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Low Genetic Diversity in *Melanaphis sacchari* Aphid Populations at the Worldwide Scale

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Abstract

Numerous studies have examined the genetic diversity and genetic structure of invading species, with contrasting results concerning the relative roles of genetic diversity and phenotypic plasticity in the success of introduced populations. Increasing evidence shows that asexual lineages of aphids are able to occupy a wide geographical and ecological range of habitats despite low genetic diversity. The anholocyclic aphid *Melanaphis sacchari* is a pest of sugarcane and sorghum which originated in the old world, was introduced into the Americas, and is now distributed worldwide. Our purpose was to assess the genetic diversity and structuring of populations of this species according to host and locality. We used 10 microsatellite markers to genotype 1333 individuals (57 samples, 42 localities, 15 countries) collected mainly on sugarcane or sorghum. Five multilocus lineages (MLL) were defined, grouping multilocus genotypes (MLG) differing by only a few mutations or scoring errors. Analysis of a 658 bp sequence of mitochondrial COI gene on 96 individuals revealed five haplotypes, with a mean divergence of only 0.19%. The distribution of MLL appeared to be strongly influenced by geography but not by host plant. Each of the five MLL grouped individuals from (A) Africa, (B) Australia, (C) South America, the Caribbean and the Indian Ocean including East Africa, (D) USA, and (E) China. The MLL A and C, with a wide geographic distribution, matched the definition of superclone. Among aphids, *M. sacchari* has one of the lowest known rates of genetic diversity for such a wide geographical distribution.

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Introduction

Range expansion of exotic species can result from either evolutionary adaptation or generalism and plasticity often associated with a change in niche [1]. Genetic diversity is required for evolutionary adaptation, but a reduction in genetic diversity in invasive populations compared to populations in their native range is expected and often observed [2–5]. Organisms with clonal reproduction may exhibit reduced genetic diversity within populations, as better-adapted clonal genotypes expand and dominate available resources [6]. In aphids, the concept of “superclones” emerged [7] when a few asexual genotypes of the same species were able to colonize a wide geographical or ecological range of habitats [8–11]. The capacity of these populations to adapt to different conditions could be the result of a preadaptation capacity for phenotypic plasticity rather than local selection acting on genetic diversity [12,13]. What is more, this capacity may be enhanced by their high rate of reproduction and population expansion [14]. For these reasons, clonal aphids are good models to assess the ability of asexual lineages to show rapid and widespread adaptive changes to ecological conditions [15].

The old world genus *Melanaphis* van der Groot 1917 comprises around 20 species mainly associated with Poaceae, most of which originate from East Asia [16]. The sugarcane aphid *Melanaphis sacchari* (Zehntner, 1897) (Homoptera, Aphididae), which is considered to be mainly anholocyclic, is present in America, Australia, Asia and Africa. *M. sacchari* is known to be invasive in continental US [17,18] and in Central and South America [16]. The host range of this species is restricted to Poaceae [2,19]. Blackman et al. [20] hypothesised that *Melanaphis* individuals originating from sorghum or sugarcane were distinct taxa, referred to as *M. sorghi* and *M. sacchari* respectively, even though their host plant preference was not absolute. In their catalogue, Remaudi ere and Remaudi ere [21] considered *M. sorghi* to be a synonym for *M. sacchari*, but both forms were still listed as separate taxa by Blackman and Eastop [22], and it is still not clear whether *M. sacchari* constitutes a single species or a complex of sibling taxa.

In any case, *M. sacchari* is a major pest of sorghum and sugarcane. On sugarcane, it is considered to be the most common and most efficient vector of the *Sugarcane yellow leaf virus* (ScYLV), which causes yellow leaf disease [23,24], a disease of worldwide economic importance [25–27]. The aphid is also a major pest of sorghum, causing direct damage (sap feeding) and

indirect damage (sooty mould) [19]. Varietal resistance against *M. sacchari* is one of the main control tactics suggested both for sugarcane [28,29] and sorghum [19,30]. Most plant resistance to aphids is specific to a single aphid species or to a few biotypes within a species [31] and it has been demonstrated that variability exists among clonal lineages of aphids in their response to resistant cultivars [32,33]. Therefore, characterisation of the genetic diversity of aphids is critical for breeding durable and efficient resistance, which has to account for the worldwide diversity of these pests and the potential emergence of new invasive biotypes.

Based on a worldwide sample of aphids covering its area of distribution (home range as well as invasive range) and using microsatellite markers and sequencing of a fragment of the mitochondrial cytochrome c oxidase I gene (COI), the purpose of this study was to evaluate 1) the mode of reproduction of *M. sacchari*; 2) molecular evidence for the existence of sibling species; and 3) its genetic diversity and structuring according to host and locality.

