# Microsatellite flanking region similarities among different loci within insect species 

E. Meglécz*, S. J. Anderson $\dagger$, D. Bourguet $\ddagger$, R. Butcher§, A. Caldas $\uparrow$, A. Cassel-Lundhagen**, A. C. d'Acier $\ddagger$, D. A. Dawsontt, N. Faure $\ddagger \ddagger$, C. Fauvelot§§, P. Franckiๆl), G. Harper***, N. Keyghobaditt†, C. Kluetsch $\ddagger \ddagger \ddagger$, M. Muthulakshmi§§§, J. Nagaraju§§§, A. Patt $\ddagger \ddagger \ddagger$, F. Péténianๆๆाף, J.-F. Silvain $\ddagger \ddagger$, H. R. Wilcock ${ }^{* * * *}$<br>*Evolution Génome et Environnement, CASE 36, Université de Provence, Marseille, France, $\dagger$ Department of Biological Sciences, The Open University, Milton Keynes, UK; $\ddagger$ Centre de Biologie et de Gestion des Populations (CBGP), INRA, UMR-INRA-IRD-CIRAD-Agro.M, Campus International de Baillarguet, Montferrier/Lez, France; §Centre for Vectors and Vector-Borne Diseases, Faculty of Science, Mahidol University, Phaya thai, Bangkok, Thailand; $\boldsymbol{\uparrow}$ Department of Entomology, University of Maryland, USA; ** Swedish University of Agricultural Sciences, Department of Entomology, Uppsala, Sweden; $\dagger \dagger$ Department of Animal and Plant Sciences, University of Sheffield, Sheffield, UK; $\ddagger \ddagger I R D$, UR072, c/o CNRS, UPR 9034, BP1, avenue de la Terrasse, Gif-sur-Yvette Cedex, France; $\S \S$ Environmental Science, University of Bologna at Ravenna, Ravenna, Italy; $\Pi \uparrow$ IIantes et Systèmes de culture Horticoles, INRA, AgroParc, Domaine St-Paul 84914 Avignon, Cedex 9, France;***School of Applied Sciences, University of Glamorgan, Pontypridd, Mid Glamorgan; UK; $\dagger \dagger \dagger$ University of Western Ontario, Department of Biology, London, Ontario, Canada; $\ddagger \ddagger \ddagger Z o o l o g i s c h e s ~ F o r s c h u n g s m u s e u m ~ A l e x a n d e r ~ K o e n i g, ~$ Bonn, Germany; §§§Laboratory of Molecular Genetics, Centre for DNA Fingerprinting and Diagnostics, Nacharam, Hyderabad, India; $\uparrow T T E R$ Biodiversité et Environnement, Université de Provence, Marseille, France; and<br>${ }^{* * * *}$ Deceased, Department of Biological Sciences, University of Hull, Hull, UK


#### Abstract

Although microsatellites are ubiquitous in eukaryota, the number of available markers varies strongly among

Received 30 June 2006; accepted after revision 5 October 2006; first published online 7 February 2007. Correspondence: Emese Meglécz, EGEE, CASE 36, Université de Provence, 3 place Victor Hugo, F-13331, Marseille, Cedex 3, France. Tel.: +33 491106338; fax: +33 491106353; email: meglecz@up.univ-mrs.fr


taxa. This meta-analysis was conducted on 32 insect species. Sequences were obtained from two assembled whole genomes, whole genome shotgun (WGS) sequences from 10 species and screening partial genomic libraries for microsatellites from 23 species. We have demonstrated: (1) strong differences in the abundance of microsatellites among species; (2) that microsatellites within species are often grouped into families based on similarities in their flanking sequences; (3) that the proportion of microsatellites grouped into families varies strongly among taxa; and (4) that microsatellite families were significantly more often associated with transposable elements - or their remnants - than unique microsatellite sequences.

Keywords: microsatellite, flanking region, interspersed repetitive element, Lepidoptera, genome.

## Introduction

Microsatellites have been detected in a wide diversity of eukaryotes of varying complexity and genome size (Tóth et al., 2000). Although numerous studies suggest that at least a fraction of microsatellites have important functions (Li et al., 2002, and references therein), they are generally regarded as highly variable neutral markers, and are thus the most frequently used genetic markers in population biology.
Despite early reports of microsatellite abundance (Beckmann \& Soller, 1990; Beckmann \& Weber, 1992; Epplen et al., 1997), the isolation of microsatellites as useable markers appears to be more difficult in some taxa than in others (Nematoda: Fisher \& Viney, 1996; Lepidoptera: Nève \& Meglécz, 2000; Zhang, 2004; Aedes aegypti and Ixodes scapulris: Fagerberg et al., 2001). There are at least two possible, non-exclusive, explanations for this heterogeneity: (1) either the abundance of microsatellites, or (2) their localization in the genomes (e.g. coding or noncoding regions; association with repetitive DNA; extent of gene duplications or polyploidy; position on the chromosomes) significantly differ among species or higher level taxa.
Zhang (2004) has pointed out the relative scarcity of microsatellites in Lepidopteran genomes, as have Fagerberg
et al. (2001) in I. scapularis (Arachnida; Ixodida) and A. aegypti (Diptera). Both studies are based on indirect estimates, as genome data are rare for non-Dipteran (nonDrosophila) insects. The only Lepidoptera genome that has been sequenced so far is of Bombyx mori (Mita et al., 2004; Xia et al., 2004). In this species, microsatellites make up $0.31 \%$ of the genome (Prasad et al., 2005), which is rather low but not out of the range of other analysed genomes (Tóth et al., 2000). In this study, we directly compared microsatellite abundance among the 10 insect species for which Whole Genome Shotgun (WGS) sequences are available.

