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Chromosome inversions and ecological plasticity in the main African malaria mosquitoes

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Abstract

Chromosome inversions have fascinated the scientific community, mainly because of their role in the rapid adaption of different taxa to changing environments. However, the ecological traits linked to chromosome inversions have been poorly studied. Here, we investigated the roles played by 23 chromosome inversions in the adaptation of the four major African malaria mosquitoes to local environments in Africa. We studied their distribution patterns by using spatially explicit modeling and characterized the ecogeographical determinants of each inversion range. We then performed hierarchical clustering and constrained ordination analyses to assess the spatial and ecological similarities among inversions. Our results show that most inversions are environmentally structured, suggesting that they are actively involved in processes of local adaptation. Some inversions exhibited similar geographical patterns and ecological requirements among the four mosquito species, providing evidence for parallel evolution. Conversely, common inversion polymorphisms between sibling species displayed divergent ecological patterns, suggesting that they might have a different adaptive role in each species. These results are in agreement with the finding that chromosomal inversions play a role in *Anopheles* ecotypic adaptation. This study establishes a strong ecological basis for future genome-based analyses to elucidate the genetic mechanisms of local adaptation in these four mosquitoes.

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Introduction

Chromosome inversions have been considered by pioneering geneticists as the fingerprints of evolutionary processes in many different species (Krimbas and Powell 1992; Hoffmann and Rieseberg 2008). Over the last century, theoretical and experimental works have positioned chromosome inversions as key actors of chromosome architecture, local adaptation, sex evolution and speciation (Coghlan et al. 2005; Feuk et al. 2005; van Doorn and Kirkpatrick 2007; Bhutkar et al. 2008; Hoffmann and Rieseberg 2008; Sharakhova et al. 2011). Evidence that natural selection acts on chromosomal inversions has been gathered in many taxa, from mice (Lyon 2003), humans (Stefansson et al. 2005), *Drosophila* (Hoffmann et al. 2004) and monkeyflowers (Lowry and Willis 2010) to mosquitoes (Fouet et al. 2012; Ayala et al. 2013). Evolutionarily, chromosomal inversions are important because they reduce and, consequently, substantially alter recombination in heterozygotes. This characteristic together with their size (spanning hundreds or even thousands of genes) facilitate capturing favourable combinations of locally adapted alleles. Indeed, recent theoretical models (Kirkpatrick and Barton 2006; Manoukis et al. 2008; Schaeffer 2008) and ecological evidence support their role in local adaptation (Coluzzi et al. 1979b; Rodriguez-Trelles et al. 1996; Coluzzi et al. 2002; Balanya et al. 2003; Hoffmann et al. 2004; Ayala et al. 2011; Ayala et al. 2014). Although other processes like limited gene flow and genetic drift can also explain inversion clines (Dobzhansky and Wright 1943), the finding that chromosome inversion frequencies change at different collection sites was the first hint that they could contribute to local adaptation (Dobzhansky and Sturtevant 1938). Specifically, the relative frequencies of paracentric inversion polymorphisms (i.e., when a segment of a chromosome arm, which does not include the centromere, breaks and is reinserted in the reverse orientation) change in concert with the variation of different biotic and non-biotic factors. This feature allowed studying the natural selection forces that drive the evolution of inversions towards fixation (i.e., inversion frequency close to 100%) or stable polymorphism (i.e., inversion frequency stable across time) (Kirkpatrick and Barton 2006; Feder et al. 2011). However, the molecular basis of the adaptive role of these inversions remains barely known. Kirkpatrick and Kern (Kirkpatrick and Kern 2012) wrote that “the money is on the inversions” to suggest where to find genes involved in local adaptation. *Heliconius* butterflies offer a compelling example of how an inversion captures and protect several adaptive loci linked to mimicry (Joron et al. 2011). In *Drosophila*, the classical animal model for studying chromosome rearrangements, inversion frequencies and some genes within the inversion vary along environmental clines (Umina et al. 2005; Collinge et al. 2006; Kolaczowski et al. 2011; Kapun et al. 2016). In *Anopheles gambiae*, comparative genomic studies on carriers of the 2La inversion along an environmental aridity gradient led to the identification of divergent genes within the inversion (Cheng et al. 2012). These genes mainly encode signalling molecules, gustatory receptors or ion-channel genes. Interestingly, a remarkable correspondence between orthologues and their functions was observed between *Drosophila* and *An. gambiae* in comparable environmental clines, providing

evidences of potential parallel evolution (Kolaczowski et al. 2011; Cheng et al. 2012). Nevertheless, studies remain scarce. The main obstacle to phenotypic experiments lies in the difficulty of identifying the local adaptation drivers (ecological, behavioural, sexual, etc.) that model the inversion distribution in a population along a cline. For example, in sub-Saharan Africa, the *An. gambiae* 2La inversion frequency varies significantly along an aridity gradient, from complete absence in the humid rainforest of Central Africa to fixation in the arid savannas of West and East Africa (Coluzzi et al. 1985; Simard et al. 2009). In an attempt to validate the hypothesis that aridity tolerance is linked to this inversion, the thermal tolerance and desiccation resistance of 2La inversion carriers and non-carriers was tested in laboratory conditions. The results confirmed that 2La inversion carriers exhibit a higher desiccation and thermal resistance (Gray et al. 2009; Rocca et al. 2009; Fouet et al. 2012). Moreover, genome-wide expression analyses before and after exposure to thermal stress revealed that a large number of stress-linked genes were up-regulated in larvae carrying the inversion, compared with non-carriers (Cassone et al. 2011). However, only two of these up-regulated genes were in common with those found along the latitudinal cline (Cheng et al. 2012). Unfortunately, this is the only inversion where both phenotypic and molecular analyses have been carried out. Therefore, to understand the genetic mechanism of local adaptation, it is essential to identify the specific adaptive traits that shaped inversion frequencies in any species.

The availability of polytene chromosomes in *Anopheles* is a unique opportunity for studying chromosome evolution (Green and Hunt 1980; Coluzzi et al. 2002; Sharakhov et al. 2002; Sharakhova et al. 2006; Sharakhova et al. 2010a; Sharakhova et al. 2010b; Sharakhova et al. 2011; Sharakhova et al. 2013). In Africa, four *Anopheles* species have received most attention: *An. gambiae*, *An. coluzzii*, *An. arabiensis* and *An. funestus*. The first three species belong to the same complex, although *An. gambiae* and *An. coluzzii* have been only recently proposed as separate species (Coetzee et al. 2013). Until the development of molecular diagnostic tools in the 1990s, *An. arabiensis* was distinguished from the other sibling species in the *Anopheles gambiae* complex on the basis of post-zygotic barriers and five fixed inversions on chromosome X (Davidson 1964; Davidson and Hunt 1973; Coluzzi et al. 2002). These four species can thrive in a wide range of environments and live in sympatry in many sub-Saharan Africa regions (Ayala et al. 2009; Sinka et al. 2012). Their ecological plasticity and their anthropophilic habits (resting, feeding and breeding preferences) make them most efficient malaria vectors. Chromosomal rearrangements have been directly involved in *Anopheles* ecological and behavioural plasticity (Ayala et al. 2014). However, little is known about the specific environmental requirements of chromosomal inversions and how they promote local adaptation in these malaria mosquitoes.

