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# Genetic diversity of populations of Merodon aureus and M. cinereus species complexes (Diptera, Syrphidae): integrative taxonomy and implications for conservation priorities on the Balkan Peninsula 

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#### Abstract

The genetic structure of 10 populations of the Merodon aureus group from the Balkan Peninsula was examined through allozyme electrophoresis and mitochondrial DNA sequencing of the cytochrome $c$ oxidase subunit I (COI). Six diagnosable cryptic taxa were identified within the morphologically defined species M. aureus Fabricius, 1805 and M. cinereus (Fabricius, 1794), with clear separation of the populations ( ( $($. aureus $\mathrm{A}+M$. aureus B$)+$ cinereus complex $)+$ M. aureus C$)$. The parsimony analysis of COI sequence data of the aureuscinereus complex using Merodon avidus A species as an outgroup resulted in two main clades, ( $M$. aureus $\mathrm{A}+M$. aureus B$)$ and ( $(M$. aureus $\mathrm{C}+M$. cinereus $\mathrm{B}+$ M. cinereus C$)+M$. cinereus A ), which differed on average by $5.7 \%$. The observed spatial distribution of the taxonomic diversity of the group suggested that these taxa originated from a common ancestral population in the Mediterranean. Identification of genetic uniqueness and genetic endemism emphasizes the importance of molecular markers and estimation of genetic diversity in recognition of conservation units. The primary goals of the conservation measures that we propose are the protection of phylogenetic lineages within the highly diverse $M$. aureus group taxa and conservation of the genetic variation through management of important areas.


[^0]Keywords Allozymes • MtDNA • Cryptic taxa • Genetic units • Genetic diversity • Conservation • Diptera • Merodon

## Introduction

The genus Merodon is the second largest genus of Palaearctic hoverflies (Diptera, Syrphidae) with more than 50 European species (Speight 2004), and the highest diversity in the Mediterranean region. The high endemicity on the Balkan (Vujić et al. unpublished) and Iberian (Marcos Garcia et al. 2007) Peninsulas is probably due to isolation of different districts during the Pleistocene glaciations. Studying phylogenetic relationships within, and genetic distinctiveness of the genus Merodon is challenging due to the high diversity, presence of cryptic species, taxonomic uncertainties, and unclear relationships among taxa.

An example of the great morphological variability exhibited in the genus Merodon is the aureus group of taxa with bi-colored tibiae, mentioned under two names in recent literature (Šimić and Vujić 1996; Van de Weyer and Dils 2002) for the central and south-eastern Europe: Merodon aeneus Megerle in Meigen, 1822 and M. cinereus (Fabricius, 1794). In the Biosystematic Database of World Diptera (BDWD), Thompson (2005) cited Merodon aureus as a senior synonym of M. aeneus, and we accept this synonymy. There are six more synonyms of M. aureus and four synonyms for $M$. cinereus cited in Peck (1988). Morphologically similar to M. aureus is a European taxon Merodon funestus (Fabricius, 1794). Populations of M. funestus also exhibit morphological variation in the length of antennae and leg coloration.

For identification of areas of genetic endemism of taxonomically diverse and challenging taxa, such as the
genus Merodon, it is important to delineate genetic units that are geographically discrete, often characterized by non-overlapping haplotype distributions and/or occurrence of unique allozyme markers (Moritz 1994). For example, the presence of unique alleles is a marker of distinct genetic units, implying existence of cryptic taxa with reproductive incompatibility, while species-specific alleles and their combinations in spatially subdivided populations probably reflect their origin in past events of gene pool fragmentation. However, different allozyme loci can diverge at different rates and the rate of divergence can vary through time, making it difficult to allocate a time since divergence based on genetic distance (Thorpe 1982). Molecular data of both nuclear and mitochondrial genes are a valuable source of information in taxonomic and evolutionary studies in hoverflies, especially when morphological characters are insufficient or not exclusive enough for identification of morphologically cryptic taxa (e.g. Milankov et al. 2005; Rojo et al. 2006).

The first goal of this study was to identify diagnosable taxa within available populations of the aureus group occurring on the Balkan Peninsula using allozyme markers and mitochondrial DNA cytochrome $c$ oxidase subunit I gene (mtDNA COI) sequences. The second aim was to quantify genetic variability in the populations of $M$. aureus and $M$. cinereus complexes and in M. funestus as a measure of an evolutionary potential, and analyze genetic and phylogenetic relationships among the taxa. Finally, as there are no plans that focus on the management of threatened hoverflies species on the Balkan Peninsula, conservation implications for the preservation of the genetic diversity of these taxa based on our findings are discussed.

## Material and methods

## Sample collection

Samples of 10 populations of the Merodon aureus group and M. funestus were collected from five regions (Fig. 1) on the Balkan Peninsula over multiple years (population code and number of collected specimens are given in Table 1). In addition, populations of M. avidus, comprising cryptic taxa M. avidus A and M. avidus B, from five different biogeographic regions on the Balkan Peninsula were sampled to be used as an outgroup (Milankov et al. 2001) (Table 1).

Specimens were initially identified to either the aureus or cinereus complex based on the morphology. Specimens from the M. aureus and M. cinereus complexes have dark legs except for pale knees, both ends of tibiae and tarsi in part. Pale hairs on thorax and abdomen, dull scutum and rough punctuation of scutum and tergites are characteristic


Fig. 1 Map of the Balkan Peninsula. Origin of the analyzed populations: 1. Dubašnica Mountain (DUB Mt), E $21^{\circ} 59^{\prime}$, N $44^{\circ} 01^{\prime}$ (Serbia); 2. Kopaonik Mountain (KOP Mt), E $20^{\circ} 40^{\prime}$, N $43^{\circ} 15^{\prime}$ (Serbia); 3. Durmitor Mountain (DUR Mt), E $19^{\circ} 00^{\prime}$, N $43^{\circ} 11^{\prime}$ (Montenegro); 4. Morinj (MOR), E $18^{\circ} 40^{\prime}$, N $43^{\circ} 29^{\prime} 30^{\prime \prime}$ (Montenegro); 5. Prokletije Mountain (PRO Mt), E $19^{\circ} 50^{\prime}$, N $42^{\circ} 32^{\prime}$ (Serbia); 6. Šar planina Mountain (ŠAR Mt), E $21^{\circ} 05^{\prime}$, N $42^{\circ} 12^{\prime}$ (Serbia). 7. Mavrovo Lake (MAV Lk), E $20^{\circ} 44^{\prime} 30^{\prime \prime}$, N 41 $38^{\prime} 30^{\prime \prime}$ (FYR Macedonia); 8. Pindos Mountain (PIN Mt), E 20 ${ }^{\circ} 37^{\prime}$, N $39^{\circ} 14^{\prime}$ (Greece). Locality Morinj is the territory with intermixed evergreen Mediterranean maritime woodlands and maquis and Submediterranean oak woodlands at the Adriatic Sea in the Mediterranean area. Kopaonik, Durmitor, Prokletije and Sar-planina are high Dinaric mountains with different types of biomes, from deciduous woodlands at low altitudes (up to 700 m ), European coniferous boreal woodlands at the higher altitudes, to the biome of alpine and high Nordic rockgrounds pastures and snow patches in the highest zone
for the M. aureus complex, contrary to dark hairs on thorax and abdomen, shiny scutum and finer punctuation of scutum and tergites in the $M$. cinereus complex.