The use of polymerase chain reaction-based microsatellite analysis in population genetics is based upon the premise that the flanking sequences are unique for each locus (single co-dominant locus evaluation) and that they mutate much more slowly than the microsatellite repeat motif (Hancock, 1999). While there have been many studies on the mutation pattern of microsatellite core repeats (Weber \& Wong, 1993; Michalakis \& Veuille, 1996; Chakraborty et al., 1997; Schlötterer, 1998; Schug et al., 1998; Estoup \& Cornuet, 1999; Hancock, 1999; Pupko \& Graur, 1999), little is known on their origin, including the uniqueness of their flanking sequences. A recent study showed that in two butterfly species a considerable proportion of the isolated microsatellite loci cluster into families based on similarities between flanking regions of different loci (Meglécz et al., 2004). The work presented here aims to (1) test if this phenomenon is general in insects and in Lepidoptera in particular, and to (2) examine the distribution of the frequencies of these microsatellite families among taxa.

As microsatellites are generally more frequent in noncoding DNA regions (Hancock, 1995; Ramsay et al., 1999; Metzgar et al., 2000; Tóth et al., 2000; Karaoglu et al., 2005) and a large portion of noncoding DNA is repetitive in
many species (Comeron, 2001; Kidwell \& Lisch, 2001; Li et al., 2002 and references therein), we also tested the association between microsatellite families and interspersed repetitive elements.

## Results and discussion

## Microsatellite abundance

The estimation of microsatellite abundance is very difficult without genomic data. Indirect methods, such as using the frequency of positive clones, have a number of potential biases. The emergence of genomic data provides a possibility to have a clearer picture, but only for a limited number of species. Furthermore, it is difficult to directly compare the results of independent studies, as there are no standard criteria for the minimal size of the microsatellites. In this study, we have used microsatellites with at least four uninterrupted repeat units (eight for single base pairs), so that even short mono- or dinucleotide repeats are included in the analyses. In similar studies, the criteria have been both more and less stringent. For example Tóth et al. (2000) used microsatellites longer than 12 bp for a wide variety of Eukaryotes, Dieringer \& Schlötterer (2003) used at least two repetitions, Karaoglu et al. (2005) used microsatellites longer than 10 bp for fungal genomes and Prasad et al. (2005) used at least five repetitions (15 for single bases), for B. mori.

Table 1 presents the estimated proportion of pure microsatellites in the genome, for each repeat unit length, calculated from WGS and complete genome data. At first sight, our estimation seems extremely high. For example, we found that on average $\approx 16 \mathrm{~kb}$ of microsatellite is found per 1 Mb of genomic DNA in the $B$. mori genome, while this was found to be $\approx 3 \mathrm{~kb}$ in Prasad et al. (2005) for the same species. Rerunning our analyses with the same criteria as applied by Prasad et al. (2005), however, gave very similar

Table 1. Total length of the analysed genomic sequences and proportion of microsatellites observed for each repeat unit length for the complete genome and WGS data

|  | Total DNA (Mbp) | Mono- (\%) | Di- (\%) | Tri- (\%) | Tetra- (\%) | Penta- (\%) | Hexa- (\%) | All motifs (\%) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Complete genome |  |  |  |  |  |  |  |  |
| Drosophila melanogaster | 118.4 | 6.73 | 6.55 | 2.34 | 0.53 | 0.21 | 0.24 | 16.60 |
| Anopheles gambiae | 223.2 | 6.63 | 11.57 | 4.16 | 0.60 | 0.09 | 0.10 | 23.15 |
| Whole Genome Shotgun |  |  |  |  |  |  |  |  |
| Aedes aegypti | 1212.3 | 7.36 | 1.50 | 1.11 | 0.31 | 0.18 | 0.12 | 10.57 |
| Anopheles gambiae | 313.8 | 6.11 | 10.58 | 3.68 | 0.58 | 0.12 | 0.13 | 21.19 |
| Apis mellifera | 232.7 | 14.45 | 15.30 | 3.89 | 1.27 | 0.44 | 0.14 | 35.50 |
| Bombyx mori A | 393.3 | 10.03 | 3.64 | 1.39 | 0.78 | 0.16 | 0.04 | 16.06 |
| Bombyx mori B | 381.9 | 10.45 | 3.68 | 1.42 | 0.83 | 0.18 | 0.05 | 16.61 |
| Drosophila melanogaster | 132.5 | 6.58 | 6.16 | 2.22 | 0.50 | 0.45 | 0.22 | 16.13 |
| Drosophila persimilis | 175.5 | 4.65 | 9.52 | 3.24 | 0.91 | 0.65 | 0.95 | 19.93 |
| Drosophila pseudoobscura | 147.4 | 4.97 | 10.33 | 3.64 | 1.02 | 0.73 | 0.91 | 21.59 |
| Drosophila sechelia | 157.2 | 4.34 | 5.30 | 1.84 | 0.40 | 0.50 | 0.13 | 12.51 |
| Drosophila simulans | 119.3 | 4.71 | 6.01 | 2.05 | 0.43 | 0.17 | 0.13 | 13.51 |
| Drosophila yakuba | 156.8 | 5.05 | 5.80 | 2.02 | 0.42 | 0.21 | 0.33 | 13.83 |

