilla_756	KY631982
Gorilla_gorilla_761	KY631981
P_spJA7_J725	GU252027
Haemoproteus_majoris	AY099045
Haemoproteus_sp.	HM222472
Haemoproteus_spGA02Cl1	HM222486
Haemoproteus_spNA16K65	HM222487
Hepatocystis_sp. AA201_blike	JQ070951
Hepatocystis_sp.	JQ070884
Hepatocystis_spAA2012	JQ070956
HO11_Cephalophus_nigrofons	KT367819
HO13_Cephalophus_monticola	KT367833
HO613_Cephalophus_monticola	KT367836
HO9_Kinixys_erosa	KT367843
Homo_sapiens_476	KY631978
Homo_sapiens_481	KY631977
Homo_sapiens_57	KY631979
Homo_sapiens_574	KY631976
Homo_sapiens_635	KY631975
Homo_sapiens_636	KY631974
Homo_sapiens_638	KY631973
Homo_sapiens_639	KY631972
Homo_sapiens_668	KY631969
Homo_sapiens_806	KY631968
Homo_sapiens_832	KY631967
Leucocytozoon_caulleryi	AB302215
Leucocytozoon_dubreuli	AY099063
Leucocytozoon_majoris	FJ168563

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Table 4 continued

Isolate	Accession numbe
Leucocytozoon_sabrazesi	AB299369
M0278_Cephalophus_monticola	KT367834
NG238_Kinixys_erosa	KT367844
NG277_Ceratogymna_atrata	KT367825
NY195_Cephalophus_dorsalis	KT367838
Nycteria_spR_alc_C9_1	KF159720
Nycteria_spR_lan_G3_1	KF159690
Ol52_Pangolin	KT367818
OL123_Cephalophus_monticola	KT367822
OL131_Cephalophus_callipygus	KT367830
P_adleri	HM235081
P_azurophilum	AY099055
P_billcollinsi	KP875474
P_blacklocki	HM235065
P_cynomolgi	AB444126
P_falciparum_3D7	AF069605
P_gaboni	JF895307
P_gallinaceum	AF069612
P_gonderi	JF923751
P_knowlesi	JQ345504
	HM000110
P_ovale	GU723548
P_praefalciparum_MOEB	JF923761
P_reichenowi	KP875479
P_relictum	AY733090
P_spDAJ	JF923753
P_vivax	KF591834
P_atheruri	AY099054
P_giganteum	AY099053
P_vinckei_isolate_1	KJ700853
P_vinckei_isolate_2	KJ700854
P_yoelii_killicki	DQ414658
P. atheruri	HQ712051
P. cyclopsi_Hip_cy_L4_1_Schaer	KF159674
P. voltaicum_M_ang_G1_1_1	KF159671
Parahaemoproteus_spbird_sp.17	GQ141581
Parahaemoproteus_spbird_sp.19	GQ141585
Parahaemoproteus_vireonis	FJ168561
Plasmodium_spbird	GQ141574
Plasmodium_spbird_sp12	HM222485
Plasmodium_spGD2_GD201	GU252012
Plasmodium_splineage_JA01	KM598212
Polychromophilus_melanipherus_haplotype_VIII	KJ131277
Polychromophilus_murinus_haplotype_3	HM055585



Table 4 continued

Isolate	Accession number
Polychromophilus_spMin_vil_G3_2	KF159699
Polychromophilus_spPip_gran_G3_1	KF159714
Polychromophilus_spNeo_cap_G3	KF159700
Syncerus_caffer_1138	KY631953
Syncerus_caffer_1417	KY631942
Tragelaphus_eurycerus_1324	KY631950
Tragelaphus_spekii_1051	KY631961
Tragelaphus_spekii_1155	KY631959
Tragelaphus_spekii_1175	KY631958
Tragelaphus_spekii_1228	KY631957
Tragelaphus_spekii_1245	KY631956
Tragelaphus_spekii_1291	KY631955
Tragelaphus_spekii_1299	KY631954
Tragelaphus_spekii_1300	KY631952
Tragelaphus_spekii_1306	KY631951
Tragelaphus_spekii_1348	KY631948
Tragelaphus_spekii_1386	KY631946
Tragelaphus_spekii_1394	KY631945
Tragelaphus_spekii_1399	KY631944
Tragelaphus_spekii_1413	KY631943
Tragelaphus_spekii_385	KY631964
Tragelaphus_spekii_56	KY631965
U65_Podocnemis_unifilis_Peru	KF049506
Unknown_host_1036	KY631960
Unknown_host_110	KY631966
Unknown_host_512	KY631962
Unknown_host_520	KY631980
Unknown_host_649	KY631971
Unknown_host_665	KY631970
Unknown_host_819	KY631984

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Concerning the method efficiency, 30% of blood meals were obtained from 4099 hematophagous flies. This result is consistent with previous studies (*Mavoungou et al., 2008*; *Simo et al., 2012*) showing that most hematophagous flies caught using traps are often seeking hosts for a blood meal. Other methods using a dip net seem to have a better capture efficiency with more than 40% of engorged flies caught on their resting places (*Gouteux et al., 1984*). However, this method requires spending a lot of time in the field because of difficulties in finding their resting sites and catching the flies.