Allozyme analysis

All sampled populations (a total of 193 specimens) were included in the allozyme analysis in order to determine molecular markers that could be used to identify the studied taxa (Table 1). Allozyme data of the outgroup, M. avidus (Milankov et al. 2001) were used for constructing a rooted dendrogram.

Genetic variability was studied by standard 5\% polyacrylamide gel electrophoresis, following Munstermann (1979) (FUM, GPD, GPI, HAD, HK, IDH, MDH, ME, PGM, SOD) and Pasteur et al. (1988) (AAT), with slight

Table 1 Species, population code and collection location of samples included in this study

| Species | Locality | Locality code | $n$ | Sample code | GenBank accession No. | Collection date | Collectors |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| M. aureus A | Durmitor Mt. | DUR | 11 | VM351-COI <br> VM407-COI | $\begin{aligned} & \text { DQ387912 } \\ & \text { DQ387919 } \end{aligned}$ | 20/06/1998 | Milankov, V., Tanurdžić, M., Radović, D., Vujić, A. |
|  | Morinj | MOR | 27 | $\begin{aligned} & \text { VM326-COI } \\ & \text { VM354-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387906 } \\ & \text { DQ387913 } \end{aligned}$ | 16/05/1998 | Dragišić, S., Mićić R., Radenković, S., Vujić, A. |
| M. aureus B | Kopaonik Mt. | KOP | 23 | $\begin{aligned} & \text { VM333-COI } \\ & \text { VM348-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387908 } \\ & \text { DQ387910 } \end{aligned}$ | 06/06/1998 | Vujić, A. |
|  |  |  |  | $\begin{aligned} & \text { VM347-COI } \\ & \text { VM332-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387921 } \\ & \text { DQ387907 } \end{aligned}$ | 20/06/1998 | Milankov, V., Tanurdžić, M., Radović, D., Vujić, A. |
| M. aureus C | Durmitor Mt. | DUR | 21 | $\begin{aligned} & \text { VM349-COI } \\ & \text { VM366-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387911 } \\ & \text { DQ387917 } \end{aligned}$ | 28/07/1998 | Milenković, D., Vujić, A. |
|  | Morinj | MOR | 6 | $\begin{aligned} & \text { VM340-COI } \\ & \text { VM355-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387909 } \\ & \text { DQ387914 } \end{aligned}$ | 21/08/1997 | Milankov, V., Vujić, A. |
|  |  |  |  | VM428-COI | DQ387920 | 25/09/1997 | Vujić, A. |
| M. cinereus A | Kopaonik Mt. | KOP | 37 | $\begin{aligned} & \text { VM313-COI } \\ & \text { VM314-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387902 } \\ & \text { DQ387903 } \end{aligned}$ | 15/08/1997 | Milankov, V., Milenković, D., Stamenković, J., Vujić, A. |
| M. cinereus B | Durmitor Mt. | DUR | 37 | $\begin{aligned} & \text { VM315-COI } \\ & \text { VM316-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387904 } \\ & \text { DQ387905 } \end{aligned}$ | 28/07/1998 | Milenković, D., Vujić, A. |
|  | Prokletije Mt. | PRO | 4 | VM406-COI | DQ387918 | 26/07/1994 | Milankov, V., Radenković, S., Radišić, P., Radnović, D., Vujić, A. |
| M. cinereus C | Šar planina Mt. | ŠAR | 5 | $\begin{aligned} & \text { VM362-COI } \\ & \text { VM363-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387915 } \\ & \text { DQ387916 } \end{aligned}$ | 16/07/1995 | Radenković, S., Radišić, P., Vujić,A. |
| M. funestus | Morinj | MOR | 22 | $\begin{aligned} & \text { VM338-COI } \\ & \text { VM458-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387901 } \\ & \text { DQ387900 } \end{aligned}$ | $\begin{aligned} & 31 / 08 / 1997 \\ & 16 / 05 / 1998 \end{aligned}$ | Milankov, V., Vujić, A. <br> Dragišic, S., Míćić R., <br> Radenković, S., Vujić, A. |
| M. avidus $\mathrm{A}^{\mathrm{a}}$ | Morinj | MOR | 30 | $\begin{aligned} & \text { VM566-COI } \\ & \text { VM581-COI } \end{aligned}$ | DQ387897 <br> DQ387896 | 18/06/1998 <br> 31/08/1998 | Milenković, D., Vujić, A. Milankov, V., Vujić, A. |
|  | Dubašnica Mt. | DUB | 14 | - |  | 06/97; 08/97; 09/97 | Dević, D., Milankov V., <br> Milenković, D., Radenković,S., Šimić,S.,Vujić,A. |
|  | Pindos Mt. | PIN | 9 | - |  | 20/05/1997 | Radenković, S., Šimić, S.,Vujić, A. |
| M. avidus $\mathrm{B}^{\mathrm{a}}$ | Dubašnica Mt. | DUB | 21 | - |  | $\begin{aligned} & \text { 06/97; 08/97; } \\ & \text { 09/97; 07/98 } \end{aligned}$ | Dević, D., Milankov V., Milenković, D., Radenković,S., Šimić,S.,Vujić,A. |
|  | Durmitor Mt. | DUR | 9 | $\begin{aligned} & \text { VM589-COI } \\ & \text { VM605-COI } \end{aligned}$ | $\begin{aligned} & \text { DQ387898 } \\ & \text { DQ387895 } \end{aligned}$ | 20/06/199820/06/1998 | Milankov, V., Tanurdžić, M., Radović, D., Vujić, A. |
|  | Mavrovo Lake | MAV | 17 | - |  | 10/07/1998 | Radenković, S., Šimić, S.,Vujić, A. |
|  | Pindos Mnt. | PIN | 17 | - |  | 20/05/1997 | Radenković, S., Šimić, S.,Vujić, A. |

Number of specimens ( $n$ ) used for allozyme analysis and sequences amplified for molecular analysis are indicated
${ }^{\text {a }}$ Merodon avidus allozyme data were published in: Milankov et al. (2001)
modifications (Milankov 2001) (Table 2). The electrophoresis of individuals from different populations was conducted in the same gel for direct interspecific comparison. Loci were numbered and alleles marked alphabetically with respect to order of increasing anodal migration. Two
loci Sod-2 and Sod-3 were not available for comparison of populations of the aureus group and the outgroup.