Figure 1. The proportion of pure microsatellites for each repeat unit length based on WGS data. $\%$ refers to the total length of microsatellites in kbp per one Mbp of analysed DNA.

results to theirs (data not shown). It thus appears that this large discrepancy among the estimates is due to the large number of short microsatellites present in the genome of B. mori.

Nevertheless, our results enable us to compare the frequencies of microsatellites among the 10 species for which WGS data were available. We observed a three-fold variation in microsatellite frequencies among the 10 investigated insect species, with the lowest value in $A$. aegypti and the highest in Apis mellifera. Even within the Drosophila genus, the variation among species is considerable (Table 1). Running the analyses with more stringent criteria (i.e. at least five to eight uninterrupted repeats) did not change the overall pattern of variation among species (data not shown). The difference between genomes is even more pronounced when only 2-6 bp motifs are considered (Fig. 1). The proportion of mononucleotide motifs compared with other motifs is particularly high in $B$. moriand $A$. aegypti. As microsatellites with mononucleotide motifs are rarely used as markers, this introduces even greater difference among species in the abundance of potential genetic markers.

A particularly low frequency of positive clones was reported in $A$. aegypti compared with Anopheles albimanus (Fagerberg et al., 2001) and for several Lepidoptera species (Zhang, 2004 and references therein), while A. mellifera is one of the insect species for which a very large number of microsatellite markers are established (Solignac et al., 2003). Microsatellite abundance in the genome is thus probably a relevant factor in the success of microsatellite marker isolation; however, it is unlikely to be the only source of the problems specifically found in some taxa such as Lepidoptera. The low frequency of microsatellites in the genome should not lead to amplification problems and in principle, with enrichment or by increasing the number of screened clones, a sufficient number of markers could be isolated.

## Proportion of grouped sequences

Microsatellite flanking sequences were analysed species by species. Each data set was blasted against itself ( $e=1 \mathrm{E}-40$ ) and sequences were sorted into four categories based on the results of the blastn: (1) Unique if no
similarities were observed to any other sequences of the same data set; (2) Unblastable if sequence had no hits at all, not even with itself; (3) Redundant, if the identity to another sequence was higher than $95 \%$ along the whole flanking sequence; and (4) Grouped if sequences produced a significant hit with at least one different sequence and they were not redundant.

Data from complete genomes and WGS For both Drosophila melanogasterand Anopheles gambiae, the estimates of the proportion of grouped sequences were lower from complete genome sequences than from WGS (Table 2). This might be the consequence of the fact that genome assembly is the most difficult in regions with a high concentration of repetitive DNA, thus these regions might be overrepresented among the contigs of the WGS sequences. Consequently, the proportion of grouped sequences should not be taken literally from WGS data. Nevertheless, the same tendency is observed in both WGS and complete genome estimations: the genome of $A$. gambiae contains a higher proportion of microsatellite sequences grouped into families than that of $D$. melanogaster (Table 2; $\chi^{2}=4146.4, P<0.0001 ; \chi^{2}=69.1, P<0.0001$, for complete genome and WGS data, respectively).
The presence of microsatellite sequence families is clearly demonstrated in all 10 species for which WGS data were available (Table 2). The proportion of grouped sequences among WGS data sets varied largely among species ( $\chi^{2}=12997.1, P<0.0001$ ). However, as discussed above, these proportions are only rough estimates, and should be interpreted with caution. Nevertheless, it is still striking that the proportion of grouped sequences in B. mori is the highest among the investigated 10 species ( $36.1 \%$ and $34.7 \%$, respectively, for the two data sets) and it is significantly higher than the second highest proportion (23.6\% for A. aegypti; $\chi^{2}=368.5, P<0.0001 ; \chi^{2}=293.6$, $P<0.0001$, respectively).

