Ecological characteristics of *Simulium* breeding sites in West Africa

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ABSTRACT

Twenty-nine taxa of Simulium were identified amongst 527 collections of larvae and pupae from untreated rivers and streams in Liberia (362 collections in 1967-71 & 1989), Togo (125 in 1979-81), Benin (35 in 1979-81) and Ghana (5 in 1980-81). Presence or absence of associations between different taxa were used to group them into six clusters using Ward agglomerative hierarchical cluster analysis. Environmental data associated with the preimaginal habitats were then analysed in relation to the six clusters by one way ANOVA. The results revealed significant effects in determining the clusters of maximum river width (all P < 0.001 unless stated otherwise), water temperature, dry bulb air temperature, relative humidity, altitude, type of water (on a range from trickle to large river), water level, slope, current, vegetation, light conditions, discharge, length of breeding area, environs, terrain , river bed type (P < 0.01), and the supports to which the insects were attached (P < 0.01). When four non-significant contributors (wet bulb temperature, river features, height of waterfall and depth) were excluded and the reduced data-set analysed by principal components analysis (PCA), the first two principal components (PCs) accounted for 87% of the variance, with geographical features dominant in PC1 and hydrological characteristics in PC2. The analyses also revealed the ecological characteristics of each taxon's pre-imaginal

habitats, which are discussed with particular reference to members of the *Simulium damnosum* species complex, whose breeding site distributions were further analysed by canonical correspondence analysis (CCA), a method also applied to the data on non-vector species.

Keywords: Environmental variables, cluster analysis, PCA, CCA, *Simulium damnosum* complex

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1. Introduction

Blackflies of the genus *Simulium*, which includes vectors of onchocerciasis such as the *S*. *damnosum* complex, lay eggs in flowing water where their larvae and pupae develop before eclosion. As different species are found in differing categories of streams and rivers, egg-laying females presumably respond to cues in their habitats when selecting oviposition sites together with factors such as oviposition pheromones released by females that have already laid eggs in a particular micro-habitat (McCall, 1995, McCall et al., 1997). For the *S*. *damnosum* complex, the environmental cues include river size, river discharge rate, pH, conductivity and temperature (Quillévéré et al., 1976, 1977; Cheke, 2012), but to date no multivariate analyses of breeding site environmental variables for this important group of onchocerciasis vectors have been conducted, although some Central American vectors have

been studied using canonical correspondence analysis (CCA) (Millest et al., 1999). Similar studies in the tropics involving both principal components analysis (PCA) and CCA have been conducted on the blackfly faunas of Thailand (Srisuka et al., 2015) and Malaysia (Ya'cob et al., 2016a,b), with related studies using presence-absence data published for Brazil (Couceiro *et al.*, 2014) and Algeria (Chaoui Boudghane-Bendiouis et al., 2014). CCA has also been applied to environmental variables in relation to *Simulium* breeding in Europe, especially to blackfly data from the Carpathian Mountains (Illéšovà et al., 2008, 2010, Stranger et al., 2013). Such studies have revealed the significance for determining blackfly breeding habitats of variables such as altitude, water temperature, current speed, discharge slope, river width and depth, pH, conductivity, oxygen saturation and vegetation. The purpose of this retrospective study was to examine whether data on some but not all of the above variables together with some others could be used to characterise where immature stages of different species of West African Simuliidae might occur. Particular attention was paid to the various vectors of onchocerciasis found.

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We present analyses of data collected in Benin, Ghana, Liberia and Togo, consisting of information on the presence or absence of larvae and / or pupae of *Simulium* spp. in riverine breeding sites in relation to environmental characteristics of the sites surveyed in order to establish trends and relationships between the species' occurrences and the environmental variables. Regarding the vectors of onchocerciasis amongst the species identified, it is hoped that the results will help to identify the habitat types in which they may be found breeding and, in some cases, indicate the species likely to be present in a particular river. In addition, they may assist in predicting likely changes in geographical distributions in relation to climate change.