Genotype and allele frequencies were calculated directly from the observed banding patterns based on the genetic interpretation of zymograms. Calculated parameters of

Table 2 Enzymes and loci investigated (EC number and name), buffer systems and tissue extract used in electrophoresis

| Enzyme | Abb. | E.C. Number ${ }^{\text {a }}$ | Loci | Buffer ${ }^{\text {b }}$ | Tissue extract ${ }^{\text {c }}$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| aspartat amino transferases | AAT | 2.6.1.1 | Aat | TC | head |
| fumarate hydratase | FUM | 4.2.1.2. | Fum | TC | head |
| glycerol 3-phosphate dehydrogenase | GPD | 1.1.1.8. | Gpd-2 | TC | thorax |
| glucosephosphate isomerase | GPI | 5.3.1.9. | Gpi | TBE | thorax |
| b-hydroxy acid dehydrogenase | HAD | 1.1.1.30. | Had | TC | thorax |
| hexokinase | HK | 2.7.1.1. | Hk-2 | TBE | head |
|  |  |  | Hk-3 |  | thorax |
| isocitrate dehydrogenase | IDH | 1.1.1.42. | Idh-2 | TC | thorax |
| malate dehydrogenase | MDH | 1.1.1.37. | Mdh-1 | TC | head |
|  |  |  | Mdh-2 |  | head |
| malic enzyme | ME | 1.1.1.40. | Me | TBE | head |
| phosphoglucomutase | PGM | 2.7.5.1. | Pgm | TBE | thorax |
| superoxide dismutase | SOD | 1.15.1.1. | Sod-1 | TBE | thorax |
|  |  |  | Sod-2 |  |  |
|  |  |  | Sod-3 |  |  |

a EC number-Enzyme Commision
b TC buffer-1 M Tris-citric buffer $\mathrm{pH}=7.1$; TBE buffer- 1 M Tris-boric-EDTA $\mathrm{pH}=8.9$
c Added in loading buffer ( $20 \%$ sucrose, Triton X-100 [0.5\%], Tris-citrate pH 7.0 electrode buffer and trace amount of bromphenol blue tracking dye)
population genetic structure were corrected using Levene's (1949) formula for small samples. Divergence among identified genetic units within $M$. aureus and $M$. cinereus complexes was examined using pairwise Wright's $F_{S T}$ values (Weir 1996) as an estimate of the proportion of the total genetic variance. Statistical analyses of allozyme data were performed using the computer program BIOSYS-2 (Swofford and Selander 1989). Diagnostic value of allozymes was calculated after Ayala and Powell (1972). A locus was considered diagnostic if the probability for correct identification of individuals belonging to the M. aureus group was $100 \%$ (complete differentiation) or higher than $95 \%$ (frequency of a common allele in one of two species is less than 0.05). Loci with overlapping allelic frequencies of 0.15 and 0.20 (correct diagnosis with at least $85 \%$ and $80 \%$ probability, respectively), or a combination of discriminatory loci, were also used for distinguishing species.

In the pilot study of gene-enzyme varibaility, diagnostic loci and cryptic species were detected within the morphologically defined $M$. aureus and $M$. cinereus species. Based on fixed allozyme diferences associated with morphological traits (see Results), we divided sympatric populations of M. aureus into two putative taxa: M. aureus A and M. aureus C. Within morphologically defined M. aureus species, a population from Kopaonik Mt. was identified as a specific genetic unit, M. aureus B. Diagnostic allozyme loci allowed us to determine allopatric populations of M. cinereus group as the cryptic species M. cinereus A, $M$. cinereus B and $M$. cinereus C .

## DNA sequencing

We obtained a 720 bp fragment of the mtDNA COI gene, corresponding to nucleotide positions 2233-2952 in Drosophila yakuba sequence (Clary and Wolstenholme 1985). A total of 24 specimens from all 10 populations of the M. aureus group, M. funestus and four specimens of the M. avidus outgroup from two localities (Table 1) were used for sequencing (all sequences have been deposited in GenBank, Accession numbers of the analyzed specimens are listed in Table 1). DNA was extracted from legs or other parts of the fly remaining after allozyme electrophoresis using the Nucleospin Tissue DNA extraction kit (Machery-Nagel, Düren, Germany) following the manufacturer's protocols and then re-suspended in $50 \mu \mathrm{l}$ of ultra-pure water. Remains of specimens, including male genitalia, used for the morphological studies and for DNA extraction are deposited at the Finnish Museum of Natural History (Helsinki, Finland).

PCR reactions were carried out in $25 \mu \mathrm{l}$ reaction aliquots containing $2 \mu \mathrm{l}$ DNA extract, $1 \mu \mathrm{l}$ of each primer (at $10 \mathrm{pmol} / \mu \mathrm{l}$ ), $0.25 \mu \mathrm{l}$ of DNA polymerase ( $5 \mathrm{U} / \mu \mathrm{l}$ ), $2 \mu \mathrm{l}$ 2.5 mM MgCl 2 , $2.5 \mu \mathrm{l} 10 \times$ Buffer II (MBI Fermentas, St. Leon-Rot, Germany) and $4 \mu \mathrm{l} 200 \mathrm{mM}$ dNTP (GeneAmp, Applied Biosystems, Foster City, CA, USA) and ultra-pure water. Thermocycler conditions were initial denaturing at $95^{\circ} \mathrm{C} 2 \mathrm{~min}$, 29 cycles of 30 s denaturing at $94^{\circ} \mathrm{C}, 30 \mathrm{~s}$ annealing at $49^{\circ} \mathrm{C}, 2 \mathrm{~min}$ extension at $72^{\circ} \mathrm{C}$, followed by a final extension of 8 min at $72^{\circ} \mathrm{C}$. The
universally conserved primers used for amplifying and sequencing the COI fragment ( 720 bp ) were the forward primer C1-J-2183 ( $5^{\prime}$-CAA CAT TTA TTT TGA TTT TTT GG- $3^{\prime}$ ) (alias JERRY) and the reverse primer TL2-N-3014 ( $5^{\prime}$-TCC AAT GCA CTA ATC TGC CAT ATT A-3') (alias PAT) (Simon et al. 1994). PCR products were purified using the GFX PCR Purification Kit (GE Healthcare Biosciences, Little Chalfont, UK) and then sequenced (with the PCR primers) in both directions using the Big Dye Terminator Cycle Sequencing Kit (version 1.1) (Applied Biosystems, Foster City, CA, USA) at onefourth of the recommended volumes on an ABI PRISM 377 (Applied Biosystems, Foster City, CA, USA) semiautomated DNA sequencer. The sequences were edited for base-calling errors and assembled using Sequence Navigator ${ }^{\mathrm{TM}}$ (version 1.01) (Applied Biosystems, Foster City, CA, USA).

## Genetic and phylogenetic analyses

Nei's genetic distance and identity (1978) were used to compare gene frequencies among species or populations, and to construct a dendrogram using clustering with the Unweighted Pair Group Method with Arithmetic Average (UPGMA) (Sneath and Sokal 1973). Prevosti's distance (Wright 1978) was employed to estimate phylogenetic relationships through the Wagner procedure (Farris 1972) rooted tree.

Alignment of COI sequences was trivial due to lack of indels and was done by eye. Parsimony analysis was performed using NONA (Goloboff 1999) and spawn with the aid of Winclada (Nixon 2002), using heuristic search algorithm with 1000 random addition replicates (mult*1000), holding 100 trees per round (hold/100), maxtrees set to 100,000 and applying TBR branch swapping. All base positions were treated as equally weighted characters.