In this study, we investigated, from a macro-ecological perspective, the role played by 23 chromosome inversion polymorphisms in the adaptation of the four major African malaria vectors (*Anopheles gambiae*, *An. coluzzii*, *An. arabiensis* and *An. funestus*) to local environments in West-Central Africa. We studied the distribution patterns of such inversion polymorphisms by using spatially explicit modelling, and characterized the ecogeographical determinants of each inversion range. We then performed hierarchical clustering and constrained ordination analyses to assess the spatial and ecological similarities of the 23 inversions. Our results show that most inversions are environmentally structured, suggesting

that they are involved in processes of local adaptation to a wide range of habitats. The eco-geographical distribution of some inversions exhibited similar patterns among the four mosquito species, providing evidence for parallel evolution. Conversely, inversion polymorphisms that were considered to be common between sibling species displayed divergent ecological patterns, suggesting that they might have a different adaptive role in each species. These results are in agreement with the view that chromosomal inversions play a role in *Anopheles* ecotypic adaptation, by making them more easily adaptable and, therefore, more ubiquitous and potent malaria vectors.

Materials and Methods

Chromosome inversion frequency data

The full karyotypes of 35,618 mosquitoes were used in this study, among which 16% of specimens represented unpublished data (Table 1, Table S1–S4). In total, 23 polymorphic inversions in *An. gambiae*, *An. coluzzii*, *An. arabiensis* and *An. funestus* were reviewed and analysed (see Table 1). We characterized the inversions in each species, even if they were considered as common, to avoid potential biases in the analysis. Mosquito sampling was carried out in 27 countries in Africa and Saudi Arabia and included the dry savannahs of Senegal and South Africa, the Highlands of Kenya and Madagascar, and the rainforests of Cameroon and Ivory Coast (Figure 1). This vast territory is highly heterogeneous, thus increasing the genetic diversity of mosquito species (Lehmann et al. 2003; Michel et al. 2005).

Mosquitoes were collected from 1979 to 2007. We obtained the frequencies of each inversion in each sampled locality (n=1,617 villages) and for each species (Table 1). Only localities providing the frequencies of all inversions included in this study were retained. Inversions in *Anopheles* are arbitrarily classified as standard or inverted. To avoid any potential bias, we included the frequencies of both inverted and standard forms in our analyses. The geographical coordinates of the original sampling localities were recorded using a hand-held GPS receiver. Unknown geographical coordinates of localities reported in the literature were validated through the National Geospatial-Intelligence Agency [<http://geonames.nga.mil>]. For modelling purposes, we used 5 km × 5 km grid squares as territorial units on the basis of the mosquito dispersal capabilities (Costantini et al. 1996; Ayala et al. 2013). Mosquito information was then assigned to the square in which the sampling village was located.

Karyotyping analysis was identical for the four mosquito species. Briefly, ovaries of half-gravid females were dissected and stored in Carnoy's fixative solution (three parts 100% ethanol: one part glacial acetic acid, by volume). They were subsequently prepared according to standard protocols to obtain the polytene chromosomes (della Torre 1997). Paracentric inversions were identified and scored according to their respective chromosome maps (Coluzzi et al. 2002; Sharakhov et al. 2004). Only the most common polymorphic inversions were used to ensure sufficient statistical power for the subsequent analyses (Table 1).

Ecogeographical predictors

On the basis of their potential predictive power and according to previous studies on mosquitoes (Ayala et al. 2009; Costantini et al. 2009; Simard et al. 2009; Sinka et al. 2010), we selected 11 ecogeographical predictors as potential drivers of the distribution patterns of each chromosomal inversion. Predictors fell within three categories: spatial (four predictors), land (four predictors) and climate (three predictors). All information was transferred to the territorial units by using zonal statistics tools.

Spatial variables—Spatial predictors were investigated to uncover geographical trends in the distributions associated with historical events, or species – inversions – population dynamics (Real et al. 2003). We investigated four spatial predictors: latitude, longitude, the product of latitude and longitude, and distance to the equator (measured as the absolute value of latitude) of each territorial unit, to account for the data spatial structure (Legendre 1998; Kennington et al. 2006).

Land variables—The importance of land use in explaining insect distribution patterns is well known (Acevedo et al. 2010; Hortal et al. 2010), including for *Anopheles* species (Manoukis et al. 2008; Simard et al. 2009). Here, we considered the Normalized Difference Vegetation Index (NDVI) and its seasonal variations (Acevedo et al. 2010). NDVI is a measure of the amount and vigour of vegetation on the land surface directly related to soil moisture, and has been successfully used to highlight changes in land cover (Nicholson et al. 1990; Nicholson and Farrar 1994). The NDVIs were derived from a monthly imaging dataset (<http://modis.gsfc.nasa.gov/data/dataproduct/mod13.php>) over a 13-year period, from 1998 to 2010, at a spatial resolution of ~1 km. Four different NDVI-derived variables were quantified on the basis of their importance in mosquito phenology (Clements 1999): yearly mean, yearly variation, wettest quarter mean (May to October), wettest quarter variation (May to October). Variations were quantified as the variation coefficients of yearly/quarterly means in the 13-year period. We selected the wettest quarter on the basis of mosquito population dynamics (Molineaux and Gramiccia 1980) and because mosquitoes are mainly captured during this season (Lindsay et al. 1998).

Climate variables—Three predictors were selected from the range of topoclimatic predictors associated with inversion distributions in Diptera: temperature, rainfall and elevation (Petrarca et al. 2000; Hoffmann et al. 2004; Ayala et al. 2011). In our study, the mean temperature and rainfall were quantified for the wettest quarter of the year, because this is the most important period for mosquito population dynamics (Moffett et al. 2007). Data on bioclimatic variables and altitude are available from the Worldclim project database (see (Hijmans et al. 2005) for details) at a spatial resolution of ~5 km.

Statistical Analysis

Ecogeographical models—Using an inductive approach we determined the macro-ecological requirements of the chromosomal inversions at the locations where they occurred (Corsi et al. 2000). We related the frequency of each inversion to the predictors using generalized linear models (GLM) with binomial distribution (number of inversions relative to the number of sampled mosquitoes per sampling locality, see below) and a logistic link

function (Hosmer et al. 1989). To obtain the most parsimonious model for each polymorphic rearrangement we used a forward-backward stepwise model-selection procedure. All steps were assessed to decrease the Akaike Information Criterion (AIC) (Akaike 1974). Models were built on a subset of randomly selected sampling localities for each inversion (80%) and then projected to the whole study area. The remaining sampling localities (20%) were used as independent data to evaluate the predictive performance of the models. To this aim, calibration plots (Pearce and Ferrier 2000) and Pearson's correlations were employed to statistically determine the relationship between the predicted probabilities and the observed frequencies (Zheng and Agresti 2000). Bins with $n < 15$ specimens were not included in this evaluation, because this is the minimum sample size required to estimate a frequency with acceptable accuracy (Jovani and Tella 2006). We restricted the ecogeographical models to West-Central Africa to avoid the inclusion of sparsely sampled areas in eastern and southern Africa (Figure 1) where the uncertainty of the model predictions could not be properly assessed (Heikkinen et al. 2012). Finally, potential confounding effects between spatial and environmental factors can be expected in models of species spatial distribution (Dormann 2007). To rule out this modelling bias, we investigated spatial autocorrelation in the residuals by estimating the Moran's I index (a measure of global spatial autocorrelation) for each model (Table S6).