2. Methods

During field work in Liberia in the 1967-1971 period, all rivulets, streams and rivers in which Simulium spp. could breed were examined by R. Garms for the presence of larvae and pupae and, in many cases, traps were placed to collect Crustacea which might harbour pre-imaginal stages of phoretic species of blackflies, as described by Garms (1974). In Benin, Ghana and Togo only riverine sites that might harbour breeding S. damnosum s.l. were prospected by either or both of R. A. Cheke and R. Garms as part of research programmes and routine activities organised by the WHO Onchocerciasis Control Programme (OCP), but restricted to the period 1979-81 in rivers yet to be treated with insecticides. Most of these sites were accessed by helicopter and environmental data were collected simultaneously when pre-imaginal stages of *Simulium* were searched for. The collections were standardised insofar as a checklist was used for recording information; in Liberia all possible sites and substrates where immature stages of any species of Simulium might be found were searched but in Benin, Ghana and Togo searches were concentrated at positions likely to harbour members of the S. damnosum complex. The environmental data included measurements and subjective assessments of habitat characteristics (Table 1). Regrettably, no data on pH, dissolved oxygen concentrations or conductivity were obtained in sufficient amounts to be included in the analyses. For the multivariate analyses described below, missing values for environmental variables were replaced by their overall mean values.

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Data were collated from 527 sites in Liberia (362 collections in 1967-71 & 1989), Togo (125 in 1979-81), Benin (35 in 1979-81) and Ghana (5 in 1980-81). Garms (1974) provided maps of the Liberian sites and of the geographical distributions within the country of each species that he identified. See also Garms & Vajime (1975) for the *S. damnosum* complex in Liberia. The sites in Benin, Ghana and Togo were all amongst those depicted in maps published by Garms et al. (1982, 1983, 1989), Cheke & Garms (1983) and Cheke *et al.* (1987). Presence or absence associations between different taxa were used to group them into six clusters using Ward agglomerative hierarchical cluster analysis programmed in R. Environmental data associated with the pre-imaginal habitats were then analysed in relation to the clusters by one way ANOVA. When four non-significant contributors (wet bulb temperature, river features, height of waterfall and depth) were excluded, the reduced data-set was analysed by both principal components analysis (PCA) and canonical correspondence analysis (CCA; Ter Braak, 1986; Legendre & Legendre 1998, calculated as described by Oksanen et al., 2016). Regression and ordination are combined in CCA to generate a gradient that optimizes the distinctions between the species' distributions on ordination axes which are constrained as linear combinations of environmental variables (Ter Braak, 1986). Thereby, CCA results in an ordination diagram of samples, species and environmental variables optimally displaying how community composition varies in relation to the environment.

3. Results

Table 2 lists the taxa found and the countries they were recorded in. *S. sirbanum* was not recorded in Liberia in the samples for this study but the species is known to occur there (Garms *et al.* 1991). *S. sanctipauli* sensu Post 1996 and *S. soubrense* Vajime & Dunbar 1975 sensu Post 1986 were subsumed within *S. sanctipauli* s.l. in the analyses *S. ovazzae, S. berneri* and *S. lumbwanum* is are phoretic on freshwater crabs *Liberonautes latidactylus* and mayfly larvae of the genera *Elassoneuria* and *Afronurus. S. manense* was listed as *S. debegene* de Meillon 1934 by Garms (1974). *S. bovis* was not listed as occurring in Togo by Adler & Crosskey (2016), but pupae were collected there in R. Anie, 12.07.1979, and larvae and pupae in R. Na, 23.08.81.

Because the majority of the Liberian samples were collected and identified before the description of different cytospecies of the *S. damnosum* complex, we were unable to assign all such samples to taxa lower than *S. damnosum* s.l. except in a few cases such as in the St. Paul and Yah rivers where subsequent identifications provided strong indications of the taxa likely to be present. Also, for Liberia, we used a category of *S. sanctipauli* s.l. as we could not be sure which of the various members of the *S. sanctipauli* sub-complex reported in Liberia had been present at the sites and times of our collections, especially since different forms may occur sympatrically (Güzelhan & Garms 1991).