Materials and Methods

Insect samples

Here, an ‘individual’ refers to one individual aphid and a ‘sample’ refers a several individuals collected from the same host plant species in a given locality and date. The complete set of individuals (Table S1) comprised 57 samples from 42 localities in a total of 15 countries or provinces, and from five host plants: sugarcane, pearl millet, and three wild or cultivated sorghum species (*Sorghum bicolor*, *S. halepense*, *S. verticilliflorum*). The three sorghum species were considered as a single host plant, hereafter named ‘sorghum’.

Aphids were collected from wild or cultivated plants and placed in 70% ethanol in Eppendorf tubes, kept frozen at -80°C until they were processed. Only a few aphids were collected on each plant sampled to avoid collecting several individuals from the same colony.

Sampling was carried out from 2002 to 2009 by our team in Reunion Island and by colleagues in the other parts of the world (see acknowledgements). Geographic coordinates of sampling localities are provided in Table S1. No specific permissions were required for sampling aphids in these locations. The field studies did not involve endangered or protected species.

DNA extraction, genotyping and sequencing

DNA extraction. DNA was extracted using the “salting-out” protocol of Sunnucks and Hales [34]. Briefly, it consists in extracting DNA from whole aphids by crushing them in a TNES/Proteinase K buffer and precipitating DNA in ethanol. This method is simple and fast, and provided sufficient DNA for phylogeny and microsatellite PCR analyses.

Genotyping. According to their polymorphism, ten microsatellite loci (Tab. 1) were selected among the 14 previously developed by our team for *M. sacchari* [35]. PCR reactions were performed with labelled primers and multiplexed into two mixes (Type-it, standard procedure, Qjagen), and the following thermocycling protocol was used: denaturation at 95°C for 15 min, 25 denaturation cycles for 30 s at 94°C , a 1-min 30 s annealing step at 54°C , and a 30-s elongation step at 72°C . We used an ABI prism 3110 for genotyping after addition of an internal size standard for each sample (GeneScan LIZ 500, Applied Biosystems). Alleles were identified at each locus by comparison with the size standard using GeneMapper version 2.5 software (Applied Biosystems).

Sequencing. A total of 91 aphids were chosen among the worldwide sample to represent different combinations of region and host plant. COI fragments were amplified using the LCO1490 and HCO2198 primers designed by Folmer et al. [36]. PCR was carried out using the protocol of Kim and Lee [37]. PCR products were purified and sequenced by a subcontractor (Cogenics), and a consensus sequence of 658 pb was chosen for later analyses.

Data analysis

Clonal diversity analysis. Micro-Checker software [38] was run on the whole population. No evidence was found for the presence of null alleles. Any single combination of alleles was retrieved from genotyping data and arranged as unique multilocus genotypes (MLGs). Given the clonal reproduction of *M. sacchari*, we assumed that the different occurrences of the same MLG in a sample were the result of local clonal reproduction. We therefore retained a single representative of each MLG in each of the 57 samples for genetic and diversity analysis.

Using GENCLONE software [39], we computed a matrix of pairwise genetic distance between MLGs computed as the number of allelic differences between MLGs [40]. Examination of the distribution of these distances enabled us to define a threshold below which MLGs were considered to belong to the same multilocus lineage (MLL), i.e. genotypes which differed slightly due to mutation or scoring errors according to Arnaud-Haond et al. [40]. The same matrix of pairwise distances was also used to construct a minimum spanning network using HAPSTAR software [41]. On the set of identical loci within each MLL, we computed p_{sex} , the probability that the repeated MLGs originated from distinct sexual reproductive events. A p_{sex} value lower than 0.01 supported the hypothesis that MLGs originated from the same MLL [40]. To describe clonal diversity, we computed the clonal richness index as $R = (G-1)/(N-1)$, where G is the number of genotypes detected (either MLGs for R_{MLG} or MLL for R_{MLL}), and N is the number of samples [42].

Phylogenetic analysis. Sequence alignments of the COI gene were performed using Geneious software version 5.6.6 [43]. Five sequences from *M. sacchari* individuals collected in India [44] were retrieved from GenBank and added to our data. Four sequences from three species of the *Melanaphis* genus were also retrieved from GenBank and used as outgroups: *M. donacis* (referenced HQ443314), *M. bambusae* (referenced EU701747 and EU701746) and *M. japonica* (referenced GU457792). Maximum Likelihood inference performed with MEGA6 [45] was used to choose the most reliable evolutionary model of base substitution to infer the evolutionary history. Based on the AICc criterion, the best model proved to be the General Time Reversible model with gamma distribution of evolutionary rates among sites (GTR+G) [46]. The GTR+G model was then used with MEGA6 to reconstruct the subsequent phylogenetic tree through the Maximum Likelihood method, with 10,000 bootstrap replicates for branch support.