Tsetse flies provided 99% of the collected blood meals (54% by Glossina palpalis palpalis) and they are an interesting candidate as 'flying syringes'. Indeed, differently from stomoxids and tabanids, both sexes are exclusively hematophagous in tsetse flies. In addition, G. p. palpalis is considered to be an opportunistic species concerning its feeding behaviour, thus explaining the large diversity of blood meals (Clausen et al., 1998; Simo et al., 2008; Weitz, 1963). Conversely, stomoxids and tabanids show sex-specific differences in feeding behaviour and this may partly explain the smaller number of blood meals collected in these two families. In stomoxids, both sexes are



hematophagous, but males sometimes feed on nectar (*Wall and Shearer*, 1997). Moreover, the digestion of stomoxids starts more rapidly than in the other hematophagous flies (*Moffatt et al.*, 1995). Male and female tabanids feed on nectar just after their emergence as adults. Only after having been fertilized, females start sucking blood (*Mullens*, 2002). Therefore, engorged stomoxid and tabanid flies are more difficult to capture. Additionally, the lack of engorged stomoxids and tabanids could be explained by the fact that we sampled flies only at floor level. Indeed, some stomoxid species readily feed on arboreal monkeys that are mostly found higher in the tree layer (*Mayoungou et al.*, 2008).

The low rate (35%) of blood meal identifications could be explained by the degradation of host DNA during digestion in the fly midgut or by a too small blood quantity in the midgut. The stage of digestion might influence DNA degradation and the host identification efficiency. Nevertheless, the diversity of hosts we successfully identified, mainly in tsetse fly blood meals, was large, including big terrestrial (elephants) and semi-aquatic mammals (hippopotamus) and also reptiles and birds. As previously noted, the diversity of blood meals can be due to the fly high mobility, their opportunistic feeding behaviour and their frequent feeding. In our study, most blood meals were from terrestrial animals (i.e. that live primarily on the ground) and very few from arboreal species. As mentioned above, this result is potentially biased by the trophic preferences of tsetse flies and by the capture method that excluded canopy levels. Previous studies have shown that hematophagous flies sampled in canopies mainly feed on arboreal species (Mavoungou et al., 2008). Therefore, changes in trap position could broaden the range of host species analysed. We can also notice the absence of small mammals (e.g., rodents or bats) within the diversity of host vertebrates we identified. This may be explained by the trophic preferences of the flies we sampled which could have a preferential taste for large vertebrates as previously documented for tsetse flies (e.g. [Muturi et al., 2011; Späth, 2000]).

Concerning pathogen detection, we detected extant haemosporidian parasites in 8.65% of the 428 blood meals for which the host origin was successfully identified. Moreover, we also detected parasites in blood meals of unknown origin, thus increasing the number of detected parasites. Together, these results show that blood meals collected from hematophagous flies are suitable for tracking blood-borne pathogens from wild animals. Haemosporidian pathogens ingested by hematophagous flies during their blood meal can remain detectable in the fly digestive tract even after partial digestion of the blood meal. We observed congruence between the identified hosts and the detected pathogens. As expected, *P. falciparum* was detected in human blood and *P. adleri* in gorilla blood. Haemosporidian lineages are often host-specific or restricted to certain classes of vertebrate hosts. Therefore, the unknown host could be inferred from the detected haemosporidian species (*Figure 1c*). For example, the blood meal from unknown host N°110 could have originated from a *Kinixys* turtle (*Kinixys sp.*). Similarly, the blood meals from the unknown hosts N°649, 520, 665, 512 and 819 could have originated from humans (*Homo sapiens*).

The present study demonstrates the possibility to use hematophagous flies as 'flying syringes' to analyze the diversity of pathogens circulating in wildlife. We think that there is now room for improvement of the tool; for instance, by improving the methods used to identify the blood meals and the pathogens. Since DNA is likely to be degraded in many blood meals (Calvignac-Spencer et al., 2013; Schnell et al., 2012), the use of PCR systems targeting fragments of shorter size could potentially improve the performance of detection. A trial study based on 89 previously unidentified blood meals using a PCR system amplifying a shorter fragment (<150 bp) (Boessenkool et al., 2012) than the one used in the present study allowed the identification of 76% (n = 68) of the hosts (Figure 2). This represents an important gain of sensitivity. However, these primers are still not ideal for our purpose as they were designed for optimal amplification of mammal DNA and often fail to properly amplify the DNA of other classes of vertebrates. A similar PCR system targeting the entire range of vertebrates still remains to be developed. For Plasmodium, our trial for amplifying a shorter fragment of Cytb (<200 bp) using a combination of previously published primers did not increase the sensitivity. Indeed, out of 91 samples for which the blood meal was successfully identified but in which no haemosporidian infection was detected with our long Cytb PCR system, only one was shown to be positive with the short PCR system. However, it is possible that other PCR systems, more optimized, could indeed improve the sensitivity of Plasmodium detection. Another direction of improvement could be the use of high-throughput sequencing technologies on pools of blood-engarged flies or amplicons to ease the identification of both hosts and parasites (especially