Hierarchical clustering analysis—Clustering methods are a powerful tool for classifying similar objects in groups and are particularly useful for selecting species with similar biogeographical patterns (Kreft and Jetz 2010; Olivero et al. 2011) or with genetic/molecular similarities (Heard et al. 2005). We investigated similarities among the ecogeographical patterns of the polymorphic inversions in West-Central Africa. To carry out the hierarchical clustering analysis we randomly selected 1666 evaluation points in accordance with our sampling design (Legendre 1998), each of which was attributed with the probability of occurrence of each inversion according to the models' predictions.

To avoid mismatching due to inversion arrangements (standard versus inverted), we used the absolute value of the Pearson's correlation coefficient to estimate pairwise distances between the predicted probability of occurrence of each inversion. This index indicates similarity in shape between two (or more) profiles, and fits perfectly with the aim of clustering together inversions that respond similarly to ecogeographical gradients and species ranges. To maximize similarity within groups, we used the unweighted pair-group method with arithmetic averages (UPGMA) as agglomeration method. This produces less distortion from the original similarities than complete or single linkages and is consistently the best performing clustering algorithm for biogeographical classifications (Kreft and Jetz 2010). We then searched for chorotypes, defined as clusters of inversions the probabilities of occurrence of which are similarly distributed within the group and/or dissimilarly in relation to other inversions. Chorotype detection was performed as described by Olivero *et al.* (Olivero et al. 2011). A cluster was considered a quantitative chorotype if: (a) $I_H = 1$; or else (b) I_H was positive, had higher values than those of the other clusters including the distributions involved, and the proportion of "+" signs between the cluster and the most similar cluster was significantly lower than the proportion of "+" signs within the cluster (evaluated using a G-test of independence; (Sokal and Rohlf 1981)).

Canonical Ordination—The canonical correspondence analysis (CCA) is a statistical method for ordering species along canonical axes according to their (ecological) optima (Ter Braak 1987). We used this technique to relate the probability of inversion occurrence (on the evaluation points, Figure S2) with the set of ecological predictors (in this case, excluding spatial predictors to account only for ecological similarity). This allows for fairly easy ecological interpretation of inversion assemblages (Ayala et al. 2011). The statistical significance of the canonical axes and environmental predictors was assessed with permutation tests (10,000 times). To improve the CCA ecological comprehension, we plotted the major habitat boundaries according to (Olson et al. 2001). To determine the number of components to interpret from our CCA, we used the *broken stick model* (based on eigenvalues from random data) and the *Kaiser-Guttman rule* (based on the average value of eigenvalues) procedures (Jackson 1993).

All computational statistics were performed using the IBM-SPSS 19.0 software (IBM Corporation, New York, USA) and “R” v3.0.1 (R Development Core Team, <http://cran.r-project.org/>), with the addition of the “amap” (Lucas 2011), “vegan” (Oksanen et al. 2011), and ggvegan (<https://github.com/gavinsimpson/ggvegan>) libraries.

Results

Data on the inversions and *Anopheles* species retrieved from the literature and from our unpublished work and used for this study are summarized in Table 1, Figure 1 and Supplementary Material (Tables S1–S4).

Biogeographical patterns of chromosomal inversions

Using a generalized linear model approach, the frequency of each inversion was correlated with several ecogeographical predictors to calculate the inversion ecogeographical favourability throughout the study area (West-Central Africa, highlighted in Figure 1). Table 2 summarizes the predictors that were found to drive inversion frequency variations and the explained deviance (goodness-of-fit) for each model (see also Table S5). Despite the variability between models, latitude and precipitation were repeatedly the most important predictors of the inversion distribution patterns in the four mosquito species. Conversely, distance to the equator was the least significant variable in the final models. Overall, the models’ predictive performance was high. Indeed, predictions were highly correlated with the observed values in the datasets used for the independent validations (Table S5 and Figure S1). However in some cases (for instance, 2Rt in *An. funestus*), model performance was adequate according to the calibration plots, but could not be statistically evaluated due to insufficient independent data. Moreover, the inversion models did not exhibit a significant spatial autocorrelation bias as indicated by the Moran’s I value close to zero. This means that the residuals of the models for each inversion show a random spatial pattern (Table S6).

Next, the statistical models were represented in geographical space to obtain cartographic models of the expected probability of occurrence of a given inversion, on the basis of the local ecogeographical conditions (Figure 2). The predicted frequencies were subject to the local occurrence of each species and, accordingly, these maps should be interpreted as the predicted inversion frequency of the target species in a given locality.

Hierarchical clustering analysis

The predicted inversion frequencies throughout the study area were then correlated with each other to find similar eco-geographical patterns. Inversions that showed a significantly similar environmental distribution were grouped in chorotypes. We identified three significant inversion chorotypes in the study area (Figure 3). The first chorotype contained the 2Rd inversion in *An. gambiae* and *An. coluzzii*; the second contained the inversions 2Ru in *An. coluzzii* and 2Rc in *An. arabiensis*; the third comprised the inversions 2La, 2Rb and 2Rc in *An. gambiae* and *An. coluzzii*, and 2Rt, 3Ra, 3Rb and 3La in *An. funestus*. Altogether, the dendrogram highlighted important features concerning the inversion spatial patterns. Overall, chromosome inversions in *An. gambiae* and *An. coluzzii* showed a significant common spatial pattern. Moreover, chromosome inversions in *An. funestus*, *An. gambiae* and *An. coluzzii* were associated with similar environmental clines, revealing the presence of significantly correlated eco-geographical patterns. Finally, some inversions exhibited a very specific spatial pattern, such as 2Rj in *An. gambiae*.

Environmental drivers of inversions

The maximum correlations between the set of land and climate predictors (spatial predictors were excluded to account only for ecological similarity) and the inversion frequencies were then determined using a canonical correspondence analysis (CCA) method. This statistical approach allowed us to define the ecological optimum of each inversion and to represent the overall contribution of the tested ecological predictors to the chromosomal polymorphism frequency among species (Figure 4). The seven CCA axes were all statistically significant (ANOVA, $p < 0.001$). In accordance with the *broken stick model* and the *Kaiser–Guttman rule*, only the first two CCA axes, which accounted for almost 80% of the total variance in the inversion dataset, were retained (Jackson 1993). The first axis explained 65.61% (ANOVA, $F=1219$, $p < 0.001$) and the second 13.67% of the total variance (ANOVA $F= 254$, $p < 0.001$). The other five CCA axes represent 10.3%, 6.1%, 2.6, 0.9% and 0.7%, respectively. The significance of all tested ecological predictors was assessed by permutation tests (ANOVA, $p < 0.001$). The first axis could be ascribed to an aridity gradient (Figure S2A) and structured chromosomal inversions according to their tolerance to aridity, relative to the habitat boundaries. The second CCA axis represented an environmental gradient mostly influenced by vegetation productivity patterns and elevation (Figure S2B). Here, inversions with an ecological optimum in the savanna biome showed the highest frequency variations.

Discussion

This study elucidates the ecological basis of the spatially structured distribution of 23 chromosomal inversions in the four major malaria vectors in Africa. Three major outcomes emerge: i) each chromosome inversion shows a specific and unique eco-geographical pattern, possibly explaining how mosquito species can extend their habitat ranges; ii) some of the inversion polymorphisms in the four mosquito species exhibit common ecogeographical patterns, suggesting that they are involved in the adaptation to similar local pressures; and iii) some inversion polymorphisms, presumably shared by sibling species, exhibit contrasting ecological patterns, suggesting a different adaptive role. Altogether, our

results establish a strong ecological basis for future genomic studies to elucidate the genetic bases of inversion contribution to local adaptation in these four malaria vectors.