Population genetic analysis. We used GENEPOP [47] to compute population genetics parameters for each of the MLLs delimited by GENCLONE. We tested departures from Hardy–Weinberg equilibrium and heterozygote deficit and excess, and calculated population fixation index values (F_{is}). Genetic differentiation between MLLs was tested with a G test and pairwise F_{ST} were computed.

Results

Genetic and clonal diversity

We genotyped a total of 1333 aphids using the ten microsatellite markers. When we retained a single representative of each MLG in each of the 57 samples, this yielded a dataset containing 98 individuals.

Global genetic diversity was low, with 36 MLGs found (Table 1). Global clonal richness was also low, with a $R_{MLG} = 0.361$. The distribution of the pairwise number of different alleles between MLG appeared multimodal, with a first minimum located at a distance of five alleles (Figure S1). Grouping MLGs which differed by one to four alleles defined five groups. Calculation of p_{sex} on the set of identical loci within each of these five groups yielded values <0.01 , confirming that the MLGs within each group were unlikely to have derived from distinct reproductive events. We therefore considered that the five groups defined five multilocus lineages (MLLs) which grouped slightly distinct MLGs resulting from step mutations or scoring errors (Table 1, Figure 1). Considering the five MLLs, clonal richness was very low, as shown by the $R_{MLL} = 0.041$.

Phylogenetic relationship within samples

Within our 91 *M. sacchari* COI sequences, only three distinct haplotypes were observed (Figure 2). One haplotype was observed in individuals belonging to MLL-A, MLL-B or MLL-E (Figure 1, Figure 2). The second haplotype was only observed in individuals belonging to MLL-C. The third haplotype was observed in individuals belonging to MLL-D. No association of haplotypes with the host plant was observed (Figure 2). These three haplotypes differed from the two available in GenBank from five Indian samples, giving a total of five haplotypes and five nucleotide substitutions among 96 *M. sacchari* individuals. The phylogenetic tree built from a 658 bp

fragment of the COI gene clearly separated the four *Melanaphis* species with $>80\%$ bootstrap support (Figure S2). But within the *M. sacchari* sequences, the presence of distinct taxa was not supported by bootstrap analysis at the 80% threshold. Intraspecific genetic divergence in *M. sacchari* was low, with a mean pairwise divergence of 0.19% (range 0.000.61%). When the five *M. sacchari* sequences retrieved from GenBank were excluded, the mean divergence was 0.17% (range 0.000.30%). The mean divergence of *M. sacchari* sequences with the closest taxa, *M. japonica*, was 1.06% (range 0.92%–1.39%).

Standard population genetics

Plotting the results of the factorial correspondence analysis with GENETIX confirmed the grouping of the 36 MLGs in five MLLs (Figure 3). Factor 1 distinguished MLL-A and MLL-E, and a group formed by the three MLL-B, MLL-C and MLL-D. Factor 2 distinguished MLL-E from other MLLs. Factor 3 distinguished between MLL-B, MLL-C and MLL-D.

Genetic differentiation between the five MLLs was strong, with a highly significant F_{st} ranging from 0.262 to 0.694 (Table S2).

The five populations comprised by each distinct MLL differed significantly from Hardy-Weinberg equilibrium, and showed a clear signature of asexual reproduction with a significant heterozygote excess and negative F_{IS} values (Table S3).

Geographical and host distribution of MLLs

Distribution of the MLLs revealed strong geographical structuring (Figure 4). MLL-A was observed in Africa, MLL-B was restricted to Australia, MLL-C exhibited the widest distribution area (South America, the Caribbean, the Indian Ocean and East Africa), MLL-D was observed in the USA, and MLL-E was only observed in China. Kenya was the only country where two MLLs were observed simultaneously: one sample (Ken1) contained a mix