in the case of mixed blood meals or mixed infections). Finally, another way to improve the tool could be to use high-throughput multiplexed pathogen detection methods for the simultaneous testing of many samples in rapid succession. With such improvements, this approach of 'xenorsurveillance' could usefully complete recently developed methods based on the analysis of other invertebrates (carrion flies (Hoffmann et al., 2016), mosquitoes [Grubaugh et al., 2015]) and become an innovative way for the concomitant surveillance of many enzootic blood-borne pathogens, such as viruses (chikungunya, Zika), bacteria, protozoa and macro-parasites. The use of hematophagous flies as 'flying syringes' could indeed improve public health management by allowing the surveillance and early detection of zoonotic pathogens and thus prevent they spread to humans before they cause massive infections. This tool could also help to better understand the circulation in wildlife of other enzootic viruses, such as chikungunya or Zika, especially at the interface between natural/sylvan environments and, consequently, improving our knowledge of their natural history. From a broader perspective, this method could also be useful for people interested in wildlife biodiversity and conservation. Indeed, it could help monitoring the wildlife diversity within a specific region as demonstrated with other invertebrate systems (Calvignac-Spencer et al., 2013; Lee et al., 2015; Schnell et al., 2012; Schubert et al., 2015). More importantly, it could also allow detecting the emergence of new diseases in wild animals that may threaten their long-term survival.

Conclusion

Despite the significant scientific advances in the medical field, humans are still unable to predict where, when and how epidemics arise. Around 60% of emerging diseases in humans are of zoonotic origin. The progressive reduction of wild habitats will increase the contacts between humans and species that are potential reservoirs of diseases. We propose here a new non-invasive tool that can help identifying pathogens that circulate in wildlife before they spread in humans.

Materials and methods

Study sites

The fly sampling was carried out in four wildlife reserves in Gabon (*Figure 1a*): Moukalaba-Doudou National Park (MDNP; S: 2° 26′ 08"/E: 10° 25′ 18"), La Lopé National Park (LNP; S: 0° 31′ 31"/E: 11° 32′ 34"), La Lékédi Park (LP; S: 1° 45′ 32"/E: 13° 03′ 16") and Ivindo National Park (INP; N: 0° 30′ 82"/E: 12° 48′ 20"). Both MDNP and LNP are dominated by mature forests and mosaic forest-savannah. The INP is largely dominated by mature forest with some open biotopes that characterize the secondary forest. The LP is a private park dominated by large savannahs and some secondary forest and primary forest patches.

Sampling strategy

Hematophagous flies were sampled during the rainy and dry seasons between 2012 and 2014. In INP and MDNP, sampling was done during two years following a gradient of human activity from primary forest to villages. In the other parks, flies were sampled during a single year. Flies were collected by using Vavoua and Nzi traps (*Laveissiere and Grebaut, 1990*; *Acapovi et al., 2001*; *Mihok, 2002*; *Gilles et al., 2007*). The Vavoua trap, initially developed for the capture of tsetse flies was also successfully used for the capture of stomoxids at La Réunion Island (*Laveissiere and Grebaut, 1990*; *Gilles et al., 2007*). The Nzi trap was more adapted to the capture of *Glossina pallidipes* and tabanids in Africa (*Acapovi et al., 2001*; *Mihok, 2002*). In each park, we placed 24 traps (12 Vavoua and 12 Nzi) during 2 weeks per climatic season. Each trap was activated from 7:00 AM to 5:00 PM.

Identification and dissection of hematophagous flies

Freshly collected hematophagous flies were identified using a stereo-microscope and taxonomic procedure. The fly species (tsetse, stomoxids and tabanids) was determined following the determination keys of *Pollock (1982)*, *Brunhes et al., 1998, Zumpt, 1973, Garros et al. (2004)* and *Oldroyd (1973)*, on the basis of their morphological characteristics, such as size, color, wing venation structure and proboscis.



After species identification, engorged flies were dissected individually in a drop of Dulbecco's phosphate buffered saline solution (1x DPBS) to isolate blood meals from midgut. Each hematophagous fly was dissected on a slide using one forceps and one scalpel that were changed each time to avoid contaminations. Each blood meal was transferred in a 1.5-ml microtube containing 50 μ l of RNA*later* stabilization solution (Qiagen: Store at RT Tissue Collection) to stabilize and protect nucleic acids of vertebrate hosts and pathogens contained in the blood meals. Samples were kept at ambient temperature during field session and then frozen at -80°C until DNA extraction.