The role of inversions in environmental adaptation

Chromosome inversions have often been considered as major drivers in the geographical expansion of many species (Krimbas and Powell 1992), including *An. gambiae*, *An. coluzzii*, *An. arabiensis* and *An. funestus*. Our study shows that the frequency of most of the studied inversions could be explained by the tested ecogeographical gradients (Table 2). An important novelty of our study is that we could compare and pull together not only different inversions from the same species but among species. This fact allow to expand our knowledge about inversions, permitting to interpret patterns between alternative arrangements and inversions. Only for two inversions (2Rd and 2Ru in *An. coluzzii*), the models could not explain their ecogeographical variation (only 17% and 26%, respectively). On the other hand, each main ecogeographical gradient had a different weight on each inversion (Table 2, Figure 2). The CCA identified the major environmental variables that influenced the distribution of each inversion and provided evidence that chromosomal rearrangements are linked to different environmental gradients (Figure 4). To date, chromosomal rearrangements have been exclusively linked to specific ecogeographical predictors, such as latitude (Hoffmann and Rieseberg 2008), but our analyses provide a broader picture of the habitats where these inversions play an important role for local adaptation. For instance, inversion 2Rd in *An. gambiae* and *An. coluzzii* plays a robust role in the adaptation to elevation. It is not the first time that an inversion has been associated with altitudinal clines (Collinge et al. 2006). Unfortunately, previous studies on the risk of malaria transmission at high altitude in Africa did not characterize the chromosome polymorphisms of the adapted populations (Tchuinkam et al. 2010). However, the major ecological challenge for these four mosquito species is the adaptation to humid conditions. Many of the inversions in *An. coluzzii*, *An. gambiae* and *An. funestus* are thought to have a savanna origin (Ayala and Coluzzi 2005; Ayala et al. 2009; Kamali et al. 2014) and have been linked to local acclimatization to rainforest habitat (Figure 4, Figure 3 – Chorotype 3). At least for *An. coluzzii* and *An. gambiae*, breakpoints analyses have confirmed that species with the inverted 2La karyotype (arid savanna) originated from the introgression from the arid adapted species *An. arabiensis* (White et al. 2009b), predate species with the standard 2La karyotype (forest) (Sharakhov et al. 2006; Fontaine et al. 2015). This is in agreement with the crucial role played by these rearrangements in malaria expansion to rainforest habitats (Annan et al. 2007). Similarly, inversion 3Ra in *An. arabiensis* appears to be associated with the adaption to the mosaic forest-savannah biome. Despite the many evidences, we cannot conclude that all the observed inversion distribution patterns have an exclusive ecological basis. Indeed, in *Anopheles*, like in other organisms, chromosomal inversions have been correlated also with non eco-geographical factors (Hoffmann and Rieseberg 2008), such as resting behavior (Coluzzi et al. 1977; Bryan et al. 1987; Costantini et al. 1999), host preference (Coluzzi et al. 1979b; Petrarca and Beier 1992), insecticide resistance (Brooke et al. 2002) and *Plasmodium* infection (Petrarca and Beier 1992). Therefore, although ecological forces seem to be the major drivers of inversion distribution (Table 2), other behavioural and/or physiological traits could have a non-negligible effect on their frequency patterns. Another plausible explanation is that demographic forces are

responsible for the clinal patterns observed. However, several factors lead us to think that clinal variation is related to the action of natural forces (Endler 1977). Firstly, many inversions exhibit parallel clines across the continent (Table S1–S4). For instance, in *An. gambiae*, Simard *et al.*, (Simard et al. 2009) and Coluzzi *et al.*, (Coluzzi et al. 1979a) found identical clinal patterns in Cameroon and Nigeria, respectively. Secondly, another important factor is migration. Strong gene flow between populations would quickly make disappear any inversion gradient in absence of selection. According to neutral genetic markers, *An. gambiae/An. coluzzii* and *An. funestus* exhibit parallel genetic structure across Africa coherent with a common expansion (Lehmann et al. 2003; Michel et al. 2005). Both continental studies highlight the strong gene flow between natural populations of these mosquitoes, denoting weak population structure. On the other hand, scattered countrywide studies in *An. arabiensis* revealed as well important gene flow between populations of this vector (Donnelly et al. 1999; Simard et al. 2000). Finally, the large effective population size in all these mosquitoes reinforce the assumptions of one panmictic population through the continent with limited gene flow barriers (Donnelly et al. 1999; Lehmann et al. 2003; Michel et al. 2005). Overall, these evidences are the strongest indications that environmental selection forces, and not demographic forces, are responsible for the observed environmental gradients of most of our inversions.

A key feature in the evolution of chromosome polymorphisms is the additive (i.e., the sum) or epistatic (i.e., the interaction) effect of inversions (Schaeffer et al. 2003). Our hierarchical and ordination analyses revealed common environmental patterns across inversions within species (i.e., 2Rb and 2La in *An. gambiae* and *An. coluzzii*; 3Ra, 3Rb and 3La in *An. funestus*). The existence of ecotypes— chromosomal inversion forms – within *An. gambiae* (Coluzzi 1982) and *An. funestus* (Costantini et al. 1999) has been determined on the basis of stable combinations of polymorphic rearrangements. These combinations exhibit strong linkage disequilibrium (LD) and are persistent in different habitats (Costantini et al. 2009; Simard et al. 2009; Ayala et al. 2011). Theoretical models (Schaeffer 2008; Burger and Akerman 2011) and empirical evidence from *Drosophila* (Schaeffer et al. 2003) and *Heliconius* (Joron et al. 2011) support the hypothesis that epistatic effects among inversions may maintain LD in a heterogeneous environment. However, epistatic interactions between inversions will certainly interfere with the model assumption that inversion frequencies depend on ecogeographical variables, but not on the frequency of other inversions. Unfortunately, our dataset did not provide information on the entire karyotype to enable us to build models with different sets of inversions and therefore to investigate the recombinant fitness for each biome, as already done in *Heliconius* (Le Poul et al. 2014). On the other hand, the high performance of most of our models reveals that if an epistatic effect exists, it should be less important than the environmental conditions. Nevertheless, it would be important to investigate whether and how these inversions in *An. gambiae*, *An. coluzzii*, *An. arabiensis* or *An. funestus* work together to implement fitness in different habitats.