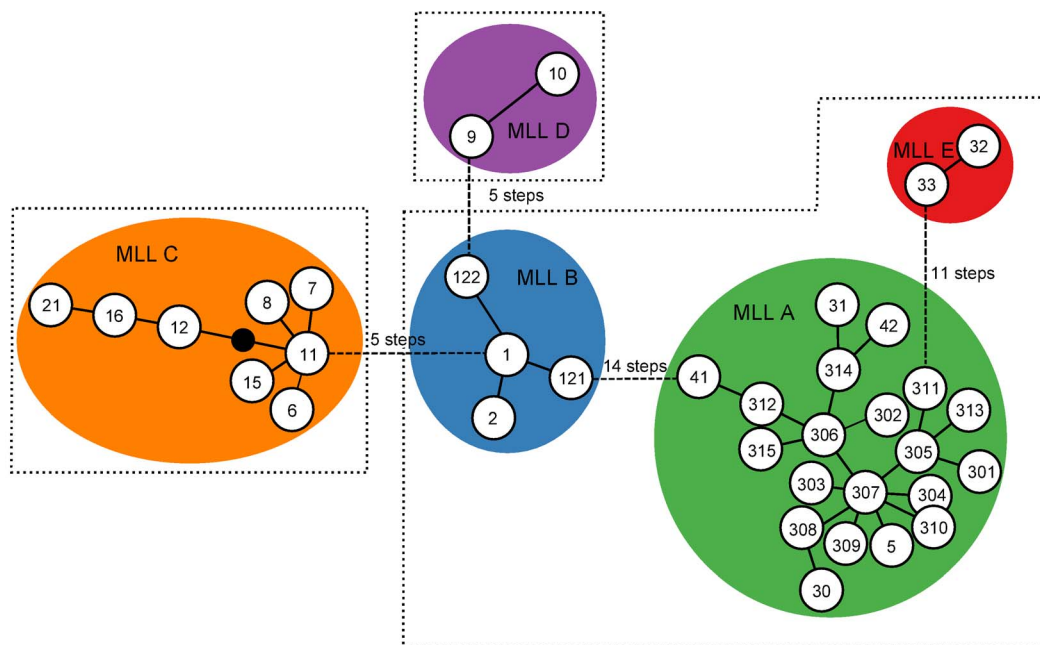


Figure 1. Minimum spanning network of *Melanaphis sacchari* microsatellite distances computed as the number of allele differences between MLGs. Each node represents one step in the network, i.e. a distance of one allele. The numbers in the circles represent MLGs according to Table 1. Coloured backgrounds represent the Multi Locus Lineages (MLLs). MLGs in the same dashed line box share the same COI haplotype.

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Table 1. Observed microsatellite Multi Locus Genotypes (MLG): allele size (bp) at each locus, distribution by host plant and in each of the five Multi Locus Lineages (MLL) defined with GENCLONE. Within MLL allelic variations are in bold.

MLL	MLG	CIR- <i>Ms</i> -G08	CIR- <i>Ms</i> -G403	CIR- <i>Ms</i> -B03	CIR- <i>Ms</i> -C08	CIR- <i>Ms</i> -G01	CIR- <i>Ms</i> -E01	CIR- <i>Ms</i> -G12	CIR- <i>Ms</i> -E03	CIR- <i>Ms</i> -D02	CIR- <i>Ms</i> -G02
MLL A	Ms5	229/229	253/253	215/215	189/203	206/210	245/247	214/216	176/176	220/228	250/316
	Ms30	229/229	253/253	215/215	189/203	204/206	245/247	214/216	176/176	220/228	250/318
	Ms31	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	228/230	254/314
	Ms41	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/226	252/314
	Ms42	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/226	254/314
	Ms301	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	248/312
	Ms302	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	248/314
	Ms303	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	250/298
	Ms304	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	250/310
	Ms305	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	250/312
	Ms306	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	250/314
	Ms307	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	250/316
	Ms308	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	250/318
	Ms309	229/229	253//253	215/215	189//203	206/ 206	245/247	214/216	176/176	220/228	250/343
	Ms310	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	250/345
	Ms311	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	252/312
	Ms312	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	252/314
	Ms313	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	254/312
	Ms314	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	254/314
	Ms315	229/229	253/253	215/215	189/203	206/ 206	245/247	214/216	176/176	220/228	256/314
MLL B	Ms1	233/233	253/259	213/215	197/199	185/206	247/247	212/216	188/193	226/232	199/199
	Ms2	233/233	253/259	213/215	197/199	185/206	247/247	212/216	188/193	226/252	199/199
	Ms121	233/233	253/259	213/215	197/199	185/206	247/247	212/216	188/191	226/232	199/199
	Ms122	233/233	253/259	213/215	197/199	185/206	247/247	212/216	188/191	226/232	199/201
MLL C	Ms6	233/233	251/259	213/213	197/199	185/210	247/247	212/216	186/193	228/234	199/199
	Ms7	233/233	251/259	213/213	197/199	185/210	247/247	204/212	186/193	228/232	199/199
	Ms8	233/233	251/259	213/213	197/199	185/212	247/247	212/216	186/193	228/232	199/199
	Ms11	233/233	251/259	213/213	197/199	185/210	247/247	212/216	186/193	228/232	199/199
	Ms12	233/233	251/259	213/213	195/197	185/210	247/247	212/216	186/193	228/232	199/205
	Ms15	233/233	251/259	213/213	197/199	185/210	247/247	212/216	186/193	228/232	199/201
	Ms16	233/233	251/259	213/213	195/197	185/210	247/247	212/216	186/193	228/232	199/203
	Ms21	233/233	251/259	213/213	195/197	185/210	247/247	212/216	186/193	228/234	199/203
MLL D	Ms9	233/233	251/259	213/213	197/199	185/206	247/247	212/216	186/188	226/234	201/201
	Ms10	233/233	251/259	213/213	197/199	185/206	247/247	212/216	186/188	226/234	201/201