DNA extraction

Samples were centrifuged at 15,000 rpm at 4°C for 10 min to remove the RNA*later* solution. Pellets were used to extract DNA using the DNeasy Blood and Tissue Kit (Qiagen) according to the manufacturer's instructions. Extracted DNA was eluted in 100 μ l of buffer AE and stored at -20° C.

Blood meal identification

The origin of blood meals was determined using the extracted DNA to amplify a 450 bp fragment of the *cytochrome b* (*Cytb*) gene using previously published primers (*Townzen et al., 2008*). PCR amplifications were performed using a GeneAmp 9700 thermal cycler (Applied Biosystems, USA) with 50 μ L reaction mixtures containing 4 μ L template DNA, 10 mM Tris-HCl (pH = 9), 50 mM KCl, 3 mM MgCl₂, 20 pmol each primer (5'CCCCTCAGAATGATATTTGTCCTCA3' and 5'CCATCCAACATC TCAGCATGATGAAA3'), 200 mM dNTP and 1 U Taq polymerase. The thermal cycling conditions consisted of 3.5 min at 95°C, 40 cycles of 30s at 95°C, 50s at 58°C, and 40s at 72°C, followed by 5 min at 72°C. When *Cytb* amplification failed, a 330 bp and/or a 660 bp fragment of the *cytochrome oxydase* subunit I (*COI*) gene was amplified using previously described primers and protocols (*Townzen et al., 2008*). All PCR-amplified products (10 μ l) were run on 1.5% agarose gels in TBE buffer, and positive samples were sent to Beckman Coulter Genomics (France) for sequencing in both directions (forward and reverse) after purification. Consensus sequences were compared with existent sequences using the NCBI nucleotide Blast search (*Altschul et al., 1990*) to determine the host species. Hosts were identified when the amplified and reference sequences showed at least 98% similarity.

Haemosporidia detection and identification

Haemosporidian parasite detection was performed in samples with identified blood meal origin and also in 100 randomly chosen samples for which blood meal origin could not be identified.

Haemosporidian parasites were detected by PCR amplification of a portion of the *Cytb* gene (~790 bp) using a nested PCR protocol, as previously published (*Ollomo et al., 2009*). PCR products were checked on 1.5% agarose gels before shipment to EUROFINS MWG (Germany) for sequencing in both directions (reverse and forward) after purification. Multiple alignments of haemosporidian sequences were done using Muscle (*Edgar, 2004*). A phylogenetic tree with the haemosporidian sequences obtained in our study and a set of reference sequences was built using Maximum likelihood (ML) methods and phylogeny.fr (*Dereeper et al., 2008*) (see *Table 4* for accession numbers). The ML model used for construction of the tree was GTR (General Time Reversible)+ Γ (Gamma distribution)+ Γ (Invariable site distribution).

Anti-contamination procedures

Several measures were taken to avoid contaminations during our manipulations. Extraction of DNA was performed at the CIRMF (Gabon) in a laboratory working on mosquitoes. The room in which extraction was performed was away from the rooms in which DNA was amplified in this lab.

DNA extracts were then sent to France at the IRD (Montpellier). There, blood meal and *Plasmo-dium* identification was performed. This lab had never worked before on *Plasmodium* from ungulates or reptiles. Amplification of host DNA was never or very rarely performed in this lab. When the work was performed, no work on *Plasmodium* has been performed in this lab for almost 4 years. In addition, the laboratory is designed to avoid contaminations. Clearly defined and separated areas are devoted for each step of the PCR process: one area is devoted to the preparation of reagents (mix PCR). Another room is dedicated to the pre-PCR manipulation (loading of native DNA). This step is done under a cabinet to avoid contamination of the sample with DNA from the operator. Finally, an



area is devoted to PCR-amplified DNA. In this area, cabinets are used to deposit the first PCR product into the reagents of the second PCR (for nested PCRs). All cabinets are equipped with UV lamps and are always decontaminated with DNA-free solutions before and after manipulations. Gloves and coats are changed when moving between the areas and plugged tips are used at all steps. Blank controls were always incorporated at all steps of the experimental procedure and were always negative.

Several observations confirm the authenticity of our results: (1) >80% of the hosts that were found have never been manipulated in our lab (hosts that are not humans or non-human primates); (2) the parasite always corresponded to the expected host (antelope parasites were always found in antelopes, human parasites in humans and gorilla parasites in gorillas). Contaminations by external DNA would have lead to random association of hosts and parasites; (3) A new lineage of parasites was discovered.