## Results

Species boundaries

## Morphological diagnostic traits

Extensive variability observed in the coloration of hairs on eyes, legs, thorax, and abdomen; and the structure of tegument on scutum and tergites, indicated possible occurrence of cryptic taxa within the analyzed species. Based on morphological characters of the studied populations of Merodon aureus and M. cinereus occurring on the Balkan Peninsula, four morphotypes could be distinguished (Table 1):

1. Merodon aureus A and B, morphologically inseparable: abdomen and mesonotum with dense, yellowreddish hairs in male; female abdomen with shorter, mixed pale and black hairs, tergites II-IV with more or less clear stripes of polinosity with pale pilosity; tegument of mesonotum and tergites with golden tomentum; hind femora pale haired; at least upper half of eyes black haired. Analyzed populations were from Durmitor Mt., Morinj (M. aureus A DUR, M. aureus A MOR) and Kopaonik Mt. (M. aureus B KOP).
2. Merodon aureus C : abdomen and mesonotum with dense, yellow-reddish hairs in males; female abdomen with shorter, mixed pale and black hairs, tergites II-IV with more or less clear pollinose stripes; tegument of mesonotum and tergites with golden tomentum, but less than in Merodon aureus A and B: hind femora with black hairs on apical third; eyes pale haired, exceptionally with a few black hairs on dorsal corner. Analyzed populations were from Durmitor Mt. (DUR) and Morinj (MOR).
3. Merodon cinereus A and C, morphologically inseparable: mesonotum with pale hairs in anterior half and black in posterior (from only few to almost completely black hairs); female abdomen with shorter, mixed pale and black hairs, tergites II-IV with more or less clear pollinose stripes; tegument of mesonotum and tergites finely punctured, shiny; tergites 2 and 3 in males pale haired; tergite 4 with at least few black hairs (in $M$ cinereus A usually completely black haired); female abdomen with shorter, mixed pale and black hairs, tergites II-IV with more or less clear pollinose stripes. Analyzed populations were from Kopaonik ( $M$ cinereus A KOP) and Šar-Planina Mts. (M. cinereus C ŠAR).
4. Merodon cinereus B: mesonotum with pale hairs in anterior half and black in posterior (from only few to almost completely black Merodon cinereus A and C; tegument of mesonotum and tergites finely punctured, shiny; tergites $2-4$ in males pale haired with two stripes of black hairs; female abdomen with shorter, mixed pale and black hairs, tergites II-IV with more or less clear pollinose stripes. Analyzed populations were from Durmitor (DUR) and Prokletije Mts (PRO).

Taxa recognised as new to science in present study will be described in a separate taxonomic paper (Vujić et al. unpublished).

## Diagnostic allozyme loci

Based on diagnostic allozyme loci and fixed allelic differences, six well-defined genetic units were recognized within the aureus and cinereus complexes. Three taxa were
delineated within the morphologically defined M. aureus species. Based on the alleles at the diagnostic loci (Aat, Had, Me and Sod-1) M. aureus C was distinguishable from M. aureus A and M. aureus B. The allele Had ${ }^{m}$ within the M. aureus complex defined a group comprising M. aureus A and M. aureus B. The cladogram resolved these taxa as sister groups. A specific allele of diagnostic significance $\left(\mathrm{Pgm}^{c}\right)$ was observed in the population of the cryptic species M. aureus B from Kopaonik Mt. (Table 3). Diagnostic allozyme loci allowed identification of allopatric populations of $M$. cinereus group as cryptic species M. cinereus A, M. cinereus B and $M$. cinereus C . Identification and delimitation of $M$. cinereus A and $M$. cinereus B was possible with a $100 \%$ probability based on the genotypes at the Had locus. Furthermore, populations could be identified based on the specific allelomorphs at the Gpi and Me loci. Alleles $G p i^{k}$ and $M e^{l}$ defined M. cinereus B PRO and M. cinereus C ŠAR populations, respectively (Table 3 ).

Unique alleles were also used to assess the spatial variability between conspecific populations and interpopulation variation. Some private alleles registered within the M. aureus complex ( $\mathrm{Me}^{j}, \mathrm{Me}^{m}, \mathrm{Pgm}^{c}, \mathrm{Fum}^{c}, \mathrm{Gpi}^{i}, \mathrm{Had}^{n}$ ) were common alleles with $M$. desuturinus Vujić, Šimić et Radenković, 1995 (Milankov et al. unpublished), M. cinereus B and/or M. funestus. Likewise, some unique alleles within the M. cinereus complex (Aat ${ }^{a}, \mathrm{Fum}^{c}, \mathrm{Gpi}^{a}, \mathrm{Had}^{n}$, Mdh-2 ${ }^{g}, M e^{i}, P g m^{h}$ and Sod-1 ${ }^{e}$ ) were shared with M. aureus A, M. aureus B, M. funestus, M. aureus C and/or M. desuturinus (Milankov et al. unpublished).

## Mitochondrial COI diagnostic haplotypes

A total of 10 haplotypes were found in 22 analyzed individuals. There were 86 variable positions defining haplotypes. Haplotypes formed two groups of the aureus and cinereus complexes that differed from one another by 38 or more nucleotide changes. One haplotype corresponded to M. aureus B (haplotype III), one to M. aureus C MOR, M. cinereus B and $M$. cinereus C (haplotype VII), one to M. cinereus A (haplotype VIII), two to M. aureus A (I and II haplotypes), two to M. funestus (IX and X haplotypes) and four to M. aureus C (IV, V, VI and VII haplotypes), which were subdivided into two main lineages (Fig. 2).

Genetic diversity and divergence

## Population genetic structure

A total of 44 allozymes were registered in all populations and across all loci. Within the aureus complex of populations, the greatest number of alleles was observed at Morinj
locality (Table 3). Among the populations from the cinereus complex, M. cinereus A KOP had the greatest number of allelomorphs (Table 3). The largest number of alleles and genotypes were identified at the Me (6) and Had (7) loci, respectively (Table 3).

Analysis of parameters of population genetic structure showed differences among the analyzed species (Table 4). The mean number of alleles per locus $(A)$ ranged from 1.1 to 1.5 , and frequency of polymorphic loci $(P)$ was the lowest in the population of $M$. cinereus C and the highest in M. aureus C and $M$. funestus. Out of 15 analyzed loci, only Mdh-1. Sod-2 and Sod-3 isozyme loci were monomorphic (with a common allele) in all 10 studied populations. Average observed heterozygosity $\left(H_{o}\right)$ in all populations was lower than expected heterozygosity $\left(H_{e}\right)$ (Table 4). Heterozygous genotypes were identified only at the Gpi, Had and Sod-1 loci, however none were registered in populations of M. aureus B and M. funestus. At the Gpi locus genotype $G p i^{a / j}$ ( 0.045 frequency) was noted in M. cinereus B DUR. A common heterozygote $H a d^{m / q}$ was detected in M. cinereus A KOP $(0.459)$ and $M$. cinereus C ŠAR (0.200) populations as well as one unique $\mathrm{Had}^{n / r}$ in M. cinereus B DUR (0.027). M. aureus A DUR and M. aureus A MOR, and M. aureus C DUR and M. aureus C MOR had species-specific heterozygotes at Sod-1 locus (Sod-1 $1^{f / h}: 0.429$ and 0.400 , Sod- $1{ }^{d / e}: 0.313$ and 0.500 , respectively). Temporal distribution of alleles at the Gpi, $M e$ (allele $G p^{k}, M e^{k}, M e^{i}$ in "summer" generation), Had, $I d h-2$ and Sod-1 loci (Had ${ }^{q}, I d h-2^{\mathrm{b}}, \operatorname{Sod}-1^{e}$ in "spring" generation) was observed in the $M$. funestus MOR population.