Table 1. Cont.

MLL	MLG	CIR-MS-G08	CIR-MS-G403	CIR-MS-B03	CIR-MS-C08	CIR-MS-G01	CIR-MS-E01	CIR-MS-G12	CIR-MS-E03	CIR-MS-D02	CIR-MS-G02
MLL E	Ms32	229/229	253/253	215/215	197/197	198/206	245/245	208/212	216/224	220/222	216/216
	Ms33	229/229	253/253	215/215	197/197	198/206	245/245	208/212	224/224	220/222	216/216

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of MLL-A and MLL-B, one sample (Ken5) contained MLL-A alone, and three samples (Ken2, Ken3, Ken4) contained MLL-B alone.

No host plant structuring was observed: in all the countries where both sorghum and sugarcane samples were collected, each MLL was found on both host plants (Table 2).

Discussion

Molecular analysis revealed a very low genetic diversity among 57 samples collected in 15 countries on two main hosts, with 36 MLGs structured in five MLLs. The distribution of MLLs was strongly structured by geography but not by the host plant (sorghum vs. sugarcane).

Sequencing the COI 'barcoding' region, a typical locus used for species discrimination and phylogeny, particularly in aphids [48,49], did not enable the detection of cryptic species in our samples. Specifically, we observed no molecular evidence for a clear separation into two species, *M. sacchari* and *M. sorghi*. We found sequence variations peaking at 0.61%, with a mean value of 0.19%, both of which are within the range of intraspecific divergence observed in the Aphididae family by Footit et al. [50] or Lee et al. [51].

Reproduction

Population genetic parameters were consistent with populations which only reproduce by apomictic parthenogenesis, as previously described by Blackman and Eastop [22]. Each population significantly differed from Hardy-Weinberg equilibrium, with a high heterozygote excess. These features are a common consequence of populations which have reproduced clonally for a long time. In a global study of genetic diversity on *Aphis gossypii*, Carletto et al. [52] obtained similar results to ours as they observed low genetic diversity, with the predominance of a few clones at the worldwide scale reproducing only by apomictic parthenogenesis. But later, evidence for sexual reproduction of *A. gossypii* was found in Iran [53]. High genetic diversity and evidence for sexual reproduction was also observed in *A. gossypii* alate spring migrants in France [54]. Similarly, in the *Brachycaudus helichrysi* (Kaltenbach) sibling species H2, a sexually reproducing population was identified in India, despite almost exclusively clonal reproduction at the worldwide scale [11]. This shows that sexual admixture can still exist in a local population even in species which are highly clonal at the worldwide scale. A holocycle has been observed in *M. sacchari* in Asia [16] and this suggests that higher genetic diversity may exist in some parts of its geographic distribution area even if our sampling did not allow us to observe it.

Geographic genetic structure

Microsatellite analyses showed that population structuring at the worldwide scale was only influenced by geography, delimiting five MLLs corresponding to five geographic zones: Africa, China, Australia, USA, and South America – Indian Ocean (including Kenya) and the Caribbean. Variation within each of the four biggest zones (excluding China where only one sample was analysed) was low, with MLGs within a zone differing by a few step mutations, which we suggest is due to one or few separate introductions. These results suggest that each of the five zones was colonized separately following the introduction of one or a few clones from the region of origin of *M. sacchari*, which the present study did not allow us to identify. At least two of the five MLLs covered a very wide geographic area and matched the pattern of a single asexual genotype with a high capacity for dispersal which