Same patterns, same causes: parallel chromosome evolution in Anopheles

The hierarchical clustering analysis showed that chorotype 3 included inversions from three different species: *An. gambiae*, *An. coluzzii* and *An. funestus*. This cluster can be interpreted as a biogeographical pattern with strong internal similarity. Moreover, the CCA confirmed

that these inversions have similar ecological gradients (Figure 4). These three species live in sympatry throughout much of their geographical range in Africa (Gillies and Coetzee 1987). In *An. gambiae* and *An. coluzzii*, the origin of the inversions 2La, 2Rb and 2Rc predates their divergence (White et al. 2009a; Lawniczak et al. 2010; Fontaine et al. 2015); therefore, we could expect similar patterns among their shared ancestral polymorphisms. On the other hand, *An. funestus* and *An. gambiae* diverged ~35 Mya (Krzywinski and Besansky 2003; Neafsey et al. 2015). Therefore, any common spatial pattern could be attributed to similar environmental pressures (Ayala et al. 2009; Sinka et al. 2010). One hypothesis is that these inversions captured similar sets of genes. Several authors found nearly perfect synteny preservation between arms when analysing whole chromosome arms (Sharakhov et al. 2002; Neafsey et al. 2015). This means that if inversions captured similar sets of genes, they should be on homologous arms. In a key paper, Sharakhova *et al.* (Sharakhova et al. 2011) investigated the non-random distribution of genes along homologous arms of malaria vectors. They found significant non-random gene combinations on *An. gambiae* 2Rb and on *An. funestus* 2Rh and 2Rd. Here, we found that inversions 2Rb and 2Rh are both strongly associated with rainforest habitat (Figure 2 and Figure 4). Unfortunately, the 2Rd inversion frequencies in *An. funestus* were too low and uneven to be included in the analysis. However, this rearrangement has been consistently associated with forest areas in Cameroon (Cohuet et al. 2005). Therefore, environmental adaptation to very humid conditions could have preserved specific gene combinations within these three inversions in the two species. Moreover, Sharakhova *et al.* (Sharakhova et al. 2011) found an almost significant ($p = 0.07$) association between 2La in *A. gambiae* and 3Rb in *An. funestus*. The very similar patterns of the main environmental gradients for 2La and 3Rb in the study area (Figure 2 and Figure 4) indicate their common role in adaptation to an aridity cline. However, these two inversions, together with 3Ra in *An. funestus*, show limited co-linearity, leaving strong doubts that they captured comparable large blocks of genes (Sharakhov et al. 2002). Nevertheless, we cannot exclude the possibility that smaller segments and/or few genes are shared within the inversions 2La and 3Ra/3Rb. Theoretically, the inversion would only need to capture one locally adapted locus to increase its frequency (Kirkpatrick and Barton 2006). The availability of the *An. gambiae*, *An. coluzzii* and *An. funestus* genomes will make possible to identify genes common to these species (Neafsey et al. 2015). In conclusion, despite the fast evolution of autosomal arms and gene shuffling, natural selection may preserve specific gene combinations within polymorphic inversions, particularly in distant species, such as *An. gambiae*-*An. coluzzii* and *An. funestus*, that are subject to similar environmental pressures. Thus, the most plausible scenario is that inversions in these species occurred independently (i.e., parallel evolution) and captured locally adapted genes in homologous arms. This knowledge may be very useful for identifying the gene signatures of local adaptation in these malaria vectors (Neafsey et al. 2015).

Common, but ecologically divergent inversions

Our analyses revealed some degree of ecogeographical divergence among hypothetically common inversions in species of the *An. gambiae* complex. Although *Anopheles arabiensis*, *An. gambiae* and *An. coluzzii* are characterized by different fixed chromosomal rearrangements on the X chromosome, they potentially share three chromosome inversions: 2La, which is fixed in *An. arabiensis* and polymorphic in *An. gambiae* and *An. coluzzii*, as

well as 2Rb and 2Rc, which are polymorphic in all three species (Table 1, (Coluzzi et al. 2002)). Our predictive models, dendrogram and CCA revealed very different ecogeographical distribution patterns for 2Rb and 2Rc between *An. gambiae*/*An. coluzzii* and *An. arabiensis* (Figure 2, Figure 3, and 4; Table 2). The most plausible explanation is that the ecological and local adaptation patterns (and thus, captured genes) of these inversions are not the same or have rapidly evolved after speciation among mosquito species. Originally, they were characterized by the presence of a banding pattern (Coluzzi et al. 2002) and therefore, a potential mismatch in breakpoint recognitions is highly possible. Moreover, there is extensive evidence for breakpoint reuse and inversion recycling in other Diptera (Ranz et al. 2007). Indeed, differences in gene expression patterns might be expected if breakpoints are different between species (Puig et al. 2004). No molecular information is available for 2Rc. On the other hand, 2Rb breakpoints have been molecularly characterized in recent years. Lobo *et al.* (Lobo et al. 2010) argued that the homozygous 2Rb inverted form has a single common origin in all three sibling species, while the homozygous standard form (2Rb+) may have arisen twice through breakpoint reuse. In our models, the inversion 2Rb+ shows contrasting ecological patterns in *An. gambiae*-*An. coluzzii* and *An. arabiensis*. On the basis of molecular data, White *et al.* (White et al. 2009b) hypothesized that introgression from *An. arabiensis* brought the 2Rb arrangement into *An. gambiae*, while introgression in the opposite direction introduced 2Rb+ into *An. arabiensis*. The occurrence of the last introgression was supported by Fontaine et al., (Fontaine et al. 2015), who confirmed the ancestral status of the standard 2Rb+ form. Recent statistical models have reinforced the hypothesis that inversions can act as “*cassettes of genes that can accelerate adaptation by crossing species boundaries*” (Kirkpatrick and Barrett 2015). On the basis of our results, we hypothesize that a new 2Rb+ inversion might have appeared through breakpoint reuse in *An. gambiae* after speciation of *An. arabiensis* (Lobo et al. 2010). This new 2Rb+ could have brought new mutations that favoured (together with other inversions, see above) the colonization of rainforest habitats by *An. gambiae*/*An. coluzzii* (Sharakhov et al. 2006). New molecular and phenotypic analyses of the inversion 2Rb+ across species and along their geographical range might confirm this last hypothesis.

Conclusions and implications for malaria control

To date, vector control is the main approach for reducing malaria transmission (Enayati and Hemingway 2010). However, environmental and behavioural diversity within vector populations constitute serious challenges to the efficacy of any malaria control strategy (Ferguson et al. 2010). Much effort is now required to understand the adaptive mechanisms of vector species for improving such control strategies (Boete 2005; Windbichler et al. 2011).

Our study reveals the potential roles played by inversion polymorphisms in the ecological success of the four major malaria mosquitoes. The importance of chromosome inversions in adaptation is attested by the strong, significant correlations between their frequencies and ecogeographical predictors, and by the strong, spatially-structured patterns identified in the study area. Therefore, inversion polymorphisms may have enabled considerable geographical expansion of these mosquitoes, with major consequences for malaria parasite transmission. Moreover, the extensive reshuffling of gene orders confirms, to some extent,