## Interspecific divergence

The greatest average genetic distance $(0.89 \pm 0.07)$ was between $M$. funestus and the group consisting of taxa of the aureus and cinereus complexes (group II in Fig. 3). The average genetic distance among the analyzed aureus and cinereus taxa was $0.22 \pm 0.07$ (group I). Within the aureus complex, two lineages were observed: closely related M. aureus A and $M$. aureus B species (average genetic distance within this lineage was $0.06 \pm 0.05$ ), and $M$. aureus $\mathrm{C}(0.33 \pm 0.02$ distance from the $M$. aureus A and M. aureus B lineage). When the outgroup was included, the highest $D$-value was detected for avidus populations and the M. funestus species comparison (cluster IV: $1.99 \pm 0.08$ ) (Fig. 3). Analysis of allozyme variability and genetic relationships among taxa of the aureus group and avidus outgroup found common alleles and genotypes at the Gpi, Hk-2, Hk-3, Mdh-2 and Pgm loci, and the M. aureus A species was the closest to subgeneric outgroup populations $(D=0.86-1.01)$. Indeed, the UPGMA

Table 3 Total number of detected alleles ( $n=$ number of analyzed individuals) and allelic frequency at variable loci in the $M$. aureus group and M. funestus (common alleles with M. avidus outgroup are underlined)

| Species | M. aureus A |  | M. aureus B KOP | M. aureus C |  | M. cinereus A KOP | M. cinereus C ŠAR | . cinereus B |  | M. funestus MOR |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Population ${ }^{\text {c }}$ | DUR | MOR |  | DUR | MOR |  |  | DUR | PRO |  |


| Locus | Allele |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Aat | $\mathrm{A}^{\text {b }}$ | - | - | - | - | - | - | - | $0.027^{\text {a }}$ | - | - |
|  | B | 1.000 | 1.000 | 1.000 | 0.200 | 0.167 | 1.000 | 1.000 | 0.973 | 1.000 | - |
|  | C | - | - | - | 0.800 | 0.833 | - | - | - | - | 1.000 |
| Fum | A | 0.909 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 0.750 | - |
|  | C | $0.091{ }^{\text {a }}$ | - | - | - | - | - | - | - | $0.250^{\text {a }}$ | 1.000 |
|  | E | - | - | - | - | - | - | - | - | - | - |
| Gpd-2 | B | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | - |
|  | $\mathrm{D}^{\text {b }}$ | - | - | - | - | - | - | - | - | - | 1.000 |
|  | H | - | - | - | - | - | - | - | - | - | - |
| $G p i$ | $\mathrm{A}^{\text {b }}$ | - | - | - | - | - | - | - | $0.023^{\text {a }}$ | - | - |
|  | I | - | - | - | - | $0.167{ }^{\text {a }}$ | - | - | - | - | 0.300 |
|  | J | 0.727 | 0.684 | 0.526 | 1 | 0.500 | 0.909 | 0.800 | $0.977{ }^{\text {a }}$ | - | 0.600 |
|  | K | 0.273 | 0.316 | 0.474 |  | $0.333{ }^{\text {a }}$ | 0.091 | 0.200 | - | $1.000^{\text {a }}$ | 0.100 |
| Had | H | - | - | - | - | - | - | - | - | - | - |
|  | M | 1.000 | 1.000 | 1.000 | - | - | 0.716 | 0.500 | - | - | - |
|  | N | - | - | - | $0.059{ }^{\text {a }}$ | - | - | - | $0.014{ }^{\text {a }}$ | - | 0.882 |
|  | Q | - | - | - | 0.941 | 1.000 | 0.284 | 0.500 | - | - | 0.118 |
|  | $\mathrm{R}^{\text {b }}$ | - | - | - | - | - | - | - | 0.986 | 1.000 | - |
| Hk-2 | $\mathrm{A}^{\text {b }}$ | - | - | - | - | - | - | - | - | - | 1.000 |
|  | C | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 |  |
| $H k-3$ | $\mathrm{A}^{\text {b }}$ | - | - | - | - | - | - | - | - | - | 1.000 |
|  | C | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 |  |
| Idh-2 | A | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 1.000 | 0.818 |
|  | $\mathrm{B}^{\text {b }}$ | - | - | - | - | - | - | - | - | - | 0.182 |
| Mdh-2 | E | 1.000 | 0.962 | 1.000 | 1.000 | 0.750 | 1.000 | 1.000 | 1.000 | 0.500 |  |
|  | G | - | $0.038{ }^{\text {a }}$ | - | - | $0.250{ }^{\text {a }}$ | - | - | - | $0.500^{\text {a }}$ |  |
|  | $\mathrm{I}^{\text {b }}$ | - | - | - | - | - | - | - | - | - | 1.000 |
| Me | I | 0.909 | 1.000 | 0.870 | - | - | 0.194 | - | - | - | 0.050 |
|  | $\mathrm{J}^{\text {b }}$ | - | - | 0.043 | - | - | - | - | - | - | - |
|  | K | $0.091{ }^{\text {a }}$ | - | 0.087 | 0.875 | 0.400 | 0.722 | - | 1.000 | 0.333 | 0.300 |
|  | L | - | - | - | 0.125 | 0.400 | 0.083 | 1.000 | - | $0.667^{\text {a }}$ | - |
|  | M | - | - | - | - | $0.200^{\text {a }}$ | - | - | - | - | 0.650 |
| Pgm | $\mathrm{C}^{\text {b }}$ | - | - | 1.000 | - | - | - | - | - | - | - |
|  | F | 1.000 | 1.000 | - | $0.267^{\text {a }}$ | - | 0.568 | 1.000 | 1.000 | 1.000 | 1.000 |
|  | H | - | - | - | 0.733 | 1.000 | 0.432 | - | - | - | - |
| Sod-1 | D | - | - | - | 0.281 | 0.250 | 1.000 | 1.000 | 1.000 | 0.667 | - |
|  | E | - | - | - | 0.719 | 0.750 | - | - | - | $0.333{ }^{\text {a }}$ | 0.053 |
|  | F | 0.857 | 0.523 | 1.000 | - | - | - | - | - | - | 0.947 |
|  | $\mathrm{H}^{\text {b }}$ | 0.143 | 0.477 | - | - | - | - | - | - | - | - |
| \# Alleles |  | 17 | 18 | 18 | 20 | 22 | 20 | 17 | 18 | 19 | 22 |
| ( $n$ ) |  | 11 | 17 | 23 | 21 | 6 | 37 | 5 | 37 | 4 | 22 |

[^1]

Fig. 2 The most parsimonious tree based on mitochondrial data, length 158 steps, $\mathrm{CI}=0.87, \mathrm{RI}=0.96$. Filled circles $=$ nonhomoplasious changes, open circles $=$ homoplasious changes
dendrogram using Nei's Genetic Distance (1978) and Wagner procedure dendrogram using Prevosti's distance (Figs. 4, 5) supported the closest relationship of M. aureus A and $M$. aureus B species pair: ( ( $($. aureus $\mathrm{A}+M$. aureus B) + cinereus complex) + M. aureus C), while M. funestus was the most divergent species in comparison with taxa of the aureus and cinereus complexes.