Figure 1 shows results of the cluster analysis using the entire data-set, stopped at a level of six clusters and shows how the various species were associated within these different clusters. Table 3 summarises salient features of each cluster showing for each one the dominant members of the *S. damnosum* complex and the dominant non-damnosum species, together with differences between those found in Liberia only and those in the other three countries only. Cluster 2 contained species only found in Liberia and, similarly, very few of the species in clusters 5 and 6 ever occurred there. A one-way ANOVA was then performed for each river variable against the cluster codes. The results revealed significant effects (all P < 0.001 unless stated otherwise) in determining the clusters of maximum river width , water temperature , dry bulb air temperature , relative humidity (P < 0.01), altitude , type of water (on a range from trickle to large river, water level, , slope , current , vegetation , light conditions , discharge , length of breeding area , environs , terrain river bed type (P < 0.01) and the supports to which the insects were attached (P < 0.01).

Four of the river variables (wet bulb temperature, river features, height of waterfall and depth) were non-significant contributors. These were then excluded and the reduced data-set analysed by principal components analysis (PCA). The first two principal components (PCs) accounted for 87% of the variance, with geographical features dominant in PC1 and

hydrological characteristics in PC2 (Fig. 2A), which can be related to the species clusters on the PCA axes depicted in Fig. 2B.

When CCA was applied to data only for the various members of the *S. damnosum* complex involved then the CCA1 (x) axis was dominated by temperature effects and the CCA2 (y) axis mainly by hydrological characteristics of the rivers (Fig. 3A). With this in mind, interpretation of the CCA plot for the vector species (Fig. 3B) is instructive as it clearly shows the separation of the Liberian *S. sanctipauli* sub-complex according to high river discharge rates and other hydrological parameters as the top-most along the CCA2 axis and *S. yahense* according to temperature far to the right on the CCA1 axis. Other separations follow CCA2 according to hydrology, with close alignments of *S. sirbanum*, the Beffa form of *S. soubrense* and *S. damnosum* s.str., species which are often sympatric, separated from *S. squamosum* which is grouped not far from the Djodji form of *S. sanctipauli* (the two were often sympatric before the latter was extirpated, see Cheke et al. 2008).

CCA results for the environmental variables of the non-vector species have CCA1 again dominated by temperature with hydrological features notable in CCA2 (Fig 4A). Species isolated in the ordination diagram, apparently lacking non-vector congeners with similar ecological requirements, include *S. tridens*, *S. kenyae*, *S. futaense*, *S. berneri*, *S. adersi* and *S. hargreavesi*, while most of the remainder are clumped near to each other (Fig. 4B).

4. Discussion

Given the 17 environmental variables that were statistically significant contributors to the separation of the clusters, it is clear that breeding site choice in the Simuliidae is governed by a multiplicity of factors. The results also provided evidence for substantial niche separations

between species, as shown by the means of variables according to species (Tables 4 & 5) and the results of both the PCAs and the CCAs. The first two principal components (PCs) accounted for 87% of the variance, with geographical features dominant in PC1 and hydrological characteristics in PC2. In the CCA, temperature dominated CCA1 and hydrological features CCA2.

Ever since the first description of different members of the *Simulium damnosum* complex by Vajime & Dunbar (1975), it was realised that they had different ecological adaptations, typically considered as being forest for the *S. sanctipauli* sub-complex and *S. yahense*, and savanna for *S. damnosum* s.str., *S. sirbanum* and *S. sudanense* (not encountered in this study), with *S. squamosum* mostly found in forested highlands and *S. soubrense* in forest-savanna mosaic habitats. In addition the *S. sanctipauli* sub-complex was usually in large rivers, with *S. yahense* and *S. mengense* (occurs only in Cameroon and the Central African Republic and so absent from this study) in smaller rivers and streams. Although Quillévéré et al. (1976, 1977) characterised and compared river features of savanna and forest-dwelling species, there have been few other quantitative studies of the ecological requirements of the different species and none using the multivariate statistical methods applied in this investigation. The results presented here confirmed and quantified niche separations between different *Simulium* species and, in particular, those for the *S. damnosum* complex.

Results of the CCA for the *S. damnosum* complex (Fig. 3) revealed clear separations in particular for *S. sanctipauli* s.l. and *S. yahense* from the others, which in turn were split along a hydrological gradient reflecting their known ecological adaptations to rivers of different discharges and widths with temperature, for instance, defining *S. yahense* preferences. Ya'cob et al. (2016b) also found temperature, river width and discharge to be dominant in their CCAs.