Trial study to amplify shorter PCR fragments

Since DNA is likely to be degraded in many of our samples, the use of PCR systems targeting fragments of shorter size might improve performance. To determine if this could be the case with our study system, we performed supplementary analyses using (1) a PCR system targeting a shorter fragment of the vertebrate mitochondrial DNA to identify the blood meal origin and (2) a PCR system targeting a shorter fragment of the Cytb DNA to identify the parasite. For the identification of the host, the PCR system used was the one amplifying a fragment of 150 bp of 16S as described in Boessenkool et al. (2012) and using the primers 16Smam1 and 16Smam2. This PCR system was used on blood meals that failed to be identified using our original PCR system (see the paragraph 'Blood meal identification'). A total of 89 blood meals were tested for this trial study. For the parasite, we designed new primer sets to amplify a shorter fragment of the Cytb gene of the parasite (~177 bp). This new PCR system was applied to blood meals for which the host was identified but that were negative to Plasmodium with our long PCR system (~790 bp, see Material and methods above). A total of 91 blood samples were tested. For the first round of amplifications, we used 6 μL of DNA template in a 25 µL reaction volume, containing: 12.5 µL of Mix PCR (Qiagen), 2.5 µL solution Q (Qiagen), and 4 pmol of each primer (cytb1F CTCTATTAATTTAGTTAAAGCACACTT and 454R CCWGTWGCYTGCATYTATCT). Cycling conditions were 15 min at 95°C, 30 s at 94°C, 90 s at 57°C, 90 s at 72°C (40 cycles), and 10 min at 72°C. For the second round of amplification, we used 1.5 μ L of the first PCR template in a 25 μ L reaction volume, containing 2.5 μ L of 10× buffer, 1.25 mM MgCl₂, 250 μM of each dNTP, 10 pmol of each primer (454F2 WAATTAYCCATGYCCATTRAA and Plas1rc CACCATCCACTCCATAATTCTC), and 0.1 unit Tag Platinum (Invitrogen). Cycling conditions for the second round were 5 min at 95°C, 30 s at 94°C, 30 s at 50°C, 90 s at 72°C (35 cycles), and 10 min at 72°C. The amplified products (5 μL) were run on 1.5% agarose gels in TAE buffer. The PCR-amplified products (177 bp) were used as templates for sequencing. DNA sequencing was performed by Eurofins MWG.

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References

Acapovi G, Yao Y, N'Goran E, Dia ML, Desquesnes M. 2001. Abondance relative des tabanidés dans la région des savanes de côte d'Ivoire. Revue d'élevage Et De Médecine Vétérinaire Des Pays Tropicaux 54:974–980.

Alexander DJ. 2007. An overview of the epidemiology of avian influenza. Vaccine 25:5637–5644. doi: 10.1016/j. vaccine.2006.10.051, PMID: 17126960

Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic local alignment search tool. Journal of Molecular Biology 215:403–410. doi: 10.1016/S0022-2836(05)80360-2, PMID: 2231712

Baize S, Pannetier D, Oestereich L, Rieger T, Koivogui L, Magassouba N, Soropogui B, Sow MS, Keïta S, De Clerck H, Tiffany A, Dominguez G, Loua M, Traoré A, Kolié M, Malano ER, Heleze E, Bocquin A, Mély S, Raoul H, et al. 2014. Emergence of Zaire ebola virus disease in guinea. *The New England Journal of Medicine* 371: 1418–1425. doi: 10.1056/NEJMoa1404505, PMID: 24738640

Barbazan P, Thitithanyanont A, Missé D, Dubot A, Bosc P, Luangsri N, Gonzalez JP, Kittayapong P. 2008. Detection of H5N1 avian influenza virus from mosquitoes collected in an infected poultry farm in Thailand. *Vector Borne and Zoonotic Diseases* 8:105–110. doi: 10.1089/vbz.2007.0142, PMID: 18279078

Boessenkool S, Epp LS, Haile J, Bellemain E, Edwards M, Coissac E, Willerslev E, Brochmann C. 2012. Blocking human contaminant DNA during PCR allows amplification of rare mammal species from sedimentary ancient DNA. *Molecular Ecology* 21:1806–1815. doi: 10.1111/j.1365-294X.2011.05306.x, PMID: 21988749

Boundenga L, Makanga B, Ollomo B, Gilabert A, Rougeron V, Mve-Ondo B, Arnathau C, Durand P, Moukodoum ND, Okouga AP, Delicat-Loembet L, Yacka-Mouele L, Rahola N, Leroy E, Ba CT, Renaud F, Prugnolle F, Paupy