that this type of chromosomal rearrangement is very common and a frequent local adaptation mechanism in *Anopheles* (Kamali et al. 2012). On the other hand, the converging evolution of inversions in *An. gambiae*, *An. coluzzii* and *An. funestus* may provide a suitable basis for comparative studies to identify the genes responsible for environmental adaptation. Understanding the genetic mechanisms that enable these vectors to extend their geographical range will have a profound impact on malaria epidemiology. The complete genome of *An. funestus* (Besansky 2008; Neafsey et al. 2015) will now provide considerable opportunities for comparative genomic studies with *An. gambiae* and will help elucidating the common mechanisms involved in ecological adaptation. Finally, besides the adaptive role of inversions, the recognition of ecological divergence within populations of insect vectors has a direct impact on the efficacy of any vector-borne disease control strategy (Ferguson et al. 2010). Ecological and behavioural diversification within species of the *An. gambiae* complex have expanded malaria transmission spatially and temporally, compromising the efficacy of malaria control efforts (Molineaux and Gramiccia 1980). Surveillance of genetic and ecological divergence within vector populations will ultimately lead to more effective malaria vector control interventions.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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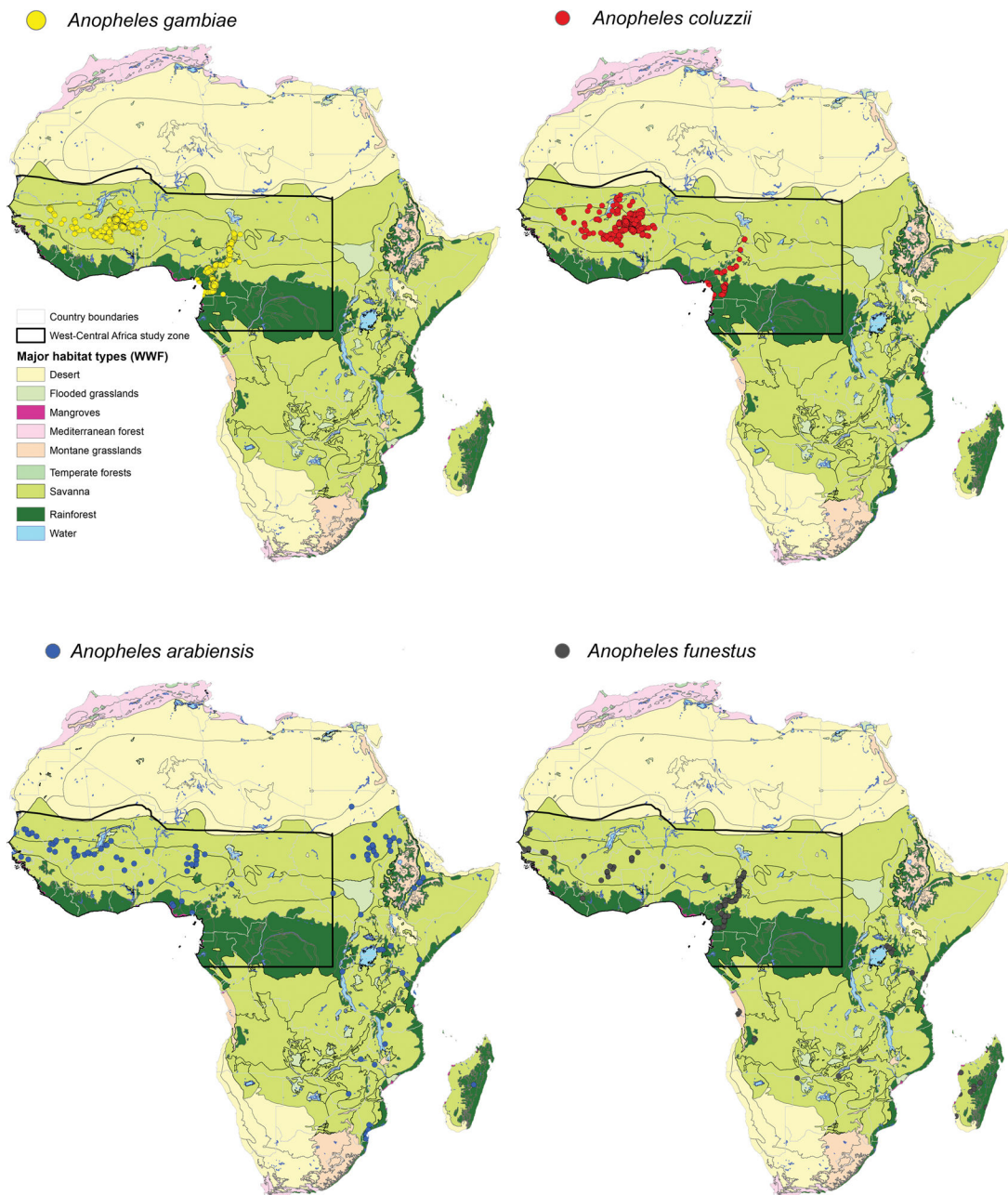


Figure 1. Sampling villages

Map of the main African habitat types (Olson et al. 2001) showing the distribution of the villages where the four *Anopheles* species were sampled: *Anopheles gambiae* (yellow dots), *An. coluzzii* (red dots), *An. arabiensis* (blue dots) and *An. funestus* (grey dots). A detailed representation of the West-Central Africa area that was selected to plot the predicted maps and carry out the similarity analyses is shown. Species' points out of the study area were used for calibration purposes (see Materials and Methods)