S. yahense was found in rivers with a mean water temperature of 24.16°C, nearly 2 degrees lower than the mean (27.65) for *S. sanctipauli* s.l., while the other species were all found in rivers with means of >28°C (Table 2). These data add to and complement river temperature ranges for different members of the *S. damnosum* complex, with only a few mean values, listed by Cheke (2012). It was not surprising that the Liberian *S. sanctipauli* s.l. was separated according to river discharge (mean 58.83 m³.sec⁻¹) and maximum river width (mean 108.42m) as the sub-complex is well known to occur predominantly in large rivers with substantial flow. Similarly, *S. yahense* which segregated on a temperature axis (see above) with slope (highest mean score of 4.09) also important, is a species of relatively cool narrow rivers (mean maximum width only 23.1m) in upland, well forested, areas and was the most likely to be found in rocky rapids (river features score the highest at 4.56). The highland habitat of *S. squamosum* was confirmed by its having the highest mean value for altitude (mean 293.75m), with *S. sanctipauli* s.l. the lowest (127.5m).

Of the non-vector species, the highland-dwelling *S. futaense* was found in the highest (mean altitude 414m) and coldest rivers (mean 21.66°C) and *S. ruficorne* in the hottest (29.25°C). *Simulium* species most often found associated with members of the *S. damnosum* complex were *S. unicornutum*, *S. cervicornutum*, *S. adersi*, *S. tridens*, *S. kenyae* and *S. hargreavesi* (Fig. 1B) and these species were separated along the CCA2 axis dominated by river features and to a lesser extent along the CCA2 axis dominated by temperature (Fig. 4B), possibly reflecting their associations with different members of the vector complex.

The statistical and computational methods used did not exist at the time the data collections began and so this was a *post hoc* study, lacking the benefit of a careful experimental design. Thus, some important measurements of variables such as dissolved oxygen, conductivity and pH were not taken, although data on hardness and carbon dioxide concentrations for some polluted and unpolluted rivers in Liberia are given by Garms et al. (1991) and physicchemical characteristics of rivers in which members of the S. damnosum complex breed were described by Grunewald (1976). Another criticism is that the samples were not taken using a randomised protocol. However, the large sample sizes counter this criticism as nearly all sites that were amenable to sampling were examined. In the OCP area, for example, it was necessary to check all rivers for any breeding by onchocerciasis vectors. In addition, the data were not all collected at the same time of the day so, given that river water temperatures in the region may vary by about 2°C during the course of a day (see Fig. 10 of Garms et al. 1991), some inaccuracy is introduced by the lack of temporal constancy. Previous studies (Quillévéré et al., 1976, 1977; Millest et al., 1999) have highlighted the importance of pH in determining species' geographical distributions but they relied on pH papers for their tests prior to the wide availability of electronic pH meters, so the accuracy of their results may not be reliable. Nevertheless, more recent studies using modern pH meters have also drawn attention to the importance of pH for Simulium breeding (Srisuka et al., 2015; Ya'cob et al., 2016a), so it is worth noting that the pH of the St.Paul river at Haindi, a major breeding site for the St.Paul form of S. soubrense, ranged from 6.9 to 7.7 between September 1967 and June 1968, when dissolved oxygen concentrations ranged from 8.3 to 10.5 mg.l⁻¹ (Bong Mining Company, unpublished data). Details of physicochemical characteristics of the River Pra in Ghana, another breeding site for S. sanctipauli s.l., were published by Wilson et al. (2005).

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The summaries of environmental data according to species (Tables 4 & 5) provide quantitative support for the qualitative summaries given for the Simuliidae of Guinea, most of which species also occur in Liberia, by Garms & Post (1967) who provided diagrams of the species' preferred river types and species' associations.

Cheke et al. (2015) used temperature data in mathematical models to predict how populations of savanna-dwelling *S. damnosum / S. sirbanum* and forest-dwelling *S. sanctipauli* in Ghana and Liberia, respectively, might change with increasing temperatures under climate change predictions and their likely consequences for future onchocerciasis transmission scenarios. The temperature data provided in this paper (Table 4) for additional members of the *S. damnosum* complex may allow similar predictions to be made for extra vector species and, also, on how they might interact competitively in the future. For instance as temperatures rise, *S. squamosum* will be more able to compete with *S. yahense* in forested areas.