Data from screening partial libraries Sequences originating from screenings of partial genomic libraries may be biased by the screening protocol applied. However, as discussed in Meglécz et al. (2004) the same protocol used in the

Table 2. Number of sequences in each category for each data set

| Species | Unique | Grouped | Redund. | UnBL. | Prop. grouped |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Complete genome |  |  |  |  |  |
| Anopheles gambiae* | 175128 | 14893 | 6141 | 108 | 0.078 |
| Anopheles gambiae $\dagger$ | 171483 | 24679 | - | 108 | 0.126 |
| Drosophila melanogaster* | 90005 | 1607 | 1835 | 58 | 0.018 |
| Drosophila melanogaster $\dagger$ | 89657 | 3790 | - | 58 | 0.041 |
| Whole Genome Shotgun |  |  |  |  |  |
| Aedes aegypti | 7432 | 2291 | 265 | 12 | 0.236 |
| Anopheles gambiae* | 8570 | 891 | 531 | 8 | 0.094 |
| Apis mellifera | 9672 | 126 | 69 | 133 | 0.013 |
| Bombyx mori ${ }^{*}$ | 6257 | 3541 | 192 | 10 | 0.361 |
| Bombyx mori B* | 6376 | 3391 | 209 | 24 | 0.347 |
| Drosophila melanogaster* | 9149 | 604 | 238 | 9 | 0.062 |
| Drosophila persimilis | 8222 | 1166 | 568 | 44 | 0.124 |
| Drosophila pseudoobscura | 9371 | 472 | 123 | 34 | 0.048 |
| Drosophila sechelia | 8770 | 738 | 475 | 17 | 0.078 |
| Drosophila simulans | 9568 | 287 | 122 | 23 | 0.029 |
| Drosophila yakuba | 8669 | 992 | 313 | 26 | 0.103 |
| Screening (Lepidoptera) |  |  |  |  |  |
| Arctia caja | 45 | 10 | 10 | 21 | 0.182 |
| Arhopala epimuta | 35 | 16 | 6 | 3 | 0.314 |
| Busseola fusca | 30 | 24 | 7 | 2 | 0.444 |
| Bombyx mori* | 26 | 9 | 0 | 2 | 0.257 |
| Coenonympha hero | 28 | 8 | 2 | 7 | 0.222 |
| Cydia pomonella | 70 | 2 | 14 | 1 | 0.028 |
| Drupadia theda | 58 | 4 | 11 | 8 | 0.065 |
| Euphydryas aurinia | 51 | 22 | 13 | 3 | 0.301 |
| Ostrinia nubilalis | 15 | 4 | 4 | 2 | 0.211 |
| Parnassius apollo | 35 | 43 | 18 | 7 | 0.551 |
| Parnassius mnemosyne | 26 | 0 | 2 | 3 | 0 |
| Parnassius smintheus | 18 | 0 | 0 | 1 | 0 |
| Polyommatus belargus | 8 | 4 | 6 | 4 | 0.333 |
| Plutella xylostella | 83 | 33 | 8 | 3 | 0.284 |
| Reissita simonyi | 28 | 4 | 3 | 0 | 0.125 |
| Zale galbanata | 8 | 0 | 1 | 0 | 0 |
| Screening (non-Lepidoptera) |  |  |  |  |  |
| Aphis fabae | 61 | 4 | 31 | 3 | 0.061 |
| Amphitmetus transversus | 26 | 9 | 9 | 0 | 0.257 |
| Culex pipens | 14 | 2 | 1 | 7 | 0.125 |
| Culex quinquefasciatus | 16 | 2 | 0 | 2 | 0.111 |
| Monolepta vincta | 30 | 2 | 2 | 0 | 0.062 |
| Plectrocnemia conspersa | 26 | 6 | 27 | 7 | 0.188 |
| Polycentropus flavomaculatus | 18 | 2 | 6 | 3 | 0.1 |

Unique = number of sequences with no similarities to any other sequences of the same data set; Grouped = number of sequences grouped into sequence families based on similarities in their flanking region; Redund. = number of redundant sequences; UnBL. = number of the sequences that did not produce any hit; Prop. Grouped = proportion of grouped sequences (Grouped/(Grouped + Unique)).
*Species with more than one data set available are in bold characters.
$\dagger$ Without the identification of redundant sequences.
same laboratory, at the same time, for different species gave strikingly different estimates of the proportion of grouped sequences (Euphydryas aurinia, Parnassius apollo, Aphis fabae). Our enlarged series of data from screenings show the same lack of association between the applied protocols and the proportions of grouped sequences. Indeed, the same protocol was used for Busseola fusca, Cydia pomonella, E. aurinia, Ostrinia nubilalis, P. apollo, A. fabae, with the percentage of grouped sequences ranging from $2.8 \%$ to $55.1 \% ~\left(\chi^{2}=75.3, P<0.0001\right.$ ), and another protocol for Drupadia theda and Arhopala epimuta, where the two species were screened simultaneously, yielded $6.5 \%$ and $31.4 \%$ grouped sequences, respectively $\left(\chi^{2}=10.3\right.$,
$P=0.0013$ ). Thus, it is unlikely that the screening protocol is responsible for the high proportion of grouped sequences in some species (Table 2).
An obvious disadvantage of the screening data is the relatively small number of sequences available for each species. By increasing the number of investigated sequences some of the sequences classified as unique can become grouped. Thus the proportion of grouped sequences depends partially on the number of sequences available. This introduces a downwards bias in the estimation of the proportion of grouped sequences in small data sets. B. mori data provide a possibility to compare estimates of the proportion of grouped sequences based on screening
vs. WGS data. As expected, the proportion of grouped sequences is lower when estimated from the screening data ( $25.7 \%$ ) than from the WGS data ( $34.7 \%$ and $36.1 \%$ for the two data sets). However, these estimates are not significantly different ( $\chi^{2}=0.882, P=0.348 ; \chi^{2}=1.222$, $P=0.267$, for the two comparisons, respectively).
In spite of the downwards bias of the proportion of grouped sequences from screening data, we demonstrated the presence of sequence families in almost all studied species apart from three species for which only a few sequences were available. Nevertheless, by decreasing the stringency of the search for sequence similarities (i.e. blast with $e=1 \mathrm{E}-10$ instead of $e=1 \mathrm{E}-40$ ) Parnassius mnemosyne and Parnassius smintheus had $6.9 \%$ and $15.8 \%$ of grouped sequences, respectively. Only Zale galbanata (with only nine sequences available) remained apparently free from grouped sequences. Thus, there appears to be a common trend that varying and often high proportion of microsatellite sequences within Lepidoptera species cluster into families. Some of the microsatellites of the seven control (non-Lepidoptera) species also grouped into families, and WGS data sets indicated that this phenomenon is indeed not restricted to Lepidoptera, but it is present in all investigated taxa (Table 2).