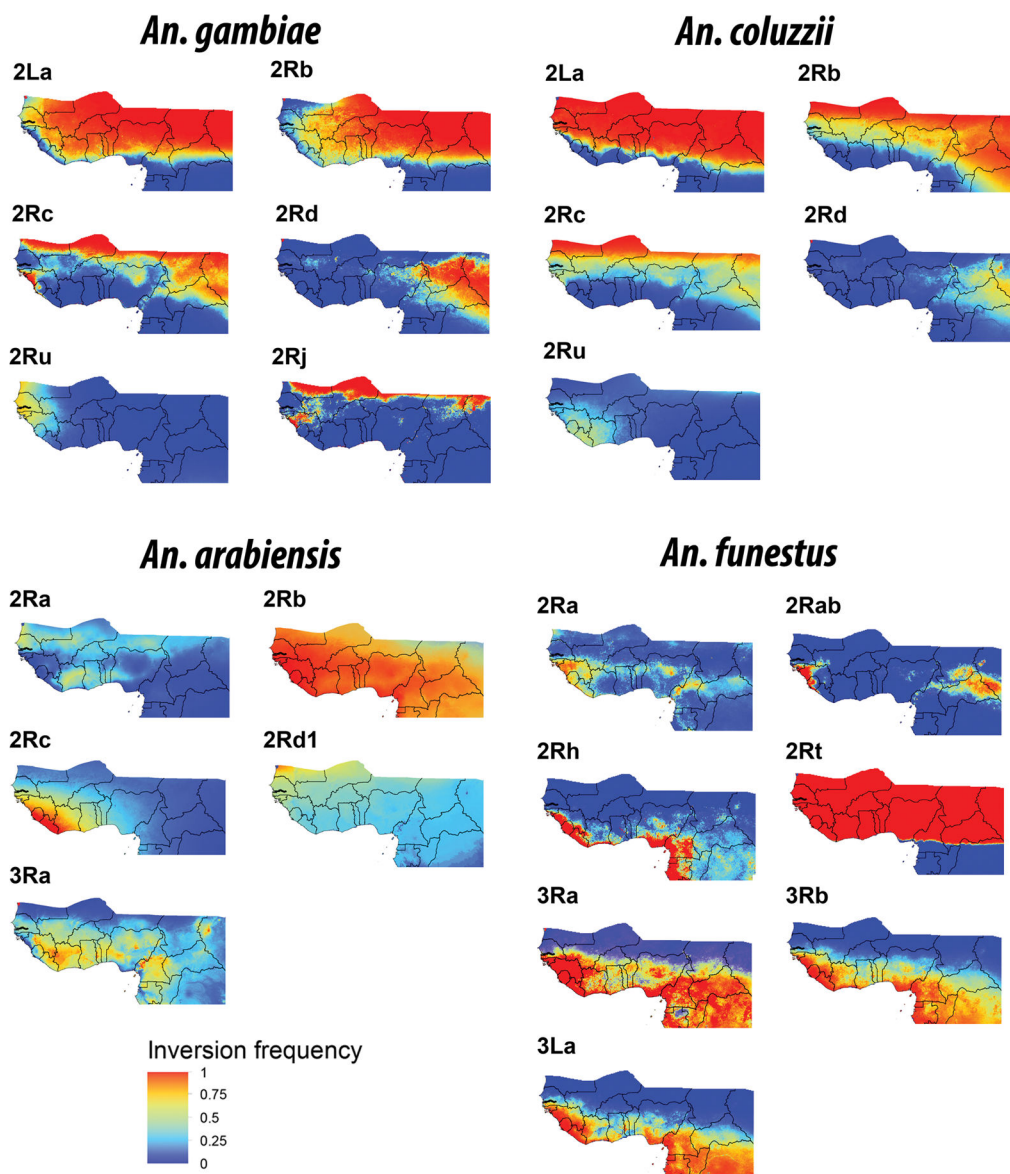


Figure 2. Maps of the predicted inversion frequencies for *Anopheles gambiae*, *An. coluzzii*, *An. arabiensis* and *An. funestus*. Predicted inversion distributions were passively plotted in the West-Central Africa study area. Blue represents a probability of 100% for the standard inversion, red represents a probability of 100% for the inverted inversion form, according to the literature data for each species (Green and Hunt 1980; Coluzzi et al. 2002). To improve their representativity, probabilities were reclassified in four classes: 0.00–0.25; 0.25–0.50; 0.50–0.75; 0.75–1.0.

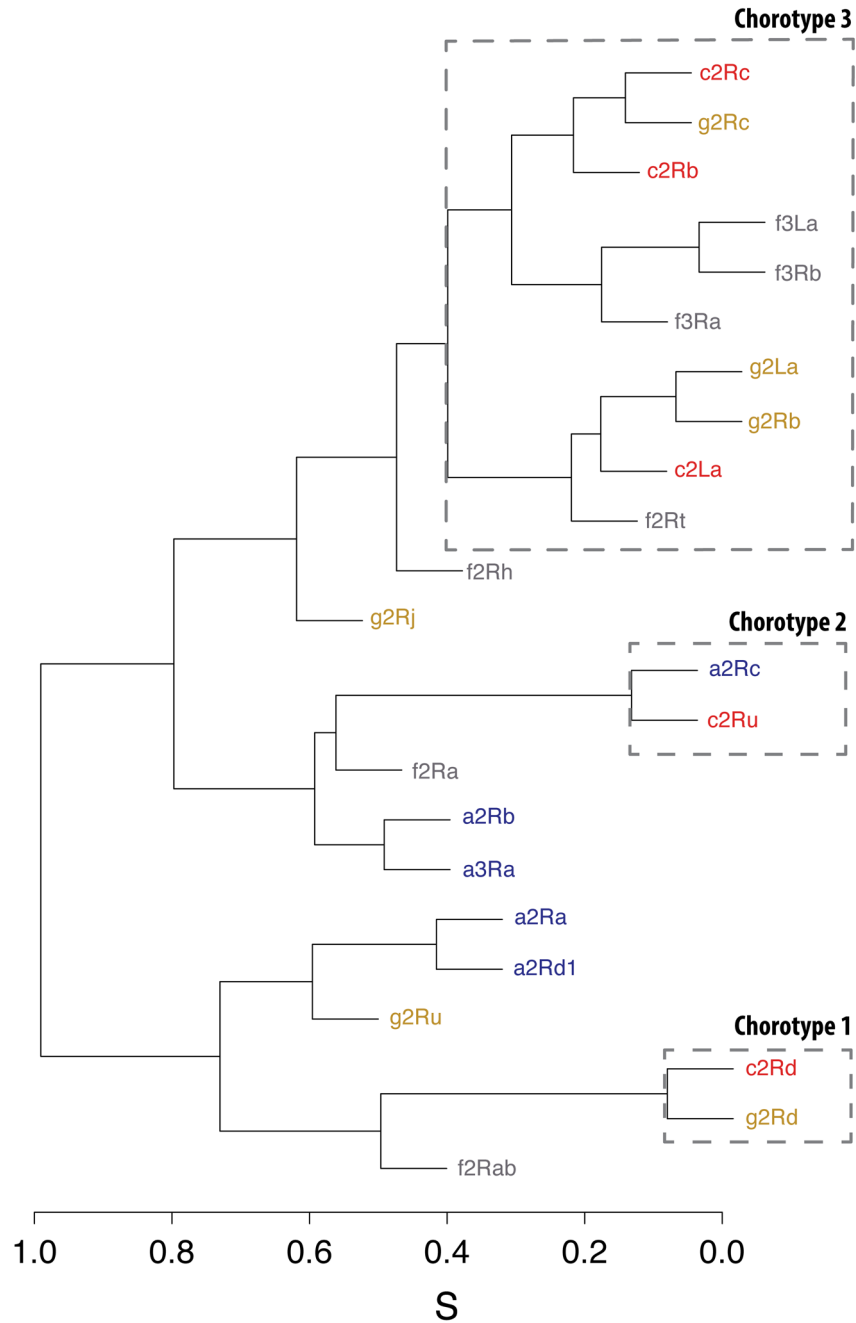


Figure 3. Dendrogram of the predicted inversion frequency distribution in West-Central Africa showing similar environmental patterns in *Anopheles* species
 Inversions included in the three chorotype clusters (Chorotype 1, Chorotype 2 and Chorotype 3) are enclosed in squares. *Anopheles* spp. are coded by letters and colours: g (yellow): *An. gambiae*; c (red): *An. coluzzii*; a (blue): *An. arabiensis*; f (grey): *An. funestus*. In x-axis, S corresponds to the Baroni-Urbani and Buser similarity index (Olivero et al. 2011). Chorotype 1 and Chorotype 2 have both a IH =1, while Chorotype 3 has a IH = 0.822 (G-test, p-value<0.001) (Olivero et al. 2011).