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Measurements						Notes					
	Unit										
River width	m										
River depth	cm										
River discharge	m ³ .sec ⁻¹ Sometimes estimated without measurements										
River water	°C										
temperature											
Air temperature (wet bulb)	°C										
Air temperature (dry bulb)	°C										
RH	%										
Height of waterfall	m										
Length of <i>Simulium</i> breeding area	m										
Altitude	m a.s.l.	•	1								
Subjective habi	tat assessments (1-5 scales; 1-3 sc	cales for light co	nditions and te	errain)	If >1 class recorded, then highest used					
	1	2	3	4	5						
Type of water	Trickle	Trickle Streamlet or stream		Medium river	Large river						
River water level	Trickle	Low	Medium	High	Inundating						
River bed	Rocky (stones > 10cm across)	Stony (mostly stones < 10cm across)	Earthen	Sandy	Muddy						
River features	Sandy island	Fish fence	Bridge	Rapids	Rocky barrier						
Slope	Inconspicuous	Moderate	Considerable	Terraced	Waterfall						
Current	Smooth	Slow	Swift	Fast	Very fast						
Vegetation	None	Plants on banks, on rocks or overgrowth on rocks	Trailing grasses or plants	Bushes	Trees						
Light conditions	Open to sun	Partially shaded	Shadowed								
Environs	Grassland	Farmland or plantation	Low bush or savanna	Forest	High forest	If >1 class recorded, then highest used unless 2 involved, then 2 used to highlight effects of man					
Terrain	Plain	Hilly	Mountainous								
Supports	Stones or rocks	Dead leaves	Sticks or roots	Trailing grasses or plants	Bushes						

 Table 1. Details of measurements made at Simulium spp. breeding sites and of subjective assessments of habitat characteristics.

Taxon	Liberia	Benin	Ghana	Togo
Simulium damnosum sensu lato	×	×	×	×
S. damnosum Theobald 1903 sensu stricto	×	×	×	X
S. sirbanum Vajime & Dunbar 1975		×		X
S. yahense Vajime & Dunbar 1975	X			X
S. sanctipauli Vajime & Dunbar 1975 sensu lato	×	×	×	×
S. sanctipauli Djodji form Surtees, Fiasorgbor, Post &			×	×
Weber 1988				
S. soubrense Beffa form Meredith, Cheke & Garms 1983		<mark>X</mark>		<mark>X</mark>
S. squamosum (Enderlein 1921) cytotaxonomic description		×	x x	<mark>×</mark>
by Vajime & Dunbar 1975				
S. tridens Freeman & de Meillon 1953	<mark>X</mark>	<mark>X</mark>		<mark>X</mark>
S. ruficorne Macquart 1838				<mark>X</mark>
S. loutetense Grenier & Ovazza 1951	×			
S. alcocki Pomeroy 1922	<mark>X</mark>			<mark>X</mark>
S. garmsi Crosskey 1969	<mark>X</mark>			
S. djallonense Roubaud & Grenier 1943	<mark>x</mark>			
S. johannae Wanson 1947	<mark>x</mark>			
S. liberiense Garms 1973	×			
S. schoutedeni Wanson 1947	×			×
. unicornutum Pomeroy 1920	×	×	×	×
S. blacklocki de Meillon 1930	×			
S. cervicornutum Pomeroy 1920	×		×	×
S. kenyae de Meillon 1940		<mark>x</mark>	× ×	<mark>×</mark>
S. adersi Pomeroy 1922	×	×		×
S. ovazzae Grenier & Mouchet 1959	×			
S. berneri Freeman 1954	×			
S. lumbwanum de Meillon 1944	×			
S. manense Elsen & Escaffre,	×			
S. futaense Garms & Post 1966	X			
S. colasbelcouri Grenier & Ovazza 1951	X			
S. hargreavesi Gibbins 1934		X		X
S. bovis de Meillon 1930				×

Table 2. Taxa recorded and the countries in which they were found.

Table 3. Features of the six different clusters showing the dominant members of the S. damnosum complex and the dominant non-damnosum species present in each cluster.