Genetic differentiation among newly defined taxa within the $M$. aureus and $M$. cinereus complexes was quantified by $F_{S T}$ parameter as well. Genetic divergence between $M$. aureus A and $M$. aureus $\mathrm{B}\left(F_{S T}=0.48\right)$ resulted from the presence of species-specific alleles at the Pgm locus $\left(F_{S T}=0.48\right)$ and variance of the allelic frequencies at $\operatorname{Sod}-1\left(F_{S T}=0.36\right)$, $M e\left(F_{S T}=0.14\right)$ and $F u m$ $\left(F_{S T}=0.05\right)$. Greater value of genetic differentiation $\left(F_{S T}=0.73\right)$ was observed on the level of the aureus complex and was caused by allelic differences of Had $\left(F_{S T}=0.95\right)$, Pgm $\left(F_{S T}=0.84\right)$, Sod-1 $\left(F_{S T}=0.76\right)$, Aat $\left(F_{S T}=0.76\right), M e\left(F_{S T}=0.66\right), G p i\left(F_{S T}=0.31\right)$ and $M d h-$ $2\left(F_{S T}=0.29\right)$ loci. Among populations of the cinereus complex the observed $F_{S T}$ value ( $F_{S T}=0.59$ ) was a result of the allelic frequency differences at $\operatorname{Had}\left(F_{S T}=0.74\right)$, Gpi $\left(F_{S T}=0.60\right), M d h-2\left(F_{S T}=0.59\right), M e\left(F_{S T}=0.51\right)$, Sod-1 $\quad\left(F_{S T}=0.40\right), \quad \operatorname{Pgm} \quad\left(F_{S T}=0.38\right) \quad$ and $\quad$ Fum ( $F_{S T}=0.26$ ) loci.

## MtDNA diversity and parsimony analysis

Sequence divergences (uncorrected $p$ divergence in \%) within and between conspecific populations ranged from $0.14 \%$ (between haplotypes I and II in M. aureus A MOR), $0.29 \%$ (IV and V in M. aureus C DUR), $0.42 \%$ (IX and X in M. funestus MOR) to $0.85 \%$ (VI and VII in M. aureus C MOR) and from $0.14 \%$ (M. aureus A) to $0.28-1.13 \%$ (M. aureus C ), respectively. Within the aureus $\mathrm{A}+$ aureus B clade there was $1.19 \%$ divergences on average. The unique haplotype of $M$. cinereus C (VIII) differed from ( $M$. aureus $\mathrm{C}+$ M. cinereus $\mathrm{B}+M$. cinereus C ) clade by $0.75 \%$ on average. Haplotype VII was the most common and widely distributed (M. aureus C MOR, M. cinereus C ŠAR, M. cinereus B DUR, M. cinereus B PRO) and differed from other members of its lineage by two (IV haplotype) to six (VI) and from another lineage by 40 (III) to 43 (II) nucleotide changes. Within the aureus group as a whole, haplotypes of M. funestus (IX, X) differed the most by 52 (II) to $71(\mathrm{~V})$ compared with the members of the aureus and cinereus complexes (Table 5). Sequence divergence between M. funestus and aureus/cinereus complexes as well as between M. avidus group and M. aureus group averaged $8.87 \%$ and $10.39 \%$, respectively (Table 5). Substitutions were most abundant at the third-codon position ( $82.80 \%$ )
Table 4 Estimates of genetic structure of the Merodon aureus group and M. funestus (DUR—Durmitor Mt, MOR—Morinj, KOP—Kopaonik Mt, ŠAR—Šar planina Mt, PRO—Prokletije Mt)

| Locus | M. aureus A |  | M. aureus B KOP | M. aureus C |  | M. cinereus AKOP | M. cinereus B |  | M. cinereus C ŠAR | M. funestus <br> MOR |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | DUR | MOR |  | DUR | MOR |  | DUR | PRO |  |  |
| Aat | - | - | - | + | + | - | x | - | - | - |
| Fum | + | - | - | - | - | - | - | + | - | - |
| Gpd-2 | - | - | - | - | - | - | - | - | - | - |
| Gpi | + | + | + | - | + | + | x | - | + | + |
| Had | - | - | - | + | - | + | x | - | + | + |
| Idh-2 | - | - | - | - | - | - | - | - | - | + |
| Mdh-2 | - | x | - | - | + | - | - | + | - | - |
| Me | + | - | + | + | + | + | - | + | - | + |
| Pgm | - | - | - | + | - | + | - | - | - | - |
| Sod-1 | + | + | - | + | + | - | - | + | - | + |
| $n$ (SE) | 9.5 (0.4) | 21.7 (1.0) | 20.2 (0.6) | 16.2 (0.2) | 4.7 (0.2) | 31.2 (1.6) | 30.9 (1.6) | 3.7 (1.3) | 4.9 (0.1) | 19.5 (0.5) |
| Mean | 15.6 |  |  | 10.45 |  |  | 17.3 |  |  |  |
| U | 2 | 1 | - | 2 | 4 | - | 3 | 5 | - | - |
| $A$ (SE) | 1.3 (0.1) | 1.2 (0.1) | 1.3 (0.2) | 1.3 (0.1) | 1.5 (0.2) | 1.3 (0.2) | 1.2 (0.1) | 1.3 (0.1) | 1.1 (0.1) | 1.5 (0.2) |
| Mean | 1.25 |  |  | 1.4 |  |  | 1.25 |  |  |  |
| $P_{(0.95)}$ | 0.267 | 0.133 | 0.200 | 0.333 | 0.333 | 0.267 | 0.000 | 0.267 | 0.133 | 0.333 |
| Mean | 0.200 |  |  | 0.333 |  |  | 0.134 |  |  |  |
| $P_{(0.99)}$ | 0.267 | 0.200 | 0.200 | 0.333 | 0.333 | 0.267 | 0.200 | 0.267 | 0.133 | 0.333 |
| Mean | 0.234 |  |  | 0.333 |  |  | 0.234 |  |  |  |
| $H_{0}$ (SE) | $\begin{aligned} & 0.019 \\ & (0.019) \end{aligned}$ | $\begin{aligned} & 0.027 \\ & (0.027) \end{aligned}$ | $\begin{aligned} & 0.057 \\ & (0.036) \end{aligned}$ | $\begin{aligned} & 0.021 \\ & (0.021) \end{aligned}$ | $\begin{aligned} & 0.033 \\ & (0.033) \end{aligned}$ | $\begin{aligned} & 0.031 \\ & \quad(0.031) \end{aligned}$ | $\begin{aligned} & 0.005 \\ & (0.003) \end{aligned}$ | 0.000 | $\begin{aligned} & 0.013 \\ & (0.013) \end{aligned}$ | 0.000 |
| Mean | 0.023 |  |  | 0.027 |  |  | 0.003 |  |  |  |
| $H_{e}(\mathrm{SE})$ | $\begin{aligned} & 0.065 \\ & (0.032) \end{aligned}$ | $\begin{aligned} & 0.067 \\ & (0.042) \end{aligned}$ | $\begin{aligned} & 0.057 \\ & (0.036) \end{aligned}$ | $\begin{aligned} & 0.096 \\ & (0.040) \end{aligned}$ | $\begin{aligned} & 0.152 \\ & (0.062) \end{aligned}$ | $\begin{aligned} & 0.100 \\ & (0.047) \end{aligned}$ | $\begin{aligned} & 0.008 \\ & (0.005) \end{aligned}$ | $\begin{aligned} & 0.118 \\ & (0.052) \end{aligned}$ | $\begin{aligned} & 0.055 \\ & (0.038) \end{aligned}$ | $\begin{aligned} & 0.109 \\ & (0.048) \end{aligned}$ |
| Mean | 0.066 |  |  | 0.124 |  |  | 0.063 |  |  |  |