GenBank entries of microsatellites Of the 415 insect species for which microsatellite sequences have been submitted to GenBank, $13 \%$ had at least one pair of grouped sequences. Note that redundant sequences were identified and discarded as before, in order to avoid a further possible bias of different teams of researchers working on the same species and finding the same microsatellite sequences. By selecting only species that have at least $10,20,30$ or 40 sequences available in GenBank, the proportion of species having at least two grouped sequences increased strongly ( $32 \%, 53 \%, 71 \%, 81 \%$, respectively). Undoubtedly, GenBank microsatellite entries are not representative of whole genomes as in most cases researchers submit only polymorphic loci for which successful amplification protocols were established. Furthermore, the number of sequences is very low for most of the species (i.e. only 58 insect species had a minimum of 20 microsatellite entries). In spite of these factors, which introduce a very strong downwards bias in the proportion of grouped sequences, a high proportion of species representing diverse insect orders had microsatellite sequence families.

Grouped vs. redundant sequences For all data sets, the same approach was used to identify grouped and redundant sequences. This is a necessary step for data originating from both screening and whole genome shotgun sequencing, in order to eliminate alleles of the same locus or multiple copies of the same allele. Our choice of the $95 \%$ cut-off is based on previously published results (for
discussion see Meglécz et al., 2004). Although, the complete genome sequences are nonredundant, we used the same method for identification of redundant sequences in order to obtain results comparable with the other data. Running the analysis with or without the identification of redundant sequences for complete genomes suggests that our procedure of eliminating redundant sequences and setting the cut-off limit to $95 \%$ is rather conservative. Indeed, a considerable portion of the sequences classified as redundant are in fact not redundant but represent different loci grouped into families. As a result, the proportions of grouped sequences within species are underestimated. However, as our principal aim was to compare the proportion of grouped sequences among the different data sets, the cutoff value is of little importance as long as it is identical for all analyses. Furthermore, demonstrating the presence of microsatellite sequence families with a conservative method gives an even stronger evidence of their occurrences. While using this conservative method, we managed to demonstrate the presence of microsatellite sequence families based on interlocus similarities of the flanking regions in a wide variety of insect species from data of diverse origins. Although the quantification of this phenomenon is difficult, our results indicate that the proportion of grouped microsatellites varies considerably among species and taxa. In particular, data from both partial genomic screens and WGS indicate that in many Lepidoptera the proportion of grouped sequences is particularly high (approximately $20 \%$ or more; Fig. 2).

## Association between microsatellites and interspersed repetitive elements

All sequences were screened for the presence of repetitive elements. When testing D. melanogaster microsatellites from the complete genome data set against the Drosophila repetitive elements bank, $6.4 \%$ of the sequences showed significant similarities to diverse repetitive elements. This proportion, however, was more than 20 times higher among grouped sequences ( $83.5 \%$ ), than among unique sequences $\left(3.5 \% ; \chi^{2}=21315 ; P<0.0001\right)$. Testing D. melanogaster microsatellite sequences against the Anopheles repetitive DNA bank gave similar but much less striking results: 1.1\% sequences showed similarities to repetitive elements in total, $1.9 \%$ among grouped sequences, $0.8 \%$ among uniques. Nevertheless, these proportions remained significantly different ( $\chi^{2}=19.4 ; P<0.0001$ ).