Cluster Number	Most abundant <i>S.</i> <i>damnosum</i> complex <mark>taxa</mark>	S. damnosum s.l. abundance	Most abundant non- <i>damnosum</i> <mark>species</mark>	Found in <mark>Liberia</mark>	Found in Benin, Ghana, Togo	
<mark>1</mark>	sanctipauli	medium	kenyae	yes	yes	
<mark>2</mark>	yahense	medium	<mark>cervicornutum</mark>	yes	no	
<mark>3</mark>	yahense	low	unicornutum	yes	yes	
<mark>4</mark>	yahense	low	<mark>schoutedeni</mark>	yes	yes	
<mark>5</mark>	<mark>squamosum</mark>	medium	adersi	no	yes	
<mark>6</mark>	sanctipauli, damnosum s.str, soubrense Beffa form	high	adersi	no	yes	

Table 4. Mean values of measured environmental variables according to *Simulium* taxon. Figures in parentheses are standard errors of the mean (SEM).

<i>Simulium</i> taxon	River width (max) (m)	Water Temp. (°C)	Air Temp Wet (°C)	Air Temp Dry (°C)	RH (%)	Altitude (m)	
damnosum s.l.	52.87 (3.85)	26.35 (0.16)	24.17 (0.15)	27.78 (0.20)	84.62 (1.12)	234.57 (13.73)	
sanctipauli s.l.	108.42 (8.62)	27.65 (0.23)	25.09 (0.41)	28.72 (0.34)	82.18 (2.48)	127.50 (22.36)	
damnosum s.str.	59.37 (6.03)	28.39 (0.22)	23.00 (-)	29.58 (0.30)	86.00 (-)	254.29 (22.66)	
sirbanum	78.89 (14.21)	28.13 (0.33)	-	29.53 (0.42)	-	250.00 (-)	
squamosum	32.02 (3.03)	28.05 (0.29)	-	29.71 (0.30)	-	293.75 (35.70)	
yahense	23.10 (2.99)	24.16 (0.16)	24.01 (0.16)	26.15 (0.24)	86.08 (1.26)	253.32 (17.61)	
sancti.Djodji	33.25 (11.21)	28.47 (0.66)	-	29.86 (0.90)	-	-	
soubrense Beffa	80.46 (10.46)	28.27 (0.27)	-	29.24 (0.45)	-	240.00 (60.00)	
alcocki	4.87 (0.65)	24.38 (0.25)	24.01 (0.34)	25.35 (0.44)	91.84 (1.55)	217.08 (37.31)	
garmsi	18.30 (3.86)	24.27 (0.14)	24.44 (0.17)	26.07 (0.24)	89.04 (1.10)	174.79 (13.12)	
djallonense	7.67 (1.52)	23.96 (0.19)	24.49 (0.19)	25.83 (0.31)	91.42 (1.22)	237.05 (31.50)	
liberiense	9.11 (1.74)	23.69 (0.19)	24.37 (0.34)	25.27 (0.43)	93.66 (1.18)	207.58 (23.17)	
schoutedeni	20.53 (2.13)	24.21 (0.12)	24.22 (0.14)	26.08 (0.20)	87.56 (1.04)	185.45 (12.84)	
cervicornutum	15.54 (1.07)	24.47 (0.14)	24.28 (0.15)	26.30 (0.22)	87.52 (1.12)	198.98 (14.18)	
unicornutum	25.78 (3.49)	24.72 (0.14)	24.28 (0.13)	26.46 (0.20)	88.06 (0.88)	197.36 (11.32)	
blacklocki	7.38 (1.92)	24.41 (0.20)	25.24 (0.41)	27.17 (0.61)	87.17 (2.52)	110.00 (14.51)	
kenyae	84.25 (9.36)	25.24 (0.29)	23.88 (0.23)	26.47 (0.34)	84.55 (1.57)	219.22 (22.62)	
ruficorne	8.00 (2.00)	29.25 (0.75)	-	-	-	-	
manense	9.50 (0.50)	24.17 (0.41)	23.67 (0.88)	25.13 (0.19)	90.67 (4.91)	196.67 (27.28)	
loutetense	6.77 (1.01)	23.96 (0.16)	24.16 (0.17)	25.66 (0.23)	90.17 (1.07)	248.63 (19.42)	
colasbelcouri	5.17 (1.28)	23.21 (0.90)	23.87 (0.99)	25.23 (0.30)	90.67 (5.40)	241.67 (30.81)	
futaense	21.00 (9.00)	21.66 (1.10)	21.62 (1.13)	23.40 (1.04)	88.20 (6.36)	414.00 (17.49)	
ovazzae	14.33 (3.50)	22.82 (0.58)	23.58 (0.53)	25.46 (0.70)	87.55 (3.84)	370.00 (44.66)	
berneri	35.83 (6.64)	24.51 (0.69)	24.12 (0.41)	26.14 (0.96)	88.11 (5.91)	357.14 (43.08)	
lumbwanum	30.00 (-)	24.35 (1.15)	24.00 (-)	24.80 (-)	95.00 (-)	50.00 (-)	
johannae	22.40 (7.95)	23.53 (0.54)	23.92 (0.85)	25.02 (0.93)	92.83 (1.60)	201.67 (64.16)	
tridens	124.00 (46.65)	27.14 (1.75)	27.00 (-)	30.67 (1.88)	60.00 (-)	10.00 (-)	
adersi	48.70 (5.54)	28.50 (0.23)	24.00 (-)	29.88 (0.29)	62.00 (-)	210.00 (10.00)	
hargreavesi	23.42 (4.36)	27.60 (0.39)	-	29.16 (0.63)	-	-	
bovis	50.00 (30.00)	26.25 (1.25)	-	-	-	-	