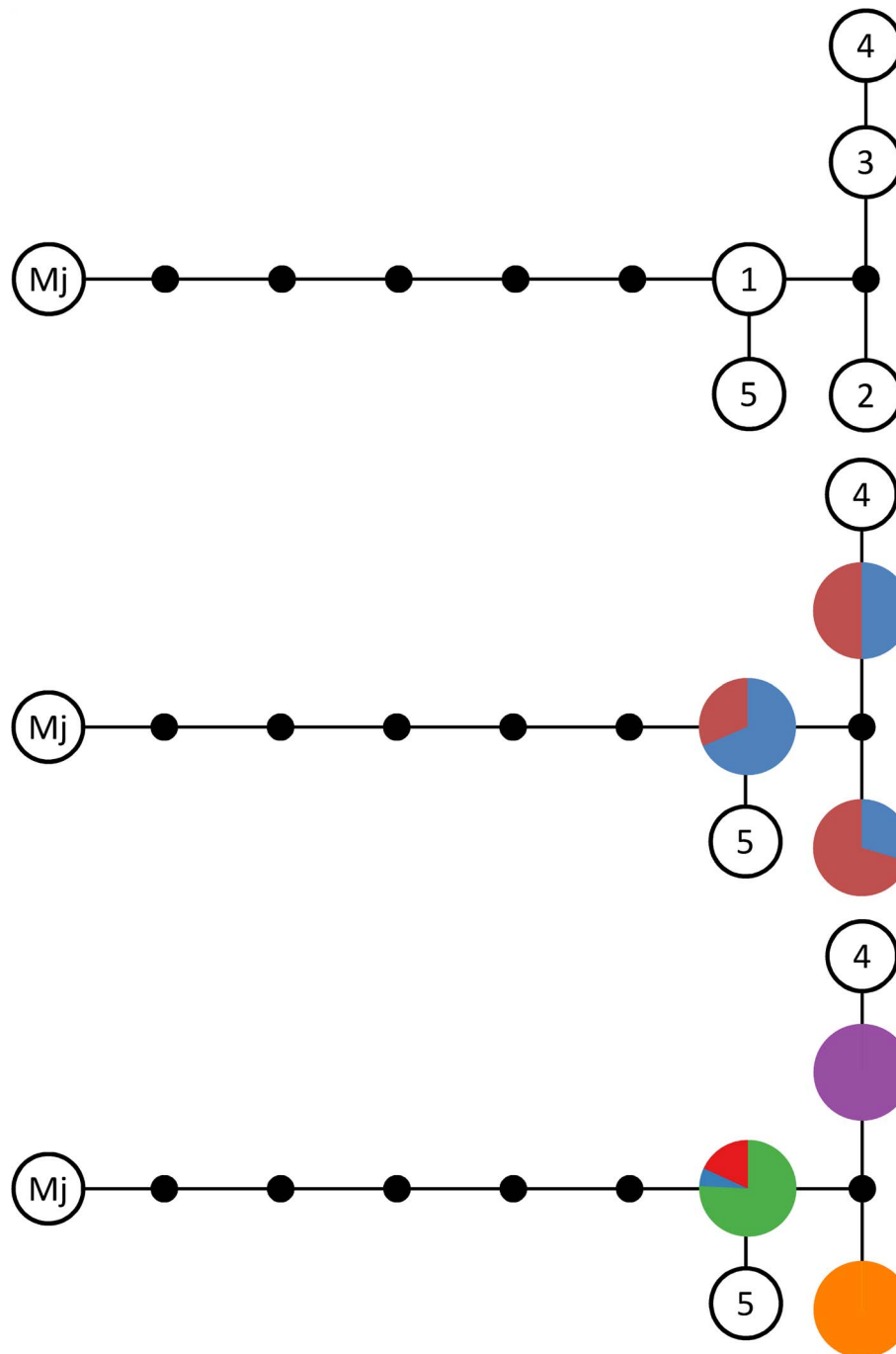


Figure 2. COI haplotype network (top), in which *Melanaphis sacchari* COI sequences originating from the present study are numbered from 1 to 3. *M. sacchari* GenBank COI sequences from India [44] are numbered 4 (JX051388, JX05189, JX051390) and 5 (HQ112185, JX051402). Mj = *Melanaphis japonica* COI sequence from GenBank (GU457792). Distribution as a function of host plant (middle): sorghum (blue) vs. sugarcane (red). Distribution as a function of MLL (bottom): A (green), B (blue), C (yellow), D (violin), E (red).
doi:10.1371/journal.pone.0106067.g002

would have spread across a large area: MLL-A was observed in West and East Africa, and MLL-C was observed in South America, the Indian Ocean, East Africa, and the Caribbean. Both MLLs match the definition of “superclone” [7], characterised by geographically and ecologically widespread distribution, which has already been documented in several aphid species [9,10,11,55,56].