- C. 2016. Haemosporidian parasites of antelopes and other vertebrates from Gabon, central africa. *PLoS One* **11**:e0148958. doi: 10.1371/journal.pone.0148958, PMID: 26863304
- **Brunhes J**, Cuisance D, Geoffroy B, Hervy J. 1998. Les glossines ou mouches Tsé-tsé. Logiciel d'identification et d'enseignement. In: *Eds ORSTOM*. France: Montpellier.
- **Burt FJ**, Rolph MS, Rulli NE, Mahalingam S, Heise MT. 2012. Chikungunya: a re-emerging virus. *The Lancet* **379**: 662–671. doi: 10.1016/S0140-6736(11)60281-X, PMID: 22100854
- Calvignac-Spencer S, Leendertz FH, Gilbert MT, Schubert G. 2013. An invertebrate stomach's view on vertebrate ecology: certain invertebrates could be used as "vertebrate samplers" and deliver DNA-based information on many aspects of vertebrate ecology. BioEssays: News and Reviews in Molecular, Cellular and Developmental Biology 35:1004–1013. doi: 10.1002/bies.201300060, PMID: 23913504
- Clausen PH, Adeyemi I, Bauer B, Breloeer M, Salchow F, Staak C. 1998. Host preferences of tsetse (Diptera: glossinidae) based on bloodmeal identifications. *Medical and Veterinary Entomology* **12**:169–180. doi: 10. 1046/j.1365-2915.1998.00097.x, PMID: 9622371
- Daszak P, Tabor GM, Kilpatrick AM, Epstein J, Plowright R. 2004. Conservation medicine and a new agenda for emerging diseases. Annals of the New York Academy of Sciences 1026:1–11. doi: 10.1196/annals.1307.001, PMID: 15604464
- de Wit E, van Doremalen N, Falzarano D, Munster VJ. 2016. SARS and MERS: recent insights into emerging coronaviruses. Nature Reviews. Microbiology 14:523–534. doi: 10.1038/nrmicro.2016.81, PMID: 27344959
- Dereeper A, Guignon V, Blanc G, Audic S, Buffet S, Chevenet F, Dufayard JF, Guindon S, Lefort V, Lescot M, Claverie JM, Gascuel O. 2008. Phylogeny.fr: robust phylogenetic analysis for the non-specialist. *Nucleic Acids Research* 36:W465–W469. doi: 10.1093/nar/gkn180, PMID: 18424797
- Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Research* 32:1792–1797. doi: 10.1093/nar/gkh340, PMID: 15034147
- Garros C, Gilles J, Duvallet G. 2004. Un nouveau caractère morphologique pour distinguer stomoxys calcitrans et S. niger (Diptera: muscidae): Comparaison de populations de l'Ile de la Réunion. Parasite: Journal De La Société Française De Parasitologie 11:329–332.
- Gilles J, David JF, Duvallet G, De La Rocque S, Tillard E. 2007. Efficiency of traps for stomoxys calcitrans and stomoxys niger niger on reunion island. *Medical and Veterinary Entomology* 21:65–69. doi: 10.1111/j.1365-2915.2006.00658.x, PMID: 17373948
- Gouteux J, Bois J, Laveissiere C, Couret D, Mustapha A. 1984. Ecologie des glossines en secteur pré-forestier de côte d'Ivoire. 9. Les lieux de repos. *Cahiers ORSTOM* 22:159–174.
- **Grubaugh ND**, Sharma S, Krajacich BJ, Fakoli LS, Bolay FK, Diclaro JW, Johnson WE, Ebel GD, Foy BD, Brackney DE. 2015. Xenosurveillance: a novel mosquito-based approach for examining the human-pathogen landscape. *PLoS Neglected Tropical Diseases* **9**:e0003628. doi: 10.1371/journal.pntd.0003628, PMID: 25775236
- Hoffmann C, Stockhausen M, Merkel K, Calvignac-Spencer S, Leendertz FH. 2016. Assessing the feasibility of fly based surveillance of wildlife infectious diseases. Scientific Reports 6:37952. doi: 10.1038/srep37952, PMID: 27 901062
- Humphery-Smith I, Donker G, Turzo A, Chastel C, Schmidt-Mayerova H. 1993. Evaluation of mechanical transmission of HIV by the african soft tick, *ornithodoros moubata*. *Aids* **7**:341–348. doi: 10.1097/00002030-199303000-00006, PMID: 8471196
- Jones KE, Patel NG, Levy MA, Storeygard A, Balk D, Gittleman JL, Daszak P. 2008. Global trends in emerging infectious diseases. *Nature* **451**:990–993. doi: 10.1038/nature06536, PMID: 18288193
- Kent RJ. 2009. Molecular methods for arthropod bloodmeal identification and applications to ecological and vector-borne disease studies. Molecular Ecology Resources 9:4–18. doi: 10.1111/j.1755-0998.2008.02469.x, PMID: 21564560
- Kuiken T, Leighton FA, Fouchier RA, LeDuc JW, Peiris JS, Schudel A, Stöhr K, Osterhaus AD. 2005. Public health. Pathogen surveillance in animals. Science 309:1680–1681. doi: 10.1126/science.1113310, PMID: 16150997
- Laveissiere C, Grebaut P. 1990. Recherches sur les pièges à glossines (Diptera: glossinidae). Mise au point d'un modèle économique: le piège 'Vavoua'. Tropical Medicine and Parasitology: Official Organ of Deutsche Tropenmedizinische Gesellschaft and of Deutsche Gesellschaft Für Technische Zusammenarbeit 41:185–192.
- Lee PS, Sing KW, Wilson JJ. 2015. Reading mammal diversity from flies: the persistence period of amplifiable mammal mtDNA in blowfly guts (*Chrysomya megacephala*) and a new DNA Mini-Barcode target. *PLoS One* 10: e0123871. doi: 10.1371/journal.pone.0123871, PMID: 25898278
- Mavoungou JF, Simo G, Gilles J, De Stordeur E, Duvallet G. 2008. [Ecology of stomoxyine fulies (Diptera: muscidae) in Gabon. II. blood meals analysis a nd epidemiologic consequences]. *Parasite* **15**:611–615. doi: 10. 1051/parasite/2008154611, PMID: 19202770
- Mihok S. 2002. The development of a multipurpose trap (the nzi) for tsetse and other biting flies. Bulletin of Entomological Research 92:385–403. doi: 10.1079/BER2002186, PMID: 12241564
- Moffatt MR, Blakemore D, Lehane MJ. 1995. Studies on the synthesis and secretion of trypsin in the midgut of stomoxys calcitrans. Comparative Biochemistry and Physiology Part B: Biochemistry and Molecular Biology 110: 291–300. doi: 10.1016/0305-0491(94)00155-N
- Mullens BA. 2002. Horse Flies and Deer Flies (Tabanidae). In: Medical and Veterinary Entomology. p. 263–277. doi: 10.1016/B978-012510451-7/50015-3
- Murray KA, Preston N, Allen T, Zambrana-Torrelio C, Hosseini PR, Daszak P. 2015. Global biogeography of human infectious diseases. PNAS 112:12746–12751. doi: 10.1073/pnas.1507442112, PMID: 26417098