Table 5. Mean values of subjective habitat assessments (1-5 scales in all cases, except 1-3 for light conditions and terrain) according to Simulium taxon. Blank cells refer to missing data.

Simulium	Туре	Water	River	River	Slope	Height	Current	Veget-	Light	Depth	Dischar	Length	Envir-	Terr-	Supp-
taxon	of	Level	Bed	Feat-		of		ation		(cm)	ge	of	ons	ain	orts
	Water			ures		water-					M3/sec	Breed-			
						fall						ing			
						(m)						Area			
<u>.</u>			4.96	2.00			2.56			20.60	15.65	(m)			0.54
Sa sensu lato	3.22	2.57	1.36	3.89	3.32	3.34	3.56	2.44	2.34	38.69	15.65	82.14	3.04	1.45	3.51
sanctipauli s.l.	4.20	2.86	1.20	3.83	2.85	2.50	3.76	2.82	2.82	40.03	58.83	124.73	3.09	1.22	3.71
aamnosum	3.64	2.54	1.36	3.89	2.78	2.02	4.00	3.04		26.19	1.37	105.36	2.95	1.14	3.92
s.str.	2.76	2.24	1 1 4	2 00	2.00	2.40		2 70		0.75	2 70	107 10	2.07	1.00	2 95
sirbunum	3.70	2.54	1.14	2.00	3.00	2.40		3.20		22.10	5.79	24.60	3.07	1.00	2.02
squumosum	3.17	2.51	1.50	5.09	2.05	2.00	2 55	1.04	2.25	42.00	1.79	64.09 E9.61	3.00	1.05	3.02
yunense cancti Diodii	2.50	2.45	1.20	4.50	4.09	5.70	5.55	1.04	2.25	45.09	1.45	76.67	2.09	1.52	3.20
sulicii Djuuji	2.00	2.05	1.00	2.20	2.25	2 50		2.37		25.05	1.00	146 56	3.50	1.00	3.00
alcocki	2.00	1.26	1.23	5.00	2.75	2.30	2.22	1.0/	1 02	20.85	2.02	20.61	2.50	1.05	4.00
aarmsi	2.00	2.20	1.00	2.00	3.52	1.53	3.22	1.94	2 33	20.85 13.04	2.03	30.01	2.30	1.09	2.00
diallonense	2.40	1.56	1.41	5.00	1 25	9.82	3.25	1.67	1.60	28.00	1.30	17 36	3.10	1.2.5	2.50
liheriense	2.05	2.00	1.45	5.00	4.23	2.41	3.21	1.07	1.00	33.00	0.51	47.06	3.95	1.54	3.24
schoutedeni	2.57	2.00	1.17	4 33	3 51	5.49	3 35	1.07	2.38	43.42	1 77	41.09	3.18	1 32	3.00
cervicornutum	2.42	2.34	1.43	4.28	3.56	6.32	3.40	2.05	2.31	39.79	1.56	47.73	3.07	1.52	3.30
unicornutum	2.57	2.36	1.54	4.14	3.55	3.65	3.28	2.01	2.21	38.57	1.99	47.75	3.12	1.39	3.14
blacklocki	2.00	2.38	2.33	5.00	3.46	13.13	3.08	2.69	2.60	35.71	0.05	25.56	3.63	1.38	3.54
kenyae	3.94	2.91	1.18	4.11	3.66	6.36	3.62	1.79	2.64	38.19	3.55	87.89	3.22	1.36	2.96
ruficorne	2.00	2.50	1.50	5.00	3.00			3.00				2.00	2.00	2.00	4.00
manense	1.67	2.00	1.00		5.00	20.50	4.00	1.00	1.00	50.00	0.01	25.00	5.00	2.50	2.00
loutetense	2.02	1.68	1.24	5.00	4.60	5.68	3.37	1.58	1.63	34.00	2.26	31.53	3.53	1.77	2.81
colasbelcouri	2.00	1.67	1.00		5.00	5.33	4.00	1.00	1.75	50.00	1.68	21.67	5.00	2.00	2.14
futaense	1.67	2.50	1.00		5.00	17.50	4.33	1.00	1.50		0.35	100.00	4.50	2.00	3.00
ovazzae	1.89	2.25	1.44		4.38	7.50	3.50	1.25	2.00	50.00	0.25	112.86	2.83	2.00	2.88