The low rate of genetic diversity observed in the whole geographic area covered by our study, and the lack of published

data about the dates of introduction of *M. sacchari* in the countries sampled, meant we were not able to reconstruct the invasion routes of this species. The only exception was continental USA. *M. sacchari* was first described in Hawaii in the late 19th century [57,58], and was first recorded in continental USA at the end of the 1970s in Florida [59,60] and in 2001 in Louisiana [18]. In our study, almost all individuals sampled in Louisiana and Hawaii belonged to the same MLG, Ms9, and shared the same COI

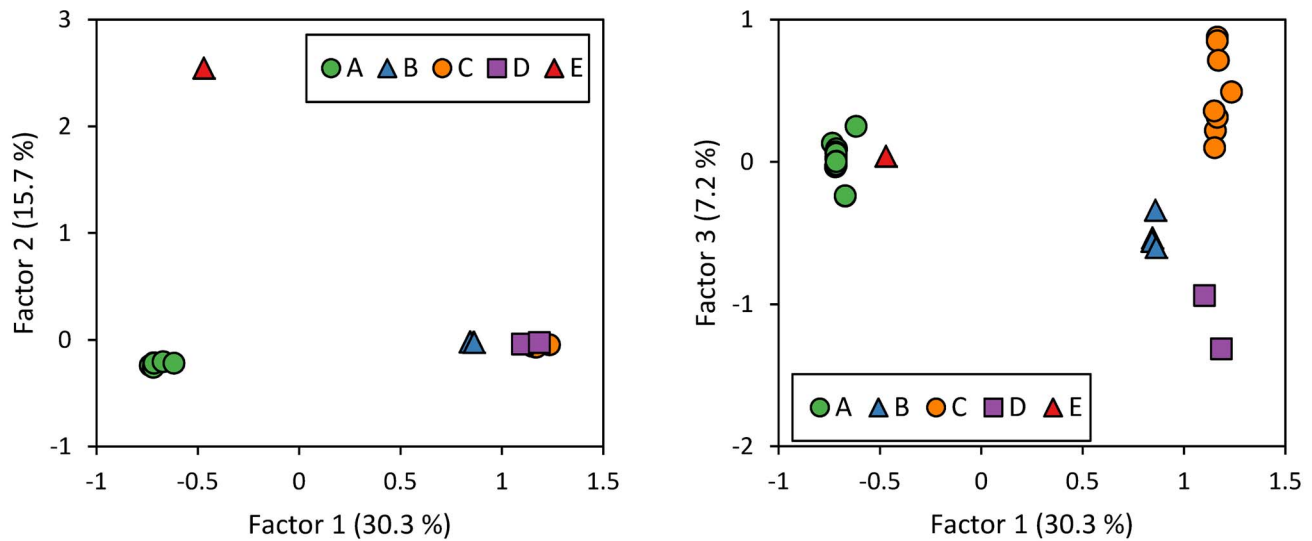


Figure 3. Factorial correspondence analysis of microsatellite data with GENETIX. Each symbol represents one of the 36 MLGs. Colours and letters refer to Multilocus Lineage (MLL) assignment with GENCLONE. doi:10.1371/journal.pone.0106067.g003

haplotype, neither of which were observed in other regions. This strongly suggests that *M. sacchari* was introduced in continental USA from Hawaii. This finding is notable, as one would expect an introduction into continental USA from either South America or the Caribbean, a shorter invasion route. However, Mondor et al. [61] emphasized that the relationship between the colonization of the Hawaiian Islands by an aphid species and its presence in continental USA was due to the high rate of commercial exchanges between the two. Here we provide an example of reverse colonization from Hawaii to continental USA. In Kenya,

MLL-C was observed in three samples from the coastal region of Kenya but was not found in a sample collected inland. This underlines the fact that, even though *M. sacchari* has been recorded in almost all areas where sugarcane is cultivated, the possibility for the expansion of some genotypes should be taken seriously, mainly its unknown potential impact on the epidemiology of the viral diseases it transmits.