$n=$ Mean sample size per locus; $\mathrm{SE}=$ standard error; $A=$ Mean number of alleles per locus; $P=$ Frequency of polymorphic loci based on the criterion of 0.95 (+); 0.99 (x); monomorphic locus $(-) ; H_{\mathrm{o}}=$ Average frequency of observed heterozygosity; $H_{e}=$ Average frequency of expected heterozygosity


Fig. 3 Genetic distance among populations of the M. aureus and $M$. cinereus complexes and $M$. funestus (black bars) and when M. avidus out group was included (patterned bars). M. funestus clearly separates from the ( $M$. aureus + M. cinereus) group (II: average $D=0.89 \pm 0.06$ ) while the average $D$ within the $M$. aureus + M. cinereus group is $0.21 \pm 0.1(\mathrm{I})$. The average distance within the M. aureus complex $(0.22 \pm 0.15)$ is larger than within the $M$. cinereus complex $(0.12 \pm 0.05)$. The average genetic distance from the outgroup M. avidus is the largest for M. funestus (IV: $1.99 \pm 0.07$ ), while the ( $M$. aureus $+M$. cinereus) group average distance from M. avidus is $1.03 \pm 0.13$ (III)


Fig. 4 UPGMA dendrogram (based on Unbiased Genetic Distance; Nei 1978) generated using allozyme data for 10 populations of seven species of the M. aureus group and M. funestus. Goodness of fit statistics: $\quad f \quad($ Farris 1972$)=2.12 ; \quad F \quad($ Prager and Wilson 1976) $=13.79 ; \%$ standard deviation (Fitch and Margoliash 1967) $=29.07 ;$ Cophenetic correlation $=0.98$
and $17.20 \%$ of mutations were at the first position. Out of a total of 93 variable sites, 16 were non-synonymous, and all of these occurred at the first-codon position.

The parsimony analysis of 23 ingroup terminals of the aureus-cinereus complex using Merodon avidus A species as outgroup (including sequence data of three additional $M$. avidus specimens, two M. avidus B and one M. avidus A) resulted in one parsimonious tree with a length of 158 steps $\mathrm{CI}=0.87, \mathrm{RI}=0.96$ (Fig. 2). The resulting tree had two main clades, (M. aureus A + M. aureus B ) and ( $(M$. aureus $\mathrm{C}+$ M. cinereus $\mathrm{B}+$ M. cinereus C$)+$ M. cinereus A$)$, that differed on average by $5.74 \%$ (uncorrected pairwise divergence).


Fig. 5 Wagner procedure dendrogram using Prevosti's distance (Wright 1978) among 10 populations of the M. aureus group, M. funestus and seven populations of the $M$. avidus outgroup generated using allozyme data. Tree rooted using outgroup method after optimization. Goodness of fit statistics: $f($ Farris 1972 $)=2.79 ; F$ (Prager and Wilson 1976) $=4.19$; \% standard deviation (Fitch and Margoliash 1967) $=12.76 ;$ Cophenetic correlation $=0.99$

## Discussion

Delimitation of genetic units
The present concept of M. aureus comprises three taxa: M. aureus A (haplotypes I and II, alleles at Had, Sod-1), M. aureus B (haplotype III, $\mathrm{Pgm}^{c}$ allele) and M. aureus C (haplotypes IV-VII, alleles at Aat, Had, Me and Sod-1) (Fig. 2; Table 3). Populations that fit the present concept of Merodon cinereus are M. cinereus A (haplotype VIII, alleles at Me locus), M. cinereus B (haplotype VII, although species-specific alleles suggest that $M$. cinereus B DUR and M. cinereus B PRO populations are likely at an early stage of speciation, in the absence of differences at mtDNA and morphological traits they are considered as conspecific for the time being) and $M$. cinereus C (haplotype VII, alleles at Me locus). However, the variability of morphological, allozyme and mtDNA data presented herein supports the notion that in delineation of the species there can be no standard level of divergence (Lipscomb et al. 2003; Will and Rubinoff 2004), and more than one source of characters should be considered in species delimitation.

The current study highlights the importance of integrating molecular and morphological characters (e.g.

Table 5 Raw fixed differences (lower matrix) and Uncorrected (" $p$ ") distance matrix for haplotypes of the M. aureus, M. cinereus complexes and M. funestus (I: M. aureus A MOR, M. aureus A DUR; II: M. aureus A MOR; III: M. aureus B KOP; IV: M. aureus C

DUR; V: M. aureus C DUR; VI: M. aureus C MOR; VII: M. aureus C MOR, M. cinereus C ŠAR, M. cinereus B DUR, M. cinereus B PRO; VIII: M. cinereus A KOP; IX: M. funestus MOR; X: M. funestus MOR)

| Haplotype | I | II | III | IV | V | VI | VII | VIII | IX | X |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| I | - | 0.14 | 1.12 | 5.77 | 5.49 | 5.91 | 5.77 | 6.03 | 7.46 | 7.61 |
| II | 1 | - | 1.27 | 5.91 | 5.63 | 6.06 | 5.91 | 6.18 | 7.32 | 7.46 |
| III | 8 | 9 | - | 5.49 | 5.21 | 5.63 | 5.59 | 5.45 | 7.75 | 7.89 |
| IV | 42 | 42 | 40 | - | 0.29 | 0.84 | 0.28 | 0.71 | 9.58 | 9.72 |
| V | 41 | 41 | 39 | 1 | - | 1.13 | 0.57 | 1.01 | 9.58 | 9.72 |
| VI | 43 | 44 | 41 | 6 | 7 | - | 0.85 | 0.86 | 9.72 | 9.86 |
| VII | 42 | 43 | 40 | 2 | 3 | 6 | - | 0.43 | 9.58 | 9.72 |
| VIII | 42 | 43 | 38 | 6 | 7 | 7 | 4 | - | 9.42 | 9.57 |
| IX | 53 | 52 | 55 | 69 | 70 | 70 | 69 | 65 | - | 0.42 |
| X | 54 | 53 | 56 | 70 | 71 | 71 | 70 | 66 | 3 | - |