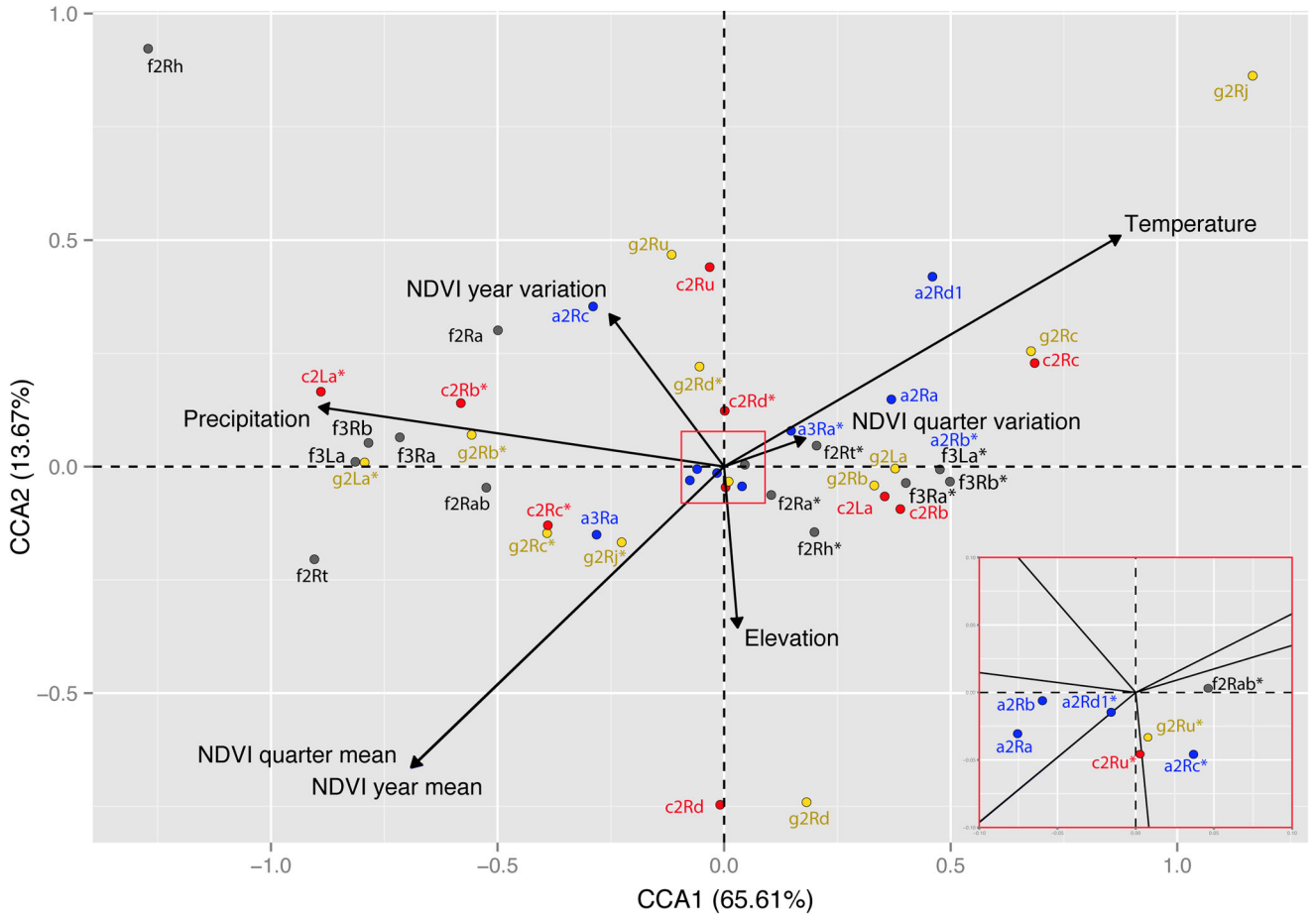


Figure 4. Canonical correspondence analysis of the inversion ecological distribution throughout West-Central Africa to highlight local adaptation patterns among chromosome inversions and mosquito species

CCA diagram showing the ordination of the chromosomal inversions (standard and inverted; red crosses indicate their ecological optima) for each species along the first two canonical axes (CCA1 and 2) that, together, explain ~80% of variance. *Anopheles* spp. are coded by letters and colours: g (yellow): *An. gambiae*; c (red): *An. coluzzii*; a (blue): *An. arabiensis*; f (grey): *An. funestus*. Asterisks represent the standard form of each inversion. Ecological predictors are passively plotted on the graph: elevation, temperature (mean temperature of the wettest quarter of the year), precipitations (mean precipitations of the wettest quarter of the year) and NDVI variables (yearly mean, yearly variation, quarterly variation and wettest quarter variation for the period included in the study). An inset of the main Figure is provided in the bottom-right corner for clarity purposes, showing the inversions encompassed in the red square.

Table 1

Summary of the chromosome inversions in the four malaria vectors *Anopheles gambiae*, *An. coluzzii*, *An. arabiensis* and *An. funestus*.

	Specimens	Villages	Inversions	Inversions names
<i>An. gambiae</i>	7949	799	6	2La; 2Rb, 2Rc, 2Rd, 2Ru, 2Rj
<i>An. coluzzii</i>	5000	528	5	2La; 2Rb, 2Rc, 2Rd, 2Ru
<i>An. arabiensis</i>	12836	125	5	2Ra, 2Rb, 2Rc, 2Rd1; 3Ra
<i>An. funestus</i>	9833	165	7	2Ra, 2Rh, 2Rab, 2Rt; 3Ra, 3Rb; 3La

Table 2

Summary of the models for each polymorphic inversion

Ecogeographical predictors selected in the final models for each inversion and *Anopheles* species. Species are coded by colour: *An. gambiae* (yellow), *An. coluzzii* (red), *An. arabiensis* (blue) and *An. funestus* (grey). Solid cells represent selected variables for each model according to AIC and open cells non-selected ones. The percentage of models in which each predictor was selected is indicated in the last row of each species. The last column represents the total explained deviance of the final model for each inversion. Lat: latitude; Long: longitude; Lat × Long: the product of latitude and longitude; abs(Lat): distance to equator expressed as absolute latitude value; Temp: mean temperature of the wettest quarter of the year; Precip: mean precipitation of the wettest quarter of the year; NDVI (normalized difference vegetation index) is numerical indicator that uses remote sensing measurements of live green vegetation (NDVI yearly mean, yearly variation, quarterly mean and variation and wettest quarter mean and variation for the period included in the study).

Inversions	Lat	Long	LatxLong	abs(Lat)	Elevation	Temp	Precip	NDVI year mean	NDVI year variation	NDVI quarter mean	NDVI quarter variation	Total
2La	■	■	■	□	■	□	■	■	■	■	■	88%
2Rb	■	■	■	□	■	□	■	■	■	■	■	81%
2Rc	■	■	■	□	■	□	■	■	■	■	■	47%
2Rd	■	■	■	□	■	□	■	■	■	■	■	72%
2Ru	■	■	■	□	■	□	■	■	■	■	■	49%
2Rj	■	■	■	□	■	□	■	■	■	■	■	70%
Total <i>An. gambiae</i>	100%	100%	83%	0%	83%	33%	100%	83%	67%	83%	67%	
2La	■	■	■	□	■	□	■	■	■	■	■	87%
2Rb	■	■	■	□	■	□	■	■	■	■	■	56%
2Rc	■	■	■	□	■	□	■	■	■	■	■	63%
2Rd	■	■	■	□	■	□	■	■	■	■	■	17%
2Ru	■	■	■	□	■	□	■	■	■	■	■	26%
Total <i>An. coluzzii</i>	100%	80%	80%	0%	60%	40%	80%	60%	60%	40%	80%	
2Ra	■	■	■	□	■	□	■	■	■	■	■	65%
2Rb	■	■	■	□	■	□	■	■	■	■	■	90%
2Rc	■	■	■	□	■	□	■	■	■	■	■	86%
2Rd	■	■	■	□	■	□	■	■	■	■	■	55%
3Ra	■	■	■	□	■	□	■	■	■	■	■	89%
Total <i>An. arabiensis</i>	80%	80%	80%	60%	60%	40%	60%	20%	40%	20%	60%	
2Ra	■	■	■	□	■	□	■	■	■	■	■	60%
2Rh	■	■	■	□	■	□	■	■	■	■	■	80%
2Rab	■	■	■	□	■	□	■	■	■	■	■	85%
2Rt	■	■	■	□	■	□	■	■	■	■	■	89%
3Ra	■	■	■	□	■	□	■	■	■	■	■	47%
3Rb	■	■	■	□	■	□	■	■	■	■	■	70%
3La	■	■	■	□	■	□	■	■	■	■	■	80%
Total <i>An. funestus</i>	100%	86%	71%	57%	71%	86%	100%	86%	71%	71%	100%	
Total	96%	87%	83%	30%	70%	52%	87%	65%	61%	57%	78%	