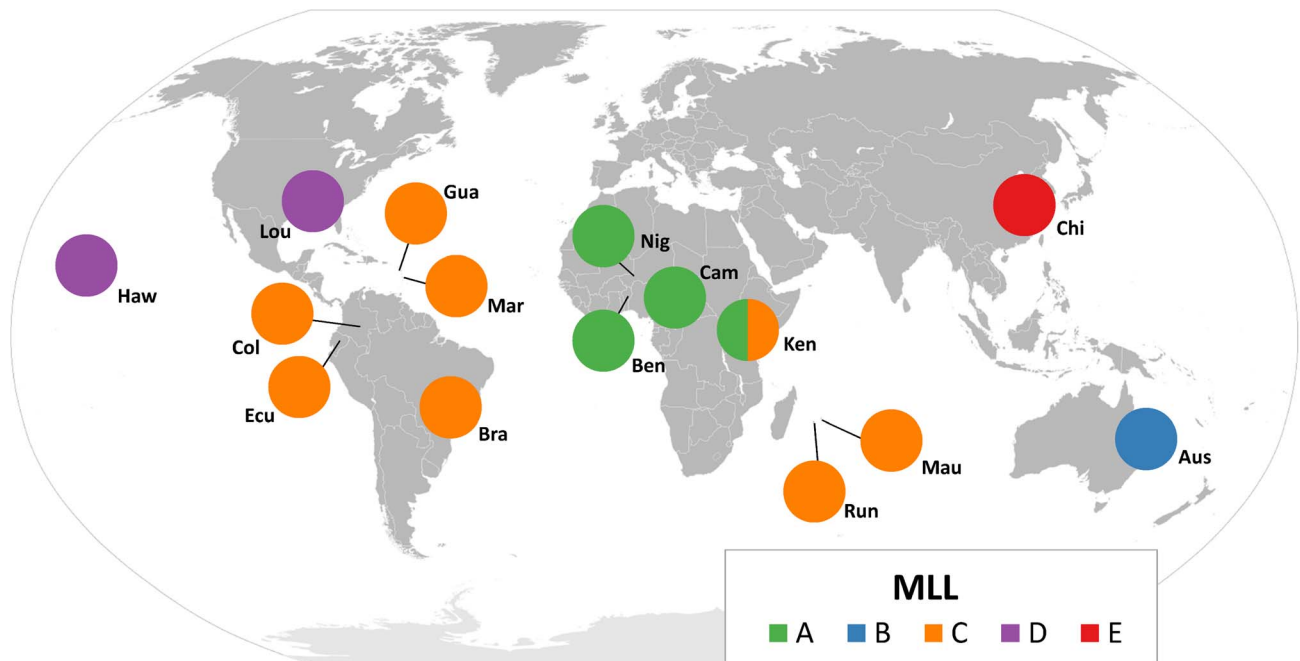


Figure 4. Relative geographical within-state distribution of Multilocus lineages (MLL). The size of circle is not proportional to the size of the sample. Aus = Australia, Bra = Brazil, Col = Columbia, Ecu = Ecuador, Gua = Guadeloupe, Haw = Hawaii, Lou = Louisiana, Mar = Martinique, Mau = Mauritius, Run = Reunion Island, Ben = Benin, Cam = Cameroon, Nig = Niger, Chi = China, Ken = Kenya. doi:10.1371/journal.pone.0106067.g004

Table 2. Distribution of the 98 individuals, a single representative of each MLG in each of the 57 samples, as a function of country/province and host plant.

State	Host plant	Number of individuals					Number of samples
		MLL-A	MLL-B	MLL-C	MLL-D	MLL-E	
Benin	sorghum	3					2
	sugarcane	6					1
	pearl millet	1					1
Cameroon	sorghum	20					5
	sorghum	2					1
Kenya	sugarcane	2		4			4
	sorghum	5					2
Niger	sugarcane		11				7
Australia	sugarcane						1
Brazil	sugarcane			2			3
Columbia	sugarcane			7			1
	sugarcane			1			5
Ecuador	sugarcane			6			3
Guadeloupe	sugarcane			3			1
Martinique	sugarcane			1			3
Mauritius	sorghum			3			5
Reunion	sugarcane			7			3
Hawaii	sugarcane				3		3
Louisiana	sorghum				1		1
	sugarcane				8		7
China	sorghum					2	1

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Supporting Information

Figure S1 Distribution of the pairwise number of different alleles between MLGs.

(PDF)

Figure S2 Molecular Phylogenetic analysis by Maximum Likelihood method with bootstrap support (10,000 replicates) using 658 bp cytochrome c oxidase subunit I sequences from 100 *Melanaphis* spp. individuals.

(PDF)

Table S1 Voucher number, sampling information, GenBank accession and SSR genotyping of individual aphids.

(XLSX)

Table S2 Genetic differentiation between MLLs: pairwise F_{ST} and significance of the G test computed with GENEPOP.

(PDF)

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Table S3 Population genetics parameters for each MLL.

(PDF)

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Author Contributions

Conceived and designed the experiments: BF SN LC HD BR. Performed the experiments: BF SM. Analyzed the data: SN BF LC HD. Contributed to the writing of the manuscript: BF SM SN HD LC BR.

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