A similar tendency was observed for $A$. gambiae microsatellites. When testing against the Anopheles repetitive elements bank the proportion of sequences showing similarities to a repetitive element is $40.1 \%$ and $3.4 \%$ for grouped and unique sequences, respectively $\left(\chi^{2}=32142\right.$; $P<0.0001$ ), while against the Drosophila bank these proportions are $0.6 \%$ and $0.3 \%\left(\chi^{2}=54.0 ; P<0.0001\right)$. Thus, for a large proportion of grouped sequences in these


Figure 2. Proportion of grouped sequences estimated from different type of data. The first letters of the three-letter codes refer to the genus the second and the third are the first two letters of the species names.
two species, associations between repetitive elements and microsatellites have been demonstrated. Furthermore, our results indicate that repetitive element banks are rather specific for each species or genus, and the sensitivity of the detection of repetitive elements is strongly reduced for species in different genera even from the same order.

Screening microsatellite sequences from WGS data for repetitive elements yielded very similar results for all Drosophila (six species) and Anopheles species as described above (data not shown). However, only $0.75 \%(n=10000)$ microsatellites sequences of $A$. aegypti showed similarities to the Anopheles repetitive bank, and for all non-Diptera species this portion was even lower. This resulted in a low power of detecting the difference between the proportion of repetitive elements among grouped and unique sequences. In spite of this, the proportion of the sequences associated with repetitive elements was still significantly higher among grouped sequences than among unique ones in B. mori (data set B; against the Anopheles bank, $\chi^{2}=9.27$; $P=0.0023$ ).

A large number of studies indicate that microsatellites are more frequent in noncoding regions than in coding ones (Wang et al., 1994; Edwards et al., 1998; Metzgar et al., 2000; Li et al., 2002; Tóth et al., 2000). Only tri- and hexanucleotide repeats seem to be more frequent in coding regions (Morgante et al., 2002), which is probably the consequence of negative selection against frameshift mutations in the coding regions (Metzgar et al., 2000). It is also known that, in species with large genomes in particular, noncoding DNA is largely composed of repetitive elements (Kidwell \& Lisch, 2001). For example, $45 \%$ of the euchromatic DNA in humans corresponds to transposable elements (Smit, 1999; Lander et al., 2001), and the size of the genome of maize has been doubled as a result of retrotransposon insertions (SanMiguel \& Bennetzen, 1998). Association between microsatellites and transposable elements has been reported in nematodes (Hoekstra et al., 1997; Johnson et al., 2006), Dipterans (Fagerberg et al., 2001; Wilder \& Hollocher, 2001), rice (Akagi et al., 2001; Temnykh et al., 2001) and barley (Ramsay et al., 1999). Similarly, our study clearly demonstrates that a large proportion of Drosophila (six species) and A. gambiae grouped microsatellites are associated with diverse transposable
elements. Furthermore, we demonstrated the specificity of the repetitive elements banks to each species/genus. This indicates that the lack of significant association between transposable elements and microsatellites for the other investigated insect species is probably the result of the scarcity of information on the transposable elements in those specific species, rather than a true lack of association. Thus, it is likely that the presence of microsatellite families is at least partially the result of the association between microsatellites and interspersed repetitive DNA elements or their remnants.

There are at least three different mechanisms that might create this kind of microsatellite-transposable element association. (1) Microsatellites arise at random in the genome, but they are more likely to be removed from coding and control regions. In this case, the association would be incidental. However, with this mechanism, most microsatellites are expected to be found near interspersed repetitive elements and thus they should become grouped. Our results do not support this hypothesis. (2) If a microsatellite arises in a transposable element, the number of microsatellite loci can be considerably amplified by transposition. (iii) Transposons might take an active role in microsatellite origin if microsatellites arise during transposition. In the two latter cases, grouped microsatellites are expected to be associated with transposable elements more often than unique ones. This is supported by our results. However, distinguishing between these two scenarios requires more detailed analyses and it is beyond the scope of this paper.

## BLAST against baculovirus sequences

Lepidoptera are commonly infected by baculoviruses (Cory \& Myers, 2003). In order to test the hypothesis of non-insect amplified DNA, all Lepidoptera microsatellite sequences were bLASTed against genomes of baculoviruses. As these analyses did not produce any significant hits, it is unlikely that virus infection is a cause of the presence of sequences families in Lepidoptera.

## Implications on the use of microsatellites as markers

Although the use of grouped microsatellites as genetic markers is not impossible (Hoekstra et al., 1997), they are unlikely to be good candidates for single locus genotyping
purposes as flanking region similarities can lead to multiple locus amplifications and unclear banding patterns. Thus the variability among species in both the proportion of grouped sequences and the abundance of microsatellites may explain the large differences in the success/effort ratio of establishing reliable microsatellite markers for Lepidoptera and more generally for insects.

We are aware of two different methods to avoid clones containing grouped sequences. van't Hof et al. (2005) have established a method to eliminate grouped sequences as a last step of the screening process prior to sequencing. An alternative approach is to isolate singlecopy DNA from the genome prior to screening for microsatellites (Elsik \& Williams, 2001), as has recently been accomplished for Aedes japonicus (Widdel et al., 2005). Once the screening is done and sequences are available, it is strongly advised to check microsatellite flanking sequences for interloci similarities prior to primer design. The program, MicroFamily (Meglécz, 2007; http://www.up. univ-mrs.fr/Local/egee/dir/meglecz/MicroFamily.html) can accomplish this task. However, if only a low number of clones/sequences are available the fact that a sequence is found to be unique, does not necessarily indicate its uniqueness in the genome. The program Primer3 for designing primers (Rozen \& Skaletsky, 2000) has a highly useful option of screening sequences for interspersed repeats using human, rodent or Drosophila repetitive element banks. However, as shown above, the sensitivity of detecting interspersed repeats from a noncongeneric repetitive bank is low.