berneri	2.88	2.86	1.38		4.43	6.00	3.86	1.17	2.33		1.52	64.17	3.17	1.50	2.86
lumbwanum	3.00		1.50		5.00		3.50	1.00	2.00		2.00	100.00			3.00
johannae	1.80	2.33	1.60		4.00	8.50	3.80	2.80		20.00	0.22	53.00	3.20	1.25	4.00
tridens	4.80	2.80	1.60	3.25	2.60		3.00	2.80			0.05	116.67	3.00	1.00	3.60
adersi	3.51	2.32	1.37	3.88	2.65	0.77	3.00	2.87		12.09	2.12	89.17	3.01	1.27	3.76
hargreavesi	2.72	2.40	1.61	4.09	3.28	3.25		3.04		0.53	3.75	45.42	3.16	1.71	3.87
bovis	4.00	3.50	1.00	4.00	3.00			3.00				100.00	2.50	1.00	4.00

Legends for Figures

Figure 1. (A) Results of cluster analysis of the full data-set (N = 527) with the six clusters used for further analyses denoted by numbered boxes. The "height" refers to the degree of similarity between groupings. (B) Distributions of species according to the six clusters (numbers 1-6) in Fig 1A for the full data-set. The heights of the bars represent the average incidence scores for assigning a species to a particular group.

Figure 2. (A) Ordination diagram for the environmental variables resulting from principal components analysis of the entire data-set. (B) Results of principal components analysis of the six species clusters derived from the complete data-set. The large dots are the centroids (means of the x and y values) and the ellipses are 80% confidence ellipses such that the centroids are 80% certain to lie within them.

Figure 3. (A) Ordination diagram for the environmental variables resulting from canonical correspondence analysis of the data on different members of the *Simulium damnosum* complex only (N = 442). (B) Ordination diagram for the onchocerciasis vectors based on canonical correspondence analysis of the different members of the *Simulium damnosum* complex only. soubr.Beffa = Beffa form of *S. soubrense*; sancti Djodji = Djodji form of *S. sanctipauli*.

Figure 4. (A). Ordination diagram for the environmental variables resulting from canonical correspondence analysis of the entire data set but excluding members of the *S. damnosum* complex and six rare species (*S. bovis, S. manense, S. colasbelcouri, S. lumbwanum, S. blacklocki, S. ruficorne*) for which data were insufficient to include in the analysis (N = 272). (B) Ordination diagram for the species resulting from canonical correspondence analysis of the entire data set but excluding members of the *S. damnosum* complex and six rare species (see under A above).