Moritz and Cicero 2004; Rubinoff and Holland 2005; Rubinoff 2006), especially when dealing with closely related species, recently diverged taxa and taxa in the progress of divergence and speciation (Avise 2000; Funk and Omland 2003). Both allozymes (diagnostic loci and a species-specific allele) and COI mtDNA (unique haplotypes) allowed determining species boundaries for morphologically inseparable M. aureus A and M. aureus B. However, while species-specific allelomorphs clearly delineated cryptic taxa within the morphologically defined M. aureus and M. cinereus species, the allozyme data were not informative enough to conclusively resolve the status of the spring and summer generations of $M$. funestus. On the other hand, morphological and mtDNA COI markers failed to discriminate some reproductively isolated but recently separated sibling taxa although they were clearly defined by allozyme markers. Relatively high gene flow might have prevented differentiation at loci not experiencing selection, which may explain low variability in morphology and sequenced mtDNA. High similarity of morphological traits in cryptic or unrecognized species is often a result of multiple factors, such as retention of ancestral morphology and convergent morphological evolution due to similar selection pressures (cf. Funk and Omland 2003). Low mtDNA diversity could reflect both recent common ancestry and stabilizing selection, or small ancestral effective population size (Hedrick et al. 2006). Additionally, longer coalescence time for mtDNA than for nuclear alleles could be results of the existence of polygynous mating systems and sex-biased dispersal (female philopatry) (Hoelzer 1997). Furthermore, introgression following hybridization and paralogy resulting from transfer of mtDNA gene copies to nucleus might limit the utility of mtDNA in recognition of species boundaries (Moritz and Cicero 2004).

Genetic and phylogenetic relationships

The observed spatial distribution of the taxonomic diversity of the aureus group suggests that the group is of Mediterranean ancestry, while relatively low differentiation among taxa indicates that lineages are at an early stage of divergence. As the larvae are internal feeders in bulbs and rhizomes (Roteray 1993), the relationship of Merodon species with bulb plants could be one of the main drivers of diversity and distribution of the genus. Like the Merodon genus, the center of diversity of bulb plants is the Mediterranean region (Mensel et al. 1965). One possible mechanism of diversification within the aureus group might be the fragmentation of a widespread ancestor and divergence in multiple glacial refugia during the Pleistocene. Another mechanism of diversification of the aureus and cinereus complexes might include founder events during recolonization of previously glaciated areas, and drift among allopatric, refugial populations. The influence of drift is supported by the very low mtDNA haplotype diversity. Using a conservative calibration factor of $1 D=5 \mathrm{Myr}$ (Nei 1987) the earliest divergence was observed for the species pair M. aureus B and . aureus $\mathrm{C}(1.76 \mathrm{Ma})$, while M. cinereus A and $M$. cinereus C appear to be the most closely related species (time of divergence estimated at 0.33 Ma ).

Sequencing data of two specimens of the M. cinereus complex from Pelister (FYR Macedonia) (Ståhls unpublished) showed a unique haplotype in a phylogeographically distant population and its closer relationships to geographically close species $M$. cinereus C and $M$. cinereus B populations than to geographically distant $M$. cinereus A . Taking into account that younger lineages typically occupy more restricted ranges than older lineages (Neigel and Avise
1993), M. aureus B and populations within the M. cinereus $\mathrm{A} / M$. cinereus C lineage seem to have diverged into genetically distinct gene pools only recently. Comparing mtDNA sequences from our study with the sequences of the members of the aureus group, M. aureus B was most closely related to the morphologically defined M. aureus species from Germany, with only one variable nucleotide site (Ståhls personal observation). Furthermore, based on the preliminary data, M. unicolor Strobl (Spain) is genetically the closest species to M. aureus B (Vujić et al. personal observation). These results indicated genetic diversity and radiation within M. aureus B lineage. We expect that the ongoing research will give us better insight into evolutionary relationships among spatially fragmented units of M. aureus B.

The UPGMA, Wagner and parsimony analyses presented a different pattern of the genetic relationship among members of the aureus group, which is not surprising considering that Wagner method and phenetic clustering differ in their algorithms and assumptions, and taking into account that nuclear and mitochondrial genomes evolve at different rates. In all runs the species pair M. aureus $\mathrm{A} / M$. aureus B was placed as a monophyletic group and M. funestus was the most distant species. However, placements of the members of M. aureus C and taxa of the cinereus complex were different, reflecting both different diversity of allozyme loci and mt gene and different pattern of evolutionary changes. In all analyses a clearly defined lineage was the $M$. cinereus A taxon, indicating independent evolution of this species. The M. aureus C, M. cinereus B and $M$. cinereus C taxa were well defined by allozyme markers. Despite the lack of the morphological divergence between $M$. cinereus A and $M$. cinereus C , allozyme loci and mtDNA showed clear divergence. In the UPGMA analysis, $M$. cinereus A and $M$. cinereus C were monophyletic, which was inconsistent with mtDNA parsimony analysis and Wagner method. On the other hand, the relationship between the two populations of M. cinereus B depended on DNA markers used. In spite of diagnostic alleles at the Gpi locus and differences in allelic frequencies, these populations shared the same COI haplotype (haplotype VII), which was likely a plesiomorph character.

## Conservation implications

Based on the observed private alleles and genotypes, unique heterozygotes, significant differences of allelic frequencies and recognized cryptic species across the full hierarchy of taxonomic and genetic diversity, it seems that allelic diversity is a suitable representation of the evolutionary potential in the aureus group, as is the case in the ruficornis (Milankov et al. in press) and avidus (Milankov et al. 2001) groups of the Merodon genus. High level of genetic variation within analyzed populations of the
M. aureus group compared to other hoverflies (e.g. Milankov et al. 2005; Ludoški et al. 2004) indicates large evolutionary potential that should be addressed through conservation management. Here we have obtained results that some loci had higher values of genetic differentiation, and we hypothesized that selection might have played a major role in creating genetic divergence among populations (under selective neutrality, all loci would be similarly affected by gene flow, historical effects and genetic drift). Observed higher level of allelic diversity at allozyme loci than in mtDNA COI gene could also be evidence of the influence of selection on the loci that are likely to be under selection or tightly linked loci. Finally, the existence of specific alleles in allochronic and sympatric populations (M. funestus MOR) points to influence of different environments driving genetic divergence among generations, and may be a sign of the intraspecific differentiation above the level of allochronic conspecific populations.

Using the available morphological and genetic data of the Merodon taxa, the present study partially resolved the taxonomic status and evolutionary relationships within the genus, and defined effective conservation units. Bearing in mind that those populations that became isolated in the remaining habitat face a risk of extinction (Bolger et al. 1997), protection of phylogenetic lineages within the sensitive aureus taxa and conservation of detected genetic variation are essential. Most of the localities of the analyzed populations are within already protected areas, such as the National Parks Durmitor, Kopaonik, Šar planina Mts. and Pindos. However, Prokletije Mt and Morinj locality have not been protected. Because of the extraordinary biodiversity, observed endemic and unique genetic diversity, Morinj locality urgently needs to be incorporated into conservation management planning. Morinj is the most threatened locality among all mentioned, due to the high degree of human encroachment, small size of diverse ecosystem patches, and small populations of many rare and endangered hoverflies species (Vujić et al. personal observation). This area was proposed for protection in 2001 but still no decision has been reached.

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[^1]:    ${ }^{\text {a }}$ Locality specific alleles within a species
    b Unique alleles allowing species delineation
    c DUR—Durmitor Mt, MOR—Morinj, KOP—Kopaonik Mt, ŠAR—Šar planina Mt, PRO—Prokletije Mt