A further problem in Lepidoptera and in many other species (Keyghobadi et al., 1999; Harper et al., 2003; Meglécz et al., 2004; Zhang, 2004; van't Hof et al., 2005) is the frequent presence of null alleles, even for apparently clearly amplifying and interpretable microsatellite loci. This problem still has to be addressed in future research. Hardison et al. (2003) have demonstrated the covariance of substitution rate and transposition frequency among different genomic regions in human and mouse genomes. One possibility is that transposition frequencies and substitution rates also covary among species, such that some species or groups of species have both a high number of transposable elements and high average mutation rates. In these species, the association between microsatellites and transposable elements can lead to many microsatellites that are not unique; but, even if amplification is successful, the high mutation rate in the flanking regions may lead to a high likelihood of null alleles.

## Conclusions

Microsatellites are used as highly polymorphic, single locus, co-dominant markers. However, our results demonstrated that in insects, a variable proportion of microsatellites
cluster into sequence families based on similarities in their flanking region. In some species, this phenomenon, in association with a relatively low abundance of microsatellites, renders the isolation of reliable microsatellite markers particularly difficult.

The association between microsatellite sequence families and interspersed repetitive elements suggests a mechanism for the creation and/or amplification of microsatellites and thus the presence of such microsatellite families, and clearly indicates that the behaviour and evolution of microsatellite sequences can only be fully understood within a larger genomic context.

## Experimental procedures

## Cleaning and extraction of sequences

Sequences from screening partial libraries Microsatellite sequence data were collected from 23 species, including 16 Lepidoptera and seven other insect species (two Diptera, two Coleoptera, two Trichoptera, one Hemiptera). For each species, all readable, microsatellite-containing, sequences resulting from one or more screenings of partial genomic libraries were analysed (Table 3). Microsatellites with motifs of 1-6 bp were identified. For the motif of a single base at least eight, for all other motifs at least four uninterrupted repetitions were necessary to retain the sequence.

Sequences were edited by (1) replacing all characters other than ACGT by $N$; (2) by deleting the extremities if they contained more than two Ns in the 10 most extreme base pairs; and (3) by removing vector and adapter contamination if the sequence produced a BLAST hit (e=1E-3) against the UniVec vector base of NCBI (ftp://ftp.ncbi.nih.gov/pub/UniVec) or the adaptors/linkers used during microsatellite isolation.

GenBank microsatellite entries All microsatellite sequences of all insects were downloaded from GenBank (10 January 2006) and treated species by species, as described above.

Whole Genome Shotgun All contigs of WGS sequencing projects were downloaded from NCBI (www.ncbi.nlm.nih.gov/ Genomes; 21 October 2005) for 10 insect species [Lepidoptera: (1) Bombyx mori (data set A: ADDK01000001-01066482, Xia et al., 2004; data set B: BAAB01000001-01213289, Mita et al., 2004); Diptera: (2) Anopheles gambiae (AAAB0100000101069724, Holt et al., 2002), (3) Aedes aegypti (AAGE0100000101655164, The Institute for Genomic Research, Broad Institute, http://msc.tigr.org/aedes/release.shtml), (4) Drosophila melanogaster (AABU01000001-01002756, Celniker et al., 2002), (5) Drosophila simulans (AAGH01000001-01025284, Drosophila simulans Sequencing Consortium, Genome Sequencing Center at Washington University School of Medicine in St Louis, http:// genome.wustl.edu/genome_group_index.cgi), (6) Drosophila pseudoobscura(AADE01000001-01012826, Richards et al., 2005), (7) Drosophila yakuba (AAEU01000001-01013569, Genome Sequencing Center at Washington University School of Medicine in St Louis, http://genome.wustl.edu/genome_group_index.cgi), (8) Drosophila persimilis (AAIZ01000001-01026813, Broad Institute, http://www.broad.mit.edu/tools/data/data-vert.html),

Table 3. Number of available sequences ( $n$ ) and their accession numbers for each species from screening partial genomic libraries

| Species | Order | $n$ | Accession nos | References |
| :--- | :--- | :--- | :--- | :--- |
| Arctia caja | Lepidoptera | 86 | AJ809344-809406 | Anderson et al. (2006) |
|  |  |  | AJ829727-829729 |  |
| Arhopala epimuta | Lepidoptera | 60 | AJ867349-867385 | DQ380801-380855 |

(9) Drosophila sechelia (AAKO01000001-01021425, Broad Institute, http://www.broad.mit.edu/tools/data/data-vert.html); Hymenoptera: (10) Apis mellifera (AADG01000001-01018946, The Honeybee Genome Sequencing Consortium, 2006).