- Muturi CN, Ouma JO, Malele II, Ngure RM, Rutto JJ, Mithöfer KM, Enyaru J, Masiga DK. 2011. Tracking the feeding patterns of tsetse flies (Glossina genus) by analysis of bloodmeals using mitochondrial cytochromes genes. PLoS One 6:e17284. doi: 10.1371/journal.pone.0017284, PMID: 21386971
- Muzari MO, Burgess GW, Skerratt LF, Jones RE, Duran TL. 2010. Host preferences of tabanid flies based on identification of blood meals by ELISA. *Veterinary Parasitology* **174**:191–198. doi: 10.1016/j.vetpar.2010.08. 040, PMID: 20884120
- Oldroyd H. 1973. Tabanidae. In: Insects and Other Arthropods of Medical Importance. p. 195-202.
- Ollomo B, Durand P, Prugnolle F, Douzery E, Arnathau C, Nkoghe D, Leroy E, Renaud F. 2009. A new malaria agent in african hominids. *PLoS Pathogens* **5**:e1000446. doi: 10.1371/journal.ppat.1000446, PMID: 19478877
- Pesapane R, Ponder M, Alexander KA. 2013. Tracking pathogen transmission at the human-wildlife interface: banded mongoose and Escherichia coli. *EcoHealth* **10**:115–128. doi: 10.1007/s10393-013-0838-2, PMID: 23612 855
- Pollock JN. 1982. Tsetse biology, systematics and distribution, techniques. In: *Training Manual for Tsetse Control Personnel.* Vol. 1. Fao. p. 1–280.
- Prugnolle F, Durand P, Neel C, Ollomo B, Ayala FJ, Arnathau C, Etienne L, Mpoudi-Ngole E, Nkoghe D, Leroy E, Delaporte E, Peeters M, Renaud F. 2010. African great apes are natural hosts of multiple related malaria species, including plasmodium falciparum. PNAS 107:1458–1463. doi: 10.1073/pnas.0914440107, PMID: 20133 889
- Marí Saéz A, Weiss S, Nowak K, Lapeyre V, Zimmermann F, Düx A, Kühl HS, Kaba M, Regnaut S, Merkel K, Sachse A, Thiesen U, Villányi L, Boesch C, Dabrowski PW, Radonić A, Nitsche A, Leendertz SA, Petterson S, Becker S, et al. 2015. Investigating the zoonotic origin of the West African Ebola epidemic. EMBO Molecular Medicine 7:17–23. doi: 10.15252/emmm.201404792, PMID: 25550396
- Santiago ML, Rodenburg CM, Kamenya S, Bibollet-Ruche F, Gao F, Bailes E, Meleth S, Soong SJ, Kilby JM, Moldoveanu Z, Fahey B, Muller MN, Ayouba A, Nerrienet E, McClure HM, Heeney JL, Pusey AE, Collins DA, Boesch C, Wrangham RW, et al. 2002. SIVcpz in wild chimpanzees. Science 295:465. doi: 10.1126/science.295.5554.465, PMID: 11799233
- Sawabe K, Hoshino K, Isawa H, Sasaki T, Hayashi T, Tsuda Y, Kurahashi H, Tanabayashi K, Hotta A, Saito T, Yamada A, Kobayashi M. 2006. Detection and isolation of highly pathogenic H5N1 avian influenza A viruses from blow flies collected in the vicinity of an infected poultry farm in Kyoto, Japan, 2004. The American Journal of Tropical Medicine and Hygiene 75:327–332. PMID: 16896143
- Schnell IB, Thomsen PF, Wilkinson N, Rasmussen M, Jensen LRD, Willerslev E, Bertelsen MF, Gilbert MTP. 2012. Screening mammal biodiversity using DNA from leeches. *Current Biology* 22:R262–R263. doi: 10.1016/j.cub. 2012.02.058, PMID: 22537625
- Schubert G, Stockhausen M, Hoffmann C, Merkel K, Vigilant L, Leendertz FH, Calvignac-Spencer S. 2015.

 Targeted detection of mammalian species using carrion fly-derived DNA. *Molecular Ecology Resources* 15:285–294. doi: 10.1111/1755-0998.12306, PMID: 25042567
- Sharp PM, Hahn BH. 2011. Origins of HIV and the AIDS pandemic. Cold Spring Harbor Perspectives in Medicine 1:a006841. doi: 10.1101/cshperspect.a006841, PMID: 22229120
- Simo G, Njiokou F, Mbida Mbida JA, Njitchouang GR, Herder S, Asonganyi T, Cuny G. 2008. Tsetse fly host preference from sleeping sickness foci in Cameroon: epidemiological implications. *Infection, Genetics and Evolution: Journal of Molecular Epidemiology and Evolutionary Genetics in Infectious Diseases* 8:34–39. doi: 10.1016/j.meegid.2007.09.005, PMID: 17977803
- Simo G, Silatsa B, Flobert N, Lutumba P, Mansinsa P, Madinga J, Manzambi E, De Deken R, Asonganyi T. 2012. Identification of different trypanosome species in the mid-guts of tsetse flies of the malanga (Kimpese) sleeping sickness focus of the democratic republic of congo. *Parasites & Vectors* 5:201. doi: 10.1186/1756-3305-5-201, PMID: 22992486
- Singh B, Daneshvar C. 2013. Human infections and detection of plasmodium knowlesi. Clinical Microbiology Reviews 26:165–184. doi: 10.1128/CMR.00079-12, PMID: 23554413
- Späth J. 2000. Feeding patterns of three sympatric tsetse species (Glossina spp.) (Diptera: glossinidae) in the preforest zone of côte d'ivoire. Acta Tropica 75:109–118. doi: 10.1016/S0001-706X(99)00096-0, PMID: 1070 8012
- Taberlet P, Coissac E, Hajibabaei M, Rieseberg LH. 2012. Environmental DNA. *Molecular Ecology* **21**:1789–1793. doi: 10.1111/j.1365-294X.2012.05542.x, PMID: 22486819
- Taylor LH, Latham SM, Woolhouse ME. 2001. Risk factors for human disease emergence. *Philosophical Transactions of the Royal Society of London. Series B, Biological Sciences* **356**:983–989. doi: 10.1098/rstb.2001. 0888, PMID: 11516376
- **Townzen JS**, Brower AV, Judd DD. 2008. Identification of mosquito bloodmeals using mitochondrial cytochrome oxidase subunit I and cytochrome b gene sequences. Medical and Veterinary Entomology **22**:386–393. doi: 10. 1111/j.1365-2915.2008.00760.x, PMID: 19120966
- Wall R, Shearer D. 1997. Veterinary Entomology. Chapman & Hall.
- Webb PA, Happ CM, Maupin GO, Johnson BJ, Ou CY, Monath TP. 1989. Potential for insect transmission of HIV: experimental exposure of cimex hemipterus and toxorhynchites amboinensis to human immunodeficiency virus. The Journal of Infectious Diseases 160:970–977. doi: 10.1093/infdis/160.6.970, PMID: 2479697
- Weitz B. 1963. The feeding habits of glossina. Bulletin of the World Health Organization 28:711–729. PMID: 13 999790
- Wikan N, Smith DR. 2016. Zika Virus: history of a newly emerging arbovirus. The Lancet. Infectious Diseases 16: e119–e126. doi: 10.1016/S1473-3099(16)30010-X, PMID: 27282424



- Wolfe ND, Daszak P, Kilpatrick AM, Burke DS. 2005. Bushmeat hunting, deforestation, and prediction of zoonoses emergence. *Emerging Infectious Diseases* **11**:1822–1827. doi: 10.3201/eid1112.040789, PMID: 164 85465
- Woolhouse M, Gaunt E. 2007. Ecological origins of novel human pathogens. *Critical Reviews in Microbiology* **33**: 231–242. doi: 10.1080/10408410701647560, PMID: 18033594
- Woolhouse ME, Howey R, Gaunt E, Reilly L, Chase-Topping M, Savill N. 2008. Temporal trends in the discovery of human viruses. *Proceedings. Biological Sciences* **275**:2111–2115. doi: 10.1098/rspb.2008.0294, PMID: 1 8505720
- **Zumpt F.** 1973. The Stomoxynae biting flies of the world. In: *Taxonomy, Biology, Economic Importance and Control Measures.*