If sequences contained more than 10 consecutive Ns , the Ns were cut out and the sequences were not joined together. Then, all microsatellites containing at least four repeat units of di-, tri-, tetra-, penta- and hexanucleotide motifs, or at least eight single nucleotide repeats, with 150 bp flanking sequence on each side were extracted. If flanking regions of different microsatellites were overlapping (i.e. when two microsatellites were separated by less than 300 bp flanking sequences), they were pooled into the same
sequence to avoid redundancy. Thus, we obtained short sequences with one or more microsatellites in each and 150 bp flanking sequence on each extremity. These were used for further analyses and we refer to each such entity as a 'sequence'. In this way, 11 pools of sequences were obtained, one from each WGS data set. From each pool, 10000 sequences were randomly selected for comparison of the flanking regions.

Complete genomes All microsatellites from the complete genome of D. melanogaster (NT_037436, NT_033779, NT_033778, NT_033777, NC_004354, NC_004353; 23 July 2005; Adams et al., 2000) and A. gambiae (NC_004818,

NT_078265, NT_078266, NT_078267, NT_078268; 23 July 2005; Severson et al., 2004) were extracted as described for WGS sequences.

## Sequence analyses

Sequences were analysed separately for each species and for each type of sequence origin (i.e. screening partial genomic libraries, GenBank entries, WGS and complete genome). In this way we treated 23 data sets from screenings, 415 data sets from GenBank entries, 11 data sets from WGS and two data sets from complete genomes. Pure microsatellite repeat regions (as defined above) were replaced by Ns in order to examine only the flanking regions in the successive analyses. Each data set was blasted against itself with $e=1 \mathrm{E}-40$, thus all bLASTn results presented in this paper represent only intraspecies comparisons. Sequences were sorted into four categories based on the results of the BLASTn. They were classified as: (1) Unique if no similarities were observed to any other sequences of the same data set; (2) unBLASTable if sequence had no hits at all, not even with itself (if the flanking region is too short or repetitive, but not a perfect microsatellite, BLAST masks the region; thus the sequence becomes uninformative and does not align even with itself); or (3) Redundant, if the identity to another sequence was higher than $95 \%$ along the whole flanking sequence. All nonredundant sequences that produced a significant hit with at least one different sequence were classified as (4) Grouped. The proportion of grouped sequences was calculated after eliminating unblastable and redundant sequences (e.g. number of grouped sequences/(number of grouped sequences + number of unique sequences)). For the whole genome sequences, both the above procedure was applied as well as the same process without identifying 'redundant' sequences.

All these operations were conducted by Perl programs written by the first author (available upon request), MICROFAMILY (Meglécz, 2007), BLASTn-2.2.10 (ftp://ftp.ncbi.nih.gov/blast/executable/) and clustalW1. 83 (Higgins et al., 1991).

Repetitive sequence banks We scanned all the sequences of our data sets against databases of repetitive sequences of $D$. melanogaster and $A$. gambiae in order to detect the presence of interspersed repetitive elements. This analysis was performed using the program RepeatMasker2 (A.F.A. Smit and P. Green, unpublished data, http://repeatmasker.genome.washington.edu/RM/ webrepeatmaskerhelp.html) and its databases using default sensitivity parameters. The proportion of grouped vs. unique sequences that showed similarities to repetitive elements were compared by $\chi^{2}$ tests conducted by the R stats package, version 2.0.1 (R Development Core Team, 2004).

Comparison with baculovirus sequences All Lepidoptera microsatellite sequence data sets were BLASTed against the 19 genomes of baculoviruses available in GenBank (downloaded 13 December 2005):
NC_007383.1, Trichoplusia ni SNPV virus;
NC_007151.1, Chrysodeixis chalcites nucleopolyhedrovirus;
NC_005137.2, Choristoneura fumiferana defective nucleopolyhedrovirus;
NC_004778.3, Choristoneura fumiferana MNPV; NC_004117.1, Mamestra configurata nucleopolyhedrovirus B;
NC_003094.2, Helicoverpa armigera nuclear polyhedrosis virus; NC_004690.1 Adoxophyes honmai nucleopolyhedrovirus;

NC_005068.1, Cryptophlebia leucotreta granulovirus;
NC_001962.1, Bombyx mori nucleopolyhedrovirus;
NC_002169.1, Spodoptera exigua nucleopolyhedrovirus;
NC_005038.1, Adoxophyes orana granulovirus;
NC_004156.1, Helothis zea virus 1;
NC_004062.1, Phthorimaea operculella granulovirus;
NC_003102.1, Spodoptera litura nucleopolyhedrovirus;
NC_002816.1, Cydia pomonella granulovirus;
NC_002654.1, Heliocoverpa armigera nucleopolyhedrovirus G4;
NC_001982.1, Helicoverpa armigera stunt virus RNA 2;
NC_001973.1, Lymantria dispar nucleopolyhedrovirus;
NC_001623.1, Autographa californica nucleopolyhedrovirus